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The Code of Federal Regulations is sold by the Superintendent of Documents. Prices of new books are listed in the first FEDERAL REGISTER issue of each week.

RECOVERY ACCOUNTABILITY AND TRANSPARENCY BOARD

4 CFR Part 200

RIN 0430-AA00

Implementation of Privacy Act of 1974

AGENCY: Recovery Accountability and Transparency Board.

ACTION: Final rule.

SUMMARY: This document institutes the Recovery Accountability and Transparency Board's (Board) final rule implementing a set of procedural regulations under the Privacy Act of 1974 (Privacy Act or the Act), Public Law 93-579, 5 U.S.C. 552a. These regulations have been written to conform to the statutory provisions of the Act. They are intended to expedite the processing of Privacy Act requests received by the Board and to ensure the proper dissemination of information to the public.

DATES: Effective November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Jennifer Dure, General Counsel, (202) 254-7900.

SUPPLEMENTARY INFORMATION: The proposed rule was published in the *Federal Register* on August 3, 2009 (74 FR 38363) for a public comment period to end on October 2, 2009. This rule sets forth the procedures to be used by members of the public when requesting records from the Board under the Privacy Act. It also establishes a timeframe for responses, a fee schedule for copying records, and charges for obtaining information, when applicable.

Public Comment

The Board received one comment on the proposed rule requesting an explanation concerning the differences between the proposed Privacy Act and Freedom of Information Act (FOIA) rules regarding what is procedurally

required in order for an individual to request access to records, in the custody of the Board. A discussion of the comment and the Board's response are set forth below.

Comments on the Proposed Rule and Explanation

Under the Board's proposed Privacy Act rules, all requests should include, among other things, the requesters full name, address, and telephone number. Requests for Privacy Act records may be made in writing, by fax, by telephone, or in person. The commenter contends that there are additional and more stringent requirements placed on a requester who requests access to his or her records in person. More specifically, such a requester must contact the Board's office at least one week before the desired appointment date. In addition, before a requester can review his or her records, the requester must provide proof of identification. Identification should be a valid copy of one of the following: A government ID, a driver's license, a passport, or other current identification that contains both an address and a picture of the requester.

According to the commenter, the process for requesting records under the Board's proposed FOIA rules "seem[s] quite simplified." Under the proposed FOIA rules (74 FR 38366), all requests for records must include the requester's full name, address, and telephone number. Such a request can be made in writing, via e-mail, or via fax. The commenter correctly points out that the proposed FOIA rule does not provide the option of an in-person request. The commenter concluded that the differences in treatment of requesters for access to the Board's Privacy Act records seem unnecessary, especially with respect to the identification information required of a requester seeking information in person.

The commenter correctly points out the difference between the proposed Privacy Act and FOIA rules, but there is a reason for the difference between them which stems from the laws at issue. Briefly, a Privacy Act request is a request from an individual seeking to review and/or make corrections to federal records, maintained and retrieved in an approved system of records, which are about that individual—with very limited

exceptions, no one else can ask for these records. A FOIA request is a request from the general public for copies of specific records maintained by a federal agency—any member of the public can make such a request. When individuals request information about themselves contained in an approved Privacy Act system of records, the request should be handled under the Privacy Act. Requested records about an individual not contained in an approved system of records asked for under the Privacy Act will have their request processed under the FOIA, since no access rights exist under the Privacy Act.

Because the nature of a Privacy Act request is narrow and specific to an individual in an approved system of records, the Board feels that providing the additional provisions to request and examine records in person is reasonable. In addition, in order to ensure that individuals who request to examine records in person are who they claim to be, it is necessary to require that individuals provide the proper proof of identification as set forth in the proposed Privacy Act rules. This Privacy Act requirement is designed to protect requesters from having their personal information disclosed to anyone else.

Executive Order 12866

The proposed regulation does not meet the criteria for a significant regulatory action under Executive Order 12866. Therefore, review by the Office of Management and Budget is not required.

Regulatory Flexibility Act

The proposed rule adds Privacy Act regulations to 4 CFR Part 200 and will not have a significant economic impact on a substantial number of small entities.

Paperwork Reduction Act

The rule imposes no additional recording and recordkeeping requirements and is therefore exempt from the requirements of the Paperwork Reduction Act.

List of Subjects in 4 CFR Part 200

Administrative practice and procedure, Privacy, Reporting and recordkeeping requirements.

■ Therefore, the Board amends Title 4 of the Code of Federal Regulations by adding Part 200 to read as follows:

CHAPTER II—RECOVERY ACCOUNTABILITY AND TRANSPARENCY BOARD

PART 200—PRIVACY ACT OF 1974

- 200.1 Purpose and scope.
- 200.2 Definitions.
- 200.3 Privacy Act records maintained by the Board.
- 200.4 Privacy Act inquiries.
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- 200.6 Processing of requests.
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- 200.8 Appealing denials of access.
- 200.9 Requests for correction of records.
- 200.10 Disclosure of records to third parties.
- 200.11 Maintaining records of disclosures.
- 200.12 Notification of systems of Privacy Act records.
- 200.13 Privacy Act training.
- 200.14 Responsibility for maintaining adequate safeguards.
- 200.15 Systems of records covered by exemptions.
- 200.16 Mailing lists.

Authority: 5 U.S.C. 552a(f).

§ 200.1 Purpose and scope.

This part sets forth the policies and procedures of the Board regarding access to systems of records maintained by the Board under the Privacy Act, Public Law 93–579, 5 U.S.C. 552a. The provisions in the Act shall take precedence over any part of the Board's regulations in conflict with the Act. These regulations establish procedures by which an individual may exercise the rights granted by the Privacy Act to determine whether a Board system of records contains a record pertaining to him or her; to gain access to such records; and to request correction or amendment of such records. These regulations also set identification requirements and prescribe fees to be charged for copying records.

§ 200.2 Definitions.

As used in this part:

(a) *Agency* means any executive department, military department, government corporation, or other establishment in the executive branch of the federal government, including the Executive Office of the President or any independent regulatory agency;

(b) *Individual* means any citizen of the United States or an alien lawfully admitted for permanent residence;

(c) *Maintain* means to collect, use, store, or disseminate records as well as any combination of these recordkeeping functions. The term also includes exercise of control over, and therefore responsibility and accountability for, systems of records;

(d) *Record* means any item, collection, or grouping of information about an

individual that is maintained by the Board and contains the individual's name or other identifying information, such as a number or symbol assigned to the individual or his or her fingerprint, voice print, or photograph. The term includes, but is not limited to, information regarding an individual's education, financial transactions, medical history, and criminal or employment history;

(e) *System of records* means a group of records under the control of the Board from which information is retrievable by use of the name of the individual or by some number, symbol, or other identifying particular assigned to the individual;

(f) *Routine use* means, with respect to the disclosure of a record, the use of a record for a purpose that is compatible with the purpose for which it was collected;

(g) *Designated Privacy Act Officer* means the person named by the Board to administer the Board's activities in regard to the regulations in this part;

(h) *Executive Director* means the chief operating officer of the Board;

(i) *Days* means standard working days, excluding weekends and federal holidays.

§ 200.3 Privacy Act records maintained by the Board.

(a) The Board shall maintain only such information about an individual as is relevant and necessary to accomplish a purpose of the agency required by statute or by Executive Order of the President. In addition, the Board shall maintain all records that are used in making determinations about any individual with such accuracy, relevance, timeliness, and completeness as is reasonably necessary to ensure fairness to that individual in the making of any determination about him or her. However, the Board shall not be required to update retired records.

(b) The Board shall not maintain any record about any individual with respect to or describing how such individual exercises rights guaranteed by the First Amendment of the Constitution of the United States, unless expressly authorized by statute or by the subject individual, or unless pertinent to and within the scope of an authorized law enforcement activity.

§ 200.4 Privacy Act inquiries.

(a) *Inquiries regarding the contents of record systems.* Any person wanting to know whether the Board's systems of records contain a record pertaining to him or her may file an inquiry in person, by mail or by telephone.

(b) *Inquiries in person* may be submitted at the Board's headquarters

located at 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006. Inquiries should be marked "Privacy Act Inquiry" on each page of the inquiry and on the front of the envelope and directed to the Privacy Act Officer.

(c) *Inquiries by mail* may be sent to: Privacy Act Officer, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006. "Privacy Act Inquiry" should be written on the envelope and each page of the inquiry.

(d) *Telephone inquiries* may be made by calling the Board's Privacy Act Officer at (202) 254–7900.

§ 200.5 Requests for access to records.

(a) All requests for records should include the following information:

(1) Full name, address, and telephone number of requester.

(2) The system of records containing the desired information.

(3) Any other information that the requester believes would help locate the record.

(b) *Requests in writing.* A person may request access to his or her own records in writing by addressing a letter to: Privacy Act Officer, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006.

(c) *Requests by fax.* A person may request access to his or her records by facsimile at (202) 254–7970.

(d) *Requests by phone.* A person may request access to his or her records by calling the Privacy Act Officer at (202) 254–7900.

(e) *Requests in person.* Any person may examine and request copies of his or her own records on the Board's premises. The requester should contact the Board's office at least one week before the desired appointment date. This request may be made to the Privacy Act Officer in writing or by calling (202) 254–7900. Before viewing the records, proof of identification must be provided. The identification should be a valid copy of one of the following:

(1) A government ID;

(2) A driver's license;

(3) A passport; or

(4) Other current identification that contains both an address and a picture of the requester.

§ 200.6 Processing of requests.

Upon receipt of a request for information, the Privacy Act Officer will ascertain whether the records identified by the requester exist, and whether they are subject to any exemption under § 200.15. If the records exist and are not subject to exemption, the Privacy Act Officer will provide the information.

(a) *Requests in writing, including those sent by fax.* Within five working days of receiving the request, the Privacy Act Officer will acknowledge its receipt and will advise the requester of any additional information that may be needed. Within 15 working days of receiving the request, the Privacy Act Officer will send the requested information or will explain to the requester why additional time is needed for a response.

(b) *Requests in person or by telephone.* Within 15 days of the initial request, the Privacy Act Officer will contact the requester and arrange an appointment at a mutually agreeable time when the record can be examined. The requester may be accompanied by no more than one person. In such case, the requestor must inform the Privacy Act Officer that a second individual will be present and must sign a statement authorizing disclosure of the records to that person. The statement will be kept with the requester's records. At the appointment, the requester will be asked to present identification as stated in § 200.5(e).

(c) *Excluded information.* If a request is received for information compiled in reasonable anticipation of litigation, the Privacy Act Officer will inform the requester that the information is not subject to release under the Privacy Act (see 5 U.S.C. 552a(d)(5)).

§ 200.7 Fees.

A fee will not be charged for searching, reviewing, or making corrections to records. A fee for copying will be assessed at the same rate established for the Freedom of Information Act requests. Duplication fees for paper copies of a record will be 10 cents per page for black and white and 20 cents per page for color. For all other forms of duplication, the Board will charge the direct costs of producing the copy. However, the first 100 pages of black-and-white copying or its equivalent will be free of charge.

§ 200.8 Appealing denials of access.

(a) If access to records is denied by the Privacy Act Officer, the requester may file an appeal in writing. The appeal should be directed to Executive Director, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006.

(b) The appeal letter must specify the denied records that are still sought, and state why denial by the Privacy Act Officer is erroneous.

(c) The Executive Director or his or her designee will respond to appeals within 20 working days of the receipt of

the appeal letter. The appeal determination will explain the basis of the decision to deny or grant the appeal.

§ 200.9 Requests for correction of records.

(a) *Correction requests.* Any person is entitled to request correction of his or her record(s) covered under the Act. The request must be made in writing and should be addressed to Privacy Act Officer, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006. The letter should clearly identify the corrections desired. In most circumstances, an edited copy of the record will be acceptable for this purpose.

(b) *Initial response.* Receipt of a correction request will be acknowledged by the Privacy Act Officer in writing within five working days. The Privacy Act Officer will provide a letter to the requester within 20 working days stating whether the request for correction has been granted or denied. If the Privacy Act Officer denies any part of the correction request, the reasons for the denial will be provided to the requester.

§ 200.10 Disclosure of records to third parties.

(a) The Board will not disclose any record that is contained in a system of records to any person or agency, except with a written request by or with the prior written consent of the individual whose record is requested, unless disclosure of the record is:

(1) Required by an employee or agent of the Board in the performance of his/her official duties.

(2) Required under the provisions of the Freedom of Information Act (5 U.S.C. 552). Records required to be made available by the Freedom of Information Act will be released in response to a request in accordance with the Board's regulation published at 4 CFR Part 201.

(3) For a routine use as published in the annual notice in the **Federal Register**.

(4) To the Census Bureau for planning or carrying out a census, survey, or related activities pursuant to the provisions of Title 13 of the United States Code.

(5) To a recipient who has provided the Board with adequate advance written assurance that the record will be used solely as a statistical research or reporting record and that the record is to be transferred in a form that is not individually identifiable.

(6) To the National Archives and Records Administration as a record that has sufficient historical or other value to

warrant its continued preservation by the United States government, or for evaluation by the Archivist of the United States, or his or her designee, to determine whether the record has such value.

(7) To another agency or to an instrumentality of any governmental jurisdiction within or under the control of the United States for a civil or criminal law enforcement activity, if the activity is authorized by law, and if the head of the agency or instrumentality has made a written request to the Board for such records specifying the particular part desired and the law enforcement activity for which the record is sought. The Board also may disclose such a record to a law enforcement agency on its own initiative in situations in which criminal conduct is suspected, provided that such disclosure has been established as a routine use, or in situations in which the misconduct is directly related to the purpose for which the record is maintained.

(8) To a person pursuant to a showing of compelling circumstances affecting the health or safety of an individual if, upon such disclosure, notification is transmitted to the last known address of such individual.

(9) To either House of Congress, or to the extent of matters within its jurisdiction, any committee or subcommittee thereof, any joint committee of Congress or subcommittee of any such joint committee.

(10) To the Comptroller General, or any of his or her authorized representatives, in the course of the performance of official duties of the Government Accountability Office.

(11) Pursuant to an order of a court of competent jurisdiction. In the event that any record is disclosed under such compulsory legal process, the Board shall make reasonable efforts to notify the subject individual after the process becomes a matter of public record.

(12) To a consumer reporting agency in accordance with 31 U.S.C. 3711(e).

(b) Before disseminating any record about any individual to any person other than a Board employee, the Board shall make reasonable efforts to ensure that the records are, or at the time they were collected, accurate, complete, timely, and relevant. This paragraph (b) does not apply to disseminations made pursuant to the provisions of the Freedom of Information Act (5 U.S.C. 552) and paragraph (a)(2) of this section.

§ 200.11 Maintaining records of disclosure.

(a) The Board shall maintain a log containing the date, nature, and

purposes of each disclosure of a record to any person or agency. Such accounting also shall contain the name and address of the person or agency to whom or to which each disclosure was made. This log will not include disclosures made to Board employees or agents in the course of their official duties or pursuant to the provisions of the Freedom of Information Act (5 U.S.C. 552).

(b) An accounting of each disclosure shall be retained for at least five years after the accounting is made or for the life of the record that was disclosed, whichever is longer.

(c) The Board shall make the accounting of disclosure of a record pertaining to an individual available to that individual at his or her request. Such a request should be made in accordance with the procedures set forth in § 200.5. This paragraph (c) does not apply to disclosure made for law enforcement purposes under 5 U.S.C. 552a(b)(7) and § 200.10(a)(7).

§ 200.12 Notification of systems of Privacy Act records.

(a) Public Notice. The Board periodically reviews its systems of records and will publish information about any significant additions or changes to those systems in the **Federal Register**. Information about systems of records maintained by other agencies that are in the temporary custody of the Board will not be published. In addition, the Office of the Federal Register biennially compiles and publishes all systems of records maintained by all federal agencies, including the Board.

(b) At least 30 days before publishing additions or changes to the Board's systems of records, the Board will publish a notice of intent to amend, providing the public with an opportunity to comment on the proposed amendments to its systems of records in the **Federal Register**.

§ 200.13 Privacy Act training.

(a) The Board shall ensure that all persons involved in the design, development, operation, or maintenance of any Board systems of records are informed of all requirements necessary to protect the privacy of individuals. The Board shall ensure that all employees having access to records receive adequate training in their protection and that records have adequate and proper storage with sufficient security to ensure their privacy.

(b) All employees shall be informed of the civil remedies provided under 5 U.S.C. 552a(g)(1) and other implications

of the Privacy Act and of the fact that the Board may be subject to civil remedies for failure to comply with the provisions of the Privacy Act and the regulations in this part.

§ 200.14 Responsibility for maintaining adequate safeguards.

The Board has the responsibility for maintaining adequate technical, physical, and security safeguards to prevent unauthorized disclosure or destruction of manual and automated records systems. These security safeguards shall apply to all systems of records in which identifiable personal data are processed or maintained, including all reports and output from such systems of records that contain identifiable personal information. Such safeguards must be sufficient to prevent negligent, accidental, or unintentional disclosure, modification, or destruction of any personal records or data; must minimize, to the extent practicable, the risk that skilled technicians or knowledgeable persons could improperly obtain access to modify or destroy such records or data; and shall further ensure against such casual entry by unskilled persons without official reasons for access to such records or data.

(a) *Manual systems.* (1) Records contained in a system of records as defined in this part may be used, held, or stored only where facilities are adequate to prevent unauthorized access by persons within or outside the Board.

(2) Access to and use of a system of records shall be permitted only to persons whose duties require such access to the information for routine uses or for such other uses as may be provided in this part.

(3) Other than for access by employees or agents of the Board, access to records within a system of records shall be permitted only to the individual to whom the record pertains or upon his or her written request.

(4) The Board shall ensure that all persons whose duties require access to and use of records contained in a system of records are adequately trained to protect the security and privacy of such records.

(5) The disposal and destruction of identifiable personal data records shall be done by shredding and in accordance with rules promulgated by the Archivist of the United States.

(b) *Automated systems.* (1) Identifiable personal information may be processed, stored, or maintained by automated data systems only where facilities or conditions are adequate to prevent unauthorized access to such systems in any form.

(2) Access to and use of identifiable personal data associated with automated data systems shall be limited to those persons whose duties require such access. Proper control of personal data in any form associated with automated data systems shall be maintained at all times, including maintenance of accountability records showing disposition of input and output documents.

(3) All persons whose duties require access to processing and maintenance of identifiable personal data and automated systems shall be adequately trained in the security and privacy of personal data.

(4) The disposal and disposition of identifiable personal data and automated systems shall be done by shredding, burning, or, in the case of electronic records, by degaussing or by overwriting with the appropriate security software, in accordance with regulations of the Archivist of the United States or other appropriate authority.

§ 200.15 Systems of records covered by exemptions.

The Board currently has no exempt systems of records.

§ 200.16 Mailing lists.

The Board shall not sell or rent an individual's name and/or address unless such action is specifically authorized by law. This section shall not be construed to require the withholding of names and addresses otherwise permitted to be made public.

Ivan J. Flores,

Paralegal Specialist, Recovery Accountability and Transparency Board.

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RECOVERY ACCOUNTABILITY AND TRANSPARENCY BOARD

4 CFR Part 201

RIN 0430-AA01

Rule Implementing the Freedom of Information Act

AGENCY: Recovery Accountability and Transparency Board.

ACTION: Final rule.

SUMMARY: This document institutes the Recovery Accountability and Transparency Board's (Board) final rule implementing a set of procedural regulations under the Freedom of Information Act (FOIA) in accordance with 5 U.S.C. 552, and Public Law 104-231, the Electronic Freedom of

Information Act Amendments of 1996. These regulations have been written to conform to the statutory provisions of the Acts, to expedite the processing of FOIA requests received by the Board, and to ensure the proper dissemination of information to the public.

DATES: Effective November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Jennifer Dure, General Counsel, (202) 254-7900.

SUPPLEMENTARY INFORMATION: The proposed rule was published in the **Federal Register** on August 3, 2009 (74 FR 38366) for a public comment period to end on October 2, 2009. This rule sets forth the procedures for members of the public to request records from the Board under both the FOIA and the Electronic Freedom of Information Act Amendments of 1996. The rule also sets forth the procedures that the Board will use when responding to such requests. It sets forth the time frames for responses and the current fee schedule for any applicable charges for information. The rule also supplies information about Board materials available to the public through the Board's Web site.

Public Comment

The Board received comments from two organizations. One commenter requested an explanation concerning the differences between the proposed FOIA and Privacy Act rules regarding what is procedurally required in order for an individual to request access to records in the custody of the Board. Under the Board's proposed Privacy Act rule, all requests should include, among other things, the requester's full name, address, and telephone number. Requests for Privacy Act records may be made in writing, by fax, by telephone, or *in person*. The commenter contends that there are additional and more stringent requirements placed on a requester who requests access to his or her records in person. More specifically, such a requester must contact the Board's office at least one week before the desired appointment date. In addition, before a requester can review his or her records, the requester must provide proof of identification. Identification should be a valid copy of one of the following: A government ID, a driver's license, a passport, or other current identification that contains both an address and a picture of the requester.

According to the commenter, the process for requesting records under the Board's proposed FOIA rule "seem[s] quite simplified." Under the proposed FOIA rule, all requests for records must

include the requester's full name, address, and telephone number. Such a request can be made in writing, via e-mail, or via fax. The commenter correctly points out that the proposed FOIA rule does not provide the option of an *in-person request*. The commenter concluded that the differences in treatment of requesters for access to the Board's records seem unnecessary, especially with respect to the identification information required of a requester seeking information in person.

The commenter correctly points out the difference between the proposed Privacy Act and FOIA rules, but there is a reason for the difference between them which stems from the laws at issue. Briefly, a Privacy Act request is a request from an individual seeking to review and/or make corrections to federal records, maintained and retrieved in an approved system of records, which are about that individual—with very limited exceptions, no one else can ask for these records. A FOIA request is a request from the general public for copies of specific records maintained by a federal agency—any member of the public can make such a request. When individuals request information about themselves contained in an approved Privacy Act system of records, the request should be handled under the Privacy Act. Requested records about an individual not contained in an approved system of records asked for under the Privacy Act will have their request processed under the FOIA, since no access rights exist under the Privacy Act.

Because the nature of a Privacy Act request is narrow and specific to an individual in an approved system of records, the Board feels that providing the additional provisions to request and examine records in person is reasonable. In addition, in order to ensure that individuals who request to examine records in person are who they claim to be, it is necessary to require that individuals provide the proper proof of identification as set forth in the proposed Privacy Act rules. This Privacy Act requirement is designed to protect requesters from having their personal information disclosed to anyone else.

The other commenter raised concerns regarding the Board's proposed definition of "agency records" under § 201.2. The Board's proposed FOIA rule defines "agency record" as "materials that are in the control of the Board and associated with Board business, as follows: (i) Materials produced by the Board. (ii) Materials produced by staff for the Board. (iii) Materials distributed by presenters at a

Board meeting or Board Committee meeting." The commenter feels that the proposed definition is too narrow. The Board agrees and has therefore modified its definition in a way as to leave open the types of information that may be considered "agency records."¹

The same commenter raised concerns regarding § 201.3 of the Board's proposed rule—publicly available documents and the electronic reading room. More specifically, § 201.3(b)(6) of the Board's proposed rule provides that "[r]ecords available electronically on the Board's Web site include * * * [c]opies of records repeatedly released in response to FOIA requests." The commenter is concerned that this provision suggests that the Board will make available a narrower category of records than what is required under FOIA. To alleviate any confusion as to whether the Board will track the law, the Board has modified § 201.3(b)(6) to track the language used in the DOJ Guidance, reflecting its intention to comply with the requirement of proactive disclosure and make records of public interest available prior to receiving frequent requests for such information.

Finally, the same commenter raised concerns regarding § 201.14(c) of the Board's proposed rule—appeals and exhaustion of administrative remedies. The commenter feels that the Board has misstated FOIA regarding when a FOIA requester may bring a lawsuit in federal court to challenge an agency's response to his or her FOIA request. The Board feels that proposed § 201.14(c) and (e) confuse the administrative appeals/judicial review issues and therefore withdraws both provisions in their entirety. The Board believes that the case law on this matter—referenced in the U.S. Department of Justice's "Guide to the Freedom of Information Act,"—speaks for itself.² As revised, the Board's regulations provide requesters with sufficient procedural information to ensure the proper review of requests.

Executive Order No. 12866

These proposed regulations do not meet the criteria for a significant regulatory action under Executive Order 12866. Thus, review by the Office of Management and Budget is not required.

Regulatory Flexibility Act

These proposed regulations will not have a significant economic impact on

¹ The Board will follow the U.S. Department of Justice's "Guide to the Freedom of Information Act" in determining what constitutes an agency record. See U.S. Department of Justice "Guide to the Freedom of Information Act," (2009), at 33.

² See *id.* at 97.

a substantial number of small entities. Therefore, a regulatory flexibility analysis as provided by the Regulatory Flexibility Act, as amended, is not required.

Paperwork Reduction Act

These proposed regulations impose no additional reporting and recordkeeping requirements. Therefore, clearance by the Office of Management and Budget is not required.

List of Subjects in 4 CFR Part 201

Administrative practice and procedure; Freedom of Information; Reporting and recordkeeping requirements.

■ Therefore, the Board amends Title 4 of the Code of Federal Regulations by adding Part 201 to read as follows:

CHAPTER II—RECOVERY ACCOUNTABILITY AND TRANSPARENCY BOARD

PART 201—PUBLIC INFORMATION AND REQUESTS

- Sec.
- 201.1 Scope.
 - 201.2 Definitions.
 - 201.3 Publicly available documents and electronic reading room.
 - 201.4 Board records exempt from public disclosure.
 - 201.5 Requests for Board records.
 - 201.6 Responsibility, form, and content of responses.
 - 201.7 Time of responses to requests.
 - 201.8 Fees.
 - 201.9 Restrictions on charging fees.
 - 201.10 Notice of anticipated fees.
 - 201.11 Requirements for waiver or reduction of fees.
 - 201.12 Denials.
 - 201.13 Business information.
 - 201.14 Appeals.
 - 201.15 Preservation of records.
 - 201.16 Other rights and services.
 - 201.17 How to track a FOIA request.

Authority: 5 U.S.C. 301, 5 U.S.C. 552 as amended; Executive Order 12600, 3 CFR, 1987 Comp., p. 235.

§ 201.1 Scope.

This part sets forth the policies and procedures of the Recovery Accountability and Transparency Board (Board) regarding public access to documents under the Freedom of Information Act (FOIA or the Act), 5 U.S.C. 552. The provisions in the Act shall take precedence over any part of the Board's regulations in conflict with the Act. This part gives the procedures the public may use to inspect and obtain copies of Board records under the FOIA, including administrative procedures which must be exhausted before a requestor invokes the jurisdiction of an

appropriate United States District Court for the Board's failure to respond to a proper request within the statutory time limits, for a denial of Board records or challenges to the adequacy of a search, or for denial of fee waiver.

§ 201.2 Definitions.

For words used in this document, unless the context indicates otherwise, singular includes the plural, plural includes the singular, present tense includes the future tense, and words of one gender include the other gender.

(a)(1) *Agency records*—Materials that are in the control of the Board and associated with Board business, including:

(i) Materials produced by the Board.

(ii) Materials produced by staff for the Board.

(iii) Materials distributed by presenters at a Board meeting or Board Committee meeting.

(2) All references to records include the entire record and/or any part of the record.

(b) *Board*—The Recovery Accountability and Transparency Board.

(c) *Chairman*—The Chairman of the Board is designated or appointed by the President.

(d) *Designated FOIA Officer*—The person designated to administer the Board's activities in regard to the regulations in this part. The FOIA Officer shall be:

(1) The Board officer having custody of, or responsibility for, agency records in the possession of the Board.

(2) The Board officer having responsibility for authorizing or denying production of records from requests filed under the FOIA.

(e) *Executive Director*—The chief operating officer of the Board.

(f) *Member*—An individual appointed to serve on the Board pursuant to Title XV, Subtitle B of the American Recovery and Reinvestment Act of 2009 (Pub. L. 111–5).

(g) *Days*—Standard working days, excluding weekends and federal holidays.

§ 201.3 Publicly available documents and electronic reading room.

(a) Many Board records are available electronically at the Board's Web site (<http://www.recovery.gov>).

(b) Records available electronically on the Board's Web site include:

(1) The rules and regulations of the Board.

(2) Statements of policy adopted by the Board.

(3) Board reports to the President and Congress, including the Committees on Appropriations of the Senate and House of Representatives.

(4) Congressional Testimony of the Chairman of the Board.

(5) Biographical information about the Chairman and other Board members.

(6) Copies of records frequently requested and released in response to FOIA requests.

(c) The cost of copying information available in the Board office shall be imposed in accordance with the provisions of § 201.8.

§ 201.4 Board records exempt from public disclosure.

5 U.S.C. 552 provides that the requirements of the FOIA do not apply to matters that are:

(a) Specifically authorized under the criteria established by an executive order to be kept secret in the interest of national defense or foreign policy and are in fact properly classified pursuant to such an executive order.

(b) Related solely to the internal personnel rules and practices of the Board.

(c) Specifically exempted from disclosure by another federal statute, provided that such statute:

(1) Requires that records are withheld from the public in such a manner that leaves no discretion on the issue; or

(2) Establishes criteria for withholding or refers to particular types of matters to be withheld.

(d) Trade secrets, and commercial or financial information obtained from a person and privileged or confidential.

(e) Interagency or intra-agency memoranda or letters that would not be available by law to a party other than an agency in litigation with the Board.

(f) Personnel, medical, or similar files that disclosing would constitute a clearly unwarranted invasion of personal privacy.

(g) Records or information compiled for law enforcement purposes, but only to the extent that the production of such law enforcement records of information:

(1) Could reasonably be expected to interfere with enforcement proceedings;

(2) Would deprive a person of a right to a fair trial or an impartial adjudication;

(3) Could reasonably be expected to constitute an unwarranted invasion of personal privacy;

(4) Could reasonably be expected to disclose the identity of any confidential source, including a state, local, or foreign agency or authority, or any private institution which furnished information on a confidential basis, and in the case of a record or information compiled by a criminal law enforcement agency in the course of a criminal investigation or by an agency conducting a lawful security

intelligence investigation, information furnished by a confidential source;

(5) Would disclose techniques and procedures for law enforcement investigations or prosecutions, or would disclose guidelines for law enforcement investigations or prosecutions if such disclosure could reasonably be expected to risk circumvention of the law; or

(6) Could reasonably be expected to endanger the life or physical safety of any individual.

(h) Contained in or related to examination, operating, or condition reports, prepared by, on behalf of, or for the use of an agency responsible for the regulation or supervision of financial institutions.

(i) Geological and geophysical information and data, including maps, concerning wells.

§ 201.5 Requests for Board records.

(a) To request Board records, you may:

(1) Write: FOIA Officer, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006;

(2) Send a request via e-mail at FOIA@ratb.gov; or

(3) Fax: (202) 254-7970.

(b) When requesting records under this section you must state, in writing:

- (1) Your full name,
- (2) Address,
- (3) Telephone number, and
- (4) At your option, electronic mail address.

(c) When making a request for records about a person, Privacy Act regulations also may apply. Please check the regulations for additional requirements before submitting a request. When making a request for records about someone other than yourself, you must include either:

(1) Written authorization signed by the person permitting you to see the records; or

(2) Proof that the individual is deceased (e.g., a death certificate or obituary).

(d) A request will be considered received for purposes of § 201.7 on the date that it is received by the Board's FOIA office. For prompt handling, write "Freedom of Information Act Request" on the letter and envelope or in the subject line of the e-mail request or fax.

(e) Each request must clearly describe the desired records in sufficient detail to enable Board personnel to locate them with reasonable effort. Response to requests may be delayed if the records are not clearly described.

(f) Whenever possible, requests should include specific information about each record sought, such as date,

title or name, author, recipient, and subject.

(g) If the FOIA Officer determines that the request does not clearly describe the records sought, he or she will either advise you of the additional information needed to locate the record or otherwise state why the request is insufficient. You will then be given the opportunity to provide additional information or to modify your request.

(h) Submitting a FOIA request shall be considered a commitment by the requestor to pay applicable fees required under § 201.8 unless the requestor seeks a waiver of fees. When making a request, you may specify a willingness to pay fees up to a specific amount.

(i) The FOIA does not require the Board to:

(1) Compile or create records solely for the purpose of satisfying a request for records.

(2) Provide records not yet in existence, even if such records may be expected to come into existence at some time in the future.

(3) Restore records destroyed or otherwise disposed of, except that the FOIA Officer must notify the requestor that the records have been destroyed or otherwise disposed of.

§ 201.6 Responsibility, form, and content of response.

The Board's Executive Director or his/her designated FOIA Officer is authorized to grant or deny any request for a record and determine appropriate fees. When determining which records are responsive to a request, the Board will include only records in its possession as of the date of the request.

(a) If no records are responsive to the request, the FOIA Officer will notify the requestor in writing.

(b) When the FOIA Officer denies a request in whole or in part, he/she will notify the requestor in writing. The response will be signed by the FOIA Officer and will include:

(1) The name and title or position of the person making the denial;

(2) A brief statement of the reasons for the denial, including the FOIA exemption(s) that the FOIA Officer has relied on in denying the request; and

(3) A statement that the denial may be appealed under § 201.14 and a description of the requirements of that section.

(c) *Referrals.* When a request for a record not created by the Board is received, the Board shall refer the requestor to the issuing agency in writing, providing the address of the agency contact and the section(s) referred.

(d) *Timing of responses to requests sent to other agencies.* The Board shall

provide, within the FOIA deadline, responses only to those parts of the request not referred.

(e) *Agreements on referrals.* The Board may make agreements with other agencies to eliminate the need for referrals for particular types of records.

§ 201.7 Timing of responses to requests.

(a) *General.* The Board shall normally respond to requests in the order of their receipt.

(b) *Acknowledgement of requests.* On receipt of a request, the Board shall send an acknowledgement letter or an e-mail confirming the requestor's agreement to pay fees under § 201.8 and providing a request number for future reference.

(c) *Time limits for responding to FOIA requests.* The Board shall make an initial determination to grant or deny a request for records within 20 days (excluding Saturday, Sunday and holidays) after the date of receipt of the request, as described in § 201.5(d), except as stated in paragraph (f) of this section. Once the Board determines whether it can grant a request entirely or in part, it shall notify the requestor in writing. The Board shall advise the requestor of any fees to be charged under § 201.8 and shall disclose records promptly on payment of the fees. Records disclosed in part shall be marked or annotated to show the amount of information deleted unless doing so would harm an interest protected by an applicable exemption. The location of the information deleted also shall be indicated on the record when technically feasible.

(d) *Unusual circumstances.* (1) If the statutory time limits for processing a request cannot be met because of "unusual circumstances" as defined in the FOIA (5 U.S.C. 552(6)(B)(iii)), the Board shall promptly notify the requestor in writing, explaining the circumstances and giving the date by which the request can be completed or if the Board cannot complete the request. If the extension is for more than 10 working days, the Board shall provide the requestor with an opportunity to:

(i) Modify the request so that it can be processed within the time limit; or

(ii) Arrange an alternative time period for processing the original request.

(2) If the Board believes that multiple requests submitted by a requestor or by requestors acting in concert constitute a single request that would otherwise involve unusual circumstances, and if the requests involve clearly related matters, they may be aggregated. Multiple requests involving unrelated matters will not be aggregated.

(e) *Expedited processing.* (1) Requests and appeals shall be taken out of order and given expedited processing whenever it is determined that they involve:

(i) Circumstances that could reasonably be expected to pose an imminent threat to the life or physical safety of an individual; or

(ii) An urgency to inform the public about an actual or alleged activity if made by a person primarily engaged in disseminating information.

(2) Requests for expedited processing may be made either at the time of the initial request or at a later time.

(3) Requests for expedited processing must include a statement explaining in detail the basis for requesting expedited processing. For example, a requestor under § 201.8 must establish that his/her professional activity is news reporting, although it need not be his/her sole occupation. The requestor also must establish a particular urgency to inform the public about government activity involved in the request, beyond the public's right to know about government activity generally.

(4) Within 10 calendar days of receipt of a request for expedited processing, the Board shall decide whether to grant the request and notify the requestor of its decision. If a request for expedited treatment is granted, the request shall be processed as soon as practicable. If a request for expedited processing is denied, an appeal of that decision shall be acted on expeditiously.

(f) *Tolling of time limits.* (1) The Board may toll the 20-day time period to:

(i) Make one request for additional information from the requester; or

(ii) Clarify the applicability or amount of any fees, if necessary, with the requester.

(2) The tolling period ends upon the Board's receipt of information from the requester or resolution of the fee issue.

§ 201.8 Fees.

(a) *General.* The Board shall charge for processing requests under the FOIA in accordance with paragraph (c) of this section, except where fees are limited under § 201.9 or where a waiver or reduction of fees is granted under § 201.11. Fees must be paid before the copies of records are sent. Fees may be paid by check or money order payable to the Treasury of the United States.

(b) *Definitions for this section.* (1) *Commercial use request*—A request from, or on behalf of, a person who seeks information for a purpose that furthers his/her commercial, trade, or profit interests including furthering those interests through litigation. The

Board shall try to determine the use to which a record will be put. When the Board believes that a request is for commercial use either because of the nature of the request or because the Board has cause to doubt the stated use, the Board shall ask the requestor for clarification.

(2) *Direct costs*—Expenses that the Board incurs in searching for, duplicating, and reviewing records in response to a request. Direct costs include the full salary of the employee performing the work and the cost of duplication of the records. Overhead expenses, such as the cost of space, heating, and lighting, are not included.

(3) *Duplication*—Making a copy of a record or the information in the record, to respond to a request. Copies can be in paper, electronic, or other format. The Board shall honor a requestor's preference for format if the record is readily reproducible in that format at a reasonable cost.

(4) *Educational institution*—A public or private undergraduate, graduate, professional or vocational school that has a program of scholarly research. For a request to be in this category, a requestor must show that the request is authorized by and made under the auspices of the qualifying institution and that the records will be used for scholarly research.

(5) *Noncommercial scientific institution*—An institution that is not operated on a commercial basis, as defined in paragraph (b)(1) of this section and is operated solely for conducting scientific research that does not promote any particular product or industry. For a request to be in this category, the requestor must show that the request is authorized and made under the auspices of the qualifying institution and that the records will be used for further scientific research.

(6) *Representative of the news media*—Any person who, or entity that, gathers information of potential interest to a segment of the public, uses editorial skills to turn the raw materials into a distinct work, and distributes that work to an audience. A freelance journalist shall be regarded as working for a news media entity if the person can demonstrate a solid basis for expecting publication through that entity, whether or not the journalist is actually employed by that entity. A publication contract is one example of a basis for expecting publication that ordinarily would satisfy this standard. The Board may consider past publication records of the requester in determining whether he or she qualifies as a "representative of the news media."

(7) *Review*—Examining a record to determine whether any part of it is exempt from disclosure, and processing a record for disclosure. Review costs are recoverable even if a record is not disclosed. Review time includes time spent considering any formal objection to disclosure made by a business submitter under § 201.13 but does not include time spent resolving general legal or policy issues regarding the application of exemptions.

(8) *Search*—The process of looking for and retrieving records, including page-by-page or line-by-line identification of information within records and reasonable efforts to locate and retrieve information from records maintained in electronic form. The Board shall ensure that searches are done in the most efficient and least expensive way that is reasonably possible.

(c) *Fees.* In responding to FOIA requests, the Board shall charge the following fees unless a waiver or a reduction of fees has been granted under § 201.11.

(1) *Search.* (i) Search fees shall be charged for all requests subject to the limitations of § 201.9. The Board may charge for time spent searching even if no responsive record is located, or if the record(s) located are withheld as exempt from disclosure.

(ii) For each quarter hour spent by clerical personnel in searching for and retrieving a requested record, the fee will be \$5. If a search and retrieval requires the use of professional personnel, the fee will be \$8 for each quarter hour. If the time of managerial personnel is required, the fee will be \$10 for each quarter hour.

(iii) For computer searches for records, requestors will be charged the direct costs of conducting the search although certain requestors (*see* § 201.9(a)) will be charged no search fee and certain other requestors (*see* § 201.9(b)) will be entitled to two hours of manual search time without charge. Direct costs include the cost of operating a computer for the search time for requested records and the operator salary for the search.

(2) *Duplication.* Duplication fees for paper copies of a record will be 10 cents per page for black and white and 20 cents per page for color. For all other forms of duplication, the Board shall charge the direct costs of producing the copy. All charges are subject to the limitations of §§ 201.9 and 201.11.

(3) *Review.* When a commercial-use request is made, review fees shall be charged as stated in paragraph (c)(1) of this section. These fees apply only to the initial record review, when the Board determines whether an

exemption applies to a particular record. Charges shall not be imposed for review at the administrative appeal level if an exemption is applied. However, records withheld under an exemption that is subsequently determined not to apply may be reviewed again to determine whether any other exemption not previously considered applies. The costs of that review shall be charged. All review fees shall be charged at the same rates as those charged in paragraph (c)(1) of this section.

§ 201.9 Restrictions on charging fees.

(a) When determining search or review fees:

(1) No search fee shall be charged for requests by educational institutions, noncommercial scientific institutions, or representatives of the news media.

(2) The Board shall provide without charge to all but commercial users:

(i) The first 100 pages of black and white duplication (or the cost equivalent); and

(ii) The first two hours of search by a clerical staff member (or the cost equivalent).

(3) When the total fee for a request will be \$14.00 or less for any request, no fee shall be charged.

(b) The Board will not assess search and/or duplication fees, as applicable, if it fails to respond to a requester's FOIA request within the time limits specified under 4 CFR 201.7, and no "unusual" circumstances (as defined in 5 U.S.C. 552(a)(6)(B) and 4 CFR 201.7(d)) or "exceptional" circumstances (as defined in 5 U.S.C. 552(a)(6)(C)) apply to the processing of the request.

§ 201.10 Notice of anticipated fees.

(a) *General.* The Board shall advise the requestor in writing of any applicable fees. If only a part of the fee can be estimated readily, the Board shall advise the requestor that this may be only a part of the total fee. After the requestor has been sent a fee estimate, the request shall not be considered received until the requestor makes a firm commitment to pay the anticipated total fee. Any such agreement must be made by the requestor in writing and must be received within 60 days of the Board's notice. If the requestor does not provide a firm commitment to pay the anticipated fee within 60 days of the notice, the request shall be closed. The requestor may be given an opportunity to work with the Board to change the request and lower the cost.

(b) *Charges for other services.* When the Board chooses as a matter of administrative discretion to provide a special service, such as certifying that

records are true copies or sending them by other than ordinary mail, the Board shall pay the costs of providing the services unless previous arrangements have been made with the requestor.

(c) *Charging interest.* The Board may charge interest on any unpaid bill starting on the 31st day following the date of billing. Interest charges shall be assessed at the rate provided in 31 U.S.C. 3717 and shall accrue from the date of the billing until payment is received by the Board. The Board shall follow the provisions of the Debt Collection Act of 1982 (Pub. L. 97-365, 96 Stat. 1749), as amended.

(d) *Aggregating requests.* If the Board reasonably believes that a requestor or a group of requestors acting together is trying to divide a request into a series of smaller requests for the purpose of avoiding fees, the Board may aggregate the requests and charge accordingly. The Board shall assume that multiple requests of the same type made within a 30-day period have been made in order to avoid fees. If requests are separated by a longer period, the Board shall aggregate them only if there is a solid basis for determining that aggregation is warranted. Multiple requests involving unrelated matters shall not be aggregated.

(e) *Advance payments.* When a requestor has previously failed to pay promptly a properly charged FOIA fee to the Board or another agency, the Board shall require proof that full payment has been made to that agency before it begins to process that requestor's FOIA request. The Board shall also require advance payment of the full amount of the anticipated fee. When advance payment is required, the request is not considered received until payment has been made.

§ 201.11 Requirements for waiver or reduction of fees.

(a) Fees for processing your request may be waived if you meet the criteria listed in paragraph (b) of this section. The burden is on you to justify entitlement to a fee waiver. Requests for fee waivers are decided on a case-by-case basis. The fact that you have received a fee waiver in the past does not mean you are automatically entitled to a fee waiver for every request you may submit, because the essential element of any fee waiver determination is whether the release of the particular documents sought in the request will likely contribute significantly to public understanding of the operations or activities of the government. The Board will rely on the fee waiver justification you have submitted in your request letter. If you do not submit sufficient

justification, your fee waiver request will be denied. The Board may, at its discretion, communicate with you to request additional information if necessary. However, the Board must make a determination on the fee waiver request within the statutory time limit, even if the Board has not received such additional information. In certain circumstances, a partial fee waiver may be appropriate, if some, but not all, of the requested records are likely to contribute significantly to public understanding of the operations and activities of the government.

(b) The Board will waive fees (in whole or part) if disclosure of all or part of the information is in the public interest because its release:

(1) Is likely to contribute significantly to public understanding of the operations or activities of the government; and

(2) Is not primarily in the commercial interest of the requester.

§ 201.12 Denials.

(a) When denying a request in any respect, the Board shall notify the requestor of that determination in writing. The types of denials include:

(1) Denials of requests, including a determination:

(i) To withhold any requested record in whole or in part;

(ii) That a requested record does not exist or cannot be located;

(iii) That a record is not readily reproducible in the form or format sought;

(iv) That what has been requested is not a record subject to the FOIA; and

(v) That the material requested is not a Board record (e.g., material produced by another agency or organization).

(2) A determination on any disputed fee matter, including a denial of a request for a fee waiver.

(3) A denial of a request for expedited processing.

(b) The denial letter shall be signed by the FOIA Officer or designee and shall include all of the following:

(1) The name and title of the person responsible for the denial.

(2) A brief statement of the reason(s) for the denial, including any FOIA exemptions applied in denying the request.

(3) An estimate of the volume of records withheld, in number of pages or in some other reasonable form of estimation. This estimate does not need to be provided if it would harm an interest protected by an applicable exemption.

(4) A statement that the denial may be appealed under § 201.14 and a description of the requirements of § 201.14.

§ 201.13 Business information.

(a) *In general.* Business information obtained by the Board from a submitter shall be disclosed under the FOIA only under this section.

(b) *Definitions.* For purposes of this section:

(1) Business information—commercial or financial records obtained by the Board that may be protected from disclosure under Exemption 4 of the FOIA.

(2) Submitter—any person or entity from which the Board obtains business records, either directly or indirectly. The term includes but is not limited to corporations and state, local, tribal, and foreign governments.

(c) *Designation of business information.* Submitters of business information shall designate any part of the record considered to be protected from disclosure under Exemption 4 of the FOIA by appropriately marking the material. This may be done either at the time the record is submitted or at a reasonable time thereafter. This designation lasts for 10 years after submittal unless the submitter requests and provides justification for a longer period.

(d) *Notice to submitters.* The Board shall provide a business submitter with prompt written notice of any FOIA request or appeal that seeks its business information under paragraph (e) of this section, except as provided in paragraph (h) of this section, to give the submitter an opportunity to object to that disclosure under paragraph (f) of this section. The notice shall either describe the records requested or include copies of the records.

(e) *Required notice.* The Board shall give notice of a FOIA request seeking business information when:

(1) The submitter has designated that the information is considered protected from disclosure under Exemption 4 of the FOIA; or

(2) The Board has reason to believe that the information may be protected from disclosure under Exemption 4 of the FOIA.

(f)(1) *Objecting to disclosure.* A submitter shall have 30 days to respond to the notice described in paragraph (d) of this section. If a submitter has an objection to disclosure, it is required to submit a detailed written statement including:

(i) All grounds for withholding any of the information under any exemption of the FOIA, and

(ii) In the case of Exemption 4, the reason why the information is a trade secret, commercial, or financial information that is privileged or confidential.

(2) If a submitter fails to respond to the notice in paragraph (d) of this section within 30 days, the Board shall assume that the submitter has no objection to disclosure. The Board shall not consider information not received by the Board until after a disclosure decision has been made. Information provided by a submitter under this paragraph might itself be subject to disclosure under the FOIA.

(g) *Notice of intent to disclose.* The Board shall consider a submitter's objections and specific grounds for nondisclosure in deciding whether to disclose the business records. Whenever the Board decides to disclose business records over the objection of a submitter, it shall give the submitter written notice, that will include:

(1) A statement of the reason(s) the submitter's objections were not sustained;

(2) A description of the business records to be disclosed; and

(3) A specified disclosure date at a reasonable time subsequent to the notice.

(h) *Exceptions to notice requirements.* The notice requirements in paragraphs (d) and (g) of this section shall not apply if:

(1) The Board determines that the information should not be disclosed;

(2) The information has been published legally or has been officially made available to the public;

(3) Disclosure of the information is required by another statute or by a regulation issued in accordance with Executive Order 12600 (3 CFR, 1987 Comp., p. 235); or

(4) The objection made by the submitter under paragraph (f) of this section appears frivolous. In such a case, the Board shall promptly notify the submitter of its decision using the guidelines in paragraph (g) of this section.

(i) *Notice of FOIA lawsuit.* When a requestor files a lawsuit seeking to compel the disclosure of business information, the Board shall promptly notify the submitter.

(j) *Corresponding notice to requestors.* When the Board provides a submitter with either notice and an opportunity to object to disclosure under paragraph (d) of this section or with its intent to disclose requested information under paragraph (g) of this section, the Board also shall notify the requestor(s). When a submitter files a lawsuit seeking to prevent the disclosure of business information, the Board shall notify the requestor(s).

§ 201.14 Appeals.

(a)(1) *Appeals of adverse determinations.* If you are dissatisfied with the Board's response to your request, you may appeal to the Board's Executive Director:

(i) By mail to: Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006;

(ii) By e-mail to: FOIA@ratb.gov; or

(iii) By fax to: 202-254-7970.

(2) The appeal must be in writing and must be received within 30 days of the date of the Board's response. The appeal letter, e-mail or fax may include as much or as little related information as you wish, as long as it clearly identifies the Board determination that you are appealing, including the assigned request number, if known. For prompt handling, please mark your appeal "Freedom of Information Act Appeal."

(b) *Responses to appeals.* Requestors shall be notified in writing of the decision on the appeal. A decision affirming an adverse determination shall include a statement of the reason(s) for the affirmation, including any FOIA exemption(s) applied, and shall include the FOIA provisions for court review of the decision. If the adverse determination is reversed or modified on appeal, the request shall be reprocessed in accordance with that appeal decision.

(d) *Denial of appeal.* An adverse determination by the Executive Director shall be the final action of the Board.

§ 201.15 Preservation of records.

The Board shall preserve all correspondence pertaining to the requests that it receives under this subpart, as well as copies of all requested records, until disposition or destruction is authorized by title 44 of the United States Code of the National Archives and Records Administration's General Records Schedule 14. Records will not be disposed of while they are the subject of a pending request, appeal, or lawsuit.

§ 201.16 Other rights and services.

Nothing in this part shall be construed to entitle any person, as a right, to any service or to the disclosure of any record to which such person is entitled under the FOIA.

§ 201.17 How to track a FOIA request.

(a) *Tracking number.* The Board will issue a tracking number to all FOIA requesters within 5 days of the receipt of the request (as described in § 201.7(b)). The tracking number will be sent via electronic mail if the requester has provided an electronic mail address.

Otherwise, the Board will mail the tracking number to the requester's physical address, as provided in the FOIA request.

(b) *Status of request.* FOIA requesters may check the status of their FOIA request(s) by contacting the FOIA Officer at FOIA@ratb.gov or (202) 254-7900.

Ivan J. Flores,

Paralegal Specialist, Recovery Accountability and Transparency Board.

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DEPARTMENT OF THE TREASURY

Office of the Comptroller of the Currency

12 CFR Part 3

[Docket ID OCC-2009-0018]

RIN 1557-AD25

FEDERAL RESERVE SYSTEM

12 CFR Parts 208 and 225

[Regulations H and Y; Docket No. R-1361]

FEDERAL DEPOSIT INSURANCE CORPORATION

12 CFR Part 325

RIN 3064-AD42

DEPARTMENT OF THE TREASURY

Office of Thrift Supervision

12 CFR Part 567

[No. OTS-2009-0020]

RIN 1550-AC34

Risk-Based Capital Guidelines; Capital Adequacy Guidelines; Capital Maintenance; Capital—Residential Mortgage Loans Modified Pursuant to the Home Affordable Mortgage Program

AGENCY: Office of the Comptroller of the Currency, Department of the Treasury; Board of Governors of the Federal Reserve System; Federal Deposit Insurance Corporation; and Office of Thrift Supervision, Department of the Treasury (the agencies).

ACTION: Final rule.

SUMMARY: The agencies have adopted a final rule to allow banks, savings associations, and bank holding companies (collectively, banking organizations) to risk weight for purposes of the agencies' capital

guidelines mortgage loans modified pursuant to the Home Affordable Mortgage Program (Program) implemented by the U.S. Department of the Treasury (Treasury) with the same risk weight assigned to the loan prior to the modification so long as the loan continues to meet other applicable prudential criteria.

DATES: The final rule becomes effective December 21, 2009.

FOR FURTHER INFORMATION CONTACT:

OCC: Margot Schwadron, Senior Risk Expert, Capital Policy Division, (202) 874-6022, or Carl Kaminski, Senior Attorney, or Ron Shimabukuro, Senior Counsel, Legislative and Regulatory Activities Division, (202) 874-5090, Office of the Comptroller of the Currency, 250 E Street, SW., Washington, DC 20219.

Board: Barbara J. Bouchard, Associate Director, (202) 452-3072, or William Tiernay, Senior Supervisory Financial Analyst, (202) 872-7579, Division of Banking Supervision and Regulation; or April Snyder, Counsel, (202) 452-3099, or Benjamin W. McDonough, Counsel, (202) 452-2036, Legal Division. For the hearing impaired only, Telecommunication Device for the Deaf (TDD), (202) 263-4869.

FDIC: Ryan Sheller, Senior Capital Markets Specialist, (202) 898-6614, Capital Markets Branch, Division of Supervision and Consumer Protection; or Mark Handzlik, Senior Attorney, (202) 898-3990, or Michael Phillips, Counsel, (202) 898-3581, Supervision Branch, Legal Division.

OTS: Teresa A. Scott, Senior Policy Analyst, (202) 906-6478, Capital Risk, or Marvin Shaw, Senior Attorney, (202) 906-6639, Legislation and Regulation Division, Office of Thrift Supervision, 1700 G Street, NW., Washington, DC 20552.

SUPPLEMENTARY INFORMATION:

Background

Under the agencies' general risk-based capital rules, loans that are fully secured by first liens on one-to-four family residential properties, that are either owner-occupied or rented, and that meet certain prudential criteria (qualifying mortgage loans) are risk-weighted at 50 percent.¹ If a banking organization holds both a first-lien and a junior-lien mortgage on the same property, and no other party holds an intervening lien, the loans are treated as a single loan secured by a first-lien mortgage and risk-weighted at 50 percent if the two loans, when

aggregated, meet the conditions to be a qualifying mortgage loan. Other junior-lien mortgage loans are risk-weighted at 100 percent.²

In general, to qualify for a 50 percent risk weight, a mortgage loan must have been made in accordance with prudent underwriting standards and may not be 90 days or more past due. Mortgage loans that do not qualify for a 50 percent risk weight are assigned a 100 percent risk weight. Each agency has additional provisions that address the risk weighting of mortgage loans. Under the OCC's general risk-based capital rules for national banks, to receive a 50 percent risk weight, a mortgage loan must "not [be] on nonaccrual or restructured."³ Under the Board's general risk-based capital rules for bank holding companies and state member banks, mortgage loans must be "performing in accordance with their original terms" and not carried in nonaccrual status in order to receive a 50 percent risk weight.⁴ Generally, mortgage loans that have been modified are considered to have been restructured (OCC), or are not considered to be performing in accordance with their original terms (Board). Therefore, under the OCC's and Board's general risk-based capital rules, such loans generally must be risk weighted at 100 percent. Under the FDIC's general risk-based capital rules, a state nonmember bank may assign a 50 percent risk weight to any modified mortgage loan, so long as the loan, as modified, is not 90 days or more past due or in nonaccrual status and meets other applicable criteria for a 50 percent risk weight.⁵ Under the OTS's general risk-based capital rules, a savings association may assign a 50 percent risk weight to any modified residential mortgage loan, so long as the loan, as modified, is not 90 days or more past due and meets other applicable criteria for a 50 percent risk weight.⁶

On June 30, 2009, the agencies published in the **Federal Register** an interim final rule (interim rule) to allow banking organizations to risk weight mortgage loans modified under the Program using the same risk weight assigned to the loan prior to the modification, so long as the loan continues to meet other applicable

² See 12 CFR Part 3, Appendix A, section 3(a)(3)(iii) (OCC); 12 CFR parts 208 and 225, Appendix A, section III.C.4. (Board); 12 CFR part 325, Appendix A, section II.C. (FDIC); and 12 CFR 567.6(1)(iv) (OTS).

³ 12 CFR Part 3, Appendix A, section 3(a)(3)(iii) (OCC).

⁴ 12 CFR parts 208 and 225, Appendix A, section III.C.3. (Board).

⁵ 12 CFR Part 325, Appendix A, section II.C. (FDIC).

⁶ 12 CFR 567.1, 12 CFR 567.6(a)(1)(iii) (OTS).

¹ See 12 CFR Part 3, Appendix A, section 3(a)(3)(iii) (OCC); 12 CFR parts 208 and 225.

prudential criteria.⁷ In many circumstances, this means that an eligible mortgage loan modified in accordance with the Program will continue to receive a 50 percent risk weight for purposes of the agencies' general risk-based capital guidelines. The agencies are now adopting the interim rule as a final rule (final rule) with changes that clarify the regulatory capital treatment of mortgage loans during the Program's trial modification period (trial period). The revisions provided under the final rule relative to the FDIC's and OTS' general risk-based capital rules are clarifying in nature.

Home Affordable Mortgage Program

On March 4, 2009, Treasury announced guidelines under the Program to promote sustainable loan modifications for homeowners at risk of losing their homes due to foreclosure.⁸ The Program provides a detailed framework for servicers to modify mortgages on owner-occupied residential properties and offers financial incentives to lenders and servicers that participate in the Program.⁹ The Program also provides financial incentives for homeowners whose mortgages are modified pursuant to Program guidelines to remain current on their mortgages after modification.¹⁰ Taken together, these incentives are intended to help responsible homeowners remain in their homes and avoid foreclosure, which is in turn intended to help ease the current downward pressures on house prices and the costs that families, communities, and the economy incur from unnecessary foreclosures.

Under the Program, Treasury has partnered with lenders and loan servicers to offer at-risk homeowners loan modifications under which the homeowners may obtain more affordable monthly mortgage payments. The Program applies to a spectrum of outstanding loans, some of which meet all of the prudential criteria under the agencies' general risk-based capital rules and receive a 50 percent risk weight and

some of which otherwise receive a 100 percent risk weight under the agencies' general risk-based capital rules.¹¹ Servicers who elect to participate in the Program are required to apply the Program guidelines to all eligible loans¹² unless explicitly prohibited by the governing pooling and servicing agreement and/or other lender servicing agreements. If a mortgage loan qualifies for modification under the Program, the Program guidelines require the lender to first reduce payments on eligible first-lien loans to an amount representing no greater than a 38 percent initial front-end debt-to-income ratio.¹³ Treasury then will match further reductions in monthly payments with the lender dollar-for-dollar to achieve a 31 percent front-end debt-to-income ratio on the first-lien mortgage.¹⁴ Borrowers whose back-end debt-to-income ratio exceeds 55 percent must agree to work with a foreclosure prevention counselor approved by the Department of Housing and Urban Development.¹⁵

In addition to the incentives for lenders, servicers are eligible for other incentive payments to encourage participation in the Program. Servicers receive an up-front servicer incentive payment of \$1,000 for each eligible first-lien modification. Lenders and servicers are eligible for one-time incentive payments of \$1,500 and \$500, respectively, for early modifications of first-lien mortgages—that is, modifications made while the borrower is still current on mortgage payments but at risk of imminent default. To encourage ongoing performance of modified loans, servicers also will

receive "Pay for Success" incentive payments of up to \$1,000 per year for up to three years for first-lien mortgages as long as borrowers remain in the Program. A borrower can likewise receive "Pay for Performance Success" incentive payments that reduce the principal balance on the borrower's first-lien mortgage up to \$1,000 per year for up to five years if the borrower remains current on monthly payments on the modified first-lien mortgage. Lenders also may receive a home price depreciation reserve payment to offset certain losses if a modified loan subsequently defaults.

For second-lien mortgages, lenders are eligible to receive incentive payments based on the difference between the interest rate on the modified first-lien mortgage and the reduced interest rate (either 1 percent or 2 percent) on the second-lien mortgage following modification.¹⁶ Servicers may receive a one-time \$500 incentive payment for successful second-lien modifications, as well as additional incentive payments of up to \$250 per year for up to three years for second-lien mortgages as long as both the modified first-lien and second-lien mortgages remain current. A borrower also may receive incentive payments of up to \$250 per year for a modified second-lien mortgage loan for up to five years for remaining current on the loan, which will be paid to reduce the unpaid principal of the first-lien mortgage. However, second-lien modification incentives only will be paid with respect to a given property if the first-lien mortgage on the property also is modified under the Program.¹⁷

Before a loan may be modified under the Program, a borrower must successfully complete a trial period of at least 90 days. During the trial period, a borrower makes payments on the eligible mortgage loan under modified terms. To complete the trial period successfully, the borrower must be current at the end of the trial period and provide certain information.¹⁸ The Program provides no incentive payments to the lender, servicer, or

⁷ See 12 CFR Part 3, Appendix A, sections 3(a)(3)(iii) and 3(a)(4) (OCC); 12 CFR parts 208 and 225, Appendix A, sections III.C.3. and III.C.4. (Board); 12 CFR part 325, Appendix A, section II.C. (FDIC); and 12 CFR 567.1 and 567.6 (OTS).

¹² For a mortgage to be eligible for the Program, the property securing the mortgage loan must be a one-to-four family owner-occupied property that is the primary residence of the mortgagee. The property cannot be vacant or condemned, and the mortgage must have an unpaid principal balance (prior to capitalization of arrearages) at or below the Fannie Mae conforming loan limit for the type of property.

¹³ A front-end debt-to-income ratio measures how much of the borrower's gross (pretax) monthly income is represented by the borrower's required payment on the first-lien mortgage, including real estate taxes and insurance.

¹⁴ To qualify for the Treasury match, servicers must follow an established sequence of actions (capitalize arrearages, reduce interest rate, extend term or amortization period, and then defer principal) to reduce the front-end debt-to-income ratio on the loan from 38 percent to 31 percent. Servicers may reduce principal on the loan at any stage during the modification sequence to meet affordability targets.

¹⁵ A back-end debt-to-income ratio measures how much of a borrower's gross (pretax) monthly income would go toward monthly mortgage and nonmortgage debt service obligations.

¹⁶ Participating servicers are required to follow certain steps in modifying amortizing second-lien mortgages, including reducing the interest rate to 1 percent or 2 percent. Lenders may receive an incentive payment from Treasury equal to half of the difference between (i) the interest rate on the first lien as modified and (ii) 1 percent, subject to a floor.

¹⁷ In some cases, servicers may choose to accept a lump-sum payment from Treasury to extinguish some or all of a second-lien mortgage under a pre-set formula.

¹⁸ Under the Program, borrowers in certain states with unique foreclosure law requirements (foreclosure restart states) will be considered to have failed the trial period if they are not current at the time the foreclosure sale is scheduled.

⁷ 74 FR 31160 (June 30, 2009); 74 FR 34499 (July 16, 2009) (OCC technical correction).

⁸ Further details about the Program, including Program terms and borrower eligibility criteria, are available at <http://www.makinghomeaffordable.gov>.

⁹ For ease of reference, the term "servicer" refers both to servicers that service loans held by other entities and to lenders who service loans that they hold themselves. The term "lender" refers to the beneficial owner or owners of the mortgage.

¹⁰ A separate aspect of the Program, the Home Affordable Refinance Program, also provides incentives for refinancing certain mortgage loans owned or guaranteed by Fannie Mae or Freddie Mac. This final rule does not apply to mortgage loans refinanced under the Home Affordable Refinance Program.

borrower during the trial period and no payments if the borrower does not successfully complete the trial period.

Comments on the Interim Rule

The agencies received six comments on the interim rule, one from a banking organization, four from trade groups representing the financial industry, and one from an individual. The commenters that addressed the interim final rule unanimously supported it, asserting that it is consistent with the important policy objectives of the Program and does not compromise the goals of safety and soundness. Commenters requested that the agencies clarify whether the rule's capital treatment is available for a mortgage loan that has been modified on a preliminary basis under the Program, but which still is within the trial period (and, thus, has not been permanently modified). Commenters also requested clarification regarding the circumstances under which a mortgage loan that was risk-weighted at 100 percent immediately prior to modification under the Program could receive a 50 percent risk weight. Some commenters suggested that such a loan should receive a 50 percent risk weight following completion of the trial period or following receipt of the first pay-for-performance incentive payments. Other commenters requested that the agencies clarify that a sustained period of repayment performance could include payments made after a loan had been modified under the Program. The agencies also received a comment on the interaction between private mortgage insurance and loan modifications, which was beyond the scope of the interim rule.

Based on an analysis of the comments, the agencies have modified the rule to specify that a mortgage modified on a permanent or trial basis pursuant to the Program and that was risk-weighted at 50 percent may continue to receive a 50 percent risk weight provided it meets other prudential criteria.¹⁹

As noted in the preamble to the interim rule, under the agencies' existing practice, past due and nonaccrual loans that receive a 100 percent risk weight may return to a 50 percent risk weight under certain circumstances, including after demonstration of a sustained period of repayment performance. Because borrower characteristics, such as debt

service capacity, impact a borrower's creditworthiness, the degree of appropriate reliance on a fixed period of payment performance may vary for different borrowers.²⁰ For these reasons, the agencies have not established a specific period of repayments that would constitute a "sustained period of performance" for a particular loan. The agencies confirm that a borrower's payments on a mortgage loan modified under the Program, including during the trial period, may be considered in assessing whether the borrower has demonstrated a sustained period of repayment performance.

Commenters also requested that the agencies (1) allow a banking organization to risk weight at 50 percent, rather than 100 percent, a second-lien mortgage loan that is modified under the Program if the first-lien mortgage loan on the property is owned by another entity, that first-lien mortgage is also modified under the Program, and there is no intervening lien; and (2) allow loans modified pursuant to the Program or similar programs that continue to qualify for 50 percent risk weight to be excluded from troubled debt restructurings reported in quarterly bank regulatory reports. Under the general risk-based capital rules all second-lien mortgage loans receive a 100 percent risk weight, unless the banking organization that holds the loan also holds the first lien, there is no intervening lien, and the loan meets other prudential criteria. The agencies believe this treatment is commensurate with the risks of junior positions, as lenders have limited access to collateral in the event of default. Therefore, the agencies have determined that allowing a banking organization to risk weight junior-lien mortgage loans at less than 100 percent is not appropriate other than in those circumstances already permitted by the agencies general risk-based capital rules. With respect to whether mortgage loans modified under the Program are considered troubled debt restructurings, the question of how these loans should be classified and reported will be determined under

generally accepted accounting principles.

Final Rule

Based on the above considerations, the agencies have adopted the interim rule in final form with the modification discussed above. Under the final rule as under the interim rule mortgage loans modified under the Program will retain the risk weight appropriate to the mortgage loan prior to modification, as long as other applicable prudential criteria remain satisfied. Accordingly, under the final rule, a qualifying mortgage loan appropriately risk weighted at 50 percent before modification under the Program would continue to be risk weighted at 50 percent during the trial period and after modification, provided it meets other prudential criteria. If a borrower does not successfully complete the trial period and the loan is not modified under the Program on a permanent basis, the loan would qualify for the 50 percent risk weight category if it meets the conditions to be a qualifying mortgage loan under the general risk-based capital rules. If the loan does not meet the conditions, it would receive a 100 percent risk weight. A mortgage loan appropriately risk weighted at 100 percent prior to modification under the Program would continue to be risk weighted at 100 percent during and after the trial period.

Consistent with the OCC's and the Board's general risk-based capital rules, if a mortgage loan were to become 90 days or more past due or carried in non-accrual status or otherwise restructured after being modified under the Program, the loan would be assigned a risk weight of 100 percent. Consistent with the FDIC's general risk-based capital rules, if a mortgage loan were to again be restructured after being modified under the Program, the loan could be assigned a risk weight of 50 percent provided the loan, as modified, is not 90 days or more past due or in nonaccrual status and meets the other applicable criteria for a 50 percent risk weight. Consistent with the OTS's general risk-based capital rules, if a mortgage loan were to again be restructured after being modified under the Program, the loan could be assigned a risk weight of 50 percent provided the loan, as modified, is not 90 days or more past due and meets the other applicable criteria for a 50 percent risk weight.

Additionally, in certain circumstances under the general risk-based capital rules (as with, for example, a direct credit substitute or recourse obligation), a banking organization is permitted to look through an exposure to the risk

¹⁹The agencies intended the interim rule to apply to loans modified on both a trial and permanent basis under the Program. Accordingly, the modifications to the final rule are clarifying in nature.

²⁰The instructions for the Consolidated Reports of Condition and Income (Call Report) and the Thrift Financial Report (TFR) define a sustained period of repayment performance as a period generally lasting " * * * a minimum of six months and would involve payments of cash or cash equivalents. (In returning the asset to accrual status, sustained historical repayment performance for a reasonable time prior to the restructuring may be taken into account.)" Call Reports instructions are available at <http://www.federalreserve.gov/reportforms/CategoryIndex.cfm?WhichCategory=3> and TFR instructions are available at <http://files.ots.treas.gov/4210058.pdf>.

weight of a residential mortgage loan underlying that exposure. In such cases, the banking organizations would follow the capital treatment provided for in the agencies' general risk-based capital rules, as modified by the final rule, when the underlying residential mortgage loan has been modified pursuant to the Program.

The agencies believe that treating mortgage loans modified under the Program in the manner described above is appropriate in light of the special and unique incentive features of the Program and the fact that the Program is offered by the federal government in order to achieve the public policy objective of promoting sustainable loan modifications for homeowners at risk of foreclosure in a way that balances the interests of borrowers, servicers, and lenders. As previously described, the Program requires that a borrower's front-end debt-to-income ratio on a first-lien mortgage modified under the Program be reduced to no greater than 31 percent, which should improve the borrower's ability to repay the modified loan, and, importantly, provides for Treasury to match reductions in monthly payments dollar-for-dollar to reduce the borrower's front-end debt-to-income ratio from 38 percent to 31 percent. In addition, as described above, the Program provides material financial incentives for servicers and lenders to take actions to reduce the likelihood of defaults, as well as incentives for servicers and borrowers designed to help borrowers remain current on modified loans. The structure and amount of these cash payments meaningfully align the financial incentives for servicers, lenders, and borrowers to encourage and increase the likelihood of participating borrowers remaining current on their mortgages. Each of these incentives is important to the agencies' determination with respect to the appropriate regulatory capital treatment of mortgage loans modified under the Program.

Regulatory Analysis

Regulatory Flexibility Act

The Regulatory Flexibility Act, 5 U.S.C. 601 *et seq.* (RFA), generally requires that, in connection with a notice of proposed rulemaking, an agency prepare and make available for public comment an initial regulatory flexibility analysis that describes the impact of a proposed rule on small entities.²¹ Under regulations issued by the Small Business Administration,²² a

small entity includes a commercial bank, bank holding company, or savings association with assets of \$175 million or less (a small banking organization). As of June 30, 2009, approximately 2,533 small bank holding companies, 386 small savings associations, 749 small national banks, 432 small state member banks, and 3,040 small state nonmember banks existed. As a general matter, the Board's general risk-based capital rules apply only to a bank holding company that has consolidated assets of \$500 million or more. Therefore, the changes to the Board's capital adequacy guidelines for bank holding companies will not affect small bank holding companies.

This rulemaking does not involve the issuance of a notice of proposed rulemaking and, therefore, the requirements of the RFA do not apply. However, the agencies note that the rule does not impose any additional obligations, restrictions, burdens, or reporting, recordkeeping or compliance requirements on banks or savings associations, including small banking organizations, nor does it duplicate, overlap or conflict with other federal rules. The rule also will benefit small banking organizations that are subject to the agencies' general risk-based capital rules by allowing mortgage loans modified under the Program to retain the risk weight assigned to the loan prior to the modification.

Paperwork Reduction Act

In accordance with the requirements of the Paperwork Reduction Act of 1995 (44 U.S.C. 3506), the agencies have reviewed the final rule to assess any information collections. There are no collections of information as defined by the Paperwork Reduction Act in the final rule.

OCC/OTS Executive Order 12866

Executive Order 12866 requires federal agencies to prepare a regulatory impact analysis for agency actions that are found to be "significant regulatory actions." Significant regulatory actions include, among other things, rulemakings that "have an annual effect on the economy of \$100 million or more or adversely affect in a material way the economy, a sector of the economy, productivity, competition, jobs, the environment, public health or safety, or state, local, or tribal governments or communities." The OCC and the OTS each determined that its portion of the final rule is not a significant regulatory action under Executive Order 12866.

OCC/OTS Unfunded Mandates Reform Act of 1995 Determination

The Unfunded Mandates Reform Act of 1995²³ (UMRA) requires that an agency prepare a budgetary impact statement before promulgating a rule that includes a federal mandate that may result in the expenditure by state, local, and tribal governments, in the aggregate, or by the private sector of \$100 million or more (adjusted annually for inflation) in any one year. If a budgetary impact statement is required, section 205 of the UMRA also requires an agency to identify and consider a reasonable number of regulatory alternatives before promulgating a rule. The OCC and the OTS each have determined that its final rule will not result in expenditures by state, local, and tribal governments, in the aggregate, or by the private sector, of \$100 million or more in any one year. Accordingly, neither the OCC nor the OTS has prepared a budgetary impact statement or specifically addressed the regulatory alternatives considered.

List of Subjects

12 CFR Part 3

Administrative practice and procedure, Banks, Banking, Capital, National banks, Reporting and recordkeeping requirements, Risk.

12 CFR Part 208

Confidential business information, Crime, Currency, Federal Reserve System, Mortgages, Reporting and recordkeeping requirements, Risk.

12 CFR Part 225

Administrative Practice and Procedure, Banks, banking, Federal Reserve System, Holding companies, Reporting and recordkeeping requirements, Securities.

12 CFR Part 325

Administrative practice and procedure, Banks, banking, Capital adequacy, Reporting and recordkeeping requirements, Savings associations, State nonmember banks.

12 CFR Part 567

Capital, Reporting and recordkeeping requirements, Risk, Savings associations.

²¹ See 5 U.S.C. 603(a).

²² See 13 CFR 121.201.

²³ See Public Law 104-4.

**Department of the Treasury
Office of the Comptroller of the
Currency**

12 CFR Chapter I

Authority and Issuance

■ For the reasons stated in the common preamble, the Office of the Comptroller of the Currency amends Part 3 of chapter I of Title 12, Code of Federal Regulations as follows:

PART 3—MINIMUM CAPITAL RATIOS; ISSUANCE OF DIRECTIVES

■ 1. The authority citation for part 3 continues to read as follows:

Authority: 12 U.S.C. 93a, 161, 1818, 1828(n), 1828 note, 1831n note, 1835, 3907, and 3909.

■ 2. In appendix A to Part 3, in section 3, revise paragraph (a)(3)(iii) to read as follows:

Appendix A to Part 3—Risk-Based Capital Guidelines

* * * * *

Section 3. Risk Categories/Weights for On-Balance Sheet Assets and Off-Balance Sheet Items

* * * * *

(a) * * *

(3) * * *

(iii) Loans secured by first mortgages on one-to-four family residential properties, either owner occupied or rented, provided that such loans are not otherwise 90 days or more past due, or on nonaccrual or restructured. It is presumed that such loans will meet the prudent underwriting standards. For the purposes of the risk-based capital guidelines, a loan modified on a permanent or trial basis solely pursuant to the U.S. Department of Treasury's Home Affordable Mortgage Program will not be considered to have been restructured. If a bank holds a first lien and junior lien on a one-to-four family residential property and no other party holds an intervening lien, the transaction is treated as a single loan secured by a first lien for the purposes of both determining the loan-to-value ratio and assigning a risk weight to the transaction. Furthermore, residential property loans made for the purpose of construction financing are assigned to the 100% risk category of section 3(a)(4) of this appendix A; however, these loans may be included in the 50% risk category of this section 3(a)(3) of this appendix A if they are subject to a legally binding sales contract and satisfy the requirements of section 3(a)(3)(iv) of this appendix A.

* * * * *

Board of Governors of the Federal Reserve System

12 CFR Chapter II

Authority and Issuance

■ For the reasons stated in the common preamble, the Board of Governors of Federal Reserve System amends parts 208 and 225 of Chapter II of title 12 of the Code of Federal Regulations as follows:

PART 208—MEMBERSHIP OF STATE BANKING INSTITUTIONS IN THE FEDERAL RESERVE SYSTEM (REGULATION H)

■ 3. The authority for part 208 continues to read as follows:

Authority: 12 U.S.C. 24, 36, 92a, 93a, 248(a), 248(c), 321–338a, 371d, 461, 481–486, 601, 611, 1814, 1816, 1818, 1820(d)(9), 1833(j), 1828(o), 1831, 1831o, 1831p–1, 1831r–1, 1831w, 1831x, 1835a, 1882, 2901–2907, 3105, 3310, 3331–3351, and 3905–3909; 15 U.S.C. 78b, 78l(b), 78l(i), 780–4(c)(5), 78q, 78q–1, and 78w, 1681s, 1681w, 6801, and 6805; 31 U.S.C. 5318; 42 U.S.C. 4012a, 4104a, 4104b, 4106 and 4128.

■ 4. In appendix A to part 208, revise Section III. C.3., to read as follows:

Appendix A to Part 208—Capital Adequacy Guidelines for State Member Banks: Risk-Based Measure

* * * * *

III. * * *

C. * * *

3. *Category 3: 50 percent.* This category includes loans fully secured by first liens⁴¹ on 1- to 4-family residential properties, either owner-occupied or rented, or on multifamily residential properties,⁴² that meet certain

⁴¹ If a bank holds the first and junior lien(s) on a residential property and no other party holds an intervening lien, the transaction is treated as a single loan secured by a first lien for the purposes of determining the loan-to-value ratio and assigning a risk weight.

⁴² Loans that qualify as loans secured by 1- to 4-family residential properties or multifamily residential properties are listed in the instructions to the commercial bank Call Report. In addition, for risk-based capital purposes, loans secured by 1- to 4-family residential properties include loans to builders with substantial project equity for the construction of 1- to 4-family residences that have been presold under firm contracts to purchasers who have obtained firm commitments for permanent qualifying mortgage loans and have made substantial earnest money deposits. Such loans to builders will be considered prudently underwritten only if the bank has obtained sufficient documentation that the buyer of the home intends to purchase the home (*i.e.*, has a legally binding written sales contract) and has the ability to obtain a mortgage loan sufficient to purchase the home (*i.e.*, has a firm written commitment for permanent financing of the home upon completion).

The instructions to the Call Report also discuss the treatment of loans, including multifamily housing loans, that are sold subject to a pro rata loss sharing arrangement. Such an arrangement should

be treated by the selling bank as sold (and excluded from balance sheet assets) to the extent that the sales agreement provides for the purchaser of the loan to share in any loss incurred on the loan on a pro rata basis with the selling bank. In such a transaction, from the standpoint of the selling bank, the portion of the loan that is treated as sold is not subject to the risk-based capital standards. In connection with sales of multifamily housing loans in which the purchaser of a loan shares in any loss incurred on the loan with the selling institution on other than a pro rata basis, these other loss sharing arrangements are taken into account for purposes of determining the extent to which such loans are treated by the selling bank as sold (and excluded from balance sheet assets) under the risk-based capital framework in the same as prescribed for reporting purposes in the instructions to the Call Report.

⁴³ Residential property loans that do not meet all the specified criteria or that are made for the purpose of speculative property development are placed in the 100 percent risk category.

⁴⁴ Prudent underwriting standards include a conservative ratio of the current loan balance to the value of the property. In the case of a loan secured by multifamily residential property, the loan-to-value ratio is not conservative if it exceeds 80 percent (75 percent if the loan is based on a floating interest rate). Prudent underwriting standards also dictate that a loan-to-value ratio used in the case of originating a loan to acquire a property would not be deemed conservative unless the value is based on the lower of the acquisition cost of the property or appraised (or if appropriate, evaluated) value. Otherwise, the loan-to-value ratio generally would be based upon the value of the property as determined by the most current appraisal, or if appropriate, the most current evaluation. All appraisals must be made in a manner consistent with the Federal banking agencies' real estate appraisal regulations and guidelines and with the bank's own appraisal guidelines.

criteria.⁴³ Loans included in this category must have been made in accordance with prudent underwriting standards;⁴⁴ be performing in accordance with their original terms; and not be 90 days or more past due or carried in nonaccrual status. For purposes of this 50 percent risk weight category, a loan modified on a permanent or trial basis solely pursuant to the U.S. Department of Treasury's Home Affordable Mortgage Program will be considered to be performing in accordance with its original terms. The following additional criteria must also be applied to a loan secured by a multifamily residential property that is included in this category: all principal and interest payments on the loan must have been made on time for at least the year preceding placement in this category, or in the case where the existing property owner is refinancing a loan on that property, all principal and interest payments on the loan being refinanced must have been made on time for at least the year preceding placement in this category; amortization of the principal and interest must occur over a period of not more than 30 years and the minimum original maturity for repayment of principal must not be less than 7 years; and the annual net operating income (before debt service) generated by the property during its most recent fiscal year must not be less than 120 percent of the loan's current annual debt service (115 percent if the loan is based on a floating interest rate) or, in the case of a cooperative or other not-for-profit housing project, the property must generate sufficient cash flow to provide comparable protection

to the institution. Also included in this category are privately-issued mortgage-backed securities provided that:

(1) The structure of the security meets the criteria described in section III(B)(3) above;

(2) If the security is backed by a pool of conventional mortgages, on 1- to 4-family residential or multifamily residential properties each underlying mortgage meets the criteria described above in this section for eligibility for the 50 percent risk category at the time the pool is originated;

(3) If the security is backed by privately issued mortgage-backed securities, each underlying security qualifies for the 50 percent risk category; and

(4) If the security is backed by a pool of multifamily residential mortgages, principal and interest payments on the security are not 30 days or more past due.

Privately-issued mortgage-backed securities that do not meet these criteria or that do not qualify for a lower risk weight are generally assigned to the 100 percent risk category.

Also assigned to this category are revenue (non-general obligation) bonds or similar obligations, including loans and leases, that are obligations of states or other political subdivisions of the U.S. (for example, municipal revenue bonds) or other countries of the OECD-based group, but for which the government entity is committed to repay the debt with revenues from the specific projects financed, rather than from general tax funds.

Credit equivalent amounts of derivative contracts involving standard risk obligors (that is, obligors whose loans or debt securities would be assigned to the 100 percent risk category) are included in the 50 percent category, unless they are backed by collateral or guarantees that allow them to be placed in a lower risk category.

* * * * *

PART 225—BANK HOLDING COMPANIES AND CHANGE IN BANK CONTROL (REGULATION Y)

■ 5. The authority for part 225 continues to read as follows:

Authority: 12 U.S.C. 1817(j)(13), 1818, 1828(o), 1831i, 1831p-1, 1843(c)(8), 1844(b), 1972(1), 3106, 3108, 3310, 3331-3351, 3907, and 3909; 15 U.S.C. 1681s, 1681w, 6801 and 6805.

■ 6. In Appendix A to part 225, revise section III.C.3., to read as follows:

Appendix A to Part 225—Capital Adequacy Guidelines for Bank Holding Companies: Risk-Based Measure

* * * * *

III. * * *
C. * * *

3. *Category 3: 50 percent.* This category includes loans fully secured by first liens⁴⁸ on 1- to 4-family residential properties, either

⁴⁸ If a banking organization holds the first and junior lien(s) on a residential property and no other party holds an intervening lien, the transaction is treated as a single loan secured by a first lien for the purposes of determining the loan-to-value ratio and assigning a risk weight.

owner-occupied or rented, or on multifamily residential properties,⁴⁹ that meet certain criteria.⁵⁰ Loans included in this category must have been made in accordance with prudent underwriting standards;⁵¹ be performing in accordance with their original terms; and not be 90 days or more past due or carried in nonaccrual status. For purposes of this 50 percent risk weight category, a loan modified on a permanent or trial basis solely pursuant to the U.S. Department of Treasury's Home Affordable Mortgage Program will be considered to be performing in accordance with its original terms. The following additional criteria must also be applied to a loan secured by a multifamily residential property that is included in this category: all principal and interest payments on the loan must have been made on time for at least the year preceding placement in this category, or in the case where the existing property owner is refinancing a loan on that property, all principal and interest payments on the loan being refinanced must have been made on time for at least the year preceding placement in this category; amortization of the principal and interest must occur over a period of not more than 30 years and the minimum original maturity for repayment of principal must not be less than 7 years; and the annual net operating income (before debt service) generated by the property during its most recent fiscal year must not be less than 120 percent of the loan's current annual debt service (115 percent if the loan is based on a floating interest rate) or, in the case of a

⁴⁹ Loans that qualify as loans secured by 1- to 4-family residential properties or multifamily residential properties are listed in the instructions to the FR Y-9C Report. In addition, for risk-based capital purposes, loans secured by 1- to 4-family residential properties include loans to builders with substantial project equity for the construction of 1- to 4-family residences that have been presold under firm contracts to purchasers who have obtained firm commitments for permanent qualifying mortgage loans and have made substantial earnest money deposits. Such loans to builders will be considered prudently underwritten only if the bank holding company has obtained sufficient documentation that the buyer of the home intends to purchase the home (*i.e.*, has a legally binding written sales contract) and has the ability to obtain a mortgage loan sufficient to purchase the home (*i.e.*, has a firm written commitment for permanent financing of the home upon completion).

⁵⁰ Residential property loans that do not meet all the specified criteria or that are made for the purpose of speculative property development are placed in the 100 percent risk category.

⁵¹ Prudent underwriting standards include a conservative ratio of the current loan balance to the value of the property. In the case of a loan secured by multifamily residential property, the loan-to-value ratio is not conservative if it exceeds 80 percent (75 percent if the loan is based on a floating interest rate). Prudent underwriting standards also dictate that a loan-to-value ratio used in the case of originating a loan to acquire a property would not be deemed conservative unless the value is based on the lower of the acquisition cost of the property or appraised (or if appropriate, evaluated) value. Otherwise, the loan-to-value ratio generally would be based upon the value of the property as determined by the most current appraisal, or if appropriate, the most current evaluation. All appraisals must be made in a manner consistent with the Federal banking agencies' real estate appraisal regulations and guidelines and with the banking organization's own appraisal guidelines.

cooperative or other not-for-profit housing project, the property must generate sufficient cash flow to provide comparable protection to the institution. Also included in this category are privately-issued mortgage-backed securities provided that:

(1) The structure of the security meets the criteria described in section III(B)(3) above;

(2) if the security is backed by a pool of conventional mortgages, on 1- to 4-family residential or multifamily residential properties, each underlying mortgage meets the criteria described above in this section for eligibility for the 50 percent risk category at the time the pool is originated;

(3) If the security is backed by privately-issued mortgage-backed securities, each underlying security qualifies for the 50 percent risk category; and

(4) If the security is backed by a pool of multifamily residential mortgages, principal and interest payments on the security are not 30 days or more past due. Privately-issued mortgage-backed securities that do not meet these criteria or that do not qualify for a lower risk weight are generally assigned to the 100 percent risk category.

Also assigned to this category are revenue (non-general obligation) bonds or similar obligations, including loans and leases, that are obligations of states or other political subdivisions of the U.S. (for example, municipal revenue bonds) or other countries of the OECD-based group, but for which the government entity is committed to repay the debt with revenues from the specific projects financed, rather than from general tax funds.

Credit equivalent amounts of derivative contracts involving standard risk obligors (that is, obligors whose loans or debt securities would be assigned to the 100 percent risk category) are included in the 50 percent category, unless they are backed by collateral or guarantees that allow them to be placed in a lower risk category.

* * * * *

Federal Deposit Insurance Corporation

12 CFR Chapter III

Authority for Issuance

■ For the reasons stated in the common preamble, the Federal Deposit Insurance Corporation amends Part 325 of Chapter III of Title 12, Code of the Federal Regulations as follows:

PART 325—CAPITAL MAINTENANCE

■ 7. The authority citation for part 325 continues to read as follows:

Authority: 12 U.S.C. 1815(a), 1815(b), 1816, 1818(a), 1818(b), 1818(c), 1818(t), 1819(Tenth), 1828(c), 1828(d), 1828(i), 1828(n), 1828(o), 1831o, 1835, 3907, 3909, 4808; Public Law 102-233, 105 Stat. 1761, 1789, 1790, (12 U.S.C. 1831n note); Public Law 102-242, 105 Stat. 2236, as amended by Public Law 103-325, 108 Stat. 2160, 2233 (12 U.S.C. 1828 note); Public Law 102-242, 105 Stat. 2236, 2386, as amended by Public Law 102-550, 106 Stat. 3672, 4089 (12 U.S.C. 1828 note).

■ 8. Amend Appendix A to part 325 by revising footnote 39 to read as follows:

Appendix A to Part 325—Statement of Policy on Risk-Based Capital

* * * * *
 II * * *
 C. * * *
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³⁹This category would also include a first-lien residential mortgage loan on a one-to-four family property that was appropriately assigned a 50 percent risk weight pursuant to this section immediately prior to modification (on a permanent or trial basis) under the Home Affordable Mortgage Program established by the U.S. Department of Treasury, so long as the loan, as modified, is not 90 days or more past due or in nonaccrual status and meets other applicable criteria for a 50 percent risk weight. In addition, real estate loans that do not meet all of the specified criteria or that are made for the purpose of property development are placed in the 100 percent risk category.

* * * * *

**Department of the Treasury
 Office of Thrift Supervision**

12 CFR Chapter V

■ For reasons set forth in the common preamble, the Office of Thrift Supervision amends part 567 of Chapter V of title 12 of the Code of Federal Regulations as follows:

PART 567—CAPITAL

■ 9. The authority for citation for part 567 continues to read as follows:

Authority: 12 U.S.C. 1462, 1462a, 1463, 1464, 1467a, 1828 (note)

PART 567—CAPITAL

■ 10. Section 576.1 is amended in the definition *Qualifying mortgage loan* by revising paragraph (4) to read as follows

§ 567.1 Definitions.

* * * * *
Qualifying mortgage loan
 * * * * *

(4) A loan that meets the requirements of this section prior to modification on a permanent or trial basis under the U.S. Department of Treasury's Home Affordable Mortgage Program may be included as a *qualifying mortgage loan*, so long as the loan is not 90 days or more past due.

* * * * *

Dated: November 10, 2009.

John C. Dugan,
Comptroller of Currency.

By order of the Board of Governors of the Federal Reserve System, November 12, 2009.

Jennifer J. Johnson,
Secretary of the Board.

Dated at Washington DC, this 12th day of November 2009.

Federal Deposit Insurance Corporation.

Valerie J. Best,
Assistant Executive Secretary.

Dated: October 29, 2009.

By the Office of the Thrift Supervision.

John E. Bowman,
Acting Director.

[FR Doc. E9-27776 Filed 11-19-09; 8:45 am]

BILLING CODE 6714-01-P; 6210-01-P; 4810-33-P; 6720-01-P

FEDERAL RESERVE SYSTEM

12 CFR Part 226

[Regulation Z; Docket No. R-1378]

Truth in Lending

AGENCY: Board of Governors of the Federal Reserve System.

ACTION: Interim final rule; request for public comment.

SUMMARY: The Board is publishing for public comment an interim final rule amending Regulation Z (Truth in Lending). The interim rule implements Section 131(g) of the Truth in Lending Act (TILA), which was enacted on May 20, 2009, as Section 404(a) of the Helping Families Save Their Homes Act. TILA Section 131(g) became effective immediately upon enactment and established a new requirement for notifying consumers of the sale or transfer of their mortgage loans. The purchaser or assignee that acquires the loan must provide the required disclosures in writing no later than 30 days after the date on which the loan is sold or otherwise transferred or assigned. The Board is issuing this interim rule, effective immediately upon publication, so that parties subject to the statutory requirement have guidance on how to comply. However, to allow time for any necessary operational changes, compliance with the interim final rule is optional for 60 days from the date of publication; during this period, covered persons would continue to be subject to the statute's requirements. The Board seeks comment on all aspects of the interim rule.

DATES: This interim final rule is effective November 20, 2009; however, to allow time for any necessary

operational changes, compliance with this interim final rule is optional until January 19, 2010. Comments must be received on or before January 19, 2010.

ADDRESSES: You may submit comments, identified by Docket No. R-1378, by any of the following methods:

- **Agency Web Site:** <http://www.federalreserve.gov>. Follow the instructions for submitting comments at <http://www.federalreserve.gov/generalinfo/foia/ProposedRegs.cfm>.

- **Federal eRulemaking Portal:** <http://www.regulations.gov>. Follow the instructions for submitting comments.

- **E-mail:** regs.comments@federalreserve.gov.

Include the docket number in the subject line of the message.

- **Fax:** (202) 452-3819 or (202) 452-3102.

- **Mail:** Address to Jennifer J. Johnson, Secretary, Board of Governors of the Federal Reserve System, 20th Street and Constitution Avenue, NW., Washington, DC 20551.

All public comments will be made available on the Board's Web site at <http://www.federalreserve.gov/generalinfo/foia/ProposedRegs.cfm> as submitted, unless modified for technical reasons. Accordingly, comments will not be edited to remove any identifying or contact information. Public comments may also be viewed electronically or in paper in Room MP-500 of the Board's Martin Building (20th and C Streets, NW.) between 9 a.m. and 5 p.m. on weekdays.

FOR FURTHER INFORMATION CONTACT: Paul Mondor, Senior Attorney, or Stephen Shin, Attorney; Division of Consumer and Community Affairs, Board of Governors of the Federal Reserve System, Washington, DC 20551, at (202) 452-2412 or (202) 452-3667. For users of Telecommunications Device for the Deaf (TDD) only, contact (202) 263-4869.

SUPPLEMENTARY INFORMATION:

I. Background

The Truth in Lending Act (TILA), 15 U.S.C. 1601 *et seq.*, seeks to promote the informed use of consumer credit by requiring disclosures about its costs and terms. TILA requires additional disclosures for loans secured by consumers' homes and permits consumers to rescind certain transactions that involve their principal dwelling. TILA directs the Board to prescribe regulations to carry out its purposes and specifically authorizes the Board, among other things, to issue regulations that contain such classifications, differentiations, or other provisions, or that provide for such

adjustments and exceptions for any class of transactions, that in the Board's judgment are necessary or proper to effectuate the purposes of TILA, facilitate compliance with TILA, or prevent circumvention or evasion of TILA. 15 U.S.C. 1604(a). TILA is implemented by the Board's Regulation Z, 12 CFR part 226. An Official Staff Commentary interprets the requirements of the regulation and provides guidance to creditors in applying the rules to specific transactions. See 12 CFR part 226, Supp. I.

On May 20, 2009, the Helping Families Save Their Homes Act of 2009 (the "2009 Act") was signed into law. Public Law 111-22, 123 Stat. 1632. Section 404(a) of the 2009 Act amended TILA to establish a new requirement for notifying consumers of the sale or transfer of their mortgage loans. The purchaser or assignee that acquires the loan must provide the required disclosures no later than 30 days after the date on which the loan is acquired. This provision is contained in TILA Section 131(g), 15 U.S.C. 1641(g), which applies to any consumer credit transaction secured by the principal dwelling of a consumer. Consequently, the disclosure requirements in Section 131(g) apply to both closed-end mortgage loans and open-end home equity lines of credit (HELOCs).

Section 131(g) became effective immediately upon enactment on May 20, 2009, and did not require the issuance of implementing regulations. Mortgage loans sold or transferred on or after that date became subject to the requirements of Section 131(g), and failure to comply can result in civil liability under TILA Section 130(a). See 15 U.S.C. 1640(a). Accordingly, as discussed below, the Board finds there is good cause for issuing an interim rule that is effective immediately upon publication, so that parties subject to the rule have guidance on how to interpret and comply with the statutory requirements.

Under the Real Estate Settlement Procedures Act (RESPA), consumers must be notified when the servicer of their mortgage loan has changed.¹ The 2009 Act's legislative history reflects that, in addition to the information provided under RESPA, the Congress intended to provide consumers with information about the identity of the owner of their mortgage loan. In some cases, consumers that have an extended right to rescind the loan under TILA Section 125, 15 U.S.C. 1635, can assert

that right against the purchaser or assignee. See TILA Section 131(c), 15 U.S.C. 1641(c). Among other things, the 2009 Act seeks to ensure that consumers attempting to exercise this right know the identity of the assignee and how to contact the assignee or its agent for that purpose. See 155 Cong. Rec. S5098-99 (daily ed. May 5, 2009); 155 Cong. Rec. S5173-74 (daily ed. May 6, 2009). The legislative history indicates, however, that TILA Section 131(g) was not intended to require notice when a transaction "does not involve a change in the ownership of the physical note," such as when the note holder issues mortgage-backed securities but does not transfer legal title to the loan. 155 Cong. Rec. S5099.

II. Summary of the Interim Final Rule

Consistent with the legislative intent, this interim final rule implements Section 404(a) of the 2009 Act by applying the new disclosure requirements to any person or entity that acquires ownership of an existing consumer mortgage loan, whether the acquisition occurs as a result of a purchase or other transfer or assignment. A person is covered by the rule only if the person acquires legal title to the debt obligation. Although TILA and Regulation Z generally apply only to persons to whom the obligation is initially made payable and that regularly engage in extending consumer credit, Section 404(a) and the interim final rule apply to persons that acquire mortgage loans without regard to whether they also extend consumer credit by originating mortgage loans. However, the interim final rule applies only to persons that acquire more than one mortgage loan in any 12-month period.

To comply with the interim rule, a covered person must mail or deliver the required disclosures on or before the 30th day following the date that the covered person acquired the loan. The disclosure need not be given, however, if the covered person transfers or assigns the loan to another party on or before that date. This exception seeks to prevent the confusion that could result if consumers receive outdated contact information for parties that no longer own their loan. For example, a covered person that acquires a mortgage loan on March 1 must mail or deliver the disclosures on or before March 31. However, if the covered person sells or assigns the loan to a third party on March 31 (or earlier), the covered person need not provide the disclosures, but subsequent purchasers would have to comply with the rule.

III. Legal Authority

General Rulemaking Authority

As noted above, TILA Section 105(a) directs the Board to prescribe regulations to carry out the act's purposes. 15 U.S.C. 1604(a). Section 404 of the 2009 Act became effective immediately without any requirement that the Board first issue implementing rules. Nevertheless, the Board finds that the legislative purpose of Section 404 will be furthered and its effectiveness enhanced by the issuance of rules that specify the manner in which covered persons can comply with its provisions. In addition, the Board believes that implementing regulations will facilitate covered persons' compliance with the statutory provisions.

TILA also specifically authorizes the Board, among other things, to:

- Issue regulations that contain such classifications, differentiations, or other provisions, or that provide for such adjustments and exceptions for any class of transactions, that in the Board's judgment are necessary or proper to effectuate the purposes of TILA, facilitate compliance with the act, or prevent circumvention or evasion. 15 U.S.C. 1604(a).

- Exempt from all or part of TILA any class of transactions if the Board determines that TILA coverage does not provide a meaningful benefit to consumers in the form of useful information or protection. The Board must consider factors identified in the act and publish its rationale at the time it proposes an exemption for comment. 15 U.S.C. 1604(f).

Authority To Issue Interim Final Rules Without Notice and Comment

The Administrative Procedures Act (APA), 5 U.S.C. 551 *et seq.*, generally requires public notice before promulgation of regulations. See 5 U.S.C. 553(b). Unless notice or a hearing is specifically required by statute, however, the APA also provides an exception "when the agency for good cause finds (and incorporates the finding and a brief statement of reasons therefore in the rules issued) that notice and public procedure thereon are impracticable, unnecessary, or contrary to the public interest." 5 U.S.C. 553(b)(B).

As an initial matter, neither TILA nor the 2009 Act specifically requires the Board to provide notice or a hearing with respect to this rulemaking. See TILA Section 105(a), 15 U.S.C. 1604(a). In addition, the Board finds that there is good cause to conclude that providing notice and an opportunity to comment before issuing this interim final rule

¹ RESPA is implemented by Regulation X, 24 CFR part 3500, which is issued by the Department of Housing and Urban Development (HUD).

would be impracticable and contrary to the public interest. The statutory requirements in Section 404 became effective upon enactment on May 20, 2009, as noted above. Covered persons must comply with those requirements even if the Board does not issue this interim final rule.

This interim final rule implements the requirements contained in the 2009 Act but also interprets the statutory text to resolve issues and ambiguities not directly addressed by the statute. Providing notice and opportunity for comment on these matters before issuing these rules is not in the public interest because the legislation was effective upon enactment. As a result, persons covered by Section 404(a) already must be in compliance with the law or face potential liability for violations. The Board is issuing final rules at this time so that covered persons receive immediate guidance on how they can comply with the law in a manner that effectuates its purposes and avoids potential liability. The Board's issuance of a notice of proposed rulemaking for public comment would not serve this purpose because it would not provide certainty regarding a covered person's compliance obligations until the rules were finalized. By clarifying that Section 404(a) of the 2009 Act covers persons that acquire mortgage loans even if they are not "creditors" as defined under TILA, the interim final rule also ensures that consumers will receive the notice that was intended by the legislation. Consequently, the Board finds that the use of notice and comment procedures before issuing these rules would be impracticable and would not be in the public interest. Interested parties will still have an opportunity to submit comments in response to this interim final rule.

Authority To Issue Interim Final Rules That Are Effective Immediately

This interim final rule is effective upon publication in the **Federal Register**. Institutions may rely on the rules immediately to ensure they are complying with the statutory requirements. However, to allow time for any necessary operational changes, compliance with the interim final rules is optional until January 19, 2010. During this 60-day period, institutions continue to be subject to the statute's requirements.

The APA generally requires that rules be published not less than 30 days before their effective date. See 5 U.S.C. 553(d). As with the notice and comment requirement, however, the APA provides an exception when "otherwise

provided by the agency for good cause found and published with the rule." 5 U.S.C. 553(d)(3). Similarly, Section 302 of the Riegle Community Development and Regulatory Improvement Act of 1994 generally requires that new regulations and amendments to existing regulations prescribed by a Federal banking agency, which impose additional reporting, disclosure, or other new requirements on insured depository institutions, take effect on the first day of the calendar quarter that begins on or after the date on which the regulations are published in final form.² There is an exception, however, when "the agency determines, for good cause published with the regulation, that the regulations should become effective before such time." 12 U.S.C. 4802(b)(1)(A).

The interim final rule implements statutory disclosure requirements that have been in effect since May 20, 2009. For the reasons discussed above, the Board finds there is good cause to make these rules effective immediately. These rules are intended to interpret and clarify the statutory requirements and provide compliance guidance. The Board will consider public comments on the provisions before adopting further rules.

Finally, TILA Section 105(d) generally provides that a regulation requiring any disclosure that differs from the disclosures previously required shall have an effective date no earlier than "that October 1 which follows by at least six months the date of promulgation." To the extent that the interim rule contains disclosure requirements that are already in effect under the statute, Section 105(d) does not apply. Moreover, the Board believes that the effective date mandated by the 2009 Act for the specific disclosures required under section 404 overrides the general provision in TILA Section 105(d).

IV. Section-by-Section Analysis

Section 226.39—Mortgage Transfer Disclosures

39(a) Scope

Section 226.39(a) defines the scope of the interim rule's coverage. The disclosure requirements of § 226.39 apply to any "covered person," with certain exceptions that are specified in the rule. For purposes of the rule, a "covered person" includes any natural person or organization (as defined in section 226.2(a)(22) of the regulation) that acquires more than one existing

mortgage loan in any 12-month period. Consistent with the statute, the rule applies to all consumer mortgage transactions secured by the principal dwelling of a consumer, whether the transaction is a closed-end loan or an open-end line of credit.

Generally, TILA and Regulation Z apply to parties that regularly extend consumer credit. However, Section 404(a) of the 2009 Act is not limited to persons that extend credit by originating loans. Section 404(a) imposes the disclosure duty on the "creditor that is the new owner or assignee of the debt." The Board believes that to give effect to the legislative purpose, the term "creditor" in Section 404(a) must be construed to refer to the owner of the debt following the sale, transfer or assignment, without regard to whether that party would be a "creditor" for other purposes under TILA or Regulation Z. The Board declines to limit Section 404(a) to parties that originate consumer loans because such an interpretation would exempt a significant percentage of mortgage transfers which are acquisitions by secondary market investors that do not extend consumer credit and are not "creditors" for purposes of other provisions of Regulation Z.

The Board also believes that Section 404(a) of the 2009 Act does not alter the definition of "creditor" as currently used in TILA or Regulation Z. Thus, the fact that a person purchases mortgage loans and provides disclosures under § 226.39 does not by itself make that person a "creditor" for purposes of TILA and Regulation Z (even if the disclosure provided under Section 404(a) uses the term "creditor"). Accordingly, in describing the persons subject to the requirements of § 226.39, the interim final rule uses the term "covered person" rather than the term "creditor."

Under the interim final rule, the disclosure requirements in § 226.39 apply only to persons that acquire more than one consumer mortgage transaction in any 12-month period. Generally, TILA and Regulation Z cover only parties that are regularly engaged in consumer credit transactions, who are expected to have the capacity to put systems in place to ensure compliance with the rules. There is no indication in the legislative history that Section 404 was intended to apply more broadly. For example, individual homeowners might choose to facilitate the sale of their home by providing seller financing and accepting the buyer's promissory note for a portion of the purchase price. At a later date, ownership of the debt obligation might be transferred to

² See Public Law 103-325, Title III, § 302(b), Sept. 23, 1994, 108 Stat. 2214, codified at 12 U.S.C. 4802(b).

another family member or to a trust for estate planning purposes, or might be transferred to another person if the original note holder dies. The Board believes that a formal notice under Section 404 is not needed in situations involving individual transfers because the acquiring party is likely to provide adequate information to borrowers to ensure that they know to whom the loan payments should be made.

Accordingly, to prevent undue burden on individuals under the interim rule, a person who acquires only one existing mortgage loan in any 12-month period is not a covered person. The Board intends to exclude persons who are not regularly engaged in the business of purchasing or investing in consumer mortgages loans and are involved in such transactions infrequently and would not have systems in place to comply. The Board specifically solicits comment on this definition and whether the scope of the interim final rule's coverage is appropriate, or whether a different standard should apply in determining which persons must comply with the disclosure requirement in § 226.39. For example, comment is requested on whether the Board should use the same standard that applies in determining whether a person is regularly engaged in extending consumer credit, which would limit the application of § 226.39 to persons that have acquired more than five mortgage loans in the preceding or current calendar year. *See* § 226.2(a)(17)(i), footnote 3.

To become a "covered person" subject to § 226.39, a person must become the owner of an existing mortgage loan by acquiring legal title to the debt obligation. Consequently, § 226.39 does not apply to persons who acquire only a beneficial interest in the loan or a security interest in the loan, such as when the owner of the debt obligation uses the loan as security to obtain financing and the party providing the financing obtains only a security interest in the loan. Section 226.39 also does not apply to a party that assumes the credit risk without acquiring legal title to the loans. Accordingly, an investor who purchases an interest in a pool of loans (such as mortgage-backed securities, pass-through certificates, participation interests, or real estate mortgage investment conduits) but does not directly acquire legal title in the underlying mortgage loan, is not covered by § 226.39.

The Board has received a letter from the Department of Housing and Urban Development's Office of General Counsel, in its capacity as legal counsel for the Government National Mortgage

Association (Ginnie Mae), seeking to clarify Ginnie Mae's status under Section 404(a) of the 2009 Act. Ginnie Mae guarantees securities that are collateralized by mortgage loans. HUD's letter states that, as the guarantor of these securities, Ginnie Mae obtains equitable title in the mortgage loans but further states that the issuers of the securities retain legal title to the loans that collateralize the securities. According to HUD, legal title to the loans is not conveyed to Ginnie Mae unless the issuer of the securities defaults in its obligations. If the securities issuer defaults, Ginnie Mae can immediately extinguish the securities issuer's interest in the loans and take legal title. Based on HUD's representations and legal opinion regarding Ginnie Mae's status, the Board believes that the requirements of § 226.39 do not apply to Ginnie Mae until it finds the issuer in default and acquires legal title to the loans.

Section 131(f) of TILA addresses the treatment of loan servicers under the assignee liability provisions in Section 131 as well as the provisions of Section 131(g) which were added by the 2009 Act. Under TILA section 131(f)(2), a party servicing the mortgage loan is not treated as the owner of the obligation if the obligation was assigned to the servicer solely for the administrative convenience of the servicer in servicing the obligation. Accordingly, the requirements of § 226.39 do not apply to a loan servicer in this circumstance, even if the servicer holds legal title to the loan.

Some industry representatives have requested clarification whether a disclosure under § 226.39 is required in the case of a merger, acquisition, or reorganization. The Board believes that the statute covers acquisitions that occur in these situations when ownership of the loan is transferred to a different legal entity. Accordingly, the interim final rule does not provide an exception for such transactions.

39(b) Disclosure Required

Section 226.39(b) contains the general requirement for covered persons to provide the disclosures required under Section 404 of the 2009 Act, unless the exception specified in § 226.39(c) applies. The disclosures must be mailed or delivered to the consumer on or before the 30th calendar day following the date that the covered person acquires the loan. For purposes of this requirement, the date that the covered person acquires the loan is deemed to be the acquisition date that is recognized in the books and records of the acquiring party. If there is more than one covered

person, the interim rule provides that only one disclosure shall be given; the covered persons must determine among themselves which one of them will provide the disclosure. If there is more than one consumer, a covered person may mail or deliver the disclosures to any consumer who is primarily liable on the obligation.

The transfer of ownership of a mortgage loan is subject to the disclosure requirements of this section when the acquiring party is a separate legal entity from the transferor, even if the parties are affiliated entities. However, if a covered person acquires a mortgage loan and subsequently transfers the loan to another entity, the regulation does not prohibit the two entities from combining their disclosures on a single document. Comment 39(b)-2 clarifies how two entities may comply with the rules in certain circumstances by providing a single form that covers both entities. For example, a covered person that acquires a loan on August 31 might mail a single disclosure on or before September 30 with the knowledge that it will assign the loan to another entity on October 15. The covered person could mail a single disclosure providing the required information for both entities and indicating when the subsequent transfer will occur.

39(c) Exceptions

To comply with the interim final rule, a covered person must mail or deliver the required disclosures on or before the 30th day following the date that the covered person acquired the loan. Section 226.39(c)(1) provides an exception, however, if the covered person transfers or assigns the loan to another party on or before that date. This exception is made pursuant to the Board's authority to make exceptions and exemptions under TILA Sections 105(a) and 105(f). 15 U.S.C. 1604(a), 1604(f). This exception seeks to prevent the confusion that could result if consumers receive outdated contact information for parties that no longer own their loans. For example, if a mortgage loan is originated on February 22 and the original creditor sells the loan on March 1 to a covered person, the covered person must mail or deliver the disclosures required by § 226.39 on or before March 31. However, under the exception in § 226.39(c)(1) the covered person would not be required to provide the disclosures if the loan is sold or otherwise transferred or assigned to a third party on or before March 31.

The Board specifically solicits public comment on the need for this exception and its scope. The Board believes that

this exception is necessary and proper to effectuate the purposes of Section 404 and to facilitate compliance. The Board is concerned about the potential for consumers to receive multiple disclosures, some of which contain information that is outdated and inaccurate by the time it is received. This can occur because during the normal securitization process, several legal entities may be created to serve as acquisition vehicles to hold the loan for a short period before delivering the loan to an entity that ultimately holds it for the investors. After origination, a loan might be assigned to one or more entities for only a few days before it is transferred to an entity that will hold it for a much longer time period.

The Board believes that consumers may be confused if they receive one or more notices on or around the 30th day identifying multiple parties that no longer own the loan. Consequently, the interim final rule requires notices to be provided only by a covered person that still owns the loan on the 30th day after the acquisition. Thus consumers would be likely to receive notices only from parties actually holding the loan as of that date. In contrast, notices sent by temporary holders would provide information that most consumers are unlikely to need or use and could create information overload for many consumers, thereby hindering their ability to determine which party should be contacted to address a particular concern. The Board believes that the disclosure of short-term holdings of the debt obligation that do not reflect the current ownership status at the time the consumer receives the notice would be of minimal value to consumers and does not provide meaningful disclosure consistent with the purposes of TILA or the 2009 Act. Thus, the Board believes that a regulatory exception adopted pursuant to TILA Section 105(a) would effectuate TILA's purposes and facilitate compliance.

The Board has also considered the relevant statutory factors in TILA Section 105(f). The Board believes that the Section 105(f) exemption is appropriate because the disclosure of ownership interests that are held less than the 30-day period would not provide a meaningful benefit to consumers in the form of useful information or protection. It would also complicate compliance and impose unnecessary burden and expense for persons that would be required to comply, that would not be outweighed by the benefits to consumers.³ The

Board requests comment on whether the scope of this exemption is appropriate and whether the 30-day period should be shorter or longer.

In some cases, the original creditor or owner of the mortgage loan may sell or transfer the legal title to secure business financing, pursuant to a repurchase agreement that obligates the original creditor or owner to repurchase the loan within a short period, typically a month or less. Under § 226.39(c)(2) of the interim final rule, if the original creditor or owner does not recognize the transaction as a sale of the loan on its books and records for accounting purposes, the acquiring party is not subject to the disclosure requirements of § 226.39. However, if the transferor does not repurchase the mortgage loan, the acquiring party must make the disclosures required by § 226.39 within 30 days after the date that the transaction is recognized as an acquisition in its books and records. This exception is also being adopted pursuant to the Board's authority in TILA Sections 105(a) and 105(f). As with the exception in § 226.39(c)(1), the exception for repurchase agreements in § 226.39(c)(2) seeks to prevent consumer confusion from the receipt of outdated disclosures. The Board believes that providing disclosures for the transactions covered by the exception in § 226.39(c)(2) would not provide a meaningful benefit to consumers in the form of useful information or protection. The Board also believes that the disclosure of transfers that are subject to repurchase agreements would complicate compliance and impose unnecessary burden and expense for persons that would be required to comply, that would not be outweighed by the benefits to consumers. Comment is requested on this exception, and any unintended consequences that may result.

39(d) Content of Required Disclosures

Section 226.39(d) sets forth the contents of the notice that must be

coverage of such transactions provides a meaningful benefit to consumers in light of specific factors. 15 U.S.C. 1604(f)(2). These factors, which the Board has reviewed, are (1) the amount of the loan and whether the disclosure provides a benefit to consumers who are parties to the transaction involving a loan of such amount; (2) the extent to which the requirement complicates, hinders, or makes more expensive the credit process; (3) the status of the borrower, including any related financial arrangements of the borrower, the financial sophistication of the borrower relative to the type of transaction, and the importance to the borrower of the credit, related supporting property, and coverage under TILA; (4) whether the loan is secured by the principal residence of the borrower; and (5) whether the exemption would undermine the goal of consumer protection.

provided under this section. The disclosures must identify the loan that was acquired or transferred and, consistent with the statute, contain the following: (1) The identity, address, and telephone number of the covered person that owns the mortgage loan; (2) the date of the acquisition or transfer; (3) contact information that the consumer can use to reach an agent or party having authority to act on behalf of the covered person; (4) the location of the place where the transfer of the ownership of the debt is recorded.

Identity, address, and telephone number. Section 226.39(d)(1) requires acquiring parties to provide their name, as well as their address and telephone number. Under the interim final rule, the party identified must be the covered person who owns the mortgage loan, regardless of whether another party has been appointed to service the loan or otherwise serve as the covered person's agent. The covered person has the option of also providing an electronic mail address or Internet Web site address but is not required to do so.

Section 226.39(d)(1) provides that if there is more than one covered person, the required information must be provided for each of them. The Board specifically solicits comments on the benefits of this approach, or whether the identification of multiple parties may create confusion for consumers. Should there be limits on the number of covered persons identified and, if so, what limits would be appropriate consistent with the legislative intent?

Acquisition date. Section 226.39(d)(2) requires disclosure of the date that the covered person acquired the loan. For purposes of this section, this is defined as the date of acquisition recognized in the books and records of the covered person. The Board believes that this approach provides flexibility to accommodate a variety of circumstances in which the acquisition could occur.

Agent's contact information. Under § 226.39(d)(3), a covered person must identify and provide contact information for the agent or party having authority to act on behalf of the covered person. The notice must identify one or more persons who are authorized to receive legal notices on behalf of the covered person and resolve issues concerning the consumer's payments on the loan. However, contact information for an agent is not required to be provided under § 226.39(d)(3) if the consumer can use the information provided for the covered person provided under paragraph § 226.39(d)(1) for these purposes. Thus, the interim final rule implements the disclosure requirement in Section 404 but does not

³ In exercising its exemption authority under Section 105(f), Board must determine whether

require that the owner of a loan designate an agent or other party for any specific purpose. The rule simply requires that the owner disclose contact information when there is such an agent, so that consumers can direct their inquiries to the appropriate party.

The Board recognizes that separate entities may be authorized by the owner of the loan to act on its behalf for different purposes. Identifying the party authorized to receive legal notices is intended to ensure that consumers have sufficient information to assert legal claims, including a right to rescind the loan, if applicable. However, a covered person might appoint a different agent to resolve loan servicing issues. In such cases, the covered person must provide contact information for each agent. If multiple agents are listed, the disclosure must state the extent to which the authority of each agent differs, for example, by indicating if only one of the agents is authorized to receive legal notices or only one is authorized to resolve issues concerning payments.

A covered person may comply with § 226.39(d)(3) by providing a telephone number on the written disclosure if the consumer can use the telephone number to obtain the address of the agent or other authorized person identified. This differs from the requirement in § 226.39(d)(1), which requires covered persons who acquire a loan to provide their name, address, and telephone number in all cases. The flexibility in § 226.39(d)(3) is intended to allow covered persons to use a single disclosure form that contains a nationwide toll-free telephone number, even though there may be different physical locations to which documents should be sent in different regions of the country. Comment is specifically solicited on this approach and whether both a telephone number and address for the agent or authorized representative should be required to be included on each disclosure under § 226.39(d)(3).

Comment 39(d)(3)–2 clarifies that the covered person has the option of also providing the agent's electronic mail address or internet web site address but is not required to do so.

Recording location. Section 404 requires that the disclosure state the location of the place where the transfer of ownership of the debt is recorded. When a mortgage loan is sold, however, the transfer in ownership of the debt instrument typically is not recorded in public records. The new owner's security interest in the property that secures the debt may or may not be recorded in the public land records or, if it is recorded, it may not yet be

recorded at the time the disclosure is sent.

Consistent with the statute, § 226.39(d)(4) of the interim final rules requires covered persons to disclose the location where their ownership of the debt is recorded. However, if the transfer of ownership has not been recorded in public records at the time the disclosure is provided, the covered person can comply with the rule by stating this fact. Whether or not the transfer of ownership has been recorded in public records at the time the disclosure is made, the disclosure may state that the transfer "is or may be recorded" at the specified location.

The covered person also has the option of disclosing the location where the covered person's *security interest* in the property is or may be recorded. In light of the fact that the transfer in ownership of the debt instrument usually is not recorded in public records, the Board specifically solicits comment on whether disclosure of the location where the security interest is recorded should be required.

Comment 39(d)(4)–2 clarifies that the covered person is not required to provide the postal address for the governmental office where the covered person's ownership interest is recorded or the name of the jurisdiction where the property is located. For example, it would be sufficient in all cases to disclose that the transaction is or may be recorded in the office of public land records or the recorder of deeds office "for the county or local jurisdiction where the property is located."

The Board has taken this approach after considering the relative costs and benefits of requiring that the disclosure provide more detailed information. Industry representatives have noted that this information may not be readily accessible to the acquiring party. A requirement to provide the name and address of the governmental office would require parties that provide such notices to develop and maintain a system for matching the property address to the correct governmental office, and keeping the database up to date with correct address information. The Board does not believe that this would provide substantial benefit to consumers because they presumably know the county or jurisdiction in which the property is located and can easily obtain the address of the governmental office from public directories or other sources. The Board solicits comments on the approach taken in the interim final rule and the relative costs and benefits of requiring more detailed disclosures about the

location where the lender's security interest is or may be recorded.

39(e) Optional Disclosures

Section 404 provides that the party acquiring a loan shall notify the borrower of "any other relevant information" regarding the new owner of the loan. The Board interprets this statutory language as permitting the Board to impose additional disclosure requirements to further the legislative purpose. Any additional disclosure requirements would be imposed by regulation after notice and comment. The Board does not believe that the statutory language requires covered persons to determine independently what additional information a reviewing court might subsequently determine to be legally relevant in order to avoid liability. Although the interim final rule does not contain any additional disclosure requirements, the Board solicits comment on whether the rule should include any such requirements. The Board also believes that, under the statutory language, covered persons are permitted, in their sole discretion, to include additional information that they might deem relevant or helpful to consumers, which is reflected in § 226.39(e) of the interim final rule. For example, the covered person may choose to inform consumers that the location where they should send mortgage payments has not changed.

V. Initial Regulatory Flexibility Analysis

The Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*) requires an initial and final regulatory flexibility analysis only when 15 U.S.C. 553 requires publication of a notice of proposed rulemaking. *See* 5 U.S.C. 603(a), 604(a). However, the Board has found good cause under 5 U.S.C. 553(b)(B) to conclude that, with respect to this interim final rule, publication of a notice of proposed rulemaking is impracticable and not in the public interest. Accordingly, the Board is not required to perform an initial or final regulatory flexibility analysis. Nonetheless, to solicit additional information from small entities subject to the interim final rule, the Board is publishing an initial regulatory flexibility analysis.

Based on its analysis and for the reasons stated below, the Board believes that this interim final rule will not have a significant economic impact on a substantial number of small entities. The Board invites comment on the effect of the interim final rule on small entities.

A. Reasons for the Interim Final Rule

As indicated above, the 2009 Act was signed into law on May 20, 2009. Section 404 amended TILA to establish a new requirement for notifying consumers of the sale or transfer of their mortgage loans. This requirement became effective immediately upon enactment on May 20, 2009, and did not require the issuance of implementing regulations. As discussed above, the Board believes there is good cause for an interim final rule so that parties subject to the rule have guidance on how to interpret and comply with the statutory requirements and consumers receive notices consistent with legislative intent.

Congress enacted TILA based on findings that economic stability would be enhanced and competition among consumer credit providers would be strengthened by the informed use of credit resulting from consumers' awareness of the cost of credit. One of the stated purposes of TILA is to provide a meaningful disclosure of credit terms to enable consumers to compare credit terms available in the marketplace more readily and avoid the uninformed use of credit.

B. Summary of 2009 Act

As described previously, the purchaser or assignee that acquires a loan must provide the required disclosures no later than 30 days after the date on which the loan is acquired. Section 226.39(c) of the rule provides an exception if the covered person transfers or assigns the loan to another party on or before that date. Section 226.39(d) sets forth the contents of the notice. Consistent with the statute, the interim final rule requires that the notice contain the following: (1) The identity, address, and telephone number of the covered person who owns the mortgage loan; (2) the acquisition date; (3) a mailing address and telephone number that the borrower can use to reach an agent of the covered person; and (4) the location where the covered person's interest in the property securing the loan is or may be recorded.

C. Statement of Objectives and Legal Basis

The **SUPPLEMENTARY INFORMATION** contains this information. The legal basis for the interim final rule is in TILA Sections 105(a), 105(f), 15 U.S.C. 1604(a), 1604(f). A more detailed discussion of the Board's rulemaking authority is set forth in the **SUPPLEMENTARY INFORMATION**.

D. Description of Small Entities to Which the Interim Final Rule Would Apply

The interim final rule would apply to all persons that acquire more than one existing mortgage loan in any 12-month period, other than servicers that take title solely as an administrative convenience to enable them to service the loans. The Board cannot identify with certainty the number of small entities that meet this definition. The Board can estimate, however, approximate numbers of small entities that purchase mortgage loans, as discussed below.

The Board can identify through data from Reports of Condition and Income ("call reports") approximate numbers of small depository institutions that would be subject to the interim final rules if they acquire more than one mortgage loan in a 12-month period. Approximately 16,345 depository institutions in the United States filed call report data in December of 2008, of which approximately 11,907 had total domestic assets of \$175 million or less and thus were considered small entities for purposes of the Regulatory Flexibility Act. Of 4231 banks, 565 thrifts and 7111 credit unions that filed call report data and were considered small entities, 4091 banks, 530 thrifts, and 4797 credit unions, totaling 9418 institutions, extended mortgage credit. For purposes of this analysis, thrifts include savings banks, savings and loan entities, co-operative banks and industrial banks.

The Board cannot identify with certainty the number of small non-depository institutions because they do not file call reports. Neither can the Board determine with certainty how many of the 11,907 institutions identified above as small entities acquired mortgage loans in 2008. Although an estimated 9418 such institutions extended mortgage credit, the Board recognizes that not all entities that extend mortgage credit also acquire existing mortgage loans. Moreover, the reverse is also true: there are entities that acquire existing mortgage loans but do not extend mortgage credit.

The Board has another source of information, data obtained under the Home Mortgage Disclosure Act (HMDA), 12 U.S.C. 2801 *et seq.*; 12 CFR part 203. Based on loan purchases reported for 2008 under HMDA, the Board estimates that 553 of the reporting institutions engaged in more than one mortgage acquisition. The 8388 lenders covered by HMDA in 2008 accounted for the majority, but not all, of the home lending in the United States.

Accordingly, the 553 institutions that reported loan purchases in 2008 probably do not represent all mortgage acquirers; institutions must report loan purchases only if they are required to report under HMDA based on loan originations and assets. Nevertheless, the Board's experience has been that the HMDA data are reasonably representative of the whole mortgage market.

A total of 2,921,684 loan purchases were reported under HMDA in 2008 by entities reporting more than one purchase (and thus subject to the interim final rule). Of those loan purchases, 2,773,918 were reported by depository institutions. Of those depository institution loan purchases, 2,122,288 (76.5%) were reported by large depository institutions (assets greater than \$175 million), and 651,630 (23.5%) were reported by small depository institutions (assets of \$175 million or less). Of the 553 HMDA reporters reporting more than one loan purchase, 502 were depository institutions. Of those 502 depository institutions, 387 (77.1%) were large and 115 (22.9%) were small. Those 115 small depository institutions represent just slightly less than one percent (0.97%) of the 11,907 total small institutions estimated above from call report data.

A total of 147,766 loan purchases were reported under HMDA by non-depository institutions that reported more than one loan purchase in 2008. The Board cannot tell from the HMDA data how many of those loan purchases were reported by small entities. Neither can the Board tell how many of the 51 non-depository institutions that reported those loan purchases are small entities. If the relative shares among small and large non-depository institutions do not differ significantly from those among depository institutions, however, the shares for non-depository institutions can be estimated. On that basis, the Board estimates that 12 small non-depository institutions reported 34,725 loan purchases and that 39 large non-depository institutions reported 113,041 loan purchases (estimates are rounded to whole numbers).

Using the foregoing numbers from 2008 HMDA data for depository institutions and the foregoing estimates for non-depository institutions, the Board estimates the following numbers for all entities reporting under HMDA combined: of the 2,921,684 loan purchases reported by 553 entities reporting more than one purchase, 2,235,329 (76.5%) were reported by 426 large entities (77%), and 686,355

(23.5%) were reported by 127 small entities (23%). Based on these estimates, less than one-quarter of the institutions reporting covered loan purchases under HMDA were small entities, and less than one-quarter of the covered loan purchases reported were reported by small entities.

The foregoing data are not complete in many respects. Not all depository institutions that file call reports are reporters under HMDA, and not all HMDA reporters file call reports. Further, some unknown number of entities purchase more than one mortgage loan in any 12-month period and yet file neither call reports nor HMDA data; how many of those are small entities also is unknown. Nevertheless, if one assumes that the existing data are reasonably representative of the market as a whole, they present an overall picture of minimal economic impact on small entities. For all these reasons, the Board believes that the interim final rule will not have a significant economic impact on a substantial number of small entities.

E. Projected Reporting, Recordkeeping, and Other Compliance Requirements

The compliance requirements of the interim final rules are described in the **SUPPLEMENTARY INFORMATION**. As indicated above, the Board is adopting a new disclosure rule requiring that consumers receive notice when ownership of their mortgage loan is transferred. The Board is aware that numerous covered persons are already complying with these statutory provisions, which became effective on May 20, 2009. Therefore the additional burden imposed by the Board's rule itself is likely to be minimal. Furthermore, the information required to be provided is easily obtainable by the covered person. The covered person must provide contact information for itself and any agent (but is not required to designate an agent), may use the acquisition date in its own books and records, and may generally describe the location where the covered person's interest in the property securing the mortgage loan is or may be recorded. This information generally is already required by the statute.

Based on informal surveys of industry representatives and practices in effect, the Board understands that entities are likely to designate servicers as their agents. Servicers already respond to consumer requests on the behalf of covered persons. Therefore, other than providing the notice itself, covered persons (including those who are small entities) are not likely to incur

significant burden in responding to consumer requests. Furthermore, the Board has provided an exception to the rule for mortgage owners who do not hold the loan more than 30 days. The Board believes that this exception balances the needs of consumers for information with the burdens on industry of compliance and the potential for confusion to consumers of multiple disclosures.

F. Other Federal Rules

The Board has not identified other rules that conflict with the rule. As indicated previously, under RESPA and HUD's Regulation X, consumers must be notified when the servicer of their mortgage loan has changed. Therefore, the disclosure of contact information for the agent of the owner of the mortgage loan, typically the servicer under applicable agreements, is already generally required by law. As a result of existing requirements, servicers are already subject to disclosure of their contact information and are already subject to calls regarding administration of payment information.

G. Significant Alternatives to the Interim Final Rule

As noted above, this interim final rule implements the statutory requirements of the 2009 Act that were effective on May 20, 2009. The Board has implemented these requirements to minimize burden while retaining benefits to consumers. The Board was not required to issue rules but has decided that rules are needed to clarify who is subject to the requirements and what information must be disclosed, and to ensure that consumers receive disclosures of ownership that are consistent with legislative intent. The Board welcomes comment on any significant alternatives that would minimize the impact of the interim final rule on small entities.

The Board welcomes further information and comment on any costs, compliance requirements, or changes in operating procedures arising from the application of the interim final rule to small businesses.

VI. Paperwork Reduction Act

In accordance with the Paperwork Reduction Act (PRA) of 1995 (44 U.S.C. 3506; 5 CFR part 1320 appendix A.1), the Board reviewed the interim final rule under the authority delegated to the Board by the Office of Management and Budget (OMB). The collection of information that is required by this final rule is found in 12 CFR 226.39. The Board may not conduct or sponsor, and an organization is not required to

respond to, this information collection unless the information collection displays a currently valid OMB control number. The OMB control number is 7100-0199.

This information collection is required to provide benefits for consumers and is mandatory (15 U.S.C. 1601 *et seq.*). Since the Board does not collect any information, no issue of confidentiality arises. The respondents/recordkeepers are persons or entities that acquire legal title to more than one mortgage loan in any 12-month period, including for-profit financial institutions and small businesses.

TILA and Regulation Z are intended to ensure effective disclosure of the costs and terms of credit to consumers. For closed-end loans, such as mortgage and installment loans, cost disclosures are required to be provided prior to consummation. Special disclosures are required in connection with certain products, such as reverse mortgages, certain variable-rate loans, and certain mortgages with rates and fees above specified thresholds. To ease the burden and cost of complying with Regulation Z (particularly for small entities), the Board provides model forms, which are appended to the regulation. TILA and Regulation Z also contain rules concerning credit advertising. Creditors are required to retain evidence of compliance with Regulation Z for 24 months (12 CFR 226.25), but Regulation Z does not specify the types of records that must be retained.

Under the PRA, the Board accounts for the paperwork burden associated with Regulation Z for the state member banks and other entities supervised by the Board that engage in activities covered by Regulation Z and, therefore, are respondents under the PRA. Appendix I of Regulation Z defines the institutions supervised by the Federal Reserve System as: state member banks, branches and agencies of foreign banks (other than federal branches, Federal agencies, and insured state branches of foreign banks), commercial lending companies owned or controlled by foreign banks, and organizations operating under section 25 or 25A of the Federal Reserve Act. Other Federal agencies account for the paperwork burden imposed on the entities for which they have administrative enforcement authority under TILA.

The current total annual burden to comply with the provisions of Regulation Z is estimated to be 1,011,311 hours for the 1,138 institutions supervised by the Federal Reserve that are deemed to be respondents for the purposes of the PRA.

As discussed in the preamble, the Board is adopting a new disclosure rule requiring that consumers receive notice when ownership of their mortgage loan is transferred. The new disclosure requirement will impose a one-time increase in the total annual burden under Regulation Z for respondents supervised by the Federal Reserve that engage in mortgage acquisitions. The Board estimates that 68 respondents⁴ supervised by the Federal Reserve will take, on average, 40 hours (one business week) to update their systems, internal procedure manuals, and provide training for relevant staff to comply with the new disclosure requirements in § 226.39. Accordingly, this revision is estimated to result in a one-time increase in the aggregate burden by 2,720 hours for these 68 respondents.

On a continuing basis, the Board estimates that 68 respondents supervised by the Federal Reserve would take, on average, 8 hours⁵ per month to comply with the new disclosure requirements, which would increase the ongoing aggregate burden by 6,528 hours annually for these respondents. Accordingly, the Board estimates that the new disclosure requirement will increase the total annual burden on a continuing basis for respondents supervised by the Federal Reserve from 1,011,311 to 1,017,839 hours (not including the one-time increase of 2,720 hours to implement the changes, as described above). This total estimated burden increase represents averages for all respondents supervised by the Federal Reserve. The Board expects that the amount of time required to implement each of the changes for a given institution may vary based on the size and complexity of the respondent.

The other federal financial institution supervisory agencies (the Office of the Comptroller of the Currency (OCC), the Office of Thrift Supervision (OTS), the Federal Deposit Insurance Corporation (FDIC), and the National Credit Union Administration (NCUA)) are responsible for estimating and reporting to OMB the

total paperwork burden for the domestically chartered commercial banks, thrifts, and federal credit unions and U.S. branches and agencies of foreign banks for which they have primary administrative enforcement jurisdiction under TILA Section 108(a), 15 U.S.C. 1607(a). These agencies may, but are not required to, use the Board's methodology for estimating burden. Using the Board's method, the total current estimated annual burden for the approximately 17,200 domestically chartered commercial banks, thrifts, and federal credit unions and U.S. branches and agencies of foreign banks supervised by the Board, OCC, OTS, FDIC, and NCUA under TILA would be approximately 17,765,525 hours. The final rule will impose a one-time increase in the estimated annual burden for the estimated 638 institutions thought to engage in mortgage acquisitions by 25,520 hours. On a continuing basis the annual burden would increase by 61,248 hours. The total annual burden is estimated to be 17,852,293 hours. The above estimates represent an average across all respondents and reflect variations between institutions based on their size, complexity, and practices.

The Board has a continuing interest in public opinion on its collections of information. At any time, comments regarding the burden estimate or any other aspect of this collection of information, including suggestions for enhancing the quality of information collected and ways for reducing the burden on respondent. Comments on the collection of information may be sent to: Secretary, Board of Governors of the Federal Reserve System, 20th and C Streets, NW., Washington, DC 20551; and to the Office of Management and Budget, Paperwork Reduction Project (7100-0199), Washington, DC 20503.

List of Subjects in 12 CFR Part 226

Consumer protection, Federal Reserve System, Mortgages, Reporting and recordkeeping requirements, Truth in lending.

Authority and Issuance

■ For the reasons set forth in the preamble, the Board amends Regulation Z, 12 CFR part 226, as set forth below:

PART 226—TRUTH IN LENDING (REGULATION Z)

■ 1. The authority citation for part 226 continues to read as follows:

Authority: 12 U.S.C. 3806; 15 U.S.C. 1604, 1637(c)(5), and 1639(l); Public Law 111-24 § 2, 123 Stat. 1734.

Subpart E—Special Rules for Certain Home Mortgage Transactions

■ 2. Add a new § 226.39 to Subpart E of Part 226 to read as follows:

§ 226.39 Mortgage transfer disclosures.

(a) *Scope.* The disclosure requirements of this section apply to any covered person except as otherwise provided in this section. For purposes of this section:

(1) A “covered person” means any person, as defined in § 226.2(a)(22), that becomes the owner of an existing mortgage loan by acquiring legal title to the debt obligation, whether through a purchase, assignment, or other transfer, and who acquires more than one mortgage loan in any twelve-month period. For purposes of this section, a servicer of a mortgage loan shall not be treated as the owner of the obligation if the servicer holds title to the loan or it is assigned to the servicer solely for the administrative convenience of the servicer in servicing the obligation.

(2) A “mortgage loan” means any consumer credit transaction that is secured by the principal dwelling of a consumer.

(b) *Disclosure required.* Except as provided in paragraph (c) of this section, any person that becomes a covered person as defined in this section shall mail or deliver the disclosures required by this section to the consumer on or before the 30th calendar day following the acquisition date. If there is more than one covered person, only one disclosure shall be given and the covered persons shall agree among themselves which covered person shall comply with the requirements that this section imposes on any or all of them.

(1) *Acquisition date.* For purposes of this section, the date that the covered person acquired the mortgage loan shall be the date of acquisition recognized in the books and records of the acquiring party.

(2) *Multiple consumers.* If there is more than one consumer liable on the obligation, a covered person may mail or deliver the disclosures to any consumer who is primarily liable.

(c) *Exceptions.* Notwithstanding paragraph (b) of this section, a covered person is not subject to the requirements of this section with respect to a particular mortgage loan if:

(1) The covered person sells or otherwise transfers or assigns legal title to the mortgage loan on or before the 30th calendar day following the date that the covered person acquired the mortgage loan; or

⁴ Based on loan purchases reported for 2008 under the Home Mortgage Disclosure Act (HMDA), 12 U.S.C. 2801 *et seq.*, and Regulation C (12 CFR part 203), the Board estimates that 58 of the 553 institutions engaged in such mortgage acquisitions are supervised by the Federal Reserve. Based on average Call Report data for the past four quarters, approximately 95 institutions that do not report under HMDA also would be subject to these new disclosure requirements and 10 of these institutions are supervised by the Federal Reserve.

⁵ Because financial institutions are familiar with the existing RESPA provisions which require notification to consumers when the servicer of their mortgage loan has changed, the Federal Reserve believes that implementation of requirements in § 226.39 should not be overly burdensome.

(2) The mortgage loan is transferred to the covered person in connection with a repurchase agreement and the transferor that is obligated to repurchase the loan continues to recognize the loan as an asset on its own books and records. However, if the transferor does not repurchase the mortgage loan, the acquiring party must make the disclosures required by § 226.39 within 30 days after the date that the transaction is recognized as an acquisition in its books and records.

(d) *Content of required disclosures.* The disclosures required by this section shall identify the loan that was acquired or transferred and state the following:

(1) The identity, address, and telephone number of the covered person who owns the mortgage loan. If there is more than one covered person, the information required by this paragraph shall be provided for each of them.

(2) The acquisition date recognized by the covered person.

(3) How to reach an agent or party having authority to act on behalf of the covered person (or persons), which shall identify a person (or persons) authorized to receive legal notices on behalf of the covered person and resolve issues concerning the consumer's payments on the loan. However, no information is required to be provided under this paragraph if the consumer can use the information provided under paragraph (d)(1) of this section for these purposes. If multiple persons are identified under this paragraph, the disclosure shall provide contact information for each and indicate the extent to which the authority of each agent differs. For purposes of this paragraph (d)(3), it is sufficient if the covered person provides only a telephone number provided that the consumer can use the telephone number to obtain the address for the agent or other person identified.

(4) The location where transfer of ownership of the debt to the covered person is recorded. However, if the transfer of ownership has not been recorded in public records at the time the disclosure is provided, the covered person complies with this paragraph by stating this fact.

(e) *Optional disclosures.* In addition to the information required to be disclosed under paragraph (d) of this section, a covered person may, at its option, provide any other information regarding the transaction.

■ 3. In Supplement I to Part 226, under Subpart E, a new *Section 226.39—Mortgage Transfer Disclosures* is added to read as follows:

Supplement I to Part 226—Official Staff Interpretations

* * * * *

Subpart E—Special Rules for Certain Home Mortgage Transactions

* * * * *

Section 226.39—Mortgage transfer disclosures.

39(a) Scope.

Paragraph 39(a)(1).

1. *Covered persons.* The disclosure requirements of § 226.39 apply to any “covered person” that becomes the legal owner of an existing mortgage loan, whether through a purchase, assignment, or other transfer, regardless of whether the person also meets the definition of a “creditor” in Regulation Z. The fact that a person purchases or acquires mortgage loans and provides disclosures under § 226.39 does not by itself make that person a “creditor” as defined in the regulation.

2. *Acquisition of legal title.* To become a “covered person” subject to § 226.39, a person must become the owner of an existing mortgage loan by acquiring legal title to the debt obligation. The transfer of ownership of a mortgage loan is subject to the disclosure requirements of this section when the acquiring party is a separate legal entity from the transferor, even if the parties are affiliated entities. Section 226.39 does not apply to persons who acquire only a beneficial interest in the loan or a security interest in the loan. Section 226.39 also does not apply to a party that assumes the credit risk without acquiring legal title to the loan. Thus, an investor that acquires mortgage-backed securities, pass-through certificates, or participation interests and does not directly acquire legal title in the underlying mortgage loans is not covered by this section.

3. *Loan servicers.* Pursuant to TILA Section 131(f)(2), the servicer of a mortgage loan is not treated as the owner of the obligation for purposes of § 226.39 if the servicer holds title to the loan as a result of the assignment of the obligation to the servicer solely for the administrative convenience of the servicer in servicing the obligation.

4. *Mergers, corporate acquisitions, or reorganizations.* Disclosures are required under § 226.39 when, as a result of a merger, corporate acquisition, or reorganization the ownership of a mortgage loan is transferred to a different legal entity.

Paragraph 39(a)(2).

1. *Mortgage transactions covered.* Section 226.39 applies to any consumer credit transaction secured by the principal dwelling of a consumer, which includes closed-end mortgage loans as well as home equity lines of credit.

39(b) Disclosure required.

1. *Generally.* A covered person must mail or deliver the disclosures required by § 226.39 on or before the 30th calendar day following the date that the covered person acquired the loan, unless the exception in § 226.39(c) applies. For example, if a covered person acquires a mortgage loan on March 1, the required disclosure must be mailed or

delivered on or before March 31. For purposes of this requirement, the date that the covered person acquires the loan is the acquisition date recognized in its books and records.

2. *Disclosure provided on behalf of multiple entities.* A mortgage loan may be acquired by a covered person and subsequently transferred to an affiliate or other entity that is also a covered person required to provide disclosures under § 226.39. In such cases, a single disclosure may be provided on behalf of both entities instead of providing two separate disclosures, as long as the disclosure satisfies the timing and content requirements applicable to both entities. For example, if a covered person acquires a loan on August 31 with the knowledge that it will assign the loan to another entity on October 15, the covered person could mail a single disclosure on or before September 30 which provides the required information for both entities and indicates when the subsequent transfer is expected to occur. Even though one person delegates responsibility for the disclosures to another covered person, each has a duty to ensure that disclosures related to its acquisition are accurate and provided in a timely manner.

39(c) Exceptions.

Paragraph 39(c)(1).

1. *Example.* If a mortgage loan is originated on February 22nd and the original creditor sells the loan on March 1 to a covered person, under the exception in § 226.39(c) the covered person would not be required to provide disclosures under § 226.39 if the loan is sold or otherwise transferred or assigned to another party on or before March 31.

Paragraph 39(c)(2).

1. *Repurchase agreements.* The original creditor or owner of the mortgage loan might sell or transfer legal title to the loan to secure short-term business financing under an agreement where the original creditor or owner is also obligated to repurchase the loan within a brief period, typically a month or less. If the original creditor or owner does not recognize such transactions as a sale of the loan on its own books and records for accounting purposes, the transfer of the loan in connection with such a repurchase agreement is not covered by § 226.39 and the acquiring party is not required to provide disclosures. However, if the transferor does not repurchase the mortgage loan, the acquiring party must make the disclosures required by § 226.39 within 30 days after the date that the transaction is recognized as an acquisition in its books and records.

39(d) Content of required disclosures.

1. *Identifying the loan.* The disclosures required by this section should identify the loan that was acquired or transferred. The covered person has flexibility in determining what information to provide for this purpose. For example, the covered person may identify the loan by stating the address of the mortgaged property along with the account number or other identification number previously known to the consumer, which may appear in a truncated format. Alternatively, the covered person might identify the loan by specifying the date on which the credit was extended and the original amount of the loan or credit line.

Paragraph 39(d)(1).

1. *Identification of covered person.* Section 226.39(d)(1) requires acquiring parties to provide their name, address, and telephone number. The party identified must be the covered person who owns the mortgage loan, regardless of whether another party has been appointed to service the loan or otherwise serve as the covered person's agent. In addition to providing a postal address and a telephone number, the covered person may, at its option, provide an address for receiving electronic mail or an internet web site address but is not required to do so.

Paragraph 39(d)(3).

1. *Identifying agents.* Under § 226.39(d)(3), the covered person must provide contact information for the agent or other party having authority to act on behalf of the covered person and who is authorized to receive legal notices on behalf of the covered person and resolve issues concerning the consumer's payments on the loan. Section 226.39(d)(3) does not require that a covered person designate an agent or other party, but if the consumer cannot use the covered person's contact information for these purposes the disclosure must provide contact information for an agent or other party that can address these matters. If multiple agents are listed on the disclosure, the disclosure shall state the extent to which the authority of each agent differs by indicating if only one of the agents is authorized to receive legal notices, or only one of the agents is authorized to resolve issues concerning payments. For purposes of § 226.39(d)(3), it is sufficient to provide a telephone number as the contact information provided that consumers can use the telephone number to obtain the mailing address for the agent or other person identified.

2. *Other contact information.* The covered person may also provide an agent's electronic mail address or internet web site address but is not required to do so.

Paragraph 39(d)(4).

1. *Recording location.* Section 226.39(d)(4) requires disclosure of the location where transfer of ownership of the debt to the covered person is recorded. If the transfer of ownership has not been recorded in public records at the time the disclosure is provided, the covered person complies with § 226.39(d)(4) by stating this fact. Whether or not the transfer has been recorded at the time the disclosure is made, the disclosure may state that the transfer "is or may be recorded" at the specified location.

2. *Postal address not required.* In disclosing the location where the transfer of ownership is recorded, the covered person is not required to provide a postal address for the governmental office where the covered person's ownership interest is recorded. The covered person also is not required to provide the name of the county or jurisdiction where the property is located. For example, it would be sufficient to disclose that the transaction is or may be recorded in the office of public land records or the recorder of deeds office "for the county or local jurisdiction where the property is located."

39(e) Optional disclosures.

1. *Generally.* Section 226.39(e) provides that covered persons may, at their option,

include additional information about the mortgage transaction that they consider relevant or helpful to consumers. For example, the covered person may choose to inform consumers that the location where they should send mortgage payments has not changed.

By order of the Board of Governors of the Federal Reserve System, November 13, 2009.

Jennifer J. Johnson,

Secretary of the Board.

[FR Doc. E9-27742 Filed 11-19-09; 8:45 am]

BILLING CODE 6210-01-P

DEPARTMENT OF ENERGY**Federal Energy Regulatory Commission****18 CFR Part 358**

[Docket No. RM07-1-002; Order No. 717-B]

Standards of Conduct for Transmission Providers; Order on Rehearing and Clarification

Issued November 16, 2009.

AGENCY: Federal Energy Regulatory Commission.

ACTION: Order on rehearing and clarification.

SUMMARY: The Federal Energy Regulatory Commission (Commission) issued Order No. 717-A to make even clearer the Standards of Conduct as implemented by Order No. 717. This order addresses requests for rehearing and clarification concerning paragraph 80 of Order No. 717-A and whether an employee who is not making business decisions about contract non-price terms and conditions is considered a "marketing function employee." **DATES:** *Effective Date:* This rule will become effective November 23, 2009. **FOR FURTHER INFORMATION CONTACT:** Leonard Tao, Office of the General Counsel—Energy Markets, Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426, (202) 502-8214.

SUPPLEMENTARY INFORMATION:

129 FERC ¶ 61,123

Before Commissioners: Jon Wellinghoff, Chairman; Suedeen G. Kelly, Marc Spitzer, and Philip D. Moeller.

I. Introduction

1. On October 16, 2008, the Commission issued Order No. 717 amending the Standards of Conduct for Transmission Providers (the Standards of Conduct or the Standards) to make them clearer and to refocus the rules on the areas where there is the greatest

potential for abuse.¹ On October 15, 2009, the Commission issued Order No. 717-A to address requests for rehearing and clarification of Order No. 717, largely affirming the reforms adopted in Order No. 717.² In this order, the Commission grants limited rehearing and clarification to address certain specific matters petitioners raised regarding one of the Commission's determinations in Order No. 717-A.

II. Discussion*Independent Functioning Rule: Marketing Function Employees*

2. In paragraph 80 of Order No. 717-A, the Commission stated the following:

The Commission clarifies that an employee in the legal, finance or regulatory division of a jurisdictional entity, whose intermittent day-to-day duties include the drafting and redrafting of non-price terms and conditions of, or exemptions to, umbrella agreements is a "marketing function employee." "Marketing functions" are not limited to only price terms and conditions of a contract, because non-price terms and conditions of a contract could contain information that an affiliate could use to its advantage. For example, delivery or hub locations in a contract are non-price terms that could be used to favor an affiliate. In addition, negotiated terms and conditions could affect the substantive rights of the parties. For this reason, we decline to make a generic finding to limit "marketing functions" to only price terms and conditions, but will consider waiver requests concerning an employee whose intermittent duties involve drafting non-price terms and conditions.³

Requests for Rehearing and Clarification

3. Several parties have requested expedited clarification regarding paragraph 80 of Order No. 717-A.⁴ Specifically, EEI and Western Utilities request that the Commission clarify that legal, finance, and regulatory personnel can be shared between an entity's transmission and marketing function units.⁵ Similarly, Otter Tail and Central Vermont seek clarification that lawyers, finance, and regulatory personnel may continue to provide support to

¹ *Standards of Conduct for Transmission Providers*, Order No. 717, 73 FR 63796 (Oct. 27, 2008), FERC Stats. & Regs. ¶ 31,280 (2008) ("Order No. 717").

² *Standards of Conduct for Transmission Providers*, Order No. 717-A, 74 FR 54463 (Oct. 22, 2009), FERC Stats. & Regs. ¶ 31,297 (2009) ("Order No. 717-A").

³ Order No. 717-A at P 80.

⁴ Edison Electric Institute (EEI) Oct. 30, 2009 Request for Clarification at 7; The Western Utilities Compliance Group (Western Utilities) Nov. 2, 2009 Request for Clarification at 6; Otter Tail Power Company (Otter Tail) Nov. 10, 2009 Request for Clarification at 1; Central Vermont Public Service Corporation (Central Vermont) Nov. 12, 2009 Request for Clarification at 1.

⁵ EEI at 7; Western Utilities at 6.

marketing function employees, including drafting and redrafting contract non-price terms, without being classified as marketing function employees.⁶

4. EEI also requests clarification that paragraph 80 in Order No. 717-A was “intended to convey that making business decisions about non-price terms and conditions can be a marketing function if the other ‘marketing function’ criteria are met.”⁷

5. If the Commission does not grant these requested clarifications prior to Order No. 717-A taking effect, EEL, Western Utilities, Otter Tail, and Central Vermont request that the Commission change the effective date of paragraph 80 until 90 days after the Commission issues an order addressing the merits of the issue.

Commission Determination

6. The Commission clarifies that the language in paragraph 80 of Order No. 717-A was overly broad. The Commission further clarifies that we intended to state in paragraph 80 of Order No. 717-A that an employee making business decisions about non-price terms and conditions can be considered a “marketing function employee” because that employee is actively and personally engaged in marketing functions. However, an employee who simply drafts or redrafts a contract, including non-price terms and conditions, without making business decisions is not a “marketing function employee.” In making our findings in paragraph 80 in Order No. 717-A, the Commission did not intend to depart from the finding in paragraph 131 in Order No. 717 that employees are not subject to the Independent Functioning Rule if they do not perform transmission functions or marketing functions or to depart from the following examples in P 131:

[I]f an attorney is rendering legal advice, he may consult with both transmission function employees and marketing function employees. Likewise, a risk management employee may develop risk guidelines for both transmission function employees and marketing function employees. And regulatory personnel may present before regulatory bodies filings that cover both transmission and marketing issues. Of course, all such employees would remain subject to the No Conduit Rule, and are prohibited from transmitting transmission function information to marketing function employees.⁸

7. In light of the above clarification to paragraph 80 of Order No. 717-A, we

will deny the petitioners’ request to extend the compliance date with respect to paragraph 80.

III. Document Availability

8. In addition to publishing the full text of this document in the **Federal Register**, the Commission provides all interested persons an opportunity to view and/or print the contents of this document via the Internet through FERC’s Home Page (<http://www.ferc.gov>) and in FERC’s Public Reference Room during normal business hours (8:30 a.m. to 5 p.m. Eastern time) at 888 First Street, NE., Room 2A, Washington, DC 20426.

9. From FERC’s Home Page on the Internet, this information is available on eLibrary. The full text of this document is available on eLibrary in PDF and Microsoft Word format for viewing, printing, and/or downloading. To access this document in eLibrary, type the docket number excluding the last three digits of this document in the docket number field.

10. User assistance is available for eLibrary and the FERC’s Web site during normal business hours from FERC Online Support at 202-502-6652 (toll free at 1-866-208-3676) or e-mail at ferconlinesupport@ferc.gov, or the Public Reference Room at (202) 502-8371, TTY (202) 502-8659. E-mail the Public Reference Room at public.referenceroom@ferc.gov.

IV. Effective Date

11. Changes to Order No. 717-A adopted in this order on rehearing and clarification are effective November 23, 2009.

By the Commission.

Kimberly D. Bose,

Secretary.

[FR Doc. E9-27875 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DELAWARE RIVER BASIN COMMISSION

18 CFR Part 410

Amendments to the Water Code and Comprehensive Plan To Implement a Revised Water Audit Approach To Identify and Control Water Loss

AGENCY: Delaware River Basin Commission.

ACTION: Final rule.

SUMMARY: By Resolution No. 2009-01 on March 11, 2009, the Delaware River Basin Commission (“Commission” or “DRBC”) approved amendments to its Water Code and Comprehensive Plan to

implement an updated water audit approach to identify and control water loss in the Basin.

DATES: *Effective Date:* November 20, 2009. The incorporation by reference of the publications listed in this rule is approved by the Director of the Federal Register as of November 20, 2009.

Applicability Date: Commencing January 1, 2012, the owners of water supply systems serving the public with sources or service areas located in the Delaware River Basin must implement an annual calendar year water audit program conforming to the IWA/AWWA Water Audit Methodology and corresponding AWWA guidance. Commencing January 1, 2013, reported “non-revenue water” must be computed in accordance with the new methodology and guidance.

FOR FURTHER INFORMATION CONTACT: Pamela M. Bush, Commission Secretary and Assistant General Counsel by phoning 609-883-9500 Ext. 203, or by e-mail to Pamela.Bush@drbc.state.nj.us.

SUPPLEMENTARY INFORMATION: The Delaware River Basin Commission (“Commission” or “DRBC”) is a federal-state regional agency charged with managing the water resources of the Delaware River Basin without regard to political boundaries. Its members are the governors of the four basin states—Delaware, New Jersey, New York, and Pennsylvania—and the North Atlantic Division Commander of the U.S. Army Corps of Engineers, representing the federal government.

Notice of the proposed amendments appeared in the **Federal Register** (73 FR 44945) on August 1, 2008 as well as in the Delaware Register of Regulations on September 1, 2008 (12 DE Reg. 275-278 (09/01/2008)), the New Jersey Register (40 N.J.R. 4499) on August 4, 2008, the New York State Register (page 2) on August 20, 2008 and the Pennsylvania Bulletin (38 Pa. B. 4373) on August 9, 2008.

The amendments to the Comprehensive Plan and Article 2 of the Water Code finalized by the Commission on March 11, 2009 phase in a program requiring water purveyors to perform a water audit and report their findings in accordance with a new audit structure established by the American Water Works Association (AWWA) and the International Water Association (IWA). Effective January 1, 2012, the owners of water supply systems serving the public with sources or service areas located in the Delaware River Basin must implement an annual calendar year water audit program conforming to the IWA/AWWA Water Audit Methodology and corresponding

⁶ Otter Tail at 1; Central Vermont at 1.

⁷ EEI at 7-8.

⁸ Order No. 717 at P 131.

AWWA guidance. Commencing January 1, 2013, reported “non-revenue water” must be computed in accordance with the new methodology and guidance. During the period between the effective date of the rule and ending December 31, 2011 (hereinafter, “phase-in period”) water purveyors are encouraged to implement the new methodology and guidance on a voluntary basis.

The Commission has determined that the new water audit methodology provides a rational approach that will facilitate more consistent tracking and reporting than the current approach allows. It will help water managers and regulators, including the Commission, state agencies, and utility managers, target their efforts to improve water supply efficiency, thereby reducing water withdrawals. Improving water accountability will contribute to achieving objective 1.3.C of the Water Resources Plan for the Delaware River Basin (DRBC 2004), which calls for ensuring maximum feasible efficiency of water use across all sectors.

The Commission conducted an informational meeting on the proposed amendments on September 10, 2008 and a public hearing on September 25, 2008, both in West Trenton, New Jersey. Written comment on the proposed amendments was accepted through October 3, 2008. The Commission received one written submission and no oral testimony on the proposed amendment. The agency made revisions to the proposed rule on its own initiative for clarification. A comment and response document summarizing the comments on the proposed rule and setting forth the Commission’s responses and revisions in detail was approved by the Commission simultaneously with adoption of the final rule.

The final form of the rule differs from the proposed rule in the following respects: For purposes of clarity, a definition of “non-revenue water” consistent with the AWWA definition was added to Section 2.1.6.A. of the rule. The definition of “unaccounted-for water” in the same section was amended to include a definition of “unaccounted-for water percent.” This change was made because the computation must return a percentage value so that it can be measured against the performance target of less than 15% unaccounted-for water.

The Commission also added language to establish that until use of the IWA/ AWWA Water Audit methodology becomes mandatory on January 1, 2012, DRBC’s regulatory standards for leak detection and repair (*i.e.*, measurement

and control of unaccounted-for-water), set forth in Section 2.1.6 of the Water Code, shall remain in force. System operators who voluntarily submit audits in a form consistent with the new methodology during the phase-in period are advised in the Commission’s comment and response document that non-revenue water volume expressed as a percentage of input volume will be treated as the equivalent of unaccounted-for-water, the measure applicable under the existing rule. The comment and response document explains that once the Water Audit method is introduced throughout the Delaware Basin and a body of data is available for analysis, a more meaningful measure of system performance will be established.

DRBC Resolution No. 2009–1 and a copy of the comment and response document are both available on the DRBC Web site, <http://www.drbc.net>. Resolution No. 2009–1 incorporates Article 2 of the Water Code, showing the amendments as proposed in August 2008 and as finally approved by the Commission on March 11, 2009. Copies of Resolution No. 2009–1 and the Water Code may be obtained from the Commission’s Secretary and Assistant General Counsel at the telephone number and e-mail address listed above. A charge for printing and mailing may apply.

List of Subjects in 18 CFR Part 410

Incorporation by reference, Water audit, Water pollution control, water reservoirs, Water supply, Watersheds.

■ For the reasons set forth in the preamble, the Delaware River Basin Commission amends part 410 of title 18 of the Code of Federal Regulations as follows:

PART 410—BASIN REGULATIONS; WATER CODE AND ADMINISTRATIVE MANUAL—PART III WATER QUALITY REGULATIONS

■ 1. The authority citation for part 410 continues to read:

Authority: Delaware River Basin Compact, 75 Stat. 688.

■ 2. Revise the first sentence of paragraph (c) of § 410.1 to read as follows:

§ 410.1 Basin regulations—Water Code and Administrative Manual—Part III Water Quality Regulations.

* * * * *

(c) Work, services, activities and facilities affecting the conservation, utilization, control, development or management of water resources within the Delaware River Basin are subject to

regulations contained within the Delaware River Basin Water Code with Amendments Through March 11, 2009, Printed: November 12, 2009, and the Administrative Manual—Part III Water Quality Regulations with Amendments Through July 16, 2008, Printed: September 12, 2008. * * *

* * * * *

Dated: November 12, 2009.

Pamela M. Bush,

Secretary and Assistant General Counsel.

[FR Doc. E9–27645 Filed 11–19–09; 8:45 am]

BILLING CODE 6360–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 520

[Docket No. FDA–2009–N–0665]

Oral Dosage Form New Animal Drugs; Sulfadimethoxine

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule.

SUMMARY: The Food and Drug Administration (FDA) is amending the animal drug regulations to reflect approval of an original abbreviated new animal drug application (ANADA) filed by First Priority, Inc. The ANADA provides for use of Sulfadimethoxine Soluble Powder in medicated drinking water of cattle, chickens, and turkeys for the treatment of various bacterial infections.

DATES: This rule is effective November 20, 2009.

FOR FURTHER INFORMATION CONTACT: John K. Harshman, Center for Veterinary Medicine (HFV–104), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 240–276–8197, e-mail: john.harshman@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: First Priority, Inc., 1590 Todd Farm Dr., Elgin, IL 60123, filed ANADA 200–443 for use of Sulfadimethoxine Soluble Powder in medicated drinking water of cattle, chickens, and turkeys for the treatment of various bacterial infections. First Priority, Inc.’s Sulfadimethoxine Soluble Powder is approved as a generic copy of ALBON (sulfadimethoxine) Soluble Powder, sponsored by Pfizer, Inc., under NADA 46–285. The ANADA is approved as of October 28, 2009, and 21 CFR 520.2220a are amended to reflect the approval.

In accordance with the freedom of information provisions of 21 CFR part

20 and 21 CFR 514.11(e)(2)(ii), a summary of safety and effectiveness data and information submitted to support approval of this application may be seen in the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852, between 9 a.m. and 4 p.m., Monday through Friday.

The agency has determined under 21 CFR 25.33 that this action is of a type that does not individually or cumulatively have a significant effect on the human environment. Therefore, neither an environmental assessment nor an environmental impact statement is required.

This rule does not meet the definition of "rule" in 5 U.S.C. 804(3)(A) because it is a rule of "particular applicability." Therefore, it is not subject to the congressional review requirements in 5 U.S.C. 801-808.

List of Subjects in 21 CFR Part 520

Animal drugs.

■ Therefore, under the Federal Food, Drug, and Cosmetic Act and under authority delegated to the Commissioner of Food and Drugs and redelegated to the Center for Veterinary Medicine, 21 CFR part 520 is amended as follows:

PART 520—ORAL DOSAGE FORM NEW ANIMAL DRUGS

■ 1. The authority citation for 21 CFR part 520 continues to read as follows:

Authority: 21 U.S.C. 360b.

§ 520.2220a [Amended]

■ 2. In § 520.2220a, in paragraph (a)(2), add in numerical sequence "058829".

Dated: November 16, 2009.

Bernadette Dunham,

Director, Center for Veterinary Medicine.

FR Doc. E9-27885 Filed 11-19-09; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF LABOR

Employee Benefits Security Administration

29 CFR 2550

RIN 1210-AB13

Investment Advice—Participants and Beneficiaries

AGENCY: Employee Benefits Security Administration, Labor.

ACTION: Withdrawal of final rule.

SUMMARY: This document withdraws final rules under the Employee

Retirement Income Security Act, and parallel provisions of the Internal Revenue Code of 1986, relating to the provision of investment advice to participants and beneficiaries in individual account plans, such as 401(k) plans, and beneficiaries of individual retirement accounts (and certain similar plans). Final rules were published in the **Federal Register** on January 21, 2009 (74 FR 3822). The effective and applicability dates of the final rules had been deferred until May 17, 2010, in order to permit a review of policy and legal issues raised with respect to the rules. As discussed in this Notice, the Department has determined to withdraw the final rules. The Department also intends to soon propose a revised rule limited to the application of the statutory exemption relating to investment advice.

DATES: Effective January 19, 2010, the final rule published January 21, 2009 amending 29 CFR Part 2550 (74 FR 3822), for which the effective and applicability date was delayed on March 20, 2009 (74 FR 11847), May 22, 2009 (74 FR 23951) and November 17, 2009 (74 FR 59092), is withdrawn.

FOR FURTHER INFORMATION CONTACT: Fred Wong, Office of Regulations and Interpretations, Employee Benefits Security Administration (EBSA), (202) 693-8500. This is not a toll-free number.
SUPPLEMENTARY INFORMATION:

A. Background

On January 21, 2009, the Department of Labor published final rules on the provision of investment advice to participants and beneficiaries of participant-directed individual account plans and to beneficiaries of individual retirement accounts and certain similar plans (IRAs) (74 FR 3822). The rules implement a statutory prohibited transaction exemption under ERISA Section 408(b)(14) and Sec. 408(g), and under section 4975 of the Internal Revenue Code of 1986 (Code),¹ and also contain an administrative class exemption granting additional relief. As published, these rules were to be effective on March 23, 2009. On February 4, 2009, the Department published in the **Federal Register** (74 FR 6007) an invitation for public comment on a proposed 60-day extension for the effective dates of the final rules until May 22, 2009, and a proposed conforming amendment to the applicability date of Section 2550.408g-1, in order to afford the Agency the opportunity to review legal and policy

issues relating to the final rules. The Department also invited public comments on the provisions of those rules and on the merits of rescinding, modifying or retaining the rules. In response to this invitation, the Department received 28 comment letters.² On March 20, 2009, the Department adopted the 60-day extension of the final rule's effective and applicability date. (See 74 FR 11847). In order to afford the Department additional time to consider the issues raised by commenters, the effective and applicability dates were further delayed until November 18, 2009 (74 FR 23951), and then until May 17, 2010.

B. Comments Received

A number of the commenters expressed the view that the final rule raises significant issues of law and policy, and should be withdrawn. Several of these commenters argued that the class exemption contained in the final rule permits financial interests that would cause a fiduciary adviser, and individuals providing investment advice on behalf of a fiduciary adviser, to have conflicts of interest, but does not contain conditions that would adequately mitigate such conflicts. They asserted that investment advice provided under the class exemption therefore might be tainted by the fiduciary adviser's conflicts. Other commenters expressed concerns about those provisions of the rule relating to the "fee-leveling" requirement under the statutory exemption. In particular, some opined that the Department's interpretation of the statutory exemption's fee-leveling requirement is incorrect for permitting the receipt of varying fees by an affiliate of a fiduciary adviser. As a result, they argued, a fiduciary adviser under such a fee-leveling arrangement has a conflict of interest, and the final rule does not adequately protect against investment advice that is influenced by the financial interests of the fiduciary adviser's affiliates. Commenters who advocated retention of the final rule argued that it contains strong safeguards that would protect the interests of plan participants and beneficiaries.

C. Analysis and Determination

As documented in the Department's regulatory impact analysis (RIA) of the January 2009 final regulation and class exemption, defined contribution (DC) plan participants and IRA beneficiaries

¹ These provisions were added to ERISA and the Code by the Pension Protection Act of 2006 (PPA), Public Law 109-280, 120 Stat. 780 (Aug. 17, 2006).

² These comments are available on the Department's Web site at: <http://www.dol.gov/ebsa/regs/cmt-investmentadvicefinalrule.html>.

often make costly investment errors. Those who receive and follow quality investment advice can reduce such errors and thereby reap substantial financial benefit. The Department estimated that the PPA statutory exemption as implemented by the final regulation, together with the final class exemption, would extend investment advice to 21 million previously unadvised participants and beneficiaries, generating \$13 billion in annual financial benefits at a cost of \$5 billion, for a net annual financial benefit of \$8 billion.

In arriving at its estimates, the Department assumed that on average participants and beneficiaries who are advised make investment errors at one-half the rate of those who are not. The Department further assumed that different types of investment advice arrangements on average would be equally effective: Arrangements operating without need for exemptive relief, those operating pursuant to the PPA, and those operating pursuant to the class exemption all would reduce investment errors by one-half on average.

The Department's assumptions regarding the effectiveness of different advice arrangements were subject to uncertainty, particularly as applied to its assessment of the final class exemption's effects. In the preamble to the January 2009 final regulation and class exemption the Department noted evidence that conflicts of interest, such as those that might be attendant to advice arrangements operating pursuant to the class exemption, can sometimes taint advice. Conflicted advisers pursuing their own interests, and the investment managers who compensate them, may profit at the expense of participants and beneficiaries. The conditions attached to the class exemption were intended to ensure that advisers operating pursuant to the class exemption would honor the interests of participants and beneficiaries.

As discussed earlier, a number of commenters raised legal and policy issues concerning the exemption and, in particular, questioned the adequacy of the final class exemption's conditions to mitigate the potential for investment adviser self-dealing. The Department believes that the questions raised in these comments are sufficient to cast doubt on the conditions' adequacy to mitigate advisers' conflicts. If conflicts are not mitigated advice might be tainted. Therefore the Department has set aside its previous assumption that participants and beneficiaries who follow advice delivered pursuant to the final class exemption will commit

investment errors at one-half the rate of those who are unadvised, together with its previous conclusion that the final class exemption's benefits justify its cost. Instead the Department believes that doubts as to whether the final class exemption's conditions are adequate to mitigate conflicts justify withdrawal of the final class exemption. Accordingly, the Department is withdrawing the January 2009 final rule. With regard to the statutory prohibited transaction exemption under ERISA Section 408(b)(14) and Section 408(g), and Code Section 4975, in order to address the absence of regulatory guidance that results from withdrawal of the January 2009 final rule, the Department intends to propose regulations that, upon adoption, implement those provisions. Work is currently being completed on those proposed regulations, and the Department anticipates that they will be published in the **Federal Register** shortly.

For the reasons set forth above, the publication on January 21, 2009 (74 FR 3822), of the final rule amending 29 CFR Part 2550, for which the effective and applicability date was delayed on March 20, 2009 (74 FR 11847), May 22, 2009 (74 FR 23951) and November 17, 2009, is withdrawn.

Signed at Washington, DC, this 16th day of November 2009.

Phyllis C. Borzi,

Assistant Secretary, Employee Benefits Security Administration, Department of Labor.

[FR Doc. E9-27889 Filed 11-19-09; 8:45 am]

BILLING CODE 4510-29-P

DEPARTMENT OF HOMELAND SECURITY

Coast Guard

33 CFR Part 165

[Docket No. USCG-2009-0946]

RIN 1625-AA00

Safety Zones; Blasting and Dredging Operations and Movement of Explosives, Columbia River, Portland to St. Helens, OR

AGENCY: Coast Guard, DHS.

ACTION: Temporary final rule.

SUMMARY: The Coast Guard is establishing two temporary safety zones on the Columbia River to help ensure the safety of the maritime public during blasting and dredging operations taking place near St. Helens, Oregon as well as the movement of explosives for those operations from Portland, Oregon to the

work site. The first temporary safety zone is a fixed zone around the area where the blasting and dredging operations will be taking place near St. Helens, Oregon. The second temporary safety zone is a moving zone around the barge KRS 200-6 at any time that it has explosives onboard.

DATES: This rule is effective from 12:01 a.m. on November 20, 2009 through 11:59 p.m. on February 28, 2010.

The safety zone has been enforced with actual notice since October 30, 2009.

ADDRESSES: Documents indicated in this preamble as being available in the docket are part of docket USCG-2009-0946 and are available online by going to <http://www.regulations.gov>, inserting USCG-2009-0946 in the "Keyword" box, and then clicking "Search." They are also available for inspection or copying at the Docket Management Facility (M-30), U.S. Department of Transportation, West Building Ground Floor, Room W12-140, 1200 New Jersey Avenue, SE., Washington, DC 20590, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

FOR FURTHER INFORMATION CONTACT: If you have questions on this temporary rule, call or e-mail MST1 Jaime Sayers, Waterways Management Division, U.S. Coast Guard Sector Portland; telephone 503-240-9319, e-mail Jaime.A.Sayers@uscg.mil. If you have questions on viewing the docket, call Renee V. Wright, Program Manager, Docket Operations, telephone 202-366-9826.

SUPPLEMENTARY INFORMATION:

Regulatory Information

The Coast Guard is issuing this temporary final rule without prior notice and opportunity to comment pursuant to authority under section 4(a) of the Administrative Procedure Act (APA) (5 U.S.C. 553(b)). This provision authorizes an agency to issue a rule without prior notice and opportunity to comment when the agency for good cause finds that those procedures are "impracticable, unnecessary, or contrary to the public interest." Under 5 U.S.C. 553(b)(B), the Coast Guard finds that good cause exists for not publishing a notice of proposed rulemaking (NPRM) because the publishing of an NPRM would be impracticable and contrary to public interest since immediate action is needed to ensure the public's safety during blasting and dredging operations. Delaying the implementation of the safety zone would subject the public to the hazards associated with blasting and dredging operations and the movement of explosives for those operations. The

danger posed by the large volume of marine traffic on the Columbia River makes safety zone regulations necessary to provide for the safety of construction support vessels, spectator craft and other vessels transiting the event area. For the safety concerns noted, it is in the public interest to have these regulations in effect during blasting and dredging operations. The Coast Guard will issue broadcast notice to mariners to advise vessel operators of navigational restrictions. On-scene Coast Guard and local law enforcement vessels will also provide actual notice to mariners.

Under 5 U.S.C. 553(d)(3), the Coast Guard finds that good cause exists for making this rule effective less than 30 days after publication in the **Federal Register** because to do otherwise would be contrary to the public interest of ensuring public safety during blasting and dredging operations, and immediate action is necessary to prevent possible loss of life and property.

Background and Purpose

As part of the Columbia River Deepening (Channel Improvement) Project, the Army Corps of Engineers must blast and dredge on portions of the Columbia River near St. Helens, Oregon. Due to the inherent dangers associated with blasting and dredging operations, a safety zone is necessary to help ensure the safety of the maritime public operating near the work site. The potential explosive arc for the work site of this project has been calculated to be approximately 832 feet.

The blasting and dredging operations also require the movement of explosives via barge from Portland, Oregon to the work site. Due to the inherent dangers associated with the movement of explosives, a safety zone is necessary to help ensure the safety of the maritime public operating near the barge when explosives are on board.

The project is also required to comply with applicable state laws.

Discussion of Rule

The Coast Guard is establishing two temporary safety zones. The first temporary safety zone applies to the navigable waters within a radius of 1500 feet centering on the Army Corps of Engineers Columbia River Deepening (Channel Improvement) Project work site near St. Helens, Oregon located on the Columbia River from Duck Club Light 6 across to Bachelor Island downstream to the point of Austin Point and across to Warrior Point, at 45°50'31.2" N/122°46'51.6" W; 45°50'31.2" N/122°46'51.6" W; 45°49'37.2" N/122°47'16.79" W;

45°49'47.9" N/122°47'42.00" W; 45°50'56.4" N/122°47'16.79" W (NAD 83). The second temporary safety zone applies to the navigable waters with a radius of 500 feet centering on the barge KRS 200-6 at any time that it has explosives onboard. Notice of the second safety zone will be issued via a Safety Marine Information Broadcast (SMIB) broadcast over Channel 16 and by actual notice on-site. Vessels will be able to transit the work site and/or barge safety zones with permission from the Captain of the Port, Portland or his designated representative. The Captain of the Port can be contacted at telephone number (503) 240-9310, or by radio on VHF Marine Band Radio, channel 16. The safety zones will be in effect from 12:01 a.m. on October 28, 2009 through 11:59 p.m. on February 28, 2010.

Regulatory Analyses

We developed this rule after considering numerous statutes and executive orders related to rulemaking. Below we summarize our analyses based on 13 of these statutes or executive orders.

Regulatory Planning and Review

This rule is not a significant regulatory action under section 3(f) of Executive Order 12866, Regulatory Planning and Review, and does not require an assessment of potential costs and benefits under section 6(a)(3) of that Order. The Office of Management and Budget has not reviewed it under that Order. The Coast Guard has made this determination based primarily on the fact that maritime traffic will be allowed to transit the safety zones with permission from the Captain of the Port, Portland so there should be little to no economic impact.

Small Entities

Under the Regulatory Flexibility Act (5 U.S.C. 601-612), we have considered whether this rule would have a significant economic impact on a substantial number of small entities. The term "small entities" comprises small businesses, not-for-profit organizations that are independently owned and operated and are not dominant in their fields, and governmental jurisdictions with populations of less than 50,000.

The Coast Guard certifies under 5 U.S.C. 605(b) that this rule will not have a significant economic impact on a substantial number of small entities. The following entities may be affected by this rule, some of which may be small entities: The owners and operators of vessels intending to operate, transit, or anchor in a portion of the Columbia

River from 12:01 a.m. on October 28, 2009 through 11:59 p.m. on February 28, 2010. The safety zones will not have a significant impact on a substantial number of small entities for the following reasons. Maritime traffic will be allowed to transit the safety zones with permission from the Captain of the Port, Portland or his designated representative, and the Coast Guard will make notifications via maritime advisories so mariners can adjust their plans accordingly.

Assistance for Small Entities

Under section 213(a) of the Small Business Regulatory Enforcement Fairness Act of 1996 (Pub. L. 104-121), we offer to assist small entities in understanding the rule so that they can better evaluate its effects on them and participate in the rulemaking process.

Small businesses may send comments on the actions of Federal employees who enforce, or otherwise determine compliance with, Federal regulations to the Small Business and Agriculture Regulatory Enforcement Ombudsman and the Regional Small Business Regulatory Fairness Boards. The Ombudsman evaluates these actions annually and rates each agency's responsiveness to small business. If you wish to comment on actions by employees of the Coast Guard, call 1-888-REG-FAIR (1-888-734-3247). The Coast Guard will not retaliate against small entities that question or complain about this rule or any policy or action of the Coast Guard.

Collection of Information

This rule calls for no new collection of information under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501-3520).

Federalism

A rule has implications for federalism under Executive Order 13132, Federalism, if it has a substantial direct effect on State or local governments and would either preempt State law or impose a substantial direct cost of compliance on them. We have analyzed this rule under that Order and have determined that it does not have implications for federalism.

Unfunded Mandates Reform Act

The Unfunded Mandates Reform Act of 1995 (2 U.S.C. 1531-1538) requires Federal agencies to assess the effects of their discretionary regulatory actions. In particular, the Act addresses actions that may result in the expenditure by a State, local, or tribal government, in the aggregate, or by the private sector of \$100,000,000 or more in any one year.

Though this rule will not result in such an expenditure, we do discuss the effects of this rule elsewhere in this preamble.

Taking of Private Property

This rule will not effect a taking of private property or otherwise have taking implications under Executive Order 12630, Governmental Actions and Interference with Constitutionally Protected Property Rights.

Civil Justice Reform

This rule meets applicable standards in sections 3(a) and 3(b)(2) of Executive Order 12988, Civil Justice Reform, to minimize litigation, eliminate ambiguity, and reduce burden.

Protection of Children

We have analyzed this rule under Executive Order 13045, Protection of Children from Environmental Health Risks and Safety Risks. This rule is not an economically significant rule and does not create an environmental risk to health or risk to safety that may disproportionately affect children.

Indian Tribal Governments

This rule does not have tribal implications under Executive Order 13175, Consultation and Coordination with Indian Tribal Governments, because it does not have a substantial direct effect on one or more Indian tribes, on the relationship between the Federal Government and Indian tribes, or on the distribution of power and responsibilities between the Federal Government and Indian tribes.

Energy Effects

We have analyzed this rule under Executive Order 13211, Actions Concerning Regulations That Significantly Affect Energy Supply, Distribution, or Use. We have determined that it is not a "significant energy action" under that order because it is not a "significant regulatory action" under Executive Order 12866 and is not likely to have a significant adverse effect on the supply, distribution, or use of energy. The Administrator of the Office of Information and Regulatory Affairs has not designated it as a significant energy action. Therefore, it does not require a Statement of Energy Effects under Executive Order 13211.

Technical Standards

The National Technology Transfer and Advancement Act (NTTAA) (15 U.S.C. 272 note) directs agencies to use voluntary consensus standards in their regulatory activities unless the agency provides Congress, through the Office of

Management and Budget, with an explanation of why using these standards would be inconsistent with applicable law or otherwise impractical. Voluntary consensus standards are technical standards (e.g., specifications of materials, performance, design, or operation; test methods; sampling procedures; and related management systems practices) that are developed or adopted by voluntary consensus standards bodies.

This rule does not use technical standards. Therefore, we did not consider the use of voluntary consensus standards.

Environment

We have analyzed this rule under Department of Homeland Security Management Directive 023-01 and Commandant Instruction M16475.ID, which guide the Coast Guard in complying with the National Environmental Policy Act of 1969 (NEPA) (42 U.S.C. 4321-4370f), and have concluded this action is one of a category of actions which do not individually or cumulatively have a significant effect on the human environment. This rule is categorically excluded, under figure 2-1, paragraph (34)(g), of the Instruction. This rule involves regulations establishing safety zones. An environmental analysis checklist and a categorical exclusion determination are available in the docket where indicated under

ADDRESSES.

List of Subjects in 33 CFR Part 165

Harbors, Marine safety, Navigation (water), Reporting and recordkeeping requirements, Security measures, Waterways.

■ For the reasons discussed in the preamble, the Coast Guard amends 33 CFR part 165 as follows:

PART 165—REGULATED NAVIGATION AREAS AND LIMITED ACCESS AREAS

■ 1. The authority citation for part 165 continues to read as follows:

Authority: 33 U.S.C. 1226, 1231; 46 U.S.C. Chapter 701, 3306, 3703; 50 U.S.C. 191, 195; 33 CFR 1.05-1, 6.04-1, 6.04-6, 160.5; Public Law 107-295, 116 Stat. 2064; Department of Homeland Security Delegation No. 0170.1.

■ 2. § 165.T13-114 is added to read as follows:

§ 165.T13-114 Safety Zones; Blasting and Dredging Operations and Movement of Explosives, Columbia River, Portland to St. Helens, OR

(a) *Location.* The following areas are safety zones: (1) All waters of the Columbia River from Duck Club Light 6

across to Bachelor Island downstream to the point of Austin Point and across to Warrior Point at 45°50'31.2" N/122°46'51.6" W; 45°50'31.2" N/122°46'51.6" W; 45°49'37.2" N/122°47'16.79" W; 45°49'47.9" N/122°47'42.00" W; 45°50'56.4" N/122°47'16.79" W (NAD 83). (2) All waters encompassed within a circle with a radius of 500 feet centered on the barge KRS 200-6 at any time that it has explosives onboard.

(b) *Definitions.* As used in this section, "designated representative" means a Coast Guard Patrol Commander, including a Coast Guard coxswain, petty officer, or other officer operating a Coast Guard vessel and a Federal, State, and local officer designated by or assisting the Captain of the Port (COTP) Portland in the enforcement of the safety zone.

(c) *Regulations.* In accordance with the general regulations in 33 CFR Part 165, Subpart C, no person may enter or remain in the safety zones established in paragraph (a) or bring, cause to be brought, or allow to remain in the safety zones established in paragraph (a) of this section any vehicle, vessel, or object unless authorized by the Captain of the Port, Portland or his designated representative.

(d) *Enforcement Period.* The safety zones established in paragraph (a) or this section are applicable from 12:01 a.m. on October 28, 2009 through 11:59 p.m. on February 28, 2010.

Dated: October 30, 2009.

F.G. Myer,

Captain, U.S. Coast Guard, Captain of the Port, Portland.

[FR Doc. E9-27725 Filed 11-19-09; 8:45 am]

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DEPARTMENT OF THE INTERIOR

National Park Service

36 CFR Part 7

RIN 1024-AD73

Special Regulations; Areas of the National Park System

AGENCY: National Park Service, Interior.

ACTION: Final rule.

SUMMARY: This rule governs winter visitation and certain recreational use in Yellowstone National Park for the 2009-2010 and 2010-2011 seasons. This final rule is issued to implement the Finding of No Significant Impact (FONSI) for the 2008 Winter Use Plans Environmental Assessment (2008 EA) approved October 15, 2009, and will provide

visitors a range of winter recreation opportunities that are appropriate to the national park setting and do not unacceptably impact or impair park resources or values. The rule requires that most recreational snowmobiles operating in the park meet certain NPS air and sound emissions requirements, requires that snowmobilers and snowcoach riders in Yellowstone be accompanied by a commercial guide, and sets daily entry limits on the numbers of snowmobiles and snowcoaches that may enter the park. Traveling off designated oversnow routes will remain prohibited.

DATES: The effective date for this rule is December 15, 2009.

FOR FURTHER INFORMATION CONTACT: John Sacklin, Management Assistant's Office, Yellowstone National Park, 307-344-2019.

SUPPLEMENTARY INFORMATION:

Background

The National Park Service (NPS) has been managing winter use issues in Yellowstone National Park, Grand Teton

National Park, and the John D. Rockefeller, Jr., Memorial Parkway for several decades under the guidance provided by a number of sources. The history of the issue was discussed at length in the notice for the proposed rule, 73 FR 65784 (November 5, 2008) and in the 2008 EA.

After the proposed rule was published on November 7, 2008, the U.S. District Court for the District of Wyoming issued an order reinstating the 2004 final rule on winter use in the parks, without its sunset provisions, "until such time as NPS can promulgate an acceptable rule to take its place." The NPS complied with the court order and on December 9, 2008, republished the 2004 regulation without its provisions terminating snowmobile and snowcoach use after the winter of 2006-2007. That regulation, among other things, imposed a limit of 720 snowmobiles per day for Yellowstone, required that all recreational snowmobiles in Yellowstone be accompanied by a commercial guide, and required that all recreational snowmobiles operating in

the park meet NPS air and sound emissions requirements for reducing noise and air pollution.

The NPS is promulgating this final regulation to replace the reinstated 2004 regulation. It provides that the park will be open to an appropriate level of oversnow vehicle use for the winter seasons of 2009-2010 and 2010-2011. During this time, NPS will determine a long-term strategy for Yellowstone winter use.

Rationale for the Final Rule

Overview of Winter Use Program

This rule provides for the enjoyment of the park's amenities by authorizing strictly managed snowmobile and snowcoach use in the park for the next two winter seasons. The rule is designed to be consistent with recent trends in oversnow vehicle use while a new long-term winter plan and rule are prepared. This rule allows for 318 snowmobiles per day in Yellowstone, as shown in the following chart, with an additional 50 snowmobiles allowed at Cave Falls.

Park entrance/location	Commercially guided snowmobiles	Commercially guided snowcoaches
(i) North Entrance*	12	13
(ii) West Entrance	160	34
(iii) South Entrance	114	13
(iv) East Entrance	20	2
(v) Old Faithful*	12	16
(vi) Cave Falls	50**	0

* Commercially guided snowmobile tours originating at the North Entrance and Old Faithful are currently provided solely by Xanterra Parks and Resorts. Because this concessioner is the sole provider at both of these areas, this regulation allows reallocation of snowmobiles between the North Entrance and Old Faithful as necessary, so long as the total daily number of snowmobiles originating from the two locations does not exceed 24. For example, the concessioner could operate 6 snowmobiles at Old Faithful and 18 at the North Entrance if visitor demand warranted it. This will allow the concessioner to respond to changing visitor demand for commercially guided snowmobile tours, thus enhancing the availability of visitor services in Yellowstone.

** These snowmobiles operate on an approximately 1-mile segment of road within the park where the use is incidental to other snowmobiling activities in the Caribou-Targhee National Forest. These snowmobiles do not need to be guided or to meet NPS air and sound emissions requirements.

This rule includes strict limits on the number of snowmobiles and snowcoaches allowed to operate within the park each day. Prior to the implementation of a managed winter use program in the winter of 2003-2004, an average of 795 snowmobiles entered Yellowstone each day, with peak days averaging approximately 1,400. This rule allows for 318 snowmobiles per day in Yellowstone, a reduction from the 720 snowmobiles authorized over the previous five winters (during which peak use never approached 720, and average use was about 36% of that limit).

For the past five winters, a managed winter use program has been in place. Visitors on snowmobiles must use snowmobiles that meet NPS

requirements for air and sound emissions (generally referred to in the 2008 EA as Best Available Technology (BAT)), but here referred to simply as NPS requirements to avoid confusion with use of the term best available technology under other environmental laws). Visitors must be accompanied by a commercial guide; visitors cannot snowmobile in Yellowstone without a guide. There is a daily limit on numbers of snowcoaches and snowmobiles. Speed limits are reduced in the busy travel corridors. The park is closed to oversnow vehicles (OSVs) at night. An extensive monitoring program is underway.

In the past five winters, an average of 259 snowmobiles (in an average of 35 commercially guided groups) have

travelled in the park each day, while snowcoach use averaged 31 per day. The peak day for snowmobiles was 557, while the peak day for snowcoaches was 60. During the past three winters, the park exceeded 318 snowmobiles on 63 of 252 days the park was open. This rule allows somewhat more than the recent annual average number of snowmobiles and snowcoaches to enter the park, but would not accommodate those recent higher use days for snowmobiles.

The most recent use levels indicate that the number of commercially guided snowmobile groups and the number of persons in those groups are very similar to those using commercial snowcoaches. In 2008-2009, the average number of snowmobile groups was 31 per day, while snowcoaches averaged 29 per day.

Each snowmobile group included an average of 8.9 people, while each snowcoach carried an average of 8.5 people.

Resource Impacts From Winter Use

Air quality is very good to excellent in the winter, despite frequent temperature inversions, which trap pollutants near the ground and affect air quality. NPS sound and air emission requirements, limits on numbers, and commercial guiding have all contributed to the improvements in air quality over historical (pre-2003) use. Only snowmobiles meeting NPS requirements are allowed. Currently, the snowmobiles use four-cycle engines that produce far less pollution than the two-cycle engines that were once used. Snowmobiles meeting NPS air emission requirements are very similar in their per passenger emissions to snowcoaches. Snowcoaches use more fuel on a per passenger basis than do snowmobiles. They average 2–4 miles per gallon while snowmobiles that meet NPS requirements get 20–26 miles per gallon. In addition, rough roads and soft snow conditions result in higher fuel consumption and high emissions for snowcoaches.

Winter use will have some effects on wildlife, just like every other form of visitor use of the park. Extensive studies of the behavioral responses of five species (bison, elk, bald eagle, trumpeter swan, and coyotes) to oversnow traffic showed that these animals rarely showed high-intensity responses (movement, defense postures, or flight) to approaching vehicles. The responses to normal snowmobile and snowcoach use that do occur do not cause the taking, frightening, or intentional disturbance that is prohibited by NPS regulations. Furthermore, thirty-five years of census data do not reveal any relationship between changing winter use patterns and elk or bison population dynamics. No wildlife populations are currently declining due to winter use (swan populations are declining, but this decline is being experienced regionally and due to factors unrelated to winter use in the park or region). Few animals are expected to be killed as a result of vehicle collisions. The best available information suggests negligible to minor effects for most species, with potential moderate effects for swans and eagles. Use will be well below levels previously studied by NPS wildlife biologists and well within the limits recommended by those studies. We conclude that winter use at the permitted levels does not pose a risk of unacceptable impacts or impairment to any wildlife population. All visitors

utilizing motorized oversnow vehicles travel with commercial guides, learning about and enjoying the abundant wildlife sightings.

Soundscapes are good to very good in the park. Snowmobiles that meet NPS sound requirements are noticeably quieter than traditional snowmobiles (at idle and while underway). In addition, snowmobiles with four-cycle engines that meet NPS requirements sound similar to snowcoaches in the winter and do not sound like traditional two-stroke snowmobiles. Commercial guiding further reduces sound levels and the amount of time that snowmobiles can be heard by reducing speeding and idling and by keeping the vehicles grouped. One concern is that some vehicles are too loud. However, monitoring results demonstrate that 94% of all high sound intensity events are caused by snowcoaches. Overly loud snowcoaches include both older, historic Bombardier snowcoaches that have not been modified or upgraded, as well as a number of modern snowcoaches. The NPS intends to implement sound and air emission requirements for snowcoaches in the long-term plan, subsequent to this rule, to address this concern. The percent of time that OSVs are heard has been a concern. As explained further below, however, NPS has determined that the percentage of time in which OSVs will be audible under this rule does not cause impairment or unacceptable impacts.

Based on a 2008 winter survey, NPS has found that visitors are enjoying the park, and they are satisfied with the management that is in place. Visitors will continue to find wildlife to be both wild and easily viewed. Under this rule, visitors will continue to find wildlife to be both wild and easily viewed. All visitors utilizing motorized vehicles will travel with commercial guides, learning about and enjoying the abundant wildlife sightings. A winter 2008 survey found a high level of satisfaction with soundscape conditions, wildlife, and the managed winter use program.

Personal exposure of employees to air pollutants has generally been greatly reduced from historic levels. Some monitoring from previous years indicated small exceedances of national standards for benzene and formaldehyde. The source could be snowcoaches or snowmobiles, or more likely both. Last winter's monitoring showed no exceedances of these standards.

Impairment, Unacceptable Impacts, and Appropriate Use

In addition to determining the environmental consequences of the alternatives, NPS policy requires consideration of impacts to determine whether actions would impair park resources. In managing National Park System units, the NPS may undertake actions that have both beneficial and adverse impacts on park resources and values. As the 2006 NPS Management Policies (Management Policies) explain (section 1.4.7.1), "Virtually every form of human activity that takes place within a park has some degree of effect on park resources or values, but that does not mean the impact is unacceptable or that the particular use must be disallowed." The NPS is generally prohibited by law from taking or authorizing any action that would or is likely to impair park resources or values. Impairment is an impact that, in the professional judgment of the responsible NPS manager, would harm the integrity of park resources or values, including the opportunities that otherwise would be present for the enjoyment of those resources or values. The responsible NPS manager generally has discretion to determine what impacts are allowed that would not impair park resources or values.

The NPS is also required to conserve the resources and values of the National Park System units and to prioritize the conservation of park resources over their use whenever the two are found to be in conflict. The NPS complies with this mandate by ensuring that a proposed use of the park will not result in unacceptable impacts to park resources or values, and by further allowing impacts to park resources only when allowing the impacts is appropriate to fulfill the purposes of the park and is necessary (meaning that the impacts are unavoidable and incapable of further mitigation in light of the authorized appropriate use).

Over the last five winter seasons, the park was intensively managed in order to provide heightened protection to the environment and prevent the impairment of park resources and values. As discussed in the FONSI and based on the analysis in the 2008 EA and monitoring and studies over the past five years, the NPS has determined that no impairment of park resources or values occurred during those five years.

The NPS has also determined that implementation of Alternative 2 (Selected Alternative) and the final rule would not result in unacceptable impacts or impairment to park resources or values. As disclosed in the 2008 EA,

the adverse impacts to wildlife would be negligible to minor, due to moderate levels of visitor use (with possible moderate effects on swans and eagles). Guiding would minimize most of these effects. For soundscapes, the adverse impacts would be negligible to moderate, due to audibility and maximum sound levels. Exceedances of maximum sound levels by snowcoaches will be mitigated while this rule is in place through driver education and reducing snowcoach travel speed. This will be communicated during pre-season meetings with commercial guides and outfitters, and to individual drivers during park-sponsored orientation training. Air quality impacts are forecast to be negligible because the air and sound emissions requirements and strict daily entry limits will reduce emissions. Impacts on visitor and employee health and safety in Yellowstone are expected to be moderately adverse due to possible high snowcoach noise exposure levels. Avalanche danger at Sylvan Pass also creates moderate adverse impacts. Both the noise exposure issues and the avalanche danger would be mitigated in several ways.

As described in the 2008 EA, the NPS's threshold for considering whether there could be an impairment is based on *major* (or significant) effects. The 2008 EA identified less than major effects on wildlife, natural soundscapes, and air quality for Alternative 2. Indeed, while some major effects have resulted from snowmobile or snowcoach use over the past five years—which included some days where snowmobile usage was nearly double the daily limit now adopted—the NPS has determined that none of the effects associated with that usage caused any impairment of park resources. Guided by this analysis and the professional judgment of National Park Service managers, the NPS has determined that there would be no impairment of park resources or values from implementation of the final rule.

Finally, the NPS has determined that the impacts associated with the OSV use permitted over the next two winter seasons, which are described at length in the 2008 EA, are both appropriate and necessary to fulfill the purposes of the park.

Section 1.5 of Management Policies, "Appropriate Use of the Parks," directs that the National Park Service must ensure that park uses that are allowed would not cause impairment of, or unacceptable impacts on, park resources or values. A new form of park use may be allowed within a park only after a determination has been made in the

professional judgment of the park manager that it will not result in unacceptable impacts. In addition, section 8.1.2 of the Management Policies, "Process for Determining Appropriate Uses," directs the NPS to evaluate the proposed use's consistency with applicable laws, executive orders, regulations, and policies; consistency with existing plans for public use and resource management; actual and potential effects on park resources or values; total costs to the NPS; and whether the public interest will be served. Finally, section 1.5 of the Management Policies directs park superintendents to continually monitor all park uses to prevent unanticipated and unacceptable impacts. If unanticipated and unacceptable impacts occur, section 1.5 directs the superintendent to engage in a thoughtful deliberative process to further manage or constrain the use, or discontinue it.

Environmental Assessment and Finding of No Significant Impact

The 2008 EA and the 2009 FONSI supporting this final rule contain the above-described evaluation of the permitted OSV use. In addition, they demonstrate that no unacceptable impacts are anticipated as a result of the use. Finally, the Preferred Alternative in the 2008 EA establishes a comprehensive monitoring and adaptive management plan to address any unanticipated unacceptable impacts. On this basis, the NPS has determined that the proposed OSV use permitted over the next two winter seasons is appropriate to fulfill the purposes of the park.

The NPS has also determined that the proposed OSV use permitted over the next two winter seasons is necessary to fulfill the purposes of the park. The National Park Service Organic Act directs the NPS to promote the use of the national parks by such means and measures as to conform to the fundamental purpose of said parks, which purpose includes providing for the enjoyment of the scenery, natural and historic objects, and wildlife within the parks (16 U.S.C. 1). Section 8.2 of Management Policies confirms that enjoyment of park resources and values by the people of the United States is one of the fundamental purposes of all parks. That section further states: "To provide for enjoyment of the parks, the National Park Service will encourage visitor use activities that are appropriate to the purpose for which the park was established, and are inspirational, educational, or healthful, and otherwise appropriate to the park environment; and will foster an understanding of and

appreciation for park resources and values, or will promote enjoyment through a direct association with, interaction with, or relation to park resources; and can be sustained without causing unacceptable impacts to park resources or values."

As explained in the 2008 EA, OSV use of Yellowstone National Park has been occurring since 1949, and snowmobiles have been used for 48 of the park's 137 years. Yellowstone is a large park, distances between attractions at Yellowstone are great, and some form of motorized vehicular access is needed to access various destination areas. Snowmobiles and snowcoaches are used for this purpose in the winter just as private vehicles and buses are used in the summer. They are both forms of transportation, not recreational activities unto themselves. Finally, snowmobiles and snowcoaches each provide very different experiences in that they provide varying levels of direct interaction with the park's resources and values.

The NPS received approximately 27,500 comments on the 2008 EA and 39,767 comments on the proposed rule. In many cases, the comments received on the proposed rule were very similar in content to those received on the 2008 EA. Numerous commenters expressed concerns that the Preferred Alternative and the rule, would violate the NPS Organic Act and would be inconsistent with the 2006 NPS Management Policies, among other things causing unacceptable impacts to park resources and values. The NPS believes most of these concerns are based on a belief that snowmobiles do not belong in the park, and should be replaced with snowcoaches. These concerns do not take into account recent monitoring and studies that show the nearly equal contribution of snowmobiles and snowcoaches to the concerns expressed by the commenters (and that snowcoaches are clearly the source of some concerns). Statistically, movement responses of wildlife were slightly higher for snowcoaches than for snowmobiles. Monitoring also indicates that commercially guided snowmobile groups and snowcoaches contribute similarly to the amount of time OSVs are heard. Snowcoaches also use more fuel on a per passenger basis than do snowmobiles. In short, neither OSV type provides a clear advantage with respect to environmental impacts. Recent monitoring and studies demonstrate that the regulated use of both snowcoaches and snowmobiles described in the Selected Alternative will not result in impairment of park resources or values,

nor will it result in unacceptable impacts on the park.

Air and Sound Emission Requirements

To mitigate impacts to air quality and the natural soundscape, the NPS is continuing the requirement that all recreational snowmobiles meet strict air and sound emissions requirements to operate in the park, with limited exceptions. For air emissions, all snowmobiles must achieve a 90% reduction in hydrocarbons and a 70% reduction in carbon monoxide, relative to EPA's baseline emissions assumptions for conventional two-stroke snowmobiles. For sound emissions, snowmobiles must operate at or below 73 dBA as measured at full throttle according to Society of Automotive Engineers (SAE) J192 test procedures (revised 1985). The Superintendent will maintain a list of approved snowmobile makes, models, and years of manufacture that meet NPS requirements. The certification is good for six years from the date on which a model is certified as meeting the requirements.

The NPS is continuing the requirement that began with the 2005 model year that all snowmobiles must be certified under 40 CFR part 1051 to a Family Emission Limit (FEL) no greater than 15 g/kW-hr for hydrocarbons (HC) and 120 g/kW-hr for carbon monoxide (CO). Snowmobiles must be tested on a five-mode engine dynamometer consistent with the test procedures specified by the EPA (40 CFR parts 1051 and 1065). Other test methods could be approved by the NPS.

The NPS is retaining the use of the FEL method for demonstrating compliance with its emissions requirements because it has several advantages. First, use of FEL will ensure that all individual snowmobiles entering the park achieve the NPS's emissions requirements, unless modified or damaged (under this regulation, snowmobiles which are modified in such a way as to increase air or sound emissions will not be in compliance with NPS requirements and therefore not permitted to enter the park). Use of FEL will also minimize any administrative burden on snowmobile manufacturers to demonstrate compliance with NPS requirements because they already provide FEL data to the EPA. Further, the EPA has the authority to ensure that manufacturers' emissions claims on their FEL applications are valid. EPA also requires that manufacturers conduct production line testing (PLT) to demonstrate that machines being manufactured actually meet the

certification levels. If PLT indicates that emissions exceed the FEL levels, then the manufacturer is required to take corrective action. Through EPA's ability to audit manufacturers' emissions claims, the NPS will have sufficient assurance that emissions information and documentation will be reviewed and enforced by the EPA. FEL also takes into account other factors, such as the deterioration rate of snowmobiles (some snowmobiles may produce more emissions as they age), lab-to-lab variability, test-to-test variability, and production line variance. In addition, under the EPA's regulations, all snowmobiles manufactured must be labeled with FEL air emissions information. This labeling will help to ensure that NPS emissions requirements are consistent with these labels. The use of FEL will avoid potential confusion for consumers.

The air emissions requirements for snowmobiles allowed to operate in the park should not be confused with standards adopted by the EPA in a final rule published in the **Federal Register** on November 8, 2002 (67 FR 68242). The EPA regulations require manufacturers to meet certain fleet averages for HC and CO emissions. For example, the Phase 1 standards required all snowmobile manufacturers to meet a fleet-wide average in 2007 of 275 g/kW-hr for CO and 100 g/kW-hr for HC, which represents a 30% reduction from the baseline emission rates for uncontrolled snowmobiles. Any particular make/model may emit more or less than the standard as long as the fleet average does not exceed the standard. Phase 2 and Phase 3 standards will be implemented in 2010 and 2012, respectively, effectively requiring the equivalent of a 50% reduction in both HC and CO as compared to average baseline levels. By comparison, NPS requires that all snowmobiles operating in the park meet a FEL of 120 g/kW-hr for CO and 15 g/kW-hr for HC. This means that snowmobiles operating in the park represent the cleanest that are commercially available.

To determine compliance with the sound emissions requirements, snowmobiles must be tested using SAE J192 test procedures (revised 1985; or potentially as further revised and adapted for use by NPS). The NPS recognizes that the SAE updated these test procedures in 2003; however, the changes between the 2003 and 1985 test procedures could yield different measurement results. The sound emissions requirement was initially established using 1985 test procedures (in addition to information provided by industry and modeling). To ensure

consistency in the test results, the NPS will at this time continue to use the 1985 test. The SAE J192 (revised 1985) test also allows for a tolerance of 2 dBA over the sound limit to account for variations in weather, snow conditions, and other factors. The NPS understands that an update to the 2003 J192 procedures may be underway, and the NPS will continue to evaluate these test procedures and possibly adopt them after these regulations are implemented. Other test methods could be approved by NPS on a case-by-case basis.

Snowmobiles may be tested at any barometric pressure equal to or above 23.4 inches Hg uncorrected (as measured at or near the test site). This exception to the SAE J192 test procedures maintains consistency with the testing conditions used to determine the sound requirement. This allowance for reduced barometric pressure is necessary since snowmobiles were tested at the elevation of Yellowstone National Park, where atmospheric pressure is lower than that under the SAE J192's requirements. Testing data indicate that snowmobiles test quieter at higher elevation, and therefore some snowmobiles may comply with the NPS's sound emissions requirements at higher elevations even though they do not when tests are conducted near sea level.

The NPS will annually publish a list of snowmobile makes, models, and years of manufacture that meet its emissions and sound requirements. Snowmobile manufacturers may demonstrate that snowmobiles are compliant with the air emissions requirements by submitting to the NPS a copy of their applications used to demonstrate compliance with EPA's general snowmobile regulation (indicating FEL). The NPS will accept this application information from manufacturers in support of conditionally certifying a snowmobile as meeting its air emissions requirements, pending ultimate review and certification by EPA at the same emissions levels identified in the application. Should EPA certify a snowmobile at an emission level that would no longer meet the NPS's requirements, this snowmobile would no longer be considered by NPS to be compliant with its requirements and would be phased-out according to a schedule that will be determined by the NPS to be appropriate. For sound emissions, snowmobile manufacturers may submit their existing Snowmobile Safety and Certification Committee (SSCC) sound level certification form. Under the SSCC machine safety standards program, snowmobiles are

certified by an independent testing company as complying with all SSCC safety standards, including sound standards. This regulation does not require the SSCC form specifically, as there could be other acceptable documentation in the future. The NPS will work cooperatively with the snowmobile manufacturers on appropriate documentation. The NPS intends to continue to rely on certified air and sound emissions data from the private sector rather than establish its own independent testing program. When the NPS certifies snowmobiles as meeting its requirements, NPS will announce how long that certification applies. Generally, each snowmobile model will be approved for entry into the park for six winter seasons after it is first listed. Based on NPS experience, six years represents the typical useful life of a snowmobile, and thus six years provides purchasers with a reasonable length of time where operation is allowed once a particular model is listed as being compliant. If a manufacturer recertifies a snowmobile model to NPS requirements for emissions and sound, it could be used for additional years. It is also based on EPA snowmobile emission regulations and the deterioration factors that are part of those regulations (EPA requires that if a manufacturer certifies its snowmobile will comply with EPA's emission regulations, the snowmobile will meet those regulations for a period of five years or 5,000 miles).

Individual snowmobiles modified in such a way as to increase sound and air emissions of hydrocarbons and carbon monoxide beyond the emission restrictions will be denied entry to the park. It is the responsibility of end users and guides and outfitters to ensure that their OSVs, whether snowmobiles or snowcoaches, comply with all applicable restrictions. Air and sound emission requirements for snowcoaches are described below. In Yellowstone, the requirement that all snowmobilers travel with commercial guides will assist NPS in enforcing these requirements, since businesses providing commercial guiding services in the park are responsible under their contracts with the park to ensure that their clients use only snowmobiles that meet the NPS's requirements. In addition, these businesses are required to ensure that snowmobiles used in the park are not modified in such a way as to increase sound or air emissions, and that snowmobiles are properly maintained.

Snowmobiles being operated on the Cave Falls Road, which extends approximately one mile into

Yellowstone from the adjacent national forest, will be exempt from air and sound emissions requirements. Because of the low level of impacts resulting from the light use of the Cave Falls Road, which is incidental to recreational use of the surrounding national forest, NPS has found it is not necessary to require these users to comply with requirements that address issues associated with use of the interior portions of the park.

Under concession contracts issued in 2003, 78 snowcoaches are currently authorized to operate in Yellowstone (and in the parkway between Flag Ranch and Yellowstone's South Entrance). Approximately 29 of these snowcoaches were manufactured by Bombardier and were designed specifically for oversnow travel. Those 29 snowcoaches were manufactured before 1983 and are referred to as "historic snowcoaches" for the purpose of this rulemaking. All other snowcoaches being used are passenger vans or light buses that have been converted for oversnow travel using tracks and/or skis. During the winter of 2008–2009, an average of 29 snowcoaches entered Yellowstone each day (during the prior winter, 2007–2008, an average of 35 snowcoaches entered the park each day).

As of the winter of 2009–2010, all snowcoaches must be commercially guided. These trained, knowledgeable operators help ensure that air and sound emission requirements are met, wildlife impacts are minimized, and visitor and employee safety is assured.

The University of Denver conducted winter emissions measurements in Yellowstone that involved the collection of emissions data from in-use snowcoaches and snowmobiles in February 2005 and February 2006. Results from that work indicate that snowcoaches and snowmobiles meeting NPS air emission requirements are now very similar in their per passenger emissions. This work also supports snowmobile air emissions requirements and the development of snowcoach air emission requirements. The snowcoach fleet should be modernized to reduce carbon monoxide and hydrocarbon emissions. However, road and snow conditions and low power-to-weight ratios of snowcoaches contribute considerably to air emissions. This means that even an upgraded snowcoach fleet operating in Yellowstone will have days for which fuel consumption and emission levels might be high.

In comparison with older carbureted snowcoaches, snowcoaches operating within EPA's Tier I standards are

cleaner. In 2004, EPA began phasing-in Tier II emissions standards for multi-passenger vans, and they will be fully phased-in during 2009. Tier II standards will require that vehicles be even cleaner than Tier I, and full emission controls will function more of the time.

During the duration of this temporary plan, all non-historic snowcoaches must meet air emission requirements, which will be the EPA emissions standards in effect when the vehicle was manufactured. This will be enforced by ensuring that all critical emission-related exhaust components are functioning properly. Malfunctioning critical emissions-related components must be replaced with the original equipment manufacturer (OEM) component where possible. If OEM parts are not available, aftermarket parts may be used. In general, catalysts that have exceeded their useful life must be replaced unless the operator can demonstrate the catalyst is functioning properly. Modifying or disabling a snowcoach's original pollution control equipment is prohibited except for maintenance purposes. Individual snowcoaches may be subject to periodic inspections to determine compliance with emission and sound requirements.

The restrictions on air and sound emissions in this rule are not a restriction on what manufacturers may produce but an end-use restriction on which commercially produced snowmobiles and snowcoaches may be used in the park. The NPS Organic Act (16 U.S.C. 1) authorizes the Secretary of the Interior to "promote and regulate" the use of national parks "by such means and measures as conform to the fundamental purpose of said parks * * * which purpose is to conserve the scenery and the natural and historic objects and the wild life therein and to provide for the enjoyment of the same in such manner and by such means as will leave them unimpaired for the enjoyment of future generations." Further, the Secretary is expressly authorized by 16 U.S.C. 3 to "make and publish such rules and regulations as he may deem necessary or proper for the use and management of the parks. * * *

" This exercise of the NPS Organic Act authority is not an effort by NPS to regulate manufacturers and is consistent with Section 310 of the Clean Air Act.

Since 2001, the park has been converting its own administrative fleet of snowmobiles to meet these NPS requirements. These newer machines have proven successful in fulfilling most of the NPS's administrative needs throughout the park. However, the NPS recognizes that some administrative applications, such as off-trail boundary

patrols in deep powder, towing heavy equipment or disabled sleds, search and rescue, or law enforcement uses, may require additional power beyond that supplied by currently available snowmobiles that meet the NPS's air and sound emissions requirements. In such limited cases, the NPS will sometimes need to use snowmobiles that do not meet the requirements this rule imposes upon recreational snowmobiles (which do not have these special needs because they travel only upon groomed roads as part of a tour group led by a commercial guide).

Guided Tours and Group Size

In order to mitigate impacts to natural soundscapes and wildlife, and for visitor and employee safety, all recreational snowmobiles and snowcoaches operated in Yellowstone must be led by a commercial guide, except for those snowmobiles being operated on the one-mile segment of the Cave Falls Road that extends into the park from the adjacent national forest. This guiding requirement has been found in practice to reduce conflicts with wildlife along roadways because these commercial guides are trained to lead visitors safely around the park with minimal disturbance to wildlife. Commercial guides are educated in safety, knowledgeable about park rules, and are required to exercise reasonable control over their clientele, which has reduced unsafe and illegal snowmobile use. Because of the contractual obligations to which commercial guides are subject, NPS has found this results in more effective enforcement of park rules. These guides receive rigorous multi-day training. They also are experts at interpreting the resources of the park to their clients. Commercial guides are employed by local businesses, not by NPS. Commercial guiding also tends to result in larger snowmobile parties than unguided use, which reduces the overall number of encounters with wildlife and reduces the amount of time that OSVs are audible (and, conversely, increases the interval of time that OSVs are not heard).

No more than eleven snowmobiles will be permitted in a group, including that of the guide. Except in emergency situations, guided parties must travel together and remain within a maximum distance of one-third mile of the first snowmobile in the group. These size and distance limits require that guided parties do not become separated, provide for sufficient and safe spacing between individual snowmobiles within the guided party, and allow the guide(s) to maintain control over the group to minimize the impacts on wildlife and

natural soundscapes. NPS thus expects that the continuation of the guiding requirement will facilitate compliance with park regulations and protect park resources.

Commercial snowmobile guides use a "follow-the-leader" approach, stopping often to talk with the group. They lead snowmobiles single-file through the park, using hand signals to pass information down the line from one snowmobile to the next, a system which has proven to be effective. Signals are used to warn group members about wildlife and other road hazards, indicate turns, reduce speed, and when to turn on or off the snowmobile. Further, all commercial guides are trained in basic first aid and CPR. In addition to first aid kits, they often carry satellite or cellular telephones, radios, and other equipment for emergency use. Guides are thus well-equipped to ensure that park regulations are enforced, wildlife are protected, and to provide a safer overall experience for visitors.

Since the winter of 2003–2004, all snowmobilers in Yellowstone have been led by commercial guides, resulting in considerable positive effects on visitor health and safety. Guides have been proven to be very effective at enforcing proper touring behavior, such as adherence to speed limits, staying on the groomed road surfaces, and other snowmobiling behaviors that are appropriate to safely and responsibly visit the park. Since implementation of the guiding program, there have been pronounced reductions in the number of law enforcement incidents and accidents associated with the use of snowmobiles, even when accounting for the reduced number of snowmobilers relative to historic use levels. The use of guides is also beneficial to wildlife, since guides are trained to respond appropriately when encountering wildlife.

Snowmobile and Snowcoach Routes

Snowmobiles and snowcoaches will continue to be restricted to designated oversnow routes, which are a subset of the same roads that are traveled by motor vehicles during the remainder of the year. In addition to most of the Grand Loop Road, certain side roads will be open for snowmobile use after noon, based on the successful experience of the NPS with temporal zoning on Firehole Canyon Drive. Virginia Cascades will be accessible only via ski and snowshoe.

The final rule also allows for up to 50 snowmobiles to enter Yellowstone on the Cave Falls Road, an approximately one-mile segment extending into the southwest corner of the park from the

Targhee National Forest. This short road segment does not connect to the rest of the oversnow routes in Yellowstone, and connects only to the national forest lands, which do not have air and sound requirements or guiding requirements. Use of this route is incidental to recreational use of the national forest lands, is far removed from the snowmobile use and the resulting impacts that occur within the interior of Yellowstone, and is therefore considered separately from the 318 snowmobile limit.

Snowmobile and snowcoach use in the two-mile road segment between Yellowstone's South Entrance and Flag Ranch in the John D. Rockefeller, Jr. Memorial Parkway will be governed by Yellowstone requirements (as is also discussed in the separate rule for the Parkway). That is all snowmobiles operating on this road segment must meet the commercial guiding, NPS air and sound requirements, daily use limits, and other requirements to operate in Yellowstone. Similarly, all snowcoaches operating on this road segment must meet Yellowstone requirements.

Monitoring and Adaptive Management

Scientific studies and monitoring of winter visitor use and park resources (including air quality, natural soundscapes, wildlife, employee health and safety, water quality, and visitor experience) will continue. As part of its adaptive management of winter use activities, NPS will close selected areas of the park to visitor use, including sections of roads, if these studies indicate that human presence or activities have unacceptable impacts on wildlife or other park resources that cannot otherwise be mitigated. A one-year notice will ordinarily be provided before any such closure is implemented unless immediate closure is deemed necessary to avoid impairment of park resources. The Superintendent will continue to have the authority under various provisions of this rule as well as 36 CFR 1.5 to take emergency actions to protect park resources and values.

The adaptive management program described in the 2008 EA provides park managers with a wide variety of tools to ensure that the goals and objectives of the winter use plans are being achieved. Some of the techniques available include adjustments in snowmobile or snowcoach use levels (up or down), adjustments in air and sound emissions requirements, visitor and guide education, timing of entries, and group sizes.

Adjustment to the daily entry limits for snowmobiles and snowcoaches is

one of several tools available to park managers to ensure that the goals and objectives of the winter use plan are maintained. Through adaptive management, if monitoring of use levels of snowmobiles and snowcoaches allowed under the FONSI indicates acceptable conditions, the NPS will increase use levels to the extent acceptable conditions can be maintained. Conversely, if monitoring of use levels of snowmobiles and snowcoaches allowed under the FONSI indicates unacceptable conditions, the NPS will reduce use levels to an extent that acceptable conditions can be maintained. In some cases, additional rulemaking would be required in order to adjust numbers.

The NPS is implementing a multi-year research proposal intended to specifically address the question of whether grooming of the Madison to Norris road segment in Yellowstone has led to alterations of bison movements and distribution. The question was identified in a report by Dr. Cormack Gates et al., entitled "The Ecology of Bison Movements and Distribution in and Beyond Yellowstone National Park" (2005). The research program will involve a linked series of experiments that will enable researchers to gain insight into how road grooming and other factors currently affect bison travel. The NPS has begun deploying cameras along travel routes to gain information on the relationship between road grooming and bison travel. The research program will include the analysis of existing data on GPS-collared bison, the tracking of additional GPS-collared bison, and use of the cameras, without necessitating the closure of the Gibbon Canyon road segment to public OSV travel. During the five year period, other roads or routes may be investigated to help understand the relationship between snow depth, grooming, and bison movement. For example, the Firehole Canyon Drive may be closed to oversnow travel and the Grand Loop Road gated to allow snowmobile and snowcoach travel, but not allow bison movement on the main road. Bison would then be forced to travel cross-country or along the ungroomed Firehole Canyon Road. Similarly, the Madison to Norris Road may be fenced or gated in the vicinity of the new bridge over the Gibbon River to restrict bison movement on the Madison to Norris Road and force bison to travel cross-country. Thus, bison movement in relation to snow depth may be tested without closing a main road. However, following the five years of data

gathering and analysis, the NPS, in consultation with the researchers, will consider closing the main Madison to Norris route to observe bison response. That decision will rely on the results of the data gathering and analysis and whether such a closure would be likely to yield informative data or conclusions. If implemented, such a closure would likely last several seasons.

Maintaining Entry by Sylvan Pass

Sylvan Pass will be open for oversnow travel (both motorized and non-motorized) for a limited core season, from December 22 through March 1 each year, subject to weather-related constraints and NPS capacities. A combination of avalanche mitigation techniques may be used, including risk assessment analyses as well as forecasting and helicopter- and howitzer-dispensed explosives. The NPS will continue to evaluate additional avalanche mitigation techniques and risk assessment tools in order to further improve safety and visitor access.

From March 2 to March 15, the NPS will maintain the road segment from the East Entrance to a point approximately four miles west of the entrance station to provide for opportunities for cross-country skiing and snowshoeing. Limited snowmobile and snowcoach use will be allowed in order to provide drop-offs for such purposes.

This approach both addresses the concerns of the communities and the National Park Service. The City of Cody, Wyoming, as well as Park County, Wyoming, and the State of Wyoming have clearly articulated the importance of this route to the community and the historical relationship between Cody and Yellowstone's East Entrance. They have spoken for the businesses near Yellowstone's East Entrance and how those businesses have been negatively impacted in recent years by the changing patterns of winter visitation and uncertainty regarding winter use in the park. They have stated how those businesses will continue to be adversely affected if the pass is closed to OSV travel in the winter. The community and businesses have also stated the value they place on the certainty of the road being open in the winter and the importance of that certainty to their businesses and guests. NPS acknowledges those values and concerns and has carefully weighed those considerations.

Avalanche control at Sylvan Pass has long represented a safety concern to the National Park Service. The 2000 Final Environmental Impact Statement (FEIS), the 2003 Supplemental Environmental

Impact Statement, the 2004 EA, and the 2007 FEIS all clearly identify the considerable avalanche danger on Sylvan Pass, which has been well known for many years. Approximately 20 avalanche paths cross the road at Sylvan Pass. They average over 600 feet of vertical drop, and the East Entrance Road crosses the middle of several of the paths, putting travelers at risk of being caught in an avalanche. NPS employees must cross several uncontrolled avalanche paths to reach the howitzer used for discharging those avalanches, and the howitzer is at the base of a cliff prone to both rock-fall and additional avalanche activity (the howitzer cannot be moved without compromising its ability to reach all avalanche zones). Artillery shells sometimes fail to explode on impact, and unexploded rounds remain on the slopes, presenting year-round hazards to both employees and visitors, both in Yellowstone and the Shoshone National Forest. Natural avalanches can and do occur, both before and after howitzer use. Using a helicopter instead of a howitzer also is a high-risk activity because of other risks, such as high winds, a helicopter contractor would have to incur.

The NPS may use a combination of techniques that have been used in the past (howitzer and helicopter), as well as techniques that may be available in the future. Area staff may use whichever tool is the safest and most appropriate for a given situation, with the full understanding that safety of employees and visitors comes first. Employees in the field make the operational determination when safety criteria have been met, and operations can be conducted with acceptable levels of risk. The NPS will not take unacceptable risks. When safety criteria have been met, the pass will be open; when they have not been met, the pass will remain closed. As with past winters, extended closures of the pass may occur, and the NPS will continue to provide notices of the road status.

Summary of and Responses to Public Comments

The NPS published a proposed rule on November 5, 2008 and accepted public comments through November 20, 2008. The NPS reopened the comment period on July 24, 2009 and accepted public comments through September 8, 2009. Comments were accepted through the mail, hand delivery, and through the Federal eRulemaking Portal: <http://www.regulations.gov>. A total of 39,767 comment documents were received.

1. *Comment:* The numbers of snowmobiles and snowcoaches that

should be permitted into the park should be set at numbers higher or lower than those proposed by the plan.

Response: A limit of 318 will produce an average considerably lower than those seen in recent years. With a limit of 720 over the last 5 years, snowmobile use did not average more than 300 per day. On most days, use was much lower than 300 (in January/February 2007, the average, for example, was 273), but the average was closer to 300 as a result of the higher numbers seen around Christmas 2006 and other peak days, when use rose as high as 543 per day. A limit of 318 will greatly reduce those peaks and thereby is expected to lower the overall average. For various reasons, it is not expected that the 318 daily limit will be reached during the next two winters. It will likely be difficult for all guides and outfitters to fill their allocations: different sizes of groups will probably create one or two unused snowmobiles per allocation, and last minute cancellations will probably leave some allocations unused. Also, using last winter as an example, one guide company had only 10 snowmobiles available to use, out of an allocation of 30. Thus, every day, 20 snowmobile allocations went unused. Finally, unless recent use patterns illustrated in the 2008 EA shift greatly, the 318 limit will not be reached every day or even often enough to produce an average more than 300. Also, as explained in the 2008 EA, NPS cannot allow higher numbers of snowmobile or snowcoaches to enter the park until the NPS analyzes their effects in an EIS, because higher numbers of snowmobiles and snowcoaches have the potential to create major adverse impacts. Additionally, at this time, NPS has not conducted sufficient analysis to determine whether higher numbers would cause unacceptable impacts or would otherwise be an appropriate use. In a long-term plan and EIS, alternatives with higher numbers of snowmobiles would be considered.

2. *Comment:* The method in which snowmobile limits should be set should be based on seasonal variations, adaptive management, annual maximums, high demand times (holidays), and/or concession contracts, as is the case for snowcoaches.

Response: As reflected in the analyses within the judicially vacated 2007 EIS providing for variable daily limits would have the potential to create major adverse impacts on park soundscapes, particularly on days when visitation exceeded 318 snowmobiles and 78 snowcoaches. Such impacts would have to be first analyzed in an EIS. Weekends are not necessarily the

busiest days; allowing higher visitation on weekends could deprive visitors the ability to enter on weekdays. Annual limits would provide variable daily limits as well and may result in major impacts. Such an alternative must be first analyzed in an EIS, and could be analyzed in the long-term plan and EIS. The decision includes an adaptive management program.

3. *Comment:* The NPS should phase out or ban snowmobiles, and transition to a snowcoach-only system.

Response: Current science suggests that a snowcoach-only system in Yellowstone could cause a number of impacts: major soundscape impacts, high fuel consumption, greater wildlife responses, and more damage to the snow road surface than from snowmobiles. At this time NPS has not conducted sufficient analysis to determine whether such a system would cause unacceptable impacts or would otherwise be an appropriate use. In a long-term plan and EIS, such a system would be considered.

4. *Comment:* The NPS should consider alternatives beyond the use of snowmobiles or snowcoaches, including plowing more roads in the winter to allow for vehicle use.

Response: As explained on 2008 EA pp. 2–8 to 2–9, plowing was dismissed as an alternative in this EA because doing so would add uncertainty and because many winter operators had already invested in oversnow equipment, assuming a plan similar to this one would indeed be implemented. The plowing option remains a possibility to consider in long-term winter use planning.

5. *Comment:* The current system of commercial guides should be modified to include non-commercial guides certifying individuals to lead groups, or the elimination of the requirement for a guide all together.

Response: The concept of non-commercial guiding or unguided access (both with training programs) has been analyzed in previous winter plans and will be evaluated in a long-term winter plan. Additionally, the NPS may consider the Certified Group Leader concept in its future long-term winter use planning. The NPS will consider non-commercial guides in long-term winter use planning. The interim plan will last for two winters, which is not sufficient time to design and implement pilot or test programs and study and report on their effects.

6. *Comment:* Snowmobile numbers should be regulated through variations in when and where snowmobiles can access the park, such as “snowmobile

only” days and/or limiting snowmobile use to certain areas of the park.

Response: Alternating kinds of visitation by week or day would be logistically difficult to implement and would not provide the consistency needed for effective trip planning for visitors in a short-term plan. In a long-term plan, the alternatives will consider a variety of spatial or temporal zoning as the comment suggests.

The requirement to use commercial guides has the effect of grouping all snowmobilers and many snowcoaches into certain time windows. Generally, these are two hour windows in the mornings and afternoons at the entrances and midday at Old Faithful. Outside of those periods commercial use is greatly reduced, and the opportunity to walk or ski in silence is more readily available. The NPS wishes to protect park soundscapes at all times of the winter, not just these less busy time periods. While visitors are certainly free to visit at less busy times to seek natural quiet, the NPS believes they also should be able to find it at other times. The NPS believes that adoption of the rule would offer ample opportunities for quiet.

7. *Comment:* The NPS should consider alternative elements that focus on non-motorized uses such as promoting cross country skiing, and snow shoeing.

Response: NPS will continue to facilitate non-motorized recreation and set ski tracks on the edges of snow roads. Snowshoers and cross-country skiers also have impacts on wildlife. The best available science indicates that cross-country skiers are more likely than snowmobiles to elicit a startle or flight response in wildlife as a result of their less regular use patterns and quiet approach to animals. Yellowstone is a large park, and it is 30 miles from West Yellowstone to Old Faithful and 50 miles from Mammoth Hot Springs to Old Faithful. Most visitors cannot ski or snowshoe these distances. For most visitors to enjoy locations in Yellowstone such as Old Faithful or the Grand Canyon of the Yellowstone, motorized access is necessary. Ski and snowshoe opportunities are available throughout the park, and many people access trailheads via snowmobile or snowcoach.

8. *Comment:* Only certain types of snowmobiles and/or snowcoaches with special technology should be allowed in the park.

Response: Electric snowmobiles could be used in Yellowstone under this winter use plan if they meet all other requirements. NPS is not aware of their commercial availability. Four-stroke

snowmobiles have been operated by concessioners within the park for the past six years. There are currently air and sound requirements for snowmobiles, and future requirements for snowcoaches are expected.

Snowmobiles that meet NPS air and sound requirements have considerably cleaner emissions and are quieter than snowmobiles that do not meet NPS requirements. The NPS continues to encourage snowmobiles (and snowcoaches) to employ improved technologies. NPS will continue to move towards air and sound requirements for snowcoaches, and snowcoaches will be required to adhere to noise and air emissions requirements, similar to those of snowmobiles.

9. *Comment:* The park should consider additional actions such as increasing law enforcement activities, lowering speed limits, stopping accommodation of winter use, prohibiting tours and allowing trips to set destinations only, and expansion of educational programs regarding winter use opportunities at Yellowstone.

Response: NPS will continue enforcement of its regulations. While an adjustment to speed limits may be analyzed further in the long-term winter use planning effort, a much lower speed limit would not allow access to Yellowstone's widely-spaced attractions. The NPS believes providing motorized oversnow access to the features of Yellowstone for the next two winter seasons helps fulfill the mission of the park to provide for visitor use and enjoyment of those resources. The current commercial guiding program provides an excellent way for the public to learn about the park and appropriate behavior. In the long-term plan, the NPS will evaluate alternatives that look at education programs for unguided or non-commercial guided opportunities.

10. *Comment:* The interim plan should be modified to include different timeframes for how long it would be in effect and different seasonal entry points.

Response: NPS believes the 2-year duration of the plan is necessary to provide adequate time to develop a new long-term winter use plan. In a long-term plan, the alternatives will consider a variety of spatial or temporal zoning as the comment suggests.

11. *Comment:* Winter use management should include either high fees for snowmobile use or subsidized snowcoach use.

Response: NPS will consider the fee suggestion in future long-term winter use planning.

12. *Comment:* NPS should create a lottery, permit, or reservation system to

limit winter use access, including a safety test or other educational component to assist the park in enforcement. Allocations among guides and outfitters should be fair and equal.

Response: Through the use of commercial guides, a reservation system is in place so that visitors can plan ahead for access to the park. Other allocation systems and education opportunities will be evaluated in the long-term winter use planning. The commercial guiding program has substantially assisted the park in improving compliance with park regulations.

13. *Comment:* Areas outside the park should be designated for snowmobile use, the park should be periodically shut down to allow for regeneration of the ecosystem, and snowmobiles should be required to stay on certain tracks if use is allowed in the Park.

Response: Whether areas outside the Park are also available for snowmobiling is not within the scope of this decision-making process. Snowmobiles in Yellowstone have always been restricted to park roads and have never been permitted off-road. The sheer size of Yellowstone means that more than one road is necessary to provide adequate visitor access. The No Action Alternative considered in the 2008 EA have closed the park and therefore better protected air quality. However, that alternative would have seriously limited access to much of the park for those not capable of skiing or snowshoeing long distances. Snowmobiles as well as snowcoaches offer visitors the opportunity to enjoy Yellowstone. With the requirement to use only snowmobiles that meet NPS air and sound requirements and are accompanied by a commercial guide, snowmobiles serve as a form of access to the features of Yellowstone, not a separate recreational activity.

14. *Comment:* NPS should require that winter users maintain 100 meter animal distance when stopping.

Response: The NPS requires visitors stay at least 100 yards (91 m) away from bears and wolves and at least 25 yards (23 m) away from all other animals—including bison, elk, bighorn sheep, deer, moose, and coyotes.

15. *Comment:* Snowmobiles should only be allowed for use by rangers, the disabled, or for emergency operations.

Response: Administrative use of snowmobiles is also managed by the NPS winter use plan, and as explained above, most NPS snowmobiles now meet NPS air and sound requirements. Similarly, researchers must also use snowmobiles that meet NPS air and sound requirements. Snowmobiles that

do not meet NPS air and sound requirements are used administratively only where necessary for the performance of park duties (for example, in deeper snow associated with boundary patrol).

Snowmobiles provide a different type of interaction with the park's attractions than do snowcoaches. Providing some level of access via both snowmobiles and snowcoaches provides for different kinds of enjoyment of the park's scenery and natural and historic objects and wildlife.

16. *Comment:* The interim plan should not use adaptive management to address existing park violations of NPS mandates.

Response: This rule does not authorize violations of any NPS mandates. NPS will continue enforcement of its regulations under any scenario, and the NPS will use adaptive management and monitoring results to make adjustments to the plan's implementation.

17. *Comment:* The 2004 rule should be retained, and the NPS should reaffirm its commitment to keeping Sylvan Pass open.

Response: Due to a pending appeal and other litigation related to reinstatement of the 2004 rule, relying on the reinstated 2004 rule would create substantial uncertainty regarding winter access, and NPS does not believe it is a viable option. In addition, there has been no current NEPA analysis or other determination that use at the levels authorized under that regulation is consistent with the NPS's statutory and other mandates. The findings of the 2007 EIS, as well as the court order vacating it, both suggest that those use levels are probably not consistent with those requirements. In order to help assure winter access to Yellowstone, the NPS is completing planning and rulemaking to replace the 2004 regulation reinstated by the Wyoming Court. A separate decision has been made, and separate regulations will be published, for Grand Teton National Park and the John D. Rockefeller, Jr. Memorial Parkway.

This decision continues the implementation of the Sylvan Pass Agreement (subject to weather-related constraints and NPS fiscal, staff, infrastructural, equipment, and other safety-related capacities) during this interim plan. Management of the Pass will continue to be evaluated in a long-term plan.

18. *Comment:* The NPS air and sound requirements should be eliminated so that individuals can drive their snowmobiles on park roads.

Response: The NPS continues to require snowmobiles (and encourage snowcoaches) to employ improved technologies. Eliminating the air and sound requirements could lead to a return of historical conditions, which were found in 2000 to constitute impairment of park resources. Even if such use could be authorized, it would at a minimum have to be analyzed in an EIS. This comment will be considered in the course of the long-term planning process.

19. *Comment:* The 2008 EA selected an incorrect “no-action”, as it did not represent the current level of activity.

Response: NPS disagrees. When the 2008 EA was prepared, the 2007 rule had been vacated. No snowmobile or snowcoach use would have been authorized without action by the NPS, because the authorizations in the 2004 rule had expired pursuant to the sunset date provisions. After the 2008 EA was issued, the U.S. District Court for the District of Wyoming reinstated the 2004 rule without the sunset clauses, and as a result, up to 720 snowmobiles per day were allowed for the winter of 2008–09. Due to a pending appeal, there is still uncertainty regarding that reinstatement. As explained above, there has been no current NEPA analysis or other determination that use at the levels authorized under that regulation is consistent with the NPS’s statutory and other mandates. Accordingly, the No Action Alternative analyzed in the 2008 EA represents a more logical and useful benchmark against which impacts can be compared, and therefore continues to better satisfy the purposes of the no action alternative under NEPA.

20. *Comment:* The snowcoach-only alternative was improperly dismissed.

Response: A snowcoach-only transportation system would have numerous impacts and might not be the least impacting form of transportation. While NPS agrees that preservation of resources is key to the fundamental mandate of Yellowstone and the entire National Park System, the suggestion that the Yellowstone National Park enabling statute and the NPS Organic Act mandate snowcoach use is incorrect. These acts direct the agency to protect park resources and provide for enjoyment without incurring impairment. If NPS is to provide for any sizeable visitor access to Yellowstone in the winter, motorized vehicle use is necessary, and NPS believes that a limit of 318 snowmobiles per day and 78 snowcoaches per day effectively allows the agency to protect its resources while providing for visitation during this two-winter period.

21. *Comment:* The NPS has received a larger percentage of comments from the past planning efforts supporting a transition from snowmobiles to snowcoaches.

Response: The NPS has reviewed all comments received throughout the past and present winter use planning efforts in compliance with the NEPA and other relevant laws and regulations. The NPS is mandated to consider all of these comments in order to provide the decision-maker with a fully informed environmental analysis to base their decision on. NPS cannot base its decision simply on the sheer numbers of comments in support or against snowmobile, snowcoach, or solely non-motorized winter use. Snowcoach use has slowly and steadily increased. Somewhat more visitors still prefer to visit Yellowstone via snowmobiles. Snowcoaches do facilitate conversations between guides and visitors, but the guiding requirement for snowmobiles also has a similar result. If visitors double up on snowmobiles, the cost is comparable to snowcoach tickets for multiple individuals. Snowmobiles and snowcoaches both cause similar soundscape, wildlife and air quality impacts. Snowcoaches may consume more fuel per capita than do the snowmobiles that meet NPS air and sound requirements for use in Yellowstone. As the FONSI indicates, it is no longer clear that snowcoaches are the “least impacting” oversnow vehicles.

22. *Comment:* The Park should work with surrounding communities to educate the public regarding responsible and appropriate behavior within Yellowstone National Park.

Response: The current commercial guiding program provides an excellent way for the public to learn about the park and appropriate behavior. In the long-term plan, the NPS will evaluate alternatives that look at education programs for unguided or non-commercial guided opportunities.

23. *Comment:* The NPS should provide the public and use a transparent and candid interpretation of the findings related to snowmobile impacts on park resources.

Response: The NPS has used the most current information available in preparing the 2008 EA and this decision. That information has led to a new and better understanding of the contribution of both snowmobiles and snowcoaches to impacts on park resources.

24. *Comment:* The proposed rule and impact analysis violates the NPS’s Organic Act of 1916, findings within the 2008 EA, the court ruling of the U.S.

District Court for Wyoming, other previous decisions on this issue, and other provided court precedents.

Response: As a result of the Wyoming District Court’s order, the reinstated 2004 rule was in effect for the winter of 2008–2009. This interim rule would be in effect for two winter seasons. NPS believes the two-year duration of the plan is necessary to provide adequate time to develop a new long-term winter use plan. NPS believes the rule is consistent with all applicable court decisions concerning prior winter use plans, and other applicable authorities.

25. *Comment:* The methodologies of the analyses were flawed because it did not compare the impacts of snowcoaches versus snowmobiles adequately, consider the historical precedent of snowmobile use, and used existing concessioner contracts as the basis for use numbers.

Response: The computations in the 2008 EA were based on actual field measurements in Yellowstone, not on hypothetical modeling or estimates. Given the average passenger load on snowmobiles and snowcoaches in Yellowstone and the actual fuel economies of these vehicles, snowcoaches consume more fuel per passenger than snowmobiles. As indicated by the August 2008 peer-reviewed paper, “Portable Emission Measurements of Yellowstone National Park Snowcoaches and Snowmobiles” by Gary A. Bishop, Ryan Stadtmuller, Donald H. Stedman, and John D. Ray in the Journal of the Air and Waste Management Association (59:936–942), snowcoaches and snowmobiles are very similar in the per-passenger emissions. The soundscape modeling in the 2007 EIS (which was not challenged on this issue) indicated that a snowcoach-only alternative would cause major adverse effects to soundscapes. More recent monitoring information indicates snowcoaches are audible for similar time periods as commercially guided snowmobile groups. Also work on snowcoach sound indicates that the loud coaches include some modern vehicles, as well as those historic coaches that have not been retrofitted.

26. *Comment:* The false studies like the two-stroke emission test (where they used a very old, very out of tune two-stroke engine and compared the results against a brand new fuel efficient car) are a criminal use of taxpayer money.

Response: Current snowmobile emission information was based on modern snowmobiles that meet NPS air and sound requirements. Two-stroke snowmobile air emissions information used standard EPA emission factors.

27. *Comment:* The economic baseline analysis used in the 2008 EA should be 540 snowmobiles per day, as opposed to zero.

Response: As discussed above, the No Action Alternative analyzed in the 2008 EA represents the most logical and useful benchmark against which impacts can be compared, and therefore continues to best satisfy the purposes of the no action alternative under NEPA.

As discussed below, the economic analysis in this rule used a different baseline, based on the reinstated 2004 rule and its limit of 720 snowmobiles per day.

28. *Comment:* The NPS methodology for determining a comment period was improper and does not need to relate to the winter use season.

Response: Little time was available to complete the 2008 EA, so the public comment period on the EA in 2008 was quite limited. The NPS regrets any difficulties entering comments into its Web-based public comment system, but notes that comments sent by regular mail were also accepted. The NPS also provided an additional 45-day comment period on the proposed rule and took into account all comments received on the rule and 2008 EA. Thus a full 60-day comment period was provided on the proposed action.

29. *Comment:* NPS Management Policies prohibit the impairment of park resources and values, and snowmobile use constitutes an impairment.

Response: No impairment to park resources was found for the Selected Alternative.

30. *Comment:* No limit should be established for snowmobile access until impairment of park resources has been identified and proven. The standard of how impairment is applied to soundscapes is too strict.

Response: The Organic Act charges NPS with providing for enjoyment of the national parks “by such means as will leave them unimpaired.” However, nothing in the Organic Act suggests that impairment is the only consideration that may justify imposing limitations on use. The Organic Act clearly authorizes appropriate limitations on use as needed to protect park resources and values. Recreational uses may be prohibited if they are not an appropriate use, which does not necessarily mean that they cause impairment. NPS also manages uses so as to minimize conflicts among them. The NPS Management Policies explain when recreational and other uses may be prohibited or restricted. The natural soundscape is one of the “park resources and values” that NPS is

required to conserve and protect from impairment under the NPS Organic Act.

31. *Comment:* A potential precedent may be set that would restrict un-guided automobile use inside the park during the summer.

Response: This is a winter use plan not a summer plan. Issues and concerns are different in the winter than in the summer, and this plan does not set a precedent for summer visitation.

32. *Comment:* Unacceptable impacts to park resources were not adequately addressed in the 2008 EA—more action is needed to prevent the unacceptable impacts caused by snowmobile use within the park.

Response: The NPS finds that the negligible to moderate impacts of the Selected Alternative described in the 2008 EA and FONSI do not meet the criteria described in the FONSI for either unacceptable impacts or impairment, and are therefore consistent with the NPS’s statutory requirements under the Organic Act.

33. *Comment:* Snowmobiles that meet NPS air and sound requirements are not impacting the air quality within the park and give off fewer emissions.

Response: All snowmobiles allowed into the parks (with certain minor exceptions) must meet NPS air and sound requirements. These are the cleanest snowmobiles on the market. Impacts on air quality were analyzed and discussed in the EA and FONSI.

34. *Comment:* Air quality is adversely affected by the use of snowmobiles in the park, primarily due to exhaust, and that it is the duty of the NPS to prevent adverse impacts to air quality.

Response: Alternative 1 considered in the 2008 EA would close the park to visitor oversnow vehicle use and therefore fully protect air quality. However, Alternative 1 would deny access to much of the park for those not capable of skiing or snowshoeing. The Selected Alternative would allow only snowmobiles that meet NPS air and sound requirements into the park. Recent use levels have been similar to or higher than the levels expected under the Selected Alternative, and air quality has been very good to excellent in the park. It is therefore expected to remain very good to excellent.

35. *Comment:* Snowmobiles and snowcoaches have the same impact on air quality.

Response: Snowcoach use has been carefully analyzed in the winter use plan, particularly since their impacts upon park soundscapes, wildlife, and air quality are at times greater than those of snowmobiles. As indicated by the August 2008 peer-reviewed paper, “Portable Emission Measurements of

Yellowstone National Park Snowcoaches and Snowmobiles” by Gary A. Bishop, Ryan Stadtmuller, Donald H. Stedman, and John D. Ray in the Journal of the Air and Waste Management Association (59:936–942), snowcoaches and snowmobiles are very similar in the per-passenger emissions. Snowcoaches also use more fuel than snowmobiles, even accounting for the different passenger loads.

36. *Comment:* While the NPS claims to have independent “authority and jurisdiction to administer some provisions of the Clean Air Act” in the 2008 EA, the State of Wyoming has primacy under the Clean Air Act; therefore, the NPS has no authority to rely on air quality standards to limit snowmobile access in the park.

Response: NPS agrees the States of Wyoming, Montana and Idaho play a primary role in implementation of the Clean Air Act as it affects the park. However, as the Federal Land Manager, the NPS also has responsibilities to protect air quality and air quality-related values in the park. The Clean Air Act is not the sole applicable authority. As explained above, this is an exercise of the NPS Organic Act authority over use within the park, not an effort by NPS to regulate manufacturers, and it is consistent with Section 310 of the Clean Air Act. Air quality is expected to remain very good to excellent under the rule. The Organic Act reserves ample discretion to the Park Service to determine how best to provide for enjoyment of the Park. Thus, NPS has exclusive responsibility to determine the appropriate level and type of public access into national parks; indeed, many other national parks are closed entirely to motorized access in the winter.

37. *Comment:* The analysis of air quality was flawed, since air quality monitoring was not conducted along road corridors and the range of impacts from pollution was not fully accounted for in the analysis. The analysis of air quality impacts was improper since the NPS has not properly explained how an action would have “major” impacts on air quality within the park.

Response: The 2008 EA used new impact threshold definitions in order to address exactly the sorts of issues raised by this comment. The definitions for this EA were intentionally adjusted downward to be more conservative—that is, more protective—of park resources. The definitions are not based on parkwide metrics; rather, they are based on actual monitoring data, which are gathered at the two places where oversnow vehicle use is highest, Old Faithful and West Yellowstone. The NPS used the National Ambient Air

Quality Standards (NAAQS) in assessing air quality impacts because they provide an objective standard established by the EPA in order to protect air quality and protect public health.

38. *Comment:* The compaction of snow is a benefit of snowmobile use, as it prevents erosion.

Response: Snowmobile and snowcoach use under this rule is confined to a portion of the existing road system. The area of compacted snow comprises a negligible portion of the park acreage and has a negligible effect on overall snowmelt, runoff patterns, and erosion.

39. *Comment:* National parks are for the entire public, not just for environmentalists or special interest groups.

Response: National parks are open to the general public. Winter use management is intended to address specific issues while providing opportunities for all visitors to enjoy the parks consistent with NPS legal mandates and policies.

40. *Comment:* Studies have shown that black carbon emissions have adverse effects on the snowpack and should be analyzed before a rule is enacted.

Response: Monitoring of pollution deposition in the snowpack has been underway for more than 10 years, and this concern has not been identified in Yellowstone. As indicated in the 2008 EA, this monitoring will continue.

41. *Comment:* Many snowmobile operators drive too fast in the park

Response: All snowmobiles are to be commercially guided, which generally has eliminated speeding and other past problems. This is demonstrated, among other things, by the reduction in citations for such violations.

42. *Comment:* Banning or limiting all automobiles within the park should be explored, since snowmobiles are not the only motorized type of vehicle that creates impacts.

Response: Regarding automobiles in the summer, this is not a summer use plan, but rather a winter use plan, so such decision-making is beyond the scope of the rule. In the winter, the majority of the park has long been closed to automobiles, with the roads groomed for oversnow vehicle use. Plowing the roads for automobile use will likely be analyzed in the long-term winter use plan.

43. *Comment:* Snowmobile use adversely affects human health and safety because of air pollution, snowmobile accidents and crashes, and improper snowmobile operation.

Response: Concerning health and safety, results of the most recent personal exposure monitoring from winter 2008–2009 shows no exceedances of standards. With the requirement for commercial guiding, law enforcement incidents related to snowmobile use have dropped dramatically in the past five years, as compared to the 1990s, thus indicating fewer accidents and violations.

44. *Comment:* The analysis of health and safety is flawed because NPS must utilize health and safety metrics that have reasoned basis in relevant health standards for determining major health and safety impacts resulting from snowmobile use.

Response: NPS safety managers use OSHA and NIOSH metrics for measuring exposure of employees to sound and air pollution, which are standard measures used by safety professionals in determining hazards.

45. *Comment:* Snowmobile operators use caution and are polite to other users; I did not see any blue haze.

Response: NPS monitoring has shown dramatic improvements in winter conditions relative to historical use.

46. *Comment:* The cost of continuing snowmobile use at the park, conducting studies on this matter, and maintaining the East Entrance Road would be too much for the amount of snowmobilers that currently access the park. Furthermore, keeping Sylvan Pass open is too dangerous for park staff.

Response: Winter operations in Yellowstone are expensive for snowmobile or snowcoach access. The interim plan continues to implement the Sylvan Pass Agreement reached with the City of Cody, Park County, Wyoming, and the State of Wyoming. Sylvan Pass will be open only when safety criteria have been met.

47. *Comment:* The Park's assertion that the snowcoach-only alternative would have hazardous effects on oversnow travel is erroneous.

Response: If travel were restricted to snowcoaches only, a consequent increase in such traffic would result assuming visitation levels remain anywhere near current levels. This increase could compound the problems already seen in the park with rutting and damage to snow roads from coaches. That is why the NPS is implementing size and weight restrictions on coaches.

48. *Comment:* The Park informed commercially guided snowmobile businesses that 14 snowmobiles a day would be allowed per concessioner, yet the number now being proposed has been decreased to nine per day.

Response: NPS recognizes that some visitors will not be able to take snowmobiles into Yellowstone. However, most visitors will be able to take a snowcoach instead. Some visitors may have to adjust their plans and visit the park on different days.

49. *Comment:* The Park needs the revenue from snowmobiling activities, so entrance fees would have to be increased as a result of banning snowmobiles from entering the Park. Otherwise, the entrance fees should be increased in order to increase law enforcement patrols.

Response: Decisions regarding the appropriate type of winter use and numbers of snowmobiles and snowcoaches are made without regard to entrance fee revenues. Entrance fees related to winter use are a small part of Yellowstone's overall budget and a small part of the fee revenue that Yellowstone receives. Winter use accounts for 100,000 of the approximately 3.2 million people that visit Yellowstone each year.

50. *Comment:* Law enforcement efforts would not necessarily be decreased with the commercial snowmobile guide requirement, as is stated in the 2008 EA. Snowmobile use within the park requires increased law enforcement, since many snowmobile operators do not abide by the rules and regulations of the park.

Response: The NPS has reviewed the methodology used to calculate law enforcement incidents and believes they correctly show a decrease with the implementation of the managed use program, including commercial guiding. With the managed use program, the NPS believes that many of the incidents observed in the past (for example, snowmobilers speeding or going off road) rarely occur today.

51. *Comment:* The potential banning or limitations placed on snowmobile access to the park would create adverse impacts to surrounding businesses, tourists, as well as the NPS, since snowmobile outfitters and businesses that benefit from tourism would have to increase the cost of snowmobile tours for tourists.

Response: The 2008 EA and rulemaking analyzed socioeconomic impacts using IMPLAN modeling. Though this model does not incorporate every potential factor in the socioeconomic setting, it allows an objective analysis structure that may be applied to the entire planning area and cumulative impact study area. With respect to the number of snowmobile and snowcoach entries permitted under the Selected Alternative and resulting impacts on operators and visitors, the

permitted entries (318 snowmobiles and 78 snowcoaches) represent an 8.2% increase in snowmobiles and a 123% increase in snowcoaches compared to the 2007–2008 average of 294 snowmobiles and 35 snowcoaches per day. The percentage increases represented by the Selected Alternative are even larger compared to the 2008–2009 average of 205 snowmobiles and 29 snowcoaches per day. While the 2008–2009 use likely reflects visitor uncertainty brought on by recent court decisions, NPS does not think that use levels will increase considerably over the next two years that the Selected Alternative will be in effect. This is because of the current economic slowdown and because NPS does not expect a considerable increase in use over such a short period of time.

52. *Comment:* The economic interests that currently depend on snowmobiling could switch to business ventures related to snowcoaches and the NPS needs to consider the value of the natural surroundings in their analysis, since the park does not exist to provide profit for businesses located outside the park. They may switch to business ventures related to cross country skiing and snowshoeing.

Response: Gateway communities provide services to park visitors that the NPS cannot provide or has chosen not to provide. Through the planning process, the NPS determines appropriate type of winter use and numbers of snowmobiles and snowcoaches. Through the concessions contract process, the NPS then determines the nature of the business opportunities available and provides potential concessioners the opportunity to submit bid to provide those services. Businesses may then compete to provide those services in the park. The NPS recognizes that each type of use and access (snowmobile, snowcoach, ski, snowshoe) creates impacts and the impacts must be weighed with regard to the protection of park resources while providing for visitor enjoyment.

53. *Comment:* Snowmobile use inside the park creates undesirable impacts to soundscapes within the park, disrupts the quiet serenity the park offers in the absence of snowmobiles, and may very well be inconsistent with desirable conditions.

Response: Even with sound from cumulative effects of all oversnow vehicles, NPS expects soundscapes impacts to stay within moderate levels, levels that would be fully acceptable and would be consistent with its desired conditions and with the 2006 Management Policies. NPS agrees that winter serenity is important and

believes that the level of use permitted by the Selected Alternative (by snowmobiles that meet NPS air and sound requirements, combined with snowcoach use) will result in large portions of the day without the sound of oversnow vehicles.

54. *Comment:* NPS should explain the adaptive management thresholds (primarily soundscape thresholds), consistency with other NPS mandates, obligation to conserve park resources and leave them unimpaired throughout the entire park, legal basis for considering soundscapes as a park resource, what an unacceptable impact is, and baseline in gauging the impacts on snowmobile use on soundscapes.

Response: The adaptive management thresholds are a management tool only; they do not represent the unacceptable impacts or impairment thresholds described in section 1.4 of the Management Policies. Rather, they are a conservative measure used to alert the NPS manager that additional attention to a particular park resource or value is merited. By reacting to the exceedance of a conservative adaptive management threshold, NPS can seek to ensure that no unacceptable impacts or impairment occur. Accordingly, the fact that these thresholds have been exceeded in the past in no way undermines NPS's determination that "sound from recreational oversnow vehicles [is] well within acceptable ranges."

In backcountry areas and travel corridors, the OSV impacts were essentially compared against natural ambient. That is, the natural ambient was the existing ambient (minus the low percentage of aircraft sounds). In the Old Faithful developed area, the natural ambient was not measurable due to other existing non-natural sounds (the heating and ventilating systems in buildings adjacent to the monitoring site are continuously audible).

The 2008 EA contains an explanation of the relationship between major impacts, unacceptable impacts, and impairment. NPS notes that the term "major" as used in the 2008 EA is equated with "significant" effects within the meaning of NEPA. Accordingly, if a major impact were predicted, the NPS would prepare an EIS.

For soundscapes, one of the "clear bright lines" separating acceptable impacts from unacceptable impacts is whether implementation of an alternative would unreasonably interfere with the natural soundscape, be inconsistent with Yellowstone's purposes or values, impede the attainment of Yellowstone's desired future conditions, create an unsafe or

unhealthy environment, or diminish opportunities for current or future generations.

NPS understands that this "line" does not establish a "quantitative" standard as the commenter requests. However, the intensity of many impacts, and the manner in which those impacts translate into impairment or unacceptable impacts, cannot be described quantitatively. In such instances, they must rely on qualitative standards which are based on the NPS manager's best professional judgment.

The soundscape impact threshold definitions in the 2008 EA make clear that recreational oversnow vehicle noise is a subject of this EA and rulemaking; however, overflights and administrative vehicles are clearly identified as contributing to the cumulative soundscapes impacts, with appropriate mitigations also identified.

55. *Comment:* Newer snowmobiles, specifically ones that meet NPS air and sound requirements, do not create noise pollution—a majority of the impacts to soundscapes within the park emanate from NPS contractors.

Response: Recent monitoring indicates that commercially guided snowmobile groups and snowcoaches contribute similarly to the audibility of oversnow vehicles. Early in the managed winter use program, some contractors were using snowmobiles that did not meet NPS requirements. Newer contracts are correcting this problem, and the NPS continues to move towards a requirement that NPS and concession employees only use snowmobiles that meet NPS air and sound requirements.

56. *Comment:* The soundscapes impacts presented in the 2008 EA could be mitigated through further management of snowmobiles and snowcoaches by the NPS.

Response: The NPS has only recently understood that modern snowcoaches are also significant contributors to the concerns regarding loud oversnow vehicles, and the NPS is still working on methodologies and test procedures for sound testing of snowcoaches. The lack of a stable, long-term plan has slowed implementation of snowcoach sound and air emission requirements. An individual snowcoach represents a significant investment, and snowcoaches are operating under 10-year contracts that were awarded in 2003. Therefore the NPS believes the long-term planning process should establish the test procedures and specifics of snowcoach sound and air emission requirements.

57. *Comment:* Experiences on a snowmobile could not be replaced with

a snowcoach, such as the feeling of openness, experience of the scenery, experience of the ability to access public lands.

Response: NPS recognizes that snowmobiles and snowcoaches offer different types of experiences for visitors.

58. *Comment:* Snowmobile use has a negative impact on visitor experience from the noise, exhaust, and wildlife disturbance.

Response: A visitor survey in 2008 specifically addressed soundscapes and wildlife and found a high level of visitor satisfaction.

59. *Comment:* Snowcoach use should be increased based on past visitation trends, as snowcoaches could enhance the visitor experience.

Response: Snowcoach ridership has increased (except for the winter of 2008–2009 when uncertainty and economic concerns reduced all winter use). With more snowcoaches, NPS now understands that snowmobiles and snowcoaches both contribute to air quality, soundscapes, and wildlife impacts. Snowcoach limits have not been reached (the peak day in the last three years was 60 of 78 authorized). Based on these concerns, the NPS cannot increase snowcoach numbers during this interim plan. The number of snowcoaches to be allowed will be addressed in the long-term winter use plan.

60. *Comment:* The mission and purpose of the NPS is to preserve national parks for future generations; snowmobile use is considered both consistent and inconsistent with this purpose.

Response: The NPS mission is to preserve and protect the park resources while providing for visitor enjoyment. The managed winter use program during the past five winters has allowed that to occur.

61. *Comment:* The interim rule should be finalized by November 15, 2009, so people could plan for the coming season. The opening date caveat that assumes accumulation of sufficient snow is improper.

Response: When the NPS reopened the comment period on the proposed rulemaking in July, it notified the public of its intent to have a rule in place for the upcoming winter season, so that people could plan accordingly. The December 15 opening date for oversnow vehicle access has been flexible for different types of vehicles, depending on snow accumulation. When there is insufficient snow for snowmobiles or steel-tracked snowcoaches, rubber tracked snowcoaches have been allowed.

62. *Comment:* Snowmobiles are an important historical use; any recent decline in use is not related to demand but the current litigation that has occurred.

Response: NPS believes that uncertainty brought on by litigation (and recently, the economic downturn) has contributed to reduced snowmobile numbers.

63. *Comment:* Current requirements for guided snowmobile use put experiencing the park out of the reach many visitors.

Response: Yellowstone has always been an expensive place to visit in the winter, and the NPS understands that guiding and snowmobile technology requirements can add to the cost of a visit. The northern areas of the park can be visited via wheeled vehicle, where visitors are able to view many features and wildlife from the roadside or via short walks, ski, or snowshoe trips.

64. *Comment:* The visitor use survey raises legitimacy concerns, and the survey may be biased.

Response: The survey used appropriate methodologies to help begin to understand the human dimensions of wildlife and soundscapes. The methods and draft instruments were made available for public review as part of the Paperwork Reduction Act process.

65. *Comment:* The NPS finding that there would be impacts on visitor access and circulation under Alternative 1 in the 2008 EA is incorrect because not all reasonable alternatives were considered, the beneficial impacts were not considered, and the thresholds applied did not take into consideration the expiration of the 2004 rule.

Response: For reasons explained therein, the NPS considered two alternatives in the 2008 EA: No Action, which presumed no snowmobile or snowcoach access, and the Proposed Action, which called for 318 snowmobiles and 78 snowcoaches. A wide range of alternatives was considered in the earlier 2007 EIS, including the alternative specifically recommended by the commenter (allow access only from South Entrance to Old Faithful in the winter). In the 2007 EIS, major adverse impacts were found to visitor access and experience with this alternative (3A in that document). A wide range of alternatives will be considered in the long-term plan and EIS.

66. *Comment:* Snowmobiles provide the opportunity to enjoy the scenic nature of the parks.

Response: Snowmobiles and snowcoaches each provide various opportunities for visitors to enjoy the

park, and each provides different experiences for visitors.

67. *Comment:* Snowmobile use affects the scenic quality and landscapes of the park as a result of exhaust and haze.

Response: The impacts that the commenters are describing seem to be those that were experienced before the managed winter program took effect. Snowmobiles that meet NPS air and sound requirements and snowcoaches produce similar air emissions on a per passenger basis. The blue haze no longer occurs.

68. *Comment:* The use of snowmobiles in the park is adversely impacting vegetation, including impacting critical habitat.

Response: Snowmobiles and snowcoaches have always been limited to the roads that visitors use in the summer months. Off-road travel is prohibited in the park. The NPS is not aware of any effects to vegetation as a result of snowmobile or snowcoach use.

69. *Comment:* Snowmobile use in the park disrupts wildlife during the winter months when the animals are more vulnerable from such impacts as noise. Others feel snowmobile and snowcoach use does not disturb wildlife. Referenced studies should be considered.

Response: Thousands of observations of wildlife reactions to nearby oversnow vehicles have extensively documented patterns of behavioral responses in some bird and ungulate species. Substantial changes in behavior are uncommon, and none of the observed responses suggest immediate threats to the health or welfare of these wildlife populations. Furthermore, the populations of these species within the park have either grown or remained stable during the decades in which winter use expanded dramatically. The exception—the trumpeter swan—declined throughout the region due to causes unrelated to winter use. Although important research questions remain regarding the ecological effects of winter use at Yellowstone, no compelling evidence has emerged regarding impacts to the studied wildlife populations from recent research to support dramatic reductions in winter access to the park.

The rule will continue winter use at approximately the same levels as experienced in the past five years. All winter visitors to Yellowstone will be required to travel in a guided group, whether with a commercial snowmobile guide or in a guided snowcoach. Effects on wildlife are expected to be similar to those seen in the last five years, primarily negligible to minor (with possible moderate effects to swans and eagles).

70. *Comment:* NPS findings regarding the impacts of snowcoaches and snowmobiles on wildlife are inconsistent with the recommendations of NPS biologists.

Response: As discussed in the FONSI, there have been some ambiguous and somewhat inconsistent statements in past papers on wildlife impacts. NPS has determined, however, that the Selected Alternative is consistent with the biologists' actual recommendations.

The 2008 EA states, "White et al. erred in stating winter use should be limited to 50,000 oversnow visitors. [emphasis in original] Rather, they intended that the phrase read '<50,000 over-snow vehicles'" (White 2008). White 2008 is a citation to a memo from Dr. White available at http://www.nps.gov/yell/parkmgmt/upload/correction_2006winuserpt.pdf which clarifies that the intended limit was indeed 50,000 vehicles, not visitors. Had the record actually suggested a limit of 50,000 visitors, rather than vehicles, NPS would have noted as much in its discussion of the snowcoach-only transportation system in the 2007 FEIS, which would accommodate 129,600 oversnow visitors (120 snowcoaches × 12 passengers per coach × 90 days per season).

In some reports, park wildlife biologists have recommended that oversnow use be limited to the numbers observed during the "past three years [2001–2004] of their study." One example, a memo by P.J. White of November 9, 2008, has been interpreted by some to mean that snowmobile use should be limited to no more than approximately 260 snowmobiles per day and snowcoaches be limited to no more than approximately 30 per day (which were the averages those years).

Other papers by the same authors, however, discussed a wider time frame (1999–2006) and higher levels of use. The peer-reviewed scientific journal article, "Behavioral Responses of Bison and Elk in Yellowstone to Snowmobiles and Snow Coaches" by John J. Borkowski, P.J. White, Robert A. Garrott, Troy Davis, Amanda R. Hardy and Daniel J. Reinhart. *Ecological Applications* 16(5) 2006, pp. 1911–1925) makes it clear that the monitoring period they are referring to is 1999 through 2004. Average daily oversnow vehicle use ranged from 593 per day during the 2002 winter to 178 oversnow vehicles per day in 2004. Maximum daily numbers ranged up to 1168 oversnow vehicles during the study. Cumulative oversnow vehicle entries for the winter season for the West Entrance alone ranged up to 46,885 for the winter season (data are found on page 1915 of

the paper). At the conclusion (p. 1924), the authors state:

This study documented that winter visitors traveling on OSVs were essentially confined to the groomed roads, typically behaved appropriately when viewing wildlife, and rarely approached wildlife except when animals were on or immediately adjacent to the road. These attributes have allowed elk and bison in Yellowstone to habituate somewhat to OSV recreation, commonly demonstrating no observable response, and rarely displaying "fight or flight" responses when animals were off road. Further, available data provide no evidence that levels and patterns of OSV traffic during the past 35 years adversely affected the population dynamics or demography of elk and bison. Thus, we suggest regulations restricting the levels and travel routes of OSVs during our study were effective at reducing disturbances to bison and elk below a level that would cause measurable fitness effects. We acknowledge the potential for fitness effects to develop if OSVs or other stressors become more severe or prolonged. Thus, we recommend park managers consider maintaining OSV traffic levels at or below those observed during our study [1999–2004]. Regardless, numerous studies have shown that scientific findings rarely persuade people to alter their values or beliefs (e.g., Meadow et al. 2005). Thus, we suspect that varying interpretations of the behavioral and physiological response data will continue to exist because of the diverse values and beliefs of the many constituencies of Yellowstone National Park.

The Selected Alternative maintains the restrictive regulations that reduced disturbances and maintains OSV traffic levels well below those observed from 1999–2004, and is thus fully consistent with the recommendations of this peer-reviewed article and the biologists' subsequent clarifications.

71. *Comment:* The NPS did not adequately show that major impacts to wildlife (such as the road packing/grooming impacts to bison) are avoided under the current interim winter use plan.

Response: The issue of bison use of groomed roadways is addressed in detail in the 2008 EA. Impact threshold definitions were based on the best information from NPS wildlife scientists, the 2006 Management Policies, and federal laws. The NPS notes that the Selected Alternative would result only in negligible to minor effects on park wildlife (with possible moderate effects on swans), and that wildlife monitoring will continue.

72. *Comment:* Sylvan Pass and the East Entrance are an important point of access to the Park—a higher number should be used to satisfy demand and justify keeping the East Entrance open.

Response: The NPS will honor the agreement reached with the State of

Wyoming, Park County, Wyoming, and the City of Cody regarding Sylvan Pass. To that end, 20 snowmobiles and 2 snowcoaches per day are allocated to the East Entrance.

73. *Comment:* The East Entrance and Sylvan Pass should not be used because of the costs to keep the entrance open versus the revenue generated—the funds saved by closing this area could be used for other park operations.

Response: The NPS reached an agreement with the Sylvan Pass Study Group and this plan continues to implement the agreement (which recognizes weather-related constraints and NPS fiscal, staff, infrastructural, equipment, and other safety-related capacities). Management of the pass will continue to be evaluated in a long-term plan.

74. *Comment:* The 15-day comment period on the draft rule was not sufficient time to offer comment, irrelevant of the NPS justification—this violates the intent of NEPA. Further, the NPS should have accepted email comments on this issue.

Response: The NPS provided 15 days for comment on the 2008 EA and a total of 60 days for comment on the proposed rule. The decision took into account all the comments received on the proposed rule and 2008 EA. The NPS Planning, Environment, and Public Comment (PEPC) web-based system allows for electronic submission of comments. The NPS regrets any difficulties entering comments into the PEPC system, but notes that comments sent by regular mail were also accepted.

75. *Comment:* The current interim plan did not include a full range of alternatives as required under NEPA. By changing the number of snowmobile allowed in the interim plan compared to what was previously allowed, and without providing a reasoned explanation, the NPS is not compliant with the Administrative Procedure Act (APA).

Response: As discussed in the purpose and need for the 2008 EA, this EA and rulemaking considered only those options that would have allowed the NPS to open the parks for an interim period without causing major impacts. NPS did not examine options that it knew, based on previous analyses, modeling data, or monitoring data, would cause major impacts. Such impacts must first be analyzed in an EIS. In order to ensure that some motorized access could occur for the upcoming winter, NPS proposed an approach it believed could likely be supported by a Finding of No Significant Impact, which required that

no major impacts from the decision could be experienced.

The past five years of monitoring and studies has provided the NPS with information that it did not have in earlier winter use decisions. Using current monitoring and science, the NPS is drawing different conclusions regarding winter use and the contributions of snowmobiles and snowcoaches to those impacts.

As the Supreme Court has recently clarified in *Federal Communications Commission v. Fox Television Stations* (2009), there is no heightened standard for agency policy changes. An agency need not provide a more detailed analysis for a new policy; it simply must provide the same level of reasoned analysis that should justify any agency decision. NPS has indicated the reasoning for the reduced numbers of snowmobiles in the 2008 EA.

76. *Comment:* The interim plan should have been an Environmental Impact Statement (EIS) level of analysis, as opposed to an EA, so the proposed rule is invalid. Furthermore, the level of analysis was flawed because the NPS has changed its definition of impacts between the various planning processes.

Response: The 2008 EA, which did not reveal any impacts greater than moderate, is an appropriate NEPA analysis document to support this interim winter use decision and rulemaking. The rule will continue a program which has been in place for the past five winters, and whose impacts are well understood through monitoring. While the interim plan is in place, a wider range of alternatives can be analyzed in a long-term plan and EIS.

Throughout the several recent winter use processes, NPS's desired conditions have remained the same. The definition of impacts has changed in recognition of the use of monitoring data versus modeling analysis to determine impacts. The 2007 EIS primarily used computer modeling, whereas the 2008 EA used the results from monitoring.

77. *Comment:* The interim plan/EA violated NEPA because it did not provide a proper level of analysis, would result in the impairment of park resources, and is pre-decisional because the proposed rule was released two days after the 2008 EA was available for public comment. The NPS should terminate the 2008 NEPA process.

Response: A final decision was not made in December 2008. NPS did not finalize this decision until nearly a year later, after also allowing an additional 45-day public comment period for the proposed rule. NPS sought to create an interim winter use plan that would probably not have a significant impact

on the environment, which among other things means that it would not require the preparation of an EIS. That does not mean, however, that NPS had prejudged the outcome of the process. The proposed rule called for implementing the Preferred Alternative in the 2008 EA, and the NPS solicited public comment on both. NPS issued its FONSI on October 15, 2009. That decision and this final rule took into account all the comments received on the 2008 EA and proposed rule.

78. *Comment:* There are potential inconsistencies with the NPS's previously published winter use National Environmental Policy Act (NEPA) documents. The 2008 proposed rule and the 2008 EA on which it is based do not address the bulk of EPA's written comments regarding the 2007 Final Environmental Impact Statement (EIS) for winter use plans in Yellowstone and Grand Teton National Parks. EPA has concerns with the proposed rule and has mitigation and monitoring recommendations. EPA will wait for the forthcoming EIS scoping period to revisit and clarify concerns with previous winter use analyses.

Response: The past five years of monitoring and studies have provided the NPS with information that it did not have in earlier winter use decisions. Using current monitoring and science, the NPS is drawing different conclusions regarding winter use and the contributions of snowmobiles and snowcoaches to those impacts. The definition of impacts has changed in recognition of the use of monitoring data versus modeling analysis to determine impacts. The 2007 EIS primarily used computer-based modeling, whereas the 2008 EA used monitoring.

79. *Comment:* Management should avoid unacceptable or major impacts and use a mitigated FONSI as one method to address impacts from snowmobile use.

Response: The Selected Alternative does do more than prevent unacceptable impacts: it avoids all impacts that are greater than moderate. It protects the very good to excellent air quality, minimizes impacts upon park wildlife, and protects park soundscapes. Also, the plan would implement an adaptive management program that managers could utilize to adjust visitation to protect park resources even more, if for some reason monitoring determines resources are not adequately protected during these two winter seasons. Furthermore, by reacting to the exceedance of a conservative adaptive management threshold, NPS can ensure

that no unacceptable impacts or impairment occur.

80. *Comment:* There is no evidence that my comments on previous efforts had been reviewed, so the NPS should ensure that comments submitted on the draft rule are reviewed and considered.

Response: All comments submitted on the 2008 EA and proposed rule were reviewed and considered. Comments made in prior planning processes are beyond the scope of this rule, but NPS did review and consider all timely comments in those processes and this one.

81. *Comment:* The NPS had conflicting statements about the environmentally preferred alternative between different NEPA efforts.

Response: The environmentally preferred alternative is determined by the range of alternatives that are being considered in the specific NEPA document. The 2007 EIS did not contain an alternative with the numbers of snowmobiles and snowcoaches that are in the Selected Alternative (318 and 78, respectively). Most alternatives called for more snowmobiles or snowcoaches, or had only limited portions of the park open to oversnow access. The Selected Alternative provides access to all park features in a highly managed program whose impacts are well understood.

82. *Comment:* Allowing snowmobile use is in conflict with purpose for which Yellowstone was established, the mandates of the NPS such as the National Park Service Act of 1916, and NPS Management Policies because of the impact this use has to wildlife, noise, and visitor experience.

Response: While NPS agrees that public enjoyment is part of the fundamental mandate of Yellowstone and the entire National Park System, the suggestion that the Yellowstone statute and the NPS Organic Act mandate some particular level or type of snowmobile use is incorrect.

While NPS agrees that preservation of resources is key to the fundamental mandate of Yellowstone and the entire National Park System, the suggestion that the Yellowstone statute and the NPS Organic Act mandate snowcoach use is incorrect. These acts merely direct the agency to conserve park resources and provide for enjoyment without incurring impairment. If NPS is to provide for any significant visitor access to Yellowstone in the winter, motorized vehicle use is necessary, and NPS believes that the limit of 318 snowmobiles per day and 78 snowcoaches per day is consistent with the park's mandate.

The NPS Management Policies state that "NPS managers must always seek

ways to avoid, or to minimize to the greatest extent practicable, adverse impacts on park resources and values.” (Section 1.4.3) This means that NPS managers must take reasonable, affirmative steps toward avoiding or minimizing adverse impacts, but it does not go so far so as to constrain the NPS’s discretion to allow impacts that the NPS deems necessary and appropriate to provide for the enjoyment or conservation of the park.

83. *Comment:* The scope of the interim plan was misdirected, as snowmobiles have a small impact when looking at the bigger picture.

Response: Historically, oversnow vehicle use (especially snowmobiles) caused most of the impacts associated with winter use in Yellowstone, for example, accounting for the majority of air pollution. During the past five years, with the managed use program, most of those historic issues have been addressed, and the NPS now understands that snowmobiles and snowcoaches are contributing similarly to winter use related impacts.

84. *Comment:* Because the definition of the word “natural” was misapplied by the NPS, and because snowmobiles travel along developed park highways and not off-road, the executive order that regulates off-road vehicles is not applicable and snowmobile use is not subject to special regulation.

Response: NPS recognizes that Executive Order 11644 (Use of Off-Road Vehicles on Public Lands, as amended by E.O. 11989) applies to all federal agencies that allow snowmobiling. The Executive Order defines off-road vehicle as “any motorized vehicle designed for or capable of cross-country travel * * *.” That Executive Order requires federal agencies to promulgate regulations. The NPS regulation, which is found at 36 CFR 2.18, requires promulgation of special regulations like this rule.

85. *Comment:* The desired conditions established in the 2008 EA were not subject to public review and that public comment must be solicited on these conditions.

Response: The desired conditions in the 2008 EA were similar to the desired conditions identified in the 2007, 2004, 2003 and 2000 winter use plans and have been subject to public review in all those past planning processes.

86. *Comment:* Including a winter use monitoring plan in the scope of the 2008 EA was unnecessary since oversnow motorized vehicle use should not be permitted.

Response: The winter-specific monitoring complements other monitoring programs. For example, the

park monitors atmospheric deposition (including mercury), visibility (including ozone), and fine particulates at other stations.

87. *Comment:* There are resources that the NPS needed to further analyze such as subnivian fauna and climate change.

Response: A review of long-term climate trends was presented in the 2007 EIS and will be considered in the new long-term winter use plan. Subnivian fauna were dismissed as an impact topic because snowmobile and snowcoach use is confined to paved and hard-packed gravel roads that visitors use in the summer. Impacts to subnivian fauna, which may occur elsewhere as a result of cross-country motorized use, do not occur in Yellowstone.

88. *Comment:* NPS misinterprets the Organic Act, Yellowstone Park Act, Clean Air Act, General Authorities Act, the NPS Management Policies, Executive Orders, and the Park’s Master Plan. The proposed rule is fundamentally flawed. Some argue that these laws require that snowmobiles be banned, while others argue that conservation should not predominate over recreation.

Response: While the NPS agrees that public enjoyment is part of the fundamental mandate of Yellowstone and the entire National Park System, the suggestion that the Yellowstone statute and the NPS Organic Act mandate some particular level or type of use is incorrect.

Under 36 CFR 2.18, snowmobile use is prohibited except where specific routes are designated, on terms that, among other things, are consistent with park values and do not damage park resources. That regulation implements Executive Order 11644, as amended by Executive Order 11989, which applies to all federal agencies that allow snowmobiling.

Nothing in the Organic Act suggests that impairment is the only consideration that may justify imposing limitations on use. For example, the portion of the Organic Act that charges NPS with conserving the scenery, natural and historic objects, and wildlife within the parks can also justify limitations on use.

NPS Management Policies state that “NPS managers must always seek ways to avoid, or to minimize to the greatest extent practicable, adverse impacts on park resources and values.” (section 1.4.3) This means that NPS managers must take reasonable, affirmative steps toward avoiding or minimizing adverse impacts, but it does not go so far so as to constrain the NPS’s discretion to allow impacts that the NPS deems necessary and appropriate to provide for

the enjoyment or conservation of the Park.

The NPS formulated this interim winter use plan for Yellowstone in full compliance with the appropriate laws, policies, and executive orders. The amount and type of snowmobile and snowcoach use, and the restrictions on that use, will allow visitors to enjoy the park while protecting park resources.

89. *Comment:* The proposed rule does not take into consideration the precedent related to providing non-commercial opportunities in national parks, as this action would set a precedent for banning other types of vehicles in other parks.

Response: The concept of non-commercial guiding or unguided access (both with training programs) has been analyzed in previous winter plans and will be evaluated in alternatives in a long-term plan. This is a winter plan, not a summer use plan and does not set a precedent for other seasons or types of visitor access, nor does it limit what may be studied in a long-term winter use plan.

90. *Comment:* The proposed rule is not consistent with the 2008 Wyoming Court Order, and does not provide the certainty that the order called for. The interim rule constitutes a final agency action subject to judicial review, so the NPS should not take final agency action on the interim rule.

Response: The NPS believes the interim rule is consistent with all applicable court orders.

91. *Comment:* Compared to snowmobiles, snowcoaches produce greater emissions so these snowmobiles that meet NPS air and sound requirements should be allowed in the park.

Response: As discussed above, snowmobiles and snowcoaches produce similar per-passenger emissions. NPS anticipates implementing NPS air and sound requirements for snowcoaches in the future, but not during these two winter seasons.

92. *Comment:* The plan is inaccurate because there is a lack of any measurable criteria.

Response: The adaptive management plan contains both quantitative and qualitative thresholds.

93. *Comment:* Poor air quality within the park stresses wildlife, deteriorates visitor experience, and contributes to climate change.

Response: The 2008 EA analysis looked at impacts to wildlife, soundscapes, and air quality which can directly or indirectly affect these resources. It identified minor impacts to wildlife, moderate impacts to

soundscapes, and negligible impacts to air quality.

94. *Comment:* Snowmobile use in the Park should be banned to reduce global warming, conserve oil resources, and to fight the “obesity epidemic.”

Response: Snowmobiles meeting NPS emission requirements get 20–26 miles per gallon—a fuel economy far better than traditional two-stroke snowmobiles, and similar on a per-passenger basis to snowcoaches. Skiers and snowshoers use snowmobiles and snowcoaches to access trails in the park.

95. *Comment:* The NPS overstated impacts to public and employee health and safety by analyzing the No Action Alternative.

Response: In taking a hard look at the impacts of the No Action Alternative (closing the park to guided snowmobile and snowcoach access), the NPS recognized some impacts would still occur as a result of administrative access needed to protect park resources. NPS deemed those impacts to be moderate for employee health and safety.

Changes to the Final Rule

After taking the public comments into consideration and after additional internal review, one change was made to the final rule, in addition to non-substantive editorial changes made to improve clarity of the rule. This change is as follows:

Paragraph 7.13(1)(6) has been revised to delete references to snowmobiles manufactured prior to 2004. The NPS certifies snowmobiles as meeting NPS requirements for a period of six years. Winter 2009–2010 will be the last winter model year 2004 snowmobiles that were certified as meeting NPS air and emission requirements will be allowed to operate in Yellowstone. Thus, in this final rule, previous

references to model year 2003 and earlier snowmobiles were deleted.

Summary of Economic Analysis

The results of the cost-benefit analysis indicate this regulation will have de minimis negative impacts. This determination is based on a consideration of current economic conditions, visitor trends from recent years and continued uncertainty of park policies from court decisions. In addition, this winter use plan will only be in place for a two-year interim period. In order to capture the widest range of possibilities, two scenarios were analyzed within this analysis. The “expected scenario” includes the impacts that are most likely to occur and the “maximum scenario” includes the worst possible impacts that might occur. NPS believes the expected scenario is most likely to occur. Given that, the selected alternative will not have an annual economic effect of \$100 million, and will not adversely affect an economic sector, productivity, jobs, the environment, or other units of government relative to the baseline. Additionally, the selected alternative will not impose significant impacts on small businesses.

Cost-Benefit Analysis

The baseline conditions for this regulatory action are influenced by recent court decisions. When the Environmental Analysis was issued in 2008, the 2007 winter use regulation had been vacated and the authorization for snowmobile access in the 2004 winter use regulation had expired pursuant to its sunset provision. Thus, without regulatory action by NPS at that time, no snowmobile access would have been permitted, wheeled vehicle travel would have continued on roads that had

been traditionally plowed, and the park would have been open to skiing and snowshoeing.

In November 2008 the Wyoming District Court ordered the reinstatement of the 2004 regulation, without its sunset provision, until NPS promulgates a regulation to take its place. The result of that decision was the continued authorization for snowmobile and snowcoach access as provided by the 2004 regulation. While there has been no current NEPA analysis or other determination that snowmobile use at the levels authorized under that regulation is consistent with NPS statutory and other mandates, these conditions describe baseline for purposes of this regulatory analysis.

In addition the recent economic downturn has also influenced winter use. Use in the winter of 2008–2009 dropped from the previous winter in part due to economic conditions.

NPS constructed two baseline scenarios to capture the possible range of impacts. The “expected scenario” assumes that under baseline conditions snowmobile and snowcoach use will not exceed the levels permitted under the selected alternative. Indeed, to be conservative, NPS assumed that snowmobile and snowcoach use under baseline conditions in this scenario would equal that permitted under the selected alternative. That assumption is considered most likely to hold given recent trends in snowmobile use, the current economic downturn, the short two-year interim period, and the likelihood of continued uncertainty of the public regarding the winter use plan. Given that assumption, changes in snowmobile and snowcoach use under the selected alternative will be de minimis, as indicated in Table 1.

TABLE 1—WINTER SEASON SNOWMOBILE AND SNOWCOACH USE UNDER THE EXPECTED SCENARIO

Alternative	Entries		
	Snowmobile	Snowcoach	Total
Baseline	28,620	7,020	35,640
Selected Alternative	28,620	7,020	35,640
Change	0	0	0

The “maximum scenario” assumes that under baseline conditions snowmobile and snowcoach use will match levels permitted under the 2004 regulation. That regulation permits 720 snowmobiles and 78 snowcoaches to access YNP per day. Therefore, under the maximum scenario the selected alternative would reduce snowmobile

use by 402 entries per day (720 entries per day under baseline minus 318 entries per day under the selected alternative). Snowcoach use would not be reduced (78 entries per day under baseline minus 78 entries per day under the selected alternative). Therefore, as many as 36,180 snowmobile entries would be reduced in the maximum

scenario over the 90-day winter use season. NPS does not believe the maximum scenario is likely to occur given the downward trend of snowmobile use in recent winter seasons, the current economic downturn, the short two-year interim period, and the likelihood of continued

uncertainty of the public regarding the winter use plan.

TABLE 2—WINTER SEASON SNOWMOBILE AND SNOWCOACH USE UNDER THE MAXIMUM SCENARIO

Alternative	Entries		
	Snowmobile	Snowcoach	Total
Baseline	64,800	7,020	71,820
Selected Alternative	28,620	7,020	35,640
Change	-36,180	0	-36,180

Benefits and Costs

As indicated in Tables 1 and 2, the impacts of the selected alternative to snowmobile use range from a reduction of zero to 402 entries per day, with zero being the most likely to occur. Impacts to visitors are quantified as “consumer surplus,” which includes the maximum willingness to pay for such activities minus the costs of participation. Therefore, consumer surplus measures

the net benefits of visitation. These total consumer surplus changes are presented in Table 3, including total present values over the two-year period that the regulation will be in effect.

NPS estimates that businesses will not incur impacts from the selected alternative under the expected scenario. That conclusion is based on the changes in snowmobile and snowcoach use presented in Table 1, which are considered most likely. However, in the

unlikely event that the maximum scenario would occur, negative impacts would be incurred. Those impacts would be associated with the decrease in snowmobile use presented in Table 2. These impacts are termed “producer surplus,” which are a net benefits that measure similar to the consumer surplus values accruing to visitors. Total producer surplus changes for businesses under the selected alternative are presented in Table 3.

TABLE 3—QUANTIFIED CONSUMER AND PRODUCER SURPLUS IMPACTS FOR THE SELECTED ALTERNATIVE

	Expected scenario		Maximum scenario	
	Total present value	Amortized annual value	Total present value	Amortized annual value
Discount Rate:				
3 percent	\$0	\$0	-\$31,305,000	-\$15,884,000
7 percent	0	0	-30,729,000	-15,884,000

Office of Management and Budget Circular A-4 recommends a 7 percent discount rate in general, and a 3 percent discount rate when analyzing the impacts to private consumption. Values are 2003 dollars rounded to the nearest 1,000.

It is possible for visitors who do not access the park by snowmobile or snowcoach to incur increases in consumer surplus from decreased snowmobile use. In the current analysis, the expected scenario is most likely to occur with de minimis changes in snowmobile and snowcoach use; therefore, no impacts associated with this phenomenon would likely occur. Under the maximum scenario, this phenomenon would increase the consumer surplus of visitors who do not access the park by snowmobile or snowcoach. However, given recent visitor trends and the relatively low level of snowmobile and snowcoach use contemplated under the selected alternative, it is not possible at this time to estimate any such changes in visitor use. Therefore, while recognizing that such impacts to visitors are possible under the selected alternative; NPS is unable to quantify those impacts.

In addition to the potential impacts described above, NPS believes there may be a positive impact on “passive” users under the maximum scenario.

These users are individuals who do not directly use park resources and perhaps never intend to do so. Economists refer to the values these users hold using several different terms, including non-use values, passive use values, and existence values. The underlying motivations for these values include the satisfaction of knowing that a particular resource is protected or a desire to preserve the resource for future generations. Under the maximum scenario, these passive users may be more confident that park resources are being protected, and will therefore incur benefits arising from the knowledge that park resources may be more protected by the Selected Alternative. Under the expected scenario, however, de minimis changes in snowmobile and snowcoach use would occur and with commensurate impacts to these passive users.

Other benefits that could not be quantified include the potential reduction in costs of road grooming and maintenance, winter staffing, snowmobile safety hazards, and law

enforcement. In general, decreasing snowmobile activity under the maximum scenario may allow the park to redirect resources towards other activities that will protect park resources and address park management needs. Under the expected scenario, these impacts are expected to be de minimis.

Explanation of the Selected Alternative

The Selected Alternative was chosen because it best balances winter use with protection of park resources to ensure that the impairment of, or unacceptable impacts to, park resources and values does not occur. The Selected Alternative demonstrates the NPS commitment to monitor winter use and to use the results to adjust the winter use program. The results of the monitoring program, including data obtained regarding air quality, wildlife, soundscapes, and health and safety, were used in formulating the alternatives in the 2008 EA. The Selected Alternative applies the lessons learned over the last several winters relative to commercial guiding, which demonstrated, among other

things, that 100% commercial guiding has been very successful and offers the best opportunity for achieving goals of protecting park resources and allowing balanced use of the park. Law enforcement incidents have been reduced well below historic numbers, even after taking into account reduced visitation. That reduction is attributed to the quality of the guided program.

The Selected Alternative uses strictly limited oversnow vehicle numbers, combined with air and sound emission requirements and 100% commercial guiding, to help ensure that the purpose and need for the environmental impact statement is best met. With access via snowmobile, snowcoaches, or non-motorized means, park visitors will have a range of appropriate winter recreational opportunities. With the significant restrictions built into snowmobile and snowcoach use, this plan also ensures that these recreational activities will not impair or irreparably harm park resources or values.

The Selected Alternative also supports the communities and businesses both near and far from the park and will encourage them to have an economically sustainable winter recreation program that relies on a variety of modes for access to the park in the winter. Peak snowmobile numbers allowed under the Selected Alternative are well below the historic averages, but the snowmobile and snowcoach limits should provide a viable program for winter access to the park.

Compliance With Other Laws

Regulatory Planning and Review (Executive Order 12866)

This document is a significant rule and has been reviewed by the Office of Management and Budget under Executive Order 12866.

(1) This rule will not have an effect of \$100 million or more on the economy. It will not adversely affect in a material way the economy, productivity, competition, jobs, the environment, public health or safety, or state, local, or tribal governments or communities. These conclusions are based on the report "Economic Analysis: Selected Winter Use Plan for Yellowstone National Park" (Best and Vigil, October 16, 2009).

(2) This rule will not create a serious inconsistency or otherwise interfere with an action taken or planned by another agency. Implementing actions under this rule will not interfere with plans by other agencies or local government plans, policies, or controls since this is an agency specific change.

(3) This rule does not alter the budgetary effects of entitlements, grants, user fees, or loan programs or the rights or obligations of their recipients. It only affects the use of over-snow machines within specific national parks. No grants or other forms of monetary supplement are involved.

(4) OMB has determined that this rule raises novel legal or policy issues. The issue has generated local as well as national interest on the subject in the Greater Yellowstone Area. The NPS has been the subject of numerous lawsuits regarding winter use management.

Regulatory Flexibility Act

The Department of the Interior certifies that this document will not have a significant economic effect on a substantial number of small entities under the Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*). A final Regulatory Flexibility Analysis has been conducted and contained in the report "Economic Analyses: Selected Winter Use Plan for Yellowstone National Park" (Best and Vigil, October 16, 2009).

Small Business Regulatory Enforcement Fairness Act (SBREFA)

This rule is not a major rule under 5 U.S.C. 804(2), the Small Business Regulatory Enforcement Fairness Act. This rule:

- a. Does not have an annual effect on the economy of \$100 million or more.
- b. Will not cause a major increase in costs or prices for consumers, individual industries, Federal, State, or local government agencies, or geographic regions.
- c. Does not have significant adverse effects on competition, employment, investment, productivity, innovation, or the ability of U.S.-based enterprises to compete with foreign-based enterprises. This rulemaking has no effect on methods of manufacturing or production and specifically affects the Greater Yellowstone Area, not national or U.S. based enterprises.

Unfunded Mandates Reform Act

This rule does not impose an unfunded mandate on State, local, or tribal governments or the private sector of more than \$100 million per year. The rule does not have a significant or unique effect on State, local or tribal governments or the private sector. A statement containing the information required by the Unfunded Mandates Reform Act (2 U.S.C. 1531 *et seq.*) is not required. This rule addresses public use of national park lands, and imposes no requirements on other agencies or governments.

Takings (Executive Order 12630)

Under the criteria in Executive Order 12630, this rule does not have significant takings implications. Access to private property located within or adjacent to the parks will be afforded the same access during winter as before this rule. No other property is affected.

Federalism (Executive Order 13132)

Under the criteria in Executive Order 13132, this rule does not have sufficient federalism implications to warrant the preparation of a Federalism summary impact statement. A Federalism summary impact statement is not required. It addresses public use of national park lands, and imposes no requirements on other agencies or governments.

Civil Justice Reform (E.O. 12988)

This rule complies with the requirements of Executive Order 12988. Specifically, this rule:

- (a) Meets the criteria of section 3(a) requiring that all regulations be reviewed to eliminate errors and ambiguity and be written to minimize litigation; and
- (b) Meets the criteria of section 3(b)(2) requiring that all regulations be written in clear language and contain clear legal standards.

Paperwork Reduction Act

This rule does not contain information collection requirements, and a submission under the Paperwork Reduction Act (PRA) is not required.

National Environmental Policy Act

The 2008 Winter Use Plans Environmental Assessment (2008 EA) was prepared and made available for public review and comment. A Finding of No Significant Impact (FONSI) was signed October 15, 2009. The 2008 EA and FONSI are available by contacting the Yellowstone National Park Management Assistant's Office or at <http://parkplanning.nps.gov/>.

Consultation With Indian Tribes (E.O. 13175)

Under the criteria in Executive Order 13175, we have evaluated this rule and determined that it has no potential effects on federally recognized Indian tribes.

The NPS has evaluated potential effects on federally recognized Indian tribes and have determined that there are no potential effects. Numerous tribes in the area were consulted in the development of the previous winter use planning documents. Their major concern was to reduce the adverse effects on wildlife by snowmobiles. This

rule does that through implementation of the guiding requirements and disbursement of snowmobile use through the various entrance stations.

Information Quality Act

In developing this rule we did not conduct or use a study, experiment, or survey requiring peer review under the Information Quality Act (Pub. L. 106–554).

Effects on the Energy Supply (E.O. 13211)

This rule is not a significant energy action under the definition in Executive Order 13211. A Statement of Energy Effects is not required.

Administrative Procedure Act:

Comment periods on the proposed rule were provided from November 5, 2008, through November 20, 2008, and from July 24, 2009, to September 8, 2009, for a total of 60 days.

This rule is effective on December 15, 2009. The National Park Service recognizes that new rules ordinarily go into effect thirty days after publication in the **Federal Register**. For this regulation, however, we have determined under 5 U.S.C. 553(d) and 318 DM 6.25 that this rule should be effective on December 15, 2009, the traditional date for commencement of the park's winter use season. This rule implements the winter use plans for Yellowstone and relieves the restrictions on the use of snowmobiles and snowcoaches that would exist in its absence. In addition, good cause exists for the effective date of December 15, 2009, for the following reasons:

(1) The NPS has in good faith publicly stated that the 2009–2010 winter season for Yellowstone National Park would commence on December 15, 2009, and the public and businesses have made decisions based on the widespread public knowledge of this opening date.

(2) The finding of no significant impact for this rule was signed on October 15, and was made available to the public for 30 days prior to the signing of this rule. By December 15, the public therefore will have had more than 60 days notice of the NPS decision.

(3) There would be no benefit to the public in delaying the effective date of this rule, given that there has already been substantial notice of the opening date and that the park will be open under conditions substantially similar to those in effect for the past three years, other than the reduced entry limits. The above-described harms to the public resulting from a procedural delay of this rule should therefore be avoided, and an effective date of December 15, 2009, is warranted.

Drafting Information: The primary authors of this regulation are John Sacklin, Management Assistant, Yellowstone National Park; Jason Waanders, Office of the Solicitor, and Phil Selleck, Regulations Program Manager, National Park Service, Washington DC.

List of Subjects in 36 CFR Part 7

District of Columbia, National parks, Reporting and recordkeeping requirements.

■ For the reasons given in the preamble, 36 CFR part 7 is amended as set forth below:

PART 7—SPECIAL REGULATIONS, AREAS OF THE NATIONAL PARK SYSTEM

■ 1. The authority for part 7 continues to read as follows:

Authority: 16 U.S.C. 1, 3, 9a, 462(k); Sec. 7.96 also issued under DC Code 10–137 (2001) and DC Code 50–2201 (2001).

■ 2. Amend § 7.13 by revising paragraph (l) to read as follows:

§ 7.13 Yellowstone National Park.

* * * * *

(l)(1) *What is the scope of this regulation?* The regulations contained in paragraphs (l)(2) through (l)(17) of this section apply to the use of snowcoaches and recreational snowmobiles. Except where indicated, paragraphs (l)(2) through (l)(17) do not apply to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(2) *What terms do I need to know?* The definitions in this paragraph (l)(2) also apply to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, and other non-recreational users authorized by the Superintendent.

Commercial guide means a guide who operates a snowmobile or snowcoach for a fee or compensation and is authorized to operate in the park under a concession contract. In this section, “guide” also means “commercial guide.”

Historic snowcoach means a Bombardier snowcoach manufactured in 1983 or earlier. Any other snowcoach is considered a non-historic snowcoach.

Oversnow route means that portion of the unplowed roadway located between the road shoulders and designated by snow poles or other poles, ropes, fencing, or signs erected to regulate oversnow activity. Oversnow routes include pullouts or parking areas that are groomed or marked similarly to roadways and are adjacent to designated

oversnow routes. An oversnow route may also be distinguished by the interior boundaries of the berm created by the packing and grooming of the unplowed roadway. The only motorized vehicles permitted on oversnow routes are oversnow vehicles.

Oversnow vehicle means a snowmobile, snowcoach, or other motorized vehicle that is intended for travel primarily on snow and has been authorized by the Superintendent to operate in the park. An oversnow vehicle that does not meet the definition of a snowcoach must comply with all requirements applicable to snowmobiles.

Snowcoach means a self-propelled mass transit vehicle intended for travel on snow, having a curb weight of over 1,000 pounds (450 kilograms), driven by a track or tracks and steered by skis or tracks, and having a capacity of at least 8 passengers. A snowcoach has a maximum size of 102 inches wide, plus tracks (not to exceed 110 inches overall); a maximum length of 35 feet; and a Gross Vehicle Weight Rating (GVWR) not exceeding 25,000 pounds.

Snowmobile means a self-propelled vehicle intended for travel on snow, with a curb weight of not more than 1,000 pounds (450 kg), driven by a track or tracks in contact with the snow, and which may be steered by a ski or skis in contact with the snow.

Snowplane means a self-propelled vehicle intended for oversnow travel and driven by an air-displacing propeller.

(3) *May I operate a snowmobile in Yellowstone National Park?* (i) You may operate a snowmobile in Yellowstone National Park in compliance with use limits, guiding requirements, operating hours and dates, equipment, and operating conditions established under this section. The Superintendent may establish additional operating conditions and must provide notice of those conditions in accordance with § 1.7(a) of this chapter or in the **Federal Register**.

(ii) The authority to operate a snowmobile in Yellowstone National Park established in paragraph (l)(3)(i) of this section is in effect through the winter season of 2010–2011.

(4) *May I operate a snowcoach in Yellowstone National Park?* (i) Snowcoaches may only be operated in Yellowstone National Park under a concessions contract. Snowcoach operation is subject to the conditions stated in the concessions contract and all other conditions identified in this section.

(ii) All non-historic snowcoaches must meet NPS air emissions

requirements, which mean the applicable EPA emissions standards for the vehicle that were in effect at the time it was manufactured.

(iii) All critical emission-related exhaust components (as listed in 40 CFR 86.004–25(b)(3)(iii) through (v)) must be functioning properly. Such critical emissions-related components may only be replaced with the original equipment manufacturer (OEM) component, where possible. Where OEM parts are not available, aftermarket parts may be used if they are certified not to worsen emission and sound characteristics.

(iv) Modifying or disabling a snowcoach's original pollution control equipment is prohibited except for maintenance purposes.

(v) Individual snowcoaches may be subject to periodic inspections to determine compliance with the requirements of paragraphs (l)(4)(ii) through (l)(4)(iv) of this section.

(vi) The authority to operate a snowcoach in Yellowstone National Park established in paragraph (l)(4)(i) of this section is in effect only through the winter season of 2010–2011.

(5) *Must I operate a certain model of snowmobile?* Only commercially available snowmobiles that meet NPS air and sound emissions requirements as set forth in this section may be operated in the park. The Superintendent will approve snowmobile makes, models, and years of manufacture that meet those requirements. Any snowmobile model not approved by the Superintendent may not be operated in the park.

(6) *How will the Superintendent approve snowmobile makes, models, and years of manufacture for use in the park?* (i) Beginning with the 2005 model year, all snowmobiles must be certified under 40 CFR part 1051, to a Family Emission Limit no greater than 15 g/kW-hr for hydrocarbons and to a Family Emission Limit no greater than 120 g/kW-hr for carbon monoxide.

(A) 2004 model year snowmobiles may use measured emissions levels (official emission results with no deterioration factors applied) to comply with the emission limits specified in paragraph (l)(6)(i) of this section.

(B) The snowmobile test procedures specified by EPA (40 CFR parts 1051 and 1065) must be used to measure air emissions from model year 2004 and later snowmobiles.

(ii) For sound emissions, snowmobiles must operate at or below 73 dBA as measured at full throttle according to Society of Automotive Engineers J192 test procedures (revised 1985). Snowmobiles may be tested at any barometric pressure equal to or

above 23.4 inches Hg uncorrected. The Superintendent may revise these testing procedures based on new information and/or updates to the SAE J192 testing procedures.

(iii) Snowmobiles meeting the requirements for air and sound emissions may be operated in the park for a period not exceeding 6 years from the date upon which first certified.

(iv) The Superintendent may prohibit entry into the park of any snowmobile that has been modified in a manner that may adversely affect air or sound emissions.

(v) These air and sound emissions requirements do not apply to snowmobiles being operated on the Cave Falls Road in Yellowstone.

(7) *Where may I operate my snowmobile in Yellowstone National Park?* (i) You may operate your snowmobile only upon designated oversnow routes established within the park in accordance with § 2.18(c) of this chapter. The following oversnow routes are so designated for snowmobile use through the winter of 2010–2011:

(A) The Grand Loop Road from its junction with Upper Terrace Drive to Norris Junction.

(B) Norris Junction to Canyon Junction.

(C) The Grand Loop Road from Norris Junction to Madison Junction.

(D) The West Entrance Road from the park boundary at West Yellowstone to Madison Junction.

(E) The Grand Loop Road from Madison Junction to West Thumb.

(F) The South Entrance Road from the South Entrance to West Thumb.

(G) The Grand Loop Road from West Thumb to its junction with the East Entrance Road.

(H) The East Entrance Road from Fishing Bridge Junction to the East Entrance.

(I) The Grand Loop Road from its junction with the East Entrance Road to Canyon Junction.

(J) The South Canyon Rim Drive.

(K) Lake Butte Road.

(L) In the developed areas of Madison Junction, Old Faithful, Grant Village, West Thumb, Lake, Fishing Bridge, Canyon, Indian Creek, and Norris.

(M) Firehole Canyon Drive, between noon and 9 p.m. each day.

(N) North Canyon Rim Drive, between noon and 9 p.m. each day.

(O) Riverside Drive, between noon and 9 p.m. each day.

(P) Cave Falls Road.

(ii) The Superintendent may open or close these routes, or portions thereof, for snowmobile travel after taking into consideration the location of wintering wildlife, appropriate snow cover, public

safety, avalanche conditions, and other factors. Notice of such opening or closing will be provided by one or more of the methods listed in § 1.7(a) of this chapter.

(iii) This paragraph (l)(7) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(iv) Maps detailing the designated oversnow routes will be available from Park Headquarters.

(8) *What routes are designated for snowcoach use?* (i) Authorized snowcoaches may be operated on the routes designated for snowmobile use in paragraphs (l)(7)(i)(A) through (l)(7)(i)(O) of this section. The restricted hours of snowmobile use described in paragraphs (1)(7)(i)(M) through (1)(7)(i)(O) do not apply to snowcoaches. Snowcoaches may also be operated on the following additional oversnow routes through the winter of 2010–2011:

(A) Fountain Flat Road.

(B) The Grand Loop Road from Canyon Junction to Washburn Hot Springs overlook.

(C) For rubber-tracked snowcoaches only, the Grand Loop Road from Upper Terrace Drive to the junction of the Grand Loop Road and North Entrance Road, and within the Mammoth Hot Springs developed area.

(ii) The Superintendent may open or close these oversnow routes, or portions thereof, or designate new routes for snowcoach travel after taking into consideration the location of wintering wildlife, appropriate snow cover, public safety, and other factors. Notice of such opening or closing shall be provided by one of more of the methods listed in § 1.7(a) of this chapter.

(iii) This paragraph (l)(8) also applies to non-administrative snowcoach use by NPS, contractor, or concessioner employees, and other non-recreational users authorized by the Superintendent.

(9) *Must I travel with a commercial guide while snowmobiling in Yellowstone and what other guiding requirements apply?* (i) All recreational snowmobile operators must be accompanied by a commercial guide.

(ii) Snowmobile parties must travel in a group of no more than 11 snowmobiles, including that of the guide.

(iii) Guided parties must travel together within a maximum of one-third mile of the first snowmobile in the group.

(iv) The guiding requirements described in this paragraph (l)(9) do not apply to snowmobiles being operated on the Cave Falls Road.

(10) *Are there limits established for the number of snowmobiles and snowcoaches permitted to operate in the park each day?* The number of snowmobiles and snowcoaches allowed to operate in the park each day is limited to a certain number per entrance or location. The limits are listed in the following table:

Park entrance/location	Commercially guided snowmobiles	Commercially guided snowcoaches
(i) North Entrance *	12	13
(ii) West Entrance	160	34
(iii) South Entrance	114	13
(iv) East Entrance	20	2
(v) Old Faithful *	12	16
(vi) Cave Falls	**50	0

* Commercially guided snowmobile tours originating at the North Entrance and Old Faithful are currently provided solely by Xanterra Parks and Resorts. Because this concessioner is the sole provider at both of these areas, this regulation allows reallocation of snowmobiles between the North Entrance and Old Faithful as necessary, so long as the total daily number of snowmobiles originating from the two locations does not exceed 24. For example, the concessioner could operate 6 snowmobiles at Old Faithful and 18 at the North Entrance if visitor demand warranted it. This will allow the concessioner to respond to changing visitor demand for commercially guided snowmobile tours, thus enhancing the availability of visitor services in Yellowstone.

** These snowmobiles operate on an approximately 1-mile segment of road within the park where the use is incidental to other snowmobiling activities in the Caribou-Targhee National Forest. These snowmobiles do not need to be guided or to meet NPS air and sound emissions requirements.

(11) *When may I operate my snowmobile or snowcoach?* The Superintendent will determine operating hours and dates. Except for emergency situations, any changes to operating hours will be made on an annual basis, and the public will be notified of those changes through one or more of the methods listed in § 1.7(a) of this chapter.

(12) *What other conditions apply to the operation of oversnow vehicles?* (i) The following are prohibited:

(A) Idling an oversnow vehicle for more than 5 minutes at any one time.

(B) Driving an oversnow vehicle while the driver's motor vehicle license or privilege is suspended or revoked.

(C) Allowing or permitting an unlicensed driver to operate an oversnow vehicle.

(D) Driving an oversnow vehicle in willful or wanton disregard for the safety of persons, property, or park resources or otherwise in a reckless manner.

(E) Operating an oversnow vehicle without a lighted white headlamp and red taillight.

(F) Operating an oversnow vehicle that does not have brakes in good working order.

(G) The towing of persons on skis, sleds, or other sliding devices by oversnow vehicles, except in emergency situations.

(ii) The following are required:

(A) All oversnow vehicles that stop on designated routes must pull over to the far right and next to the snow berm. Pullouts must be used where available and accessible. Oversnow vehicles may not be stopped in a hazardous location or where the view might be obscured, or operated so slowly as to interfere with the normal flow of traffic.

(B) Oversnow vehicle drivers must possess a valid motor vehicle driver's license. A learner's permit does not satisfy this requirement. The license must be carried by the driver at all times.

(C) Equipment sleds towed by a snowmobile must be pulled behind the snowmobile and fastened to the snowmobile with a rigid hitching mechanism.

(D) Snowmobiles must be properly registered and display a valid registration from a state or province in the United States or Canada, respectively.

(iii) The Superintendent may impose other terms and conditions as necessary to protect park resources, visitors, or employees. The public will be notified of any changes through one or more methods listed in § 1.7(a) of this chapter.

(iv) This paragraph (I)(12) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(13) *What conditions apply to alcohol use while operating an oversnow vehicle?* In addition to 36 CFR 4.23, the following conditions apply:

(i) Operating or being in actual physical control of an oversnow vehicle is prohibited when the driver is under 21 years of age and the alcohol concentration in the driver's blood or breath is 0.02 grams or more of alcohol per 100 milliliters of blood or 0.02 grams or more of alcohol per 210 liters of breath.

(ii) Operating or being in actual physical control of an oversnow vehicle is prohibited when the driver is a snowmobile guide or a snowcoach driver and the alcohol concentration in

the operator's blood or breath is 0.04 grams or more of alcohol per 100 milliliters of blood or 0.04 grams or more of alcohol per 210 liters of breath.

(iii) This paragraph (I)(13) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(14) *Do other NPS regulations apply to the use of oversnow vehicles?* (i) The use of oversnow vehicles in Yellowstone is subject to §§ 2.18(a) and (c), but not subject to §§ 2.18 (b), (d), (e), and 2.19(b) of this chapter.

(ii) This paragraph (I)(14) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(15) *Are there any forms of non-motorized oversnow transportation allowed in the park?* (i) Non-motorized travel consisting of skiing, skating, snowshoeing, or walking is permitted unless otherwise restricted under this section or other NPS regulations.

(ii) The Superintendent may designate areas of the park as closed, reopen such areas, or establish terms and conditions for non-motorized travel within the park in order to protect visitors, employees, or park resources. Notice will be made in accordance with § 1.7(a) of this chapter.

(iii) Dog sledding and ski-joring are prohibited.

(iv) Bicycles are prohibited on oversnow routes in Yellowstone.

(16) *May I operate a snowplane in Yellowstone National Park?* The operation of a snowplane in Yellowstone is prohibited.

(17) *Is violating any of the provisions of this section prohibited?* (i) Violating

any of the terms, conditions or requirements of paragraphs (l)(1) through (l)(16) of this section is prohibited.

(ii) Anyone who violates any of the terms, conditions or requirements of this regulation will be considered to have committed one separate offense for each term, condition or requirement that they violate.

Dated: November 16, 2009.

Thomas L. Strickland,

Assistant Secretary for Fish and Wildlife and Parks.

[FR Doc. E9-27893 Filed 11-17-09; 4:15 pm]

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DEPARTMENT OF THE INTERIOR

National Park Service

36 CFR Part 7

RIN 1024-AD82

Special Regulations; Areas of the National Park System

AGENCY: National Park Service, Interior.

ACTION: Final rule.

SUMMARY: This rule governs winter visitation and certain recreational use in Grand Teton National Park and the John D. Rockefeller, Jr. Memorial Parkway. This final rule is issued to implement the Finding of No Significant Impact (FONSI) for the 2008 Winter Use Plans Environmental Assessment (2008 EA) approved October 15, 2009, and will provide visitors a range of winter recreation opportunities that are appropriate to the national park setting, and that these activities do not unacceptably impact or impair park resources and values. The rule requires that recreational snowmobiles operating on Jackson Lake meet certain air and sound emissions requirements, and that such snowmobile use is for the sole purpose of accessing ice fishing opportunities on the lake. The rule sets daily entry limits on the numbers of snowmobiles allowed on Jackson Lake and on the Grassy Lake Road, and also designates the route between Flagg Ranch and the South Entrance of Yellowstone National Park for snowmobile and snowcoach use, subject to compliance with the daily entry limits and other requirements set out in the separate rule authorizing snowmobile and snowcoach use in Yellowstone National Park. Traveling off designated oversnow routes will remain prohibited.

DATES: The effective date for this rule is December 15, 2009.

FOR FURTHER INFORMATION CONTACT: Gary Pollock, Management Assistant, Grand Teton National Park, 307-344-3428.

SUPPLEMENTARY INFORMATION:

Background

The National Park Service (NPS) has been managing winter use issues in Yellowstone National Park, Grand Teton National Park, and the John D. Rockefeller, Jr., Memorial Parkway (the Parkway) for several decades under the guidance provided by a number of sources. The history of the issue was discussed at length in the notice for the proposed rule, 73 FR 65,784 (Nov. 5, 2008) and in the 2008 Winter Use Plans Environmental Assessment (2008 EA).

After the proposed rule was published, on November 7, 2008, the U.S. District Court for the District of Wyoming issued an order reinstating the 2004 final rule on winter use in the parks, without its sunset provisions, "until such time as NPS can promulgate an acceptable rule to take its place." The NPS complied with the court order and on December 9, 2008, republished the 2004 regulation without its provisions terminating snowmobile and snowcoach use after the winter of 2006-07.

The NPS is promulgating this final regulation to replace the reinstated 2004 regulation beginning with the winter season of 2009-2010.

The EA, FONSI, and other documents pertaining to winter use management in the parks can be found at <http://www.nps.gov/yell/planyourvisit/winteruse.htm>, and at <http://www.nps.gov/grte/parkmgmt/planning.htm>.

Rationale for the Final Rule

This rule allows for a limited amount of snowmobile use in Grand Teton and the Parkway to provide a range of appropriate winter activities while protecting the integrity of park resources. It allows for winter anglers to access ice fishing opportunities on the large expanse of Jackson Lake, and for snowmobile access from the adjacent Targhee National Forest to and from Flagg Ranch, via the Grassy Lake Road. The rule also designates the route between Flagg Ranch and the South Entrance of Yellowstone for use by snowmobiles and snowcoaches, subject to any daily entry limits, air and sound emissions, guiding, and other such requirements that apply to oversnow vehicle travel within Yellowstone. The designation is necessary since winter travel through the South Entrance of Yellowstone begins and ends at Flagg Ranch, approximately two miles south of the Yellowstone boundary.

The rule is designed to protect against the adverse impacts that occurred from the historical types and numbers of oversnow vehicle use in the Park and the Parkway. Experience over the past several winters has shown that a limited number of snowmobiles, in combination with the NPS requirements for air and sound emissions on Jackson Lake, allows for a range of appropriate visitor experiences while ensuring that the integrity of park resources and values is not harmed. The NPS found that the regulations that were in effect over the past several winter seasons resulted in quieter conditions, and that impacts on air quality, wildlife, other resources, and visitor experience were acceptable. This rule limits the daily number of snowmobiles allowed on Jackson Lake and the Grassy Lake Road in order to better protect park soundscapes and other resources, and includes requirements for snowmobile air and sound emissions. It also eliminates certain oversnow vehicle routes.

This rule is consistent with the 2006 NPS Management Policies. In managing units of the National Park System, the NPS may undertake actions that have both beneficial and adverse impacts on park resources and values. However, the NPS is generally prohibited by law from taking or authorizing any action that would or is likely to impair park resources and values. Impairment is defined in the 2006 NPS Management Policies in section 1.4.5 as an impact that, in the professional judgment of the responsible NPS manager, would harm the integrity of park resources or values, including the opportunities that otherwise would be present for the enjoyment of those resources and values.

The NPS is also required to conserve the resources and values of the National Park System units and to prioritize the conservation of park resources over their use whenever the two are found to be in conflict. The NPS complies with this mandate by ensuring that a proposed use of the parks will not result in unacceptable impacts to park resources and values, and by allowing impacts to park resources only when allowing the impacts is appropriate to fulfill the purposes of the park and is necessary (meaning that the impacts are unavoidable and incapable of further mitigation in light of the authorized appropriate use).

This rule initially limits the number of snowmobiles authorized in Grand Teton to 25 per day in order to provide access to ice fishing opportunities on the large expanse of Jackson Lake. The rule allows this limit to be adjusted upward or downward, not to exceed 40

snowmobiles per day, through a monitoring and adaptive management program. The daily limits, combined with a provision that all snowmobiles used on Jackson Lake must meet NPS air and sound emissions requirements, will mitigate impacts on park resources, including the natural soundscapes of the park. The rule also allows 25 snowmobiles per day on the Grassy Lake Road in the Parkway. This route is the easternmost portion of an approximately 40-mile route that traverses the Targhee National Forest from the vicinity of Ashton, Idaho, and which terminates at Flagg Ranch. The route serves as a connection to popular snowmobile touring opportunities in the vicinity of Island Park, Idaho, and West Yellowstone, Montana. Snowmobiling opportunities abound in the portion of the route that is within the national forest, and almost all use of the route within the Parkway is incidental to activities in the forest. In view of the low amount of use that has historically occurred on the Parkway portion of the Grassy Lake Road, the importance of ensuring that visitors to the remote portions of the national forest have access to the facilities and services at Flagg Ranch (including the ability to report emergencies and obtain food and gasoline), and to provide access from Flagg Ranch to the recreational opportunities available in the national forest, snowmobiles on the Grassy Lake Road are not required to meet the air and sound emission requirements.

Adjustment to the daily entry limits for snowmobiles through an adaptive management program is one of several tools available to park managers to ensure that the goals and objectives of the winter use plans are maintained. Through an adaptive management program, if monitoring of use levels indicates that conditions are acceptable and could accommodate greater use, the NPS may increase the daily entry limits on Jackson Lake to 40 snowmobiles per day. Conversely, if monitoring indicates unacceptable conditions, the NPS will reduce use levels to an extent that acceptable conditions can be maintained.

To mitigate impacts to air quality and the natural soundscape, the NPS is continuing the requirement that all recreational snowmobiles on Jackson Lake meet strict air and sound emissions requirements. For air emissions, all snowmobiles must achieve a 90% reduction in hydrocarbons and a 70% reduction in carbon monoxide, relative to EPA's baseline emissions assumptions for conventional two-stroke snowmobiles. For sound restrictions, snowmobiles

must operate at or below 73 dBA as measured at full throttle according to Society of Automotive Engineers (SAE) J192 test procedures (revised 1985). The Superintendent will maintain a list of approved snowmobile makes, models, and years of manufacture that meet NPS requirements.

The NPS is continuing the requirement that began with the 2005 model year that all snowmobiles must be certified under 40 CFR 1051 to a Family Emission Limit (FEL) no greater than 15 g/kW-hr for hydrocarbons (HC) and 120 g/kW-hr for carbon monoxide (CO). Snowmobiles must be tested on a five-mode engine dynamometer consistent with the test procedures specified by the EPA (40 CFR 1051 and 1065). Other test methods could be approved by the NPS.

The NPS is retaining the use of the FEL method for demonstrating compliance with its emissions requirements because it has several advantages. First, use of FEL will ensure that all individual snowmobiles entering the parks achieve the NPS's emissions requirements, unless modified or damaged (under this regulation, snowmobiles which are modified in such a way as to increase air or sound emissions will not be in compliance with NPS requirements and therefore not permitted to enter the parks). Use of FEL will also minimize any administrative burden on snowmobile manufacturers to demonstrate compliance with NPS requirements, because they already provide FEL data to the EPA. Further, the EPA has the authority to ensure that manufacturers' claims on their FEL applications are valid. EPA also requires that manufacturers conduct production line testing (PLT) to demonstrate that machines being manufactured actually meet the certification levels. If PLT indicates that emissions exceed the FEL levels, then the manufacturer is required to take corrective action. Through EPA's ability to audit manufacturers' emissions claims, the NPS will have sufficient assurance that emissions information and documentation will be reviewed and enforced by the EPA. FEL also takes into account other factors, such as the deterioration rate of snowmobiles (some snowmobiles may produce more emissions as they age), lab-to-lab variability, test-to-test variability, and production line variance. In addition, under the EPA's regulations, all snowmobiles manufactured must be labeled with FEL air emissions information. This will help to ensure that NPS emissions requirements are consistent with these

labels. The use of FEL will avoid potential confusion for consumers.

The air emissions requirements for snowmobiles allowed to operate in the park should not be confused with standards adopted by the EPA in a final rule published in the **Federal Register** on November 8, 2002 (67 FR 68242). The EPA regulations require manufacturers to meet certain fleet averages for HC and CO emissions. For example, the Phase 1 standards required all snowmobile manufacturers to meet a fleet-wide average in 2007 of 275 g/kW-hr for CO and 100 g/kW-hr for HC, which represents a 30-percent reduction from the baseline emission rates for uncontrolled snowmobiles. Any particular make/model may emit more or less than the standard as long as the fleet average does not exceed the standard. Phase 2 and Phase 3 standards will be implemented in 2010 and 2012, respectively, effectively requiring the equivalent of a 50% reduction in both HC and CO as compared to average baseline levels. By comparison, NPS requires that all snowmobiles operating in the Parks meet a FEL of 120 g/kW-hr for CO and 15 g/kW-hr for HC. This means that snowmobiles operating in the park represent the cleanest that are commercially available.

To determine compliance with the sound emissions requirements, snowmobiles must be tested using SAE J192 test procedures (revised 1985; or potentially as further revised and adapted for use by NPS). The NPS recognizes that the SAE updated these test procedures in 2003; however, the changes between the 2003 and 1985 test procedures could yield different measurement results. The sound emissions requirement was initially established using 1985 test procedures (in addition to information provided by industry and modeling). To ensure consistency in the test results, the NPS will at this time continue to use the 1985 test. The SAE J192 (revised 1985) test also allows for a tolerance of 2 dBA over the sound limit to account for variations in weather, snow conditions, and other factors. The NPS understands that an update to the 2003 J192 procedures may be underway, and the NPS will continue to evaluate these test procedures and possibly adopt them after these regulations are implemented. Other test methods could be approved by NPS on a case-by-case basis.

Snowmobiles may be tested at any barometric pressure equal to or above 23.4 inches Hg uncorrected (as measured at or near the test site). This exception to the SAE J192 test procedures maintains consistency with the testing conditions used to determine

the sound requirement. This allowance for reduced barometric pressure is necessary since snowmobiles were tested at the elevation of Yellowstone National Park, where atmospheric pressure is lower than that under the SAE J192's requirements. Testing data indicate that snowmobiles test quieter at high elevation, and therefore some snowmobiles may comply with the NPS's sound emissions requirements at higher elevations even though they do not when tests are conducted near sea level.

The NPS will annually publish a list of snowmobile makes, models, and years of manufacture that meet its emissions and sound requirements. Snowmobile manufacturers may demonstrate that snowmobiles are compliant with the air emissions requirements by submitting to the NPS a copy of their applications used to demonstrate compliance with EPA's general snowmobile regulation (indicating FEL). The NPS will accept this application information from manufacturers in support of conditionally certifying a snowmobile as meeting its air emissions requirements, pending ultimate review and certification by EPA at the same emissions levels identified in the application. Should EPA certify a snowmobile at an emission level that would no longer meet the NPS's requirements, this snowmobile would no longer be considered by NPS to be compliant with its requirements and would be phased out according to a schedule that will be determined by the NPS to be appropriate. For sound emissions, snowmobile manufacturers may submit their existing Snowmobile Safety and Certification Committee (SSCC) sound level certification form. Under the SSCC machine safety standards program, snowmobiles are certified by an independent testing company as complying with all SSCC safety standards, including sound standards. This regulation does not require the SSCC form specifically, as there could be other acceptable documentation in the future. The NPS will work cooperatively with the snowmobile manufacturers on appropriate documentation. The NPS intends to continue to rely on certified air and sound emissions data from the private sector rather than establish its own independent testing program. When the NPS certifies snowmobiles as meeting its requirements, it will announce how long that certification applies. Generally, each snowmobile model will be approved for entry into the parks for 6 winter seasons after it is

first listed. Based on NPS experience, 6 years represents the typical useful life of a snowmobile, and thus 6 years provides purchasers with a reasonable length of time where operation is allowed once a particular model is listed as being compliant. It is also based on EPA snowmobile emission regulations and the deterioration factors that are part of those regulations (EPA requires that if a manufacturer certifies its snowmobile will comply with EPA's emission regulations, the snowmobile will meet those regulations for a period of 5 years or 5,000 miles). The NPS recognizes that some privately owned snowmobiles used predominantly for ice fishing on Jackson Lake may have relatively low mileages even after 6 years of use, and therefore may not have experienced the type of deterioration that would cause them to fail NPS air and sound emissions requirements. The certification period for snowmobiles being operated on Jackson Lake will still be considered to be 6 years, but it may be extended up to a total of 10 years as long as the snowmobile's mileage does not exceed 6,000 miles.

Individual snowmobiles modified in such a way as to increase sound and air emissions of hydrocarbons (HC) and carbon monoxide (CO) beyond the proposed emission restrictions will be denied entry to the parks. It is the responsibility of end users to ensure that their oversnow vehicles, whether snowmobiles or snowcoaches, comply with all applicable restrictions.

Snowmobiles being operated on the Grassy Lake Road will not be required to meet air and sound emissions requirements regardless of whether they originate travel at Flagg Ranch or in the national forest. In light of the relatively short length of this segment and the very limited amount of snowmobile use, the NPS has determined that the impacts of this use of snowmobiles that does not meet NPS air and sound emissions requirements are acceptable.

Scientific studies and monitoring of winter visitor use and park resources will continue. If these studies indicate that human presence or activities have a substantial adverse effect on wildlife or other park resources that cannot otherwise be mitigated, as part of its adaptive management of winter use activities the NPS will close selected areas to visitor use. A one-year notice will ordinarily be provided before any such closure is implemented unless immediate closure is deemed necessary to avoid impairment of park resources. The Superintendent will continue to have the authority under 36 CFR 1.5 to take emergency actions to protect park resources or values.

Snowmobiles will continue to be restricted to designated routes, which are either roads that are traveled by motor vehicles during the remainder of the year, or in the case of Jackson Lake, by motorboats during the summer.

The NPS will close the Continental Divide Snowmobile Trail (CDST) as an oversnow vehicle route through most of Grand Teton and the Parkway. Experience over the past several winters strongly suggests that the minimal amount of use on this route would not substantially increase if it were to remain open, since much of the previous use of the CDST was associated with visitors traveling through to Yellowstone. The NPS recognizes that the guiding and air and sound emissions requirements for Yellowstone have contributed to a substantial reduction in the use of the CDST, since visitors have not been able to continue into Yellowstone without a guide and a snowmobile that meets the emissions requirements, as well as complying with the daily entrance caps.

The NPS also recognizes, however, that snowmobile access to and from the Targhee National Forest is important to some visitors. While the CDST will no longer be maintained or designated for snowmobile use, the air and sound emissions requirements for the Grassy Lake Road will be removed beginning with the 2009–2010 winter season. Snowmobilers will be able to transport their machines by trailer between Moran and Flagg Ranch using plowed roads, in order to connect to the Grassy Lake Road and the national forest lands to the west of the Parkway. The daily entry limit of 25 is sufficient to accommodate the levels of use that have typically occurred in the past and those which are reasonably foreseeable.

Summary of and Responses to Public Comments

The NPS published a proposed rule on November 5, 2008, (73 FR 65784) and initially accepted public comments through November 20, 2008. The comment period was reopened on July 24, 2009, and comments were accepted through September 8, 2009. Comments were accepted through the mail, hand delivery, and through the Federal eRulemaking Portal: <http://www.regulations.gov>. A total of 39,796 comment documents were received. Since the proposed rule was combined with rulemaking for winter use management in Yellowstone, many of the comments addressed issues primarily or entirely related to that park. The comments and responses below are those that were pertinent to the issues in Grand Teton and the Parkway.

Comments relevant to Yellowstone are included in a separate rulemaking.

1. *Comment:* The NPS should not require the use of BAT snowmobiles on Jackson Lake.

Response: The BAT requirement on Jackson Lake is important in mitigating the impacts of snowmobile use on Jackson Lake. Due to the large and unobstructed expanse of the frozen surface of the lake, sound from snowmobiles is able to propagate over long distances, and therefore could have a disproportionate impact on the natural soundscapes even at relatively low levels of use. The BAT requirement helps to mitigate that impact due to the reduced sound emissions compared to non-BAT snowmobiles, as well as the tonal qualities of the sound.

2. *Comment:* The daily limits for snowmobile use on Jackson Lake and the Grassy Lake Road are too low.

Response: The NPS recognizes that the levels of use on Jackson Lake in recent years may have been affected by the uncertainty over the winter use issue, and winter anglers may have been reluctant to purchase BAT snowmobiles. Should this reluctance diminish as a result of greater certainty regarding winter use management on Jackson Lake, some increase in BAT snowmobile use could be expected. In light of the amount of use in recent years, and after considering the historic levels of use on Jackson Lake, the NPS believes that the daily limits established will accommodate current and reasonably foreseeable future demand for snowmobile access on Jackson Lake. Similarly, the limits established for the Grassy Lake Road are sufficient to accommodate current use levels and those that are reasonably foreseeable.

3. *Comment:* Since the level of use analyzed in the Selected Alternative is considered less than a significant impact, the decision would unnecessarily restrict snowmobile use with no resulting benefit for park resources.

Response: The NPS believes that the snowmobile daily entry limits on Jackson Lake and the Grassy Lake Road are sufficient to accommodate current and reasonably foreseeable demand for use of those areas. At the same time, the NPS believes the Selected Alternative will be protective of park resources. The NPS is not obligated to increase the daily limits to a level that would be considered significant simply for the purpose of accommodating demand that may or may not be present. Should the need arise in the future, the NPS could revisit whether the daily entry limits should be adjusted.

4. *Comment:* The NPS should use the ongoing monitoring of the effects of snowmobile use to determine the appropriate number of snowmobiles that can access Jackson Lake.

Response: The Selected Alternative allows for an initial daily entry limit of 25 snowmobiles per day on Jackson Lake. This level of use is higher than the recent-years average of 3–5 snowmobiles per day, as well as the peak day of 17. The NPS believes that it is reasonable to expect that use will increase somewhat once the winter use management situation stabilizes, but does not have any information that suggests that the demand cannot be accommodated within the limits established in the Selected Alternative. Nevertheless, the decision allows for the entry limits to be adjusted up or down, not to exceed 40 per day based on monitoring and adaptive management.

5. *Comment:* The daily snowmobile limits on Jackson Lake and the Grassy Lake Road are too high in light of the low level of snowmobile use that has occurred in the Park and Parkway in recent years. The daily limits should be capped at current levels.

Response: The daily limits proposed in the Selected Alternative are low enough to ensure that no significant impacts will occur, but high enough to accommodate a reasonable amount of increased use. The NPS recognizes that the levels of use on Jackson Lake in recent years may have been affected by the uncertainty over the winter use issue, and winter anglers may have been reluctant to purchase BAT snowmobiles.

6. *Comment:* Snowmobile use should not be allowed on Jackson Lake because anglers can access the lake by non-motorized means, and because snowmobiles leave behind pollutants.

Response: Due to the large size of Jackson Lake, most of it would be inaccessible without the some form of motorized access.

7. *Comment:* Snowmobiles should not be allowed on Jackson Lake because they leave behind pollutants that enter the lake when the ice melts.

Response: The requirement that snowmobiles meet BAT requirements, combined with the daily entry limits, will result in negligible levels of pollutants entering the waters of Jackson Lake.

8. *Comment:* The Continental Divide Snowmobile Trail (CDST) through Grand Teton and the Parkway should be kept open. It is an important link between popular snowmobile touring opportunities portions of the CDST outside the park and in the Island Park/West Yellowstone areas.

Response: In recent years, use of the CDST through the parks was extremely low—in the neighborhood of 15 snowmobiles per season. Although use levels were higher than that prior to the initiation of guiding and BAT requirements in Yellowstone, the amount of use has always been modest. The amount of use on the CDST does not warrant the cost of continuing to construct and maintain the trail each year. Since BAT snowmobiles will no longer be required on the Grassy Lake Road, those wishing to complete a long-distance tour between other parts of the CDST and the Island Park/West Yellowstone areas will be able to trailer their snowmobiles through the parks to Flagg Ranch and continue traveling the trail from there.

9. *Comment:* The Continental Divide Snowmobile Trail between Moran Junction and Flagg Ranch should be discontinued.

Response: The CDST will no longer be designated or maintained for snowmobile use.

10. *Comment:* Implementation of a long-term plan for Grand Teton and the Parkway, while putting in place an interim plan for Yellowstone will cause further confusion and uncertainty for the public.

Response: Separate decisions regarding Yellowstone, Grand Teton, and the Parkway are not expected to create confusion and uncertainty. The oversnow vehicle use allowed under the Selected Alternative at Grand Teton and the Parkway is separate and distinct from that which occurs in Yellowstone. The use of snowmobiles on Jackson Lake for ice fishing has no connection with opportunities for touring Yellowstone, nor does use of the Grassy Lake Road. A long-term decision for Grand Teton and the Parkway will alleviate the existing confusion and uncertainty regarding winter use management in those two areas.

11. *Comment:* Snowmobiles should not be allowed on the Grassy Lake Road.

Response: The Grassy Lake Road within the Parkway is the easternmost segment of an approximately 40-mile route that extends from near Ashton, Idaho, to Flagg Ranch, mostly within the Targhee National Forest. Within the national forest, there are many opportunities for winter recreation, including snowmobile touring. Allowing snowmobile use on the portion of the Grassy Lake Road within the Parkway provides opportunities for visitors to the national forest to access the services available at Flagg Ranch, including emergency notification, and for visitors to access the adjacent national forest lands from Flagg Ranch.

12. *Comment:* The NPS should implement a winter shuttle service that would haul non-BAT snowmobiles through Grand Teton and Yellowstone so that tours to West Yellowstone would be possible without a long side trip through Island Park.

Response: The NPS would consider proposals from a potential service provider for such a service.

13. *Comment:* The EA did not include a sufficient range of alternatives.

Response: As discussed in the purpose and need for the 2008 EA, this EA and rulemaking considered only those options that would have allowed the NPS to open the parks for an interim period without causing major impacts. NPS did not examine options that it knew, based on previous analyses, modeling data, or monitoring data, would cause major impacts. Such impacts must first be analyzed in an EIS. In order to ensure that some motorized access could occur for the upcoming winter, NPS proposed an approach it believed could likely be supported by a Finding of No Significant Impact, which required that no major impacts from the decision could be experienced.

14. *Comment:* The decision to permanently allow snowmobile use in Grand Teton and the Parkway will impact future long-term decisions regarding winter access into Yellowstone.

Response: The snowmobile use authorized in Grand Teton and the Parkway is distinct and separate from oversnow vehicle access into Yellowstone. The plan for Grand Teton and the Parkway allows a limited amount of snowmobile access for ice fishing on Jackson Lake, and for use of the Grassy Lake Road between Flagg Ranch and the Targhee National Forest. Neither of these uses has any bearing on winter access into Yellowstone.

15. *Comment:* The comment period on the EA was too short.

Response: The EA and proposed rule were available for public comment for nearly concurrent 15-day periods ending on November 17, 2008, and November 20, 2008, respectively. The lengths of the comment periods were based on the time constraints that existed at the time, because NPS was attempting to complete a NEPA and rulemaking process in time for the 2008–2009 winter season. Subsequent events made it possible for NPS to reopen the proposed rule for an additional 45 days of public comment in July 2009. The issues are largely the same, and the NPS considered all of the comments made during both comment periods in the NEPA process.

16. *Comment:* By releasing a proposed rule contemporaneously with the 2008 EA, the NPS indicated that it had already made a decision regarding the outcome of the EA.

Response: Publication of the proposed rule did not prejudice or commit the NPS to a course of action since it was not a final rule. The rule could be altered any time prior to publication of a final rule in the **Federal Register**.

17. *Comment:* The NPS should disclose the amount of Federal funds that have been spent on winter use planning for the parks.

Response: Since 1997, the NPS has spent over \$11 million on planning for winter use management in Yellowstone, Grand Teton, and the Parkway.

18. *Comment:* The No-action alternative in the EA (Alt 1—Eliminate Motorized Recreational Oversnow Travel) was incorrect because it should have represented the “current level of activity,” meaning the daily entry limits that had been in effect the previous four winters.

Response: When the 2008 EA was prepared, the 2007 rule had been vacated. No snowmobile or snowcoach use would have been authorized without action by the NPS, because the authorizations in the 2004 rule had expired pursuant to the sunset date provisions. After the 2008 EA was issued, the U.S. District Court for the District of Wyoming reinstated the 2004 rule without the sunset clauses, and as a result, up to 720 snowmobiles per day were allowed for the winter of 2008–09. Due to a pending appeal, there is still uncertainty regarding that reinstatement. As explained above, there has been no current NEPA analysis or other determination that use at the levels authorized under that regulation is consistent with the NPS’s statutory and other mandates. Accordingly, the No Action Alternative analyzed in the 2008 EA represents a more logical and useful benchmark against which impacts can be compared, and therefore continues to better satisfy the purposes of the no action alternative under NEPA.

19. *Comment:* The NPS has no basis for reducing the number of snowmobiles allowed on Jackson Lake from 40 per day to 25.

Response: The rule provides an initial limit of 25 per day, but allows up to 40. The initial limit is sufficient to accommodate the amount of use that has been occurring in recent years, and the potentially higher limit of 40 is sufficient to accommodate reasonably foreseeable increases in fishing/snowmobiling demand.

20. *Comment:* The impacts associated with the use of snowmobiles violate the NPS Organic Act prohibition on the impairment of park resources.

Response: As described in the EA and FONSI, the impacts associated with the limited and carefully regulated use of snowmobiles under the Selected Alternative do not constitute impairment of park resources or cause unacceptable impacts.

21. *Comment:* The NPS must take into account its obligation to seek to perpetuate the best possible air quality in the parks.

Response: The limited amount of snowmobile use permitted in the Selected Alternative will have negligible impacts on the air quality of the parks.

22. *Comment:* The NPS should continue operating the parks under the 2004 rules.

Response: The 2004 rules were reinstated by the U.S. District Court for the District of Wyoming and were intended to provide guidance until such time as the NPS could promulgate new rules. Continued operation under the reinstated rule would lead to uncertainty, as a result of pending litigation in both the U.S. Court of Appeals for the Tenth Circuit and the U.S. District Court for the District of Columbia. In addition, there has been no current NEPA analysis or other determination that the use levels authorized under that regulation are consistent with the NPS’s statutory and other mandates.

23. *Comment:* The NPS should delay implementation of any new rules for one season in order to minimize confusion.

Response: The NPS is currently managing winter use activities under a 2004 rule that was reinstated by the U.S. District Court for the District of Wyoming. Continued operation under that rule would lead to uncertainty, as a result of pending litigation in both the U.S. Court of Appeals for the Tenth Circuit and the U.S. District Court for the District of Columbia. In addition, there has been no current NEPA analysis or other determination that the use levels authorized under that regulation are consistent with the NPS’s statutory and other mandates. In order to ensure that the Park and Parkway are open to oversnow vehicle use for the winter of 2009–2010, the NPS believes it is prudent to implement the Selected Alternative prior to the start of the season.

24. *Comment:* The NPS general regulation on snowmobiles, 36 CFR 2.18, requires that snowmobiles may only be allowed where they will not disturb wildlife.

Response: Winter use has some small and occasional effects on wildlife, just like every other form of visitor use of the park, but the impacts are expected to be acceptable, and are a necessary and unavoidable consequence of an appropriate use. Taking or intentionally disturbing wildlife are prohibited by the NPS's general regulations on natural resource preservation and wildlife protection, 36 CFR 2.1, 2.2. Section 2.18, which addresses the use of snowmobiles in the National Park System, requires that routes for snowmobile use may only be designated where the use will not disturb wildlife, but in doing so does not establish a different standard than sections 2.1 and 2.2. The wildlife responses to normal snowmobile use that are expected to occur do not cause the taking, frightening, or intentional disturbance that is prohibited by NPS regulations.

25. *Comment:* The NPS has not provided a reasoned explanation for its adaptive management thresholds and their consistency with its mandates.

Response: The adaptive management thresholds are a management tool only; they do not represent the unacceptable impacts or impairment thresholds described in section 1.4 of the Management Policies. Rather, they are a conservative measure used to alert the NPS manager that additional attention to a particular park resource or value is merited. By reacting to the exceedance of a conservative adaptive management threshold, NPS can seek to ensure that no unacceptable impacts or impairment occur.

26. *Comment:* The NPS should require winter users to maintain a 100 meter distance from animals when stopping.

Response: The NPS requires visitors to stay at least 100 yards (91 meters) away from bears and wolves, and at least 25 yards (21 meters) away from all other animals.

27. *Comment:* Promulgation of a long-term rule for winter use management at Grand Teton and the Parkway is a major Federal action pursuant to NEPA and should have required the preparation of an environmental impact statement.

Response: As disclosed in the 2008 environmental assessment and 2009 Finding of No Significant Impact, the Selected Alternative (and its implementation in the final rule) is not an action that normally requires the preparation of an environmental impact statement. Nor will it have a significant effect on the human environment.

Environmental impacts that could occur are limited in context and intensity, with generally adverse impacts that range from localized to widespread,

short- to long-term, and negligible to minor. As discussed in the EA, impacts would have to rise to the level of major to be considered significant, and these impacts fall well short of major impacts. There are no unmitigated adverse effects on public health, public safety, threatened or endangered species, sites or districts listed in or eligible for listing in the National Register of Historic Places, or other unique characteristics of the region. No highly uncertain or controversial impacts, unique or unknown risks, significant cumulative effects, or elements of precedence were identified. Implementation of the action will not violate any Federal, State, or local environmental protection law. Based on the foregoing, it has been determined that an EIS is not required for this project.

28. *Comment:* The NPS should allow the Wyoming Game and Fish Department to use non-BAT snowmobiles to access Jackson Lake.

Response: When the use of BAT snowmobiles is unsafe or impractical, or the use of non-BAT snowmobiles is otherwise necessary for the proper administration of the park, the Superintendent may allow NPS and other authorized parties to use non-BAT snowmobiles for specialized administrative purposes, such as law enforcement, search and rescue, or other management functions.

Changes to the Final Rule

After taking the public comments into consideration and after additional internal review, several changes were made to the final rule, in addition to non-substitutive editorial changes made to improve clarity of the rule. These changes are as follows:

First, paragraphs 7.21(a)(2) and 7.22(g)(2) for the Parkway and Grand Teton have been revised to fully describe all of the terms that are necessary to know, rather than simply referencing those terms in the corresponding section of the Yellowstone rule, § 7.13(l).

Second, paragraph 7.21(a)(4) was revised to eliminate the descriptions for snowcoach air emissions requirements. The revision does not necessarily remove emissions requirements, however, because any snowcoach use in the Parkway will only be pursuant to a concessions contract with Yellowstone National Park, and snowcoaches will be required to meet any emissions requirements in the Yellowstone regulations at § 7.13(l).

Third, paragraphs 7.21(a)(5) and (6) regarding the models of snowmobiles that may be operated in the Parkway, along with snowmobile air and sound

emissions requirements, were deleted and subsequent paragraphs were renumbered accordingly. The revision simply reflects the fact that snowmobile trips into Yellowstone through the South Entrance of the park begin and end at Flagg Ranch, and that all use of the two-mile route between Flagg Ranch and the South Entrance is in conjunction with those trips. Therefore, the final rule specifies at paragraph 7.21(a)(5) that snowmobiles traveling along the route between Flagg Ranch and the South Entrance must meet any air and sound emissions requirements and other conditions described in the Yellowstone regulations at § 7.13(l). The proposed rule did not include any air and sound emissions requirements for the Grassy Lake Road, and therefore the revision does not affect snowmobiles using that route.

Fourth, paragraph 7.21(a)(10), renumbered as 7.21(a)(8), was revised to remove the table showing snowmobile and snowcoach daily limits and simply describe those limits in the text. The revision also specifies that the daily entry limits for snowmobiles and snowcoaches on the route from Flagg Ranch to the South Entrance of Yellowstone are established in § 7.13(l).

Summary of the Economic Analysis

NPS analyzed the potential costs and benefits associated with the Selected Alternative as compared to the baseline conditions. The baseline conditions for this regulatory action are influenced by recent court decisions. When the Environmental Analysis was issued in 2008, the 2007 winter use regulation had been vacated and the authorization for snowmobile access in the 2004 winter use regulation had expired pursuant to its sunset provision. Thus, without regulatory action by NPS at that time, no snowmobile access would have been permitted, wheeled vehicle travel would have continued on roads that had been traditionally plowed, and the Park would have been open to skiing and snowshoeing.

However, in November 2008 the Wyoming District Court ordered the reinstatement of the 2004 regulation, without its sunset provision, until NPS promulgates an acceptable regulation to take its place. The result of that decision is the continued authorization for snowmobile access as provided by the 2004 regulation. While there has been no current NEPA analysis or other determination that snowmobile use at the levels authorized under that regulation is consistent with NPS statutory and other mandates, these conditions describe baseline for purposes of this regulatory analysis.

NPS has considered the Selected Alternative vis-à-vis these baseline conditions and determined that the resulting changes in winter season visitation will be de minimis. For example, in Grand Teton National Park, snowmobile use on Jackson Lake in recent years has averaged less than five machines per day. On a few days, snowmobile use has involved approximately 10 machines, and approximately 19 machines on the peak use day. On Grassy Lake Road, snowmobile use has typically been no more than five to 10 machines per day, and often less. Some increases beyond these levels may occur if uncertainty about authorized use levels is reduced by this regulatory action. However, such increases would be expected only gradually, if at all. The Selected Alternative will initially permit 25 snowmobiles per day on the Lake for ice fishing access, and as many as 40 snowmobiles per day if monitoring indicates acceptable resource impacts. Therefore, the Selected Alternative is not considered binding on snowmobile use within the Parks and any changes in visitation from the baseline conditions would likely be imperceptible.

Given that, NPS believes the incremental benefits and costs associated with the Selected Alternative are essentially zero for both visitors and businesses alike. Therefore, NPS estimates zero net benefits (benefits minus costs) as a result of the Selected Alternative.

NPS also analyzed the potential economic impacts of the Selected Alternative on small entities, considering the potential changes in business revenue that could occur under that alternative relative to the baseline conditions. As noted above, NPS believes that any changes in winter season visitation in the Parks resulting from the Selected Alternative will be de minimis. Therefore, NPS also believes that any revenue impacts on small entities will also be de minimis. Additionally, NPS notes that most of the visitors potentially affected by the Selected Alternative reside in the local area, and that these visitors do not utilize local hotels, restaurants, or other businesses to the extent that those coming from outside the GYA would. Therefore, NPS does not believe that significant impacts to a substantial number of small entities will occur as a result of the Selected Alternative. Given that, NPS has determined that the Selected Alternative will have de minimis impacts on the affordability or viability of local businesses, small or large.

Compliance With Other Laws

Regulatory Planning and Review (Executive Order 12866)

This document is not a significant rule and the Office of Management and Budget has not reviewed this rule under Executive Order 12866. We have made the assessments required by E.O. 12866 and the results are given below.

(1) This rule will not have an effect of \$100 million or more on the economy. It will not adversely affect in a material way the economy, productivity, competition, jobs, the environment, public health or safety, or State, local, or Tribal governments or communities. These conclusions are based on the report "Economic Analyses: Selected Winter Use Plan for Grand Teton National Park and John D. Rockefeller, Jr., Memorial Parkway" (Peacock, September 2009).

(2) This rule will not create a serious inconsistency or otherwise interfere with an action taken or planned by another agency. Implementing actions under this rule will not interfere with plans by other agencies or local government plans, policies, or controls since this is an agency-specific change.

(3) This rule does not alter the budgetary effects of entitlements, grants, user fees, or loan programs or the rights or obligations of their recipients. It only affects the use of snowmobiles within specific units of the National Park System. No grants or other forms of monetary supplement are involved.

(4) While the NPS has been the subject of numerous lawsuits regarding winter use management, this rule does not raise novel legal or policy issues.

Regulatory Flexibility Act

The NPS has determined that this regulatory action will not have a significant effect on a substantial number of small entities under the Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*). This conclusion is based on the report "Economic Analyses: Selected Winter Use Plan for Grand Teton National Park and John D. Rockefeller, Jr., Memorial Parkway" (Peacock, September 2009).

Small Business Regulatory Enforcement Fairness Act (SBREFA)

This rule is not a major rule under 5 U.S.C. 804(2), the Small Business Regulatory Enforcement Fairness Act. This rule:

(a) Does not have an annual effect on the economy of \$100 million or more. This conclusion is based on the report "Economic Analyses: Selected Winter Use Plan for Grand Teton National Park

and John D. Rockefeller, Jr., Memorial Parkway" (Peacock, September 2009).

(b) Will not cause a major increase in costs or prices for consumers, individual industries, Federal, State, or local government agencies, or geographic regions.

(c) Does not have significant adverse effects on competition, employment, investment, productivity, innovation, or the ability of U.S.-based enterprises to compete with foreign-based enterprises. This rulemaking has no effect on methods of manufacturing or production and specifically affects the immediate area surrounding Grand Teton National Park and the John D. Rockefeller, Jr. Memorial Parkway, not national or U.S.-based enterprises.

Unfunded Mandates Reform Act

This rule does not impose an unfunded mandate on State, local, or Tribal governments or the private sector of more than \$100 million per year. The rule does not have a significant or unique effect on State, local or Tribal governments or the private sector. It addresses public use of national park lands, and imposes no requirements on other agencies or governments.

Takings (Executive Order 12630)

Under the criteria in Executive Order 12630, this rule does not have significant takings implications. Access to private property located within or adjacent to the parks will still be afforded the same access during winter as before this rule. No other property is affected.

Federalism (Executive Order 13132)

In accordance with Executive Order 13132, the rule does not have sufficient federalism implications to warrant the preparation of a Federalism Assessment. It addresses public use of national park lands, and imposes no requirements on other agencies or governments.

Civil Justice Reform (Executive Order 12988)

This rule complies with the requirements of Executive Order 12988. Specifically, this rule:

(a) Meets the criteria of section 3(a) requiring that all regulations be reviewed to eliminate errors and ambiguity and be written to minimize litigation; and

(b) Meets the criteria of section 3(b)(2) requiring that all regulations be written in clear language and contain clear legal standards.

Paperwork Reduction Act

This rule does not contain information collection requirements,

and a submission under the Paperwork Reduction Act (PRA) is not required.

National Environmental Policy Act

This rule does not constitute a major Federal action significantly affecting the quality of the human environment. We have prepared an Environmental Assessment (Winter Use Plans Environmental Assessment, November 2008) under the National Environmental Policy Act of 1969. A Finding of No Significant Impact (FONSI) has also been completed. The EA and FONSI are available for review by contacting the Superintendent's Office at Grand Teton National Park, or can be found online at: <http://www.nps.gov/grte/parkmgmt/planning.htm>.

Consultation With Indian Tribes (E.O. 13175)

Under the criteria in Executive Order 13175, we have evaluated this rule and determined that it has no potential effects on Federally recognized Indian Tribes. Numerous Tribes in the area were consulted, however, in the development of the previous NEPA processes. Their major concern was to reduce the adverse effects on wildlife by snowmobiles. This rule does that by limiting the numbers of snowmobiles allowed and authorizing such use only in areas where wildlife is not abundant during the winter.

Information Quality Act

In developing this rule NPS did not conduct or use a study, experiment, or survey requiring peer review under the Information Quality Act (Pub. L. 106–554).

Effects on the Energy Supply (E.O. 13211)

This rule is not a significant energy action under the definition in Executive Order 13211. A Statement of Energy Effects is not required.

Administrative Procedure Act

Comment periods on the proposed rule were provided from November 5, 2008, through November 20, 2008, and from July 24, 2009, to September 8, 2009, for a total of 60 days.

This rule is effective on December 15, 2009. The National Park Service recognizes that new rules ordinarily go into effect thirty days after publication in the **Federal Register**. For this regulation, however, we have determined under 5 U.S.C. 553(d) and 318 DM 6.25 that this rule should be effective on December 15, 2009, the traditional date for commencement of the park's winter use season. This rule implements the winter use plans for

Grand Teton National Park and the John D. Rockefeller, Jr. Memorial Parkway and relieves the restrictions on the use of snowmobiles that would exist in its absence. In addition, good cause exists for the effective date of December 15, 2009, for the following reasons:

(1) The NPS has in good faith publicly stated that the 2009–2010 winter season for Grand Teton National Park and the John D. Rockefeller, Jr. Memorial Parkway would commence on December 15, 2009, and the public and businesses have made decisions based on the widespread public knowledge of this opening date.

(2) The finding of no significant impact for this rule was signed on October 15, and was made available to the public for 30 days prior to the signing of this rule. By December 15, the public therefore will have had more than 60 days notice of the NPS decision.

(3) There would be no benefit to the public in delaying the effective date of this rule, given that there has already been substantial notice of the opening date and that the park will be open under conditions substantially similar to those in effect for the past three years. The above-described harms to the public resulting from a procedural delay of this rule should therefore be avoided, and an effective date of December 15, 2009, is warranted.

Drafting Information: The primary authors of this regulation are Gary Pollock, Management Assistant, Grand Teton National Park; John Sacklin, Management Assistant, Yellowstone National Park; Jason Waanders, Office of the Solicitor, and Philip Selleck, Regulations Program Manager, National Park Service, Washington, DC.

List of Subjects in 36 CFR Part 7

District of Columbia, National Parks, Reporting and recordkeeping requirements.

■ For the reasons given in the preamble, 36 CFR Part 7 is amended as set forth below:

PART 7—SPECIAL REGULATIONS, AREAS OF THE NATIONAL PARK SYSTEM

■ 1. The authority for Part 7 continues to read as follows:

Authority: 16 U.S.C. 1, 3, 9a, 462(k); Sec. 7.96 also issued under DC Code 10–137(2001) and DC Code 50–2201 (2001).

■ 2. Revise § 7.21 to read as follows:

§ 7.21 John D. Rockefeller, Jr. Memorial Parkway.

(a)(1) *What is the scope of this section?* The regulations contained in paragraphs (a)(2) through (a)(17) of this

section apply to the use of snowcoaches and recreational snowmobiles. Except where indicated, paragraphs (a)(2) through (a)(15) do not apply to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(2) *What terms do I need to know?* The definitions in this paragraph (a)(2) also apply to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, and other non-recreational users authorized by the Superintendent.

Commercial guide means a guide who operates a snowmobile or snowcoach for a fee or compensation and is authorized to operate in the park under a concession contract. In this section, “guide” also means “commercial guide.”

Historic snowcoach means a Bombardier snowcoach manufactured in 1983 or earlier. Any other snowcoach is considered a non-historic snowcoach.

Oversnow route means that portion of the unplowed roadway located between the road shoulders and designated by snow poles or other poles, ropes, fencing, or signs erected to regulate oversnow activity. Oversnow routes include pullouts or parking areas that are groomed or marked similarly to roadways and are adjacent to designated oversnow routes. An oversnow route may also be distinguished by the interior boundaries of the berm created by the packing and grooming of the unplowed roadway. The only motorized vehicles permitted on oversnow routes are oversnow vehicles.

Oversnow vehicle means a snowmobile, snowcoach, or other motorized vehicle that is intended for travel primarily on snow and has been authorized by the Superintendent to operate in the park. An oversnow vehicle that does not meet the definition of a snowcoach must comply with all requirements applicable to snowmobiles.

Snowcoach means a self-propelled mass transit vehicle intended for travel on snow, having a curb weight of over 1,000 pounds (450 kilograms), driven by a track or tracks and steered by skis or tracks, and having a capacity of at least 8 passengers. A snowcoach has a maximum size of 102 inches wide, plus tracks (not to exceed 110 inches overall); a maximum length of 35 feet; and a Gross Vehicle Weight Rating (GVWR) not exceeding 25,000 pounds.

Snowmobile means a self-propelled vehicle intended for travel on snow, with a curb weight of not more than 1,000 pounds (450 kg), driven by a track or tracks in contact with the snow, and

which may be steered by a ski or skis in contact with the snow.

Snowplane means a self-propelled vehicle intended for oversnow travel and driven by an air-displacing propeller.

(3) *May I operate a snowmobile in the Parkway?* You may operate a snowmobile in the Parkway in compliance with use limits, guiding requirements, operating hours and dates, equipment, and operating conditions established under this section. The Superintendent may establish additional operating conditions and will provide notice of those conditions in accordance with § 1.7(a) of this chapter or in the **Federal Register**.

(4) *May I operate a snowcoach in the Parkway?* Snowcoaches may only be operated in the Parkway under a concessions contract. Snowcoach operation is subject to the conditions stated in the concessions contract and all other conditions identified in this section.

(5) *Where may I operate my snowmobile in the Parkway?* (i) You may operate your snowmobile only upon designated oversnow routes established within the Parkway in accordance with § 2.18(c) of this chapter. The following oversnow routes are so designated for snowmobile use:

(A) On U.S. Highway 89/191/287 from Flagg Ranch to the northern boundary of the Parkway.

(B) Grassy Lake Road from Flagg Ranch to the western boundary of the Parkway.

(C) Flagg Ranch developed area.

(ii) The Superintendent may open or close these routes, or portions thereof, for snowmobile travel after taking into consideration the location of wintering wildlife, appropriate snow cover, public safety, and other factors. The Superintendent will provide notice of such opening or closing by one or more of the methods listed in § 1.7(a) of this chapter.

(iii) The route described in paragraph (a)(5)(i)(A) of this section is subject to the air and sound emissions requirements, guiding requirements, and daily entry limits described in § 7.13(l) of this part.

(iv) This paragraph (a)(5) also applies to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(v) Maps detailing the designated oversnow routes will be available from Park Headquarters.

(6) *What routes are designated for snowcoach use?* (i) Authorized snowcoaches may only be operated on

the routes designated for snowmobile use in paragraphs (a)(6)(i)(A) and (C) of this section. No other routes are open to snowcoach use, except as provided in (a)(6)(ii) of this section.

(ii) The Superintendent may open or close these oversnow routes, or portions thereof, or designate new routes for snowcoach travel after taking into consideration the location of wintering wildlife, appropriate snow cover, public safety, and other factors. The Superintendent will provide notice of such opening or closing by one or more of the methods listed in § 1.7(a) of this chapter.

(iii) The routes described in paragraph (a)(6)(i) of this section are subject to the air and sound emissions requirements and daily entry limits in § 7.13(l) of this part.

(iv) This paragraph (a)(6) also applies to non-administrative snowcoach use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(7) *Must I travel with a commercial guide while snowmobiling in the Parkway?* Except as may be required under paragraph (a)(5)(iii) of this section, you are not required to use a guide while snowmobiling in the Parkway.

(8) *Are there limits established for the numbers of snowmobiles and snowcoaches permitted to operate in the Parkway each day?* (i) A limit of 25 snowmobiles per day applies to the Grassy Lake Road.

(ii) The daily entry limits for snowmobiles and snowcoaches on the route from Flagg Ranch to the South Entrance of Yellowstone are established in § 7.13(l) of this part.

(9) *When may I operate my snowmobile or snowcoach?* The Superintendent will determine operating hours and dates. Except for emergency situations, any changes to operating hours will be made on an annual basis and the public will be notified of those changes through one or more of the methods listed in § 1.7(a) of this chapter.

(10) *What other conditions apply to the operation of oversnow vehicles?* (i) The following are prohibited:

(A) Idling an oversnow vehicle more than 5 minutes at any one time.

(B) Driving an oversnow vehicle while the operator's motor vehicle license or privilege is suspended or revoked.

(C) Allowing or permitting an unlicensed driver to operate an oversnow vehicle.

(D) Driving an oversnow vehicle in willful or wanton disregard for the safety of persons, property, or parkway

resources or otherwise in a reckless manner.

(E) Operating an oversnow vehicle without a lighted white headlamp and red taillight.

(F) Operating an oversnow vehicle that does not have brakes in good working order.

(G) Towing persons on skis, sleds or other sliding devices by oversnow vehicles, except in emergency situations.

(ii) The following are required:

(A) All oversnow vehicles that stop on designated routes must pull over to the far right and next to the snow berm. Pullouts must be used where available and accessible. Oversnow vehicles may not be stopped in a hazardous location or where the view might be obscured, or operated so slowly as to interfere with the normal flow of traffic.

(B) Oversnow vehicle drivers must possess a valid motor vehicle driver's license. A learner's permit does not satisfy this requirement. The license must be carried by the driver at all times.

(C) Equipment sleds towed by a snowmobile must be pulled behind the snowmobile and fastened to the snowmobile with a rigid hitching mechanism.

(D) Snowmobiles must be properly registered and display a valid registration from the United States or Canada.

(iii) The Superintendent may impose other terms and conditions as necessary to protect park resources, visitors, or employees. The Superintendent will notify the public of any changes through one or more methods listed in § 1.7(a) of this chapter.

(iv) This paragraph (a)(10) also applies to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(11) *What conditions apply to alcohol use while operating an oversnow vehicle?* In addition to 36 CFR 4.23, the following conditions apply:

(i) Operating or being in actual physical control of an oversnow vehicle is prohibited when the driver is under 21 years of age and the alcohol concentration in the driver's blood or breath is 0.02 grams or more of alcohol per 100 milliliters of blood or 0.02 grams or more of alcohol per 210 liters of breath.

(ii) Operating or being in actual physical control of an oversnow vehicle is prohibited when the driver is a snowmobile guide or a snowcoach driver and the alcohol concentration in the operator's blood or breath is 0.04

grams or more of alcohol per 100 milliliters of blood or 0.04 grams or more of alcohol per 210 liters of breath.

(iii) This paragraph (a)(11) also applies to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(12) *Do other NPS regulations apply to the use of oversnow vehicles?* (i) The use of oversnow vehicles in the Parkway is subject to § 2.18(a), (b), and (c), but not to §§ 2.18(d), (e), and 2.19(b) of this chapter.

(ii) This paragraph (a)(12) also applies to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(13) *Are there any forms of non-motorized oversnow transportation allowed in the Parkway?* (i) Non-motorized travel consisting of skiing, skating, snowshoeing, or walking is permitted unless otherwise restricted under this section or other NPS regulations.

(ii) The Superintendent may designate areas of the Parkway as closed, reopen such areas, or establish terms and conditions for non-motorized travel within the Parkway in order to protect visitors, employees, or park resources. Notice will be made in accordance with § 1.7(a) of this chapter.

(14) *May I operate a snowplane in the Parkway?* The operation of a snowplane in the Parkway is prohibited.

(15) *Is violating any of the provisions of this section prohibited?* (i) Violating any of the terms, conditions or requirements of paragraphs (a)(3) through (a)(14) of this section is prohibited.

(ii) Anyone who violates any of the terms, conditions or requirements of this regulation will be considered to have committed one separate offense for each term, condition or requirement that they violate.

(b) [Reserved]

■ 3. Amend § 7.22, by revising paragraph (g) to read as follows:

§ 7.22 Grand Teton National Park.

* * * * *

(g)(1) *What is the scope of this section?* The regulations contained in paragraphs (g)(2) through (g)(20) of this section are intended to apply to the use of snowcoaches and recreational snowmobiles. Except where indicated, paragraphs (g)(2) through (g)(20) do not apply to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(2) *What terms do I need to know?* The definitions in this paragraph (g)(2) also apply to non-administrative oversnow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(i) *Commercial guide* means a guide who operates as a snowmobile or snowcoach guide for a fee or compensation and is authorized to operate in the park under a concession contract. In this section, "guide" also means "commercial guide."

(ii) *Historic snowcoach* means a Bombardier snowcoach manufactured in 1983 or earlier. Any other snowcoach is considered a non-historic snowcoach.

(iii) *Oversnow route* means that portion of the unplowed roadway located between the road shoulders and designated by snow poles or other poles, ropes, fencing, or signs erected to regulate oversnow activity. Oversnow routes include pullouts or parking areas that are groomed or marked similarly to roadways and are adjacent to designated oversnow routes. An oversnow route may also be distinguished by the interior boundaries of the berm created by the packing and grooming of the unplowed roadway. The only motorized vehicles permitted on oversnow routes are oversnow vehicles.

(iv) *Oversnow vehicle* means a snowmobile, snowcoach, or other motorized vehicle that is intended for travel primarily on snow and has been authorized by the Superintendent to operate in the park. An oversnow vehicle that does not meet the definition of a snowcoach must comply with all requirements applicable to snowmobiles.

(v) *Snowcoach* means a self-propelled mass transit vehicle intended for travel on snow, having a curb weight of over 1,000 pounds (450 kilograms), driven by a track or tracks and steered by skis or tracks, and having a capacity of at least 8 passengers. A snowcoach has a maximum size of 102 inches wide, plus tracks (not to exceed 110 inches overall); a maximum length of 35 feet; and a Gross Vehicle Weight Rating (GVWR) not exceeding 25,000 pounds.

(vi) *Snowmobile* means a self-propelled vehicle intended for travel on snow, with a curb weight of not more than 1,000 pounds (450 kg), driven by a track or tracks in contact with the snow, and which may be steered by a ski or skis in contact with the snow.

(vii) *Snowplane* means a self-propelled vehicle intended for oversnow travel and driven by an air-displacing propeller.

(3) *May I operate a snowmobile in Grand Teton National Park?* You may

operate a snowmobile in Grand Teton National Park in compliance with use limits, operating hours and dates, equipment, and operating conditions established under this section. The Superintendent may establish additional operating conditions and provide notice of those conditions in accordance with § 1.7(a) of this chapter or in the **Federal Register**.

(4) *May I operate a snowcoach in Grand Teton National Park?* It is prohibited to operate a snowcoach in Grand Teton National Park except as authorized by the Superintendent.

(5) *Must I operate a certain model of snowmobile in the park?* Only commercially available snowmobiles that meet NPS air and sound emissions requirements as set forth in this section may be operated in the park. The Superintendent will approve snowmobile makes, models, and years of manufacture that meet those requirements. Any snowmobile model not approved by the Superintendent may not be operated in the park.

(6) *How will the Superintendent approve snowmobile makes, models, and years of manufacture for use in Grand Teton National Park?* (i) Beginning with the 2005 model year, all snowmobiles must be certified under 40 CFR Part 1051, to a Family Emission Limit no greater than 15 g/kW-hr for hydrocarbons and to a Family Emission Limit no greater than 120 g/kW-hr for carbon monoxide.

(A) 2004 model year snowmobiles may use measured air emissions levels (official emission results with no deterioration factors applied) to comply with the air emission limits specified in paragraph (g)(6)(i) of this section.

(B) Snowmobiles manufactured before the 2004 model year may be operated only if they have shown to have air emissions no greater than the requirements identified in paragraph (g)(6)(i) of this section.

(C) The snowmobile test procedures specified by EPA (40 CFR parts 1051 and 1065) must be used to measure air emissions from model year 2004 and later snowmobiles. Equivalent procedures may be used for earlier model years.

(ii) For sound emissions, snowmobiles must operate at or below 73 dBA as measured at full throttle according to Society of Automotive Engineers J192 test procedures (revised 1985). Snowmobiles may be tested at any barometric pressure equal to or above 23.4 inches Hg uncorrected. The Superintendent may revise these testing procedures based on new information and/or updates to the SAE J192 testing procedures.

(iii) Snowmobiles meeting the requirements for air and sound emissions may be operated in the park for a period not exceeding 6 years from the date upon which first certified, except that snowmobiles being operated on Jackson Lake may continue to be operated up to 10 years, provided that these snowmobiles' mileage does not exceed 6,000 miles.

(iv) Snowmobiles will be exempt from these air and sound emissions requirements while in use to access lands authorized by paragraphs (g)(16) and (g)(18) of this section.

(v) The Superintendent may prohibit entry into the park of any snowmobile that has been modified in a manner that may adversely affect air or sound emissions.

(7) *Where may I operate my snowmobile in the park?* (i) You may operate your snowmobile upon the frozen water surface of Jackson Lake, a route established in accordance with § 2.18(c) of this chapter, under the following conditions:

(A) You are ice fishing, and licensed or otherwise permitted to fish in Wyoming;

(B) You possess the proper fishing gear; and

(C) You limit your snowmobile travel to a direct route to and from and between fishing locations on the lake.

(ii) The Superintendent may open or close this route, or portions thereof, for snowmobile travel, and may establish separate zones for motorized and non-motorized uses on Jackson Lake, after taking into consideration the location of wintering wildlife, appropriate snow cover, public safety and other factors. The Superintendent will provide notice of such opening or closing by one or more of the methods listed in § 1.7(a) of this chapter.

(iii) This paragraph (g)(7) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(iv) Maps detailing the designated oversnow route will be available from Park Headquarters.

(8) *Must I travel with a commercial guide while snowmobiling in Grand Teton National Park?* You are not required to use a guide while snowmobiling in Grand Teton National Park.

(9) *Are there limits established for the number of snowmobiles permitted to operate in the park each day?* (i) The number of snowmobiles allowed to operate in the park each day on Jackson Lake is 25.

(ii) The Superintendent may adjust this number up or down, not to exceed

a daily limit of 40 snowmobiles, after taking into consideration the location of wintering wildlife, appropriate snow cover, noise monitoring results, public safety and other factors. The Superintendent will provide notice of such adjustment by one or more of the methods listed in § 1.7(a) of this chapter.

(10) *When may I operate my snowmobile?* The Superintendent will determine operating hours and dates. Except for emergency situations, any changes to operating hours or dates will be made on an annual basis, and the public will be notified of those changes through one or more of the methods listed in § 1.7(a) of this chapter.

(11) *What other conditions apply to the operation of oversnow vehicles?* (i) The following are prohibited:

(A) Idling an oversnow vehicle more than 5 minutes at any one time.

(B) Driving an oversnow vehicle while the operator's motor vehicle license or privilege is suspended or revoked.

(C) Allowing or permitting an unlicensed driver to operate an oversnow vehicle.

(D) Driving an oversnow vehicle in willful or wanton disregard for the safety of persons, property, or park resources or otherwise in a reckless manner.

(E) Operating an oversnow vehicle without a lighted white headlamp and red taillight.

(F) Operating an oversnow vehicle that does not have brakes in good working order.

(G) The towing of persons on skis, sleds or other sliding devices by oversnow vehicles.

(ii) The following are required:

(A) All oversnow vehicles that stop on designated routes must pull over to the far right and next to the snow berm. Pullouts must be used where available and accessible. Oversnow vehicles may not be stopped in a hazardous location or where the view might be obscured, or operated so slowly as to interfere with the normal flow of traffic.

(B) Oversnow vehicle drivers must possess a valid motor vehicle driver's license. A learner's permit does not satisfy this requirement. The license must be carried by the driver at all times.

(C) Equipment sleds towed by a snowmobile must be pulled behind the snowmobile and fastened to the snowmobile with a rigid hitching mechanism.

(D) Snowmobiles must be properly registered and display a valid registration from the United States or Canada.

(iii) The Superintendent may impose other terms and conditions as necessary to protect park resources, visitors, or employees. The Superintendent will notify the public of any changes through one or more methods listed in § 1.7(a) of this chapter.

(iv) This paragraph (g)(11) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(12) *What conditions apply to alcohol use while operating an oversnow vehicle?* In addition to 36 CFR 4.23, the following conditions apply:

(i) Operating or being in actual physical control of an oversnow vehicle is prohibited when the driver is under 21 years of age and the alcohol concentration in the driver's blood or breath is 0.02 grams or more of alcohol per 100 milliliters of blood or 0.02 grams or more of alcohol per 210 liters of breath.

(ii) Operating or being in actual physical control of an oversnow vehicle is prohibited when the driver is a snowmobile guide or a snowcoach operator and the alcohol concentration in the driver's blood or breath is 0.04 grams or more of alcohol per 100 milliliters of blood or 0.04 grams or more of alcohol per 210 liters of breath.

(iii) This paragraph (g)(12) also applies to non-administrative over-snow vehicle use by NPS, contractor, or concessioner employees, or other non-recreational users authorized by the Superintendent.

(13) *Do other NPS regulations apply to the use of oversnow vehicles?* The use of oversnow vehicles in Grand Teton is subject to § 2.18(a), (b), and (c), but not subject to § 2.18(d) and (e) and § 2.19(b) of this chapter.

(14) *Are there any forms of non-motorized oversnow transportation allowed in the park?*

(i) Non-motorized travel consisting of skiing, skating, snowshoeing, or walking is permitted unless otherwise restricted under this section or other NPS regulations.

(ii) The Superintendent may designate areas of the park as closed, reopen such areas, or establish terms and conditions for non-motorized travel within the park in order to protect visitors, employees, or park resources.

(iii) Dog sledding and ski-joring are prohibited.

(15) *May I operate a snowplane in the park?* The operation of a snowplane in Grand Teton National Park is prohibited.

(16) *May I continue to access public lands via snowmobile through the park?*

Reasonable and direct access, via snowmobile, to adjacent public lands will continue to be permitted on the designated routes through the park identified in the following paragraphs (g)(16)(i) through (iv). Requirements established in this section related to air and sound emissions, daily entry limits, snowmobile operator age, guiding, and licensing do not apply on these oversnow routes. The following routes are designated for access via snowmobile to public lands:

(i) From the parking area at Shadow Mountain directly along the unplowed portion of the road to the east park boundary.

(ii) Along the unplowed portion of the Ditch Creek Road directly to the east park boundary.

(iii) The Continental Divide Snowmobile Trail (CDST) along U.S. 26/287 from the east park boundary to a point approximately 2 miles east of Moran Junction. If necessary for the proper administration of visitor use and resource protection, the Superintendent may extend this designated route to the Moran Entrance Station.

(iv) The Superintendent may designate additional routes if necessary to provide access to other adjacent public lands.

(17) *For what purpose may I use the routes designated in paragraph (g)(16) of this section?* You may only use those routes designated in paragraph (g)(16) of this section to gain direct access to public lands adjacent to the park boundary.

(18) *May I continue to access private property within or adjacent to the park via snowmobile?* The Superintendent may establish reasonable and direct snowmobile access routes to the inholding or to private property adjacent to park boundaries for which other routes or means of access are not reasonably available. Requirements established in this section related to air and sound emissions, snowmobile operator age, licensing, and guiding do not apply on these oversnow routes. The following routes are designated for access to private properties within or adjacent to the park:

(i) From the Antelope Flats Road off U.S. 26/89/191 to private lands in the Craighead Subdivision.

(ii) The unplowed portion of the Teton Park Road to the piece of land commonly referred to as the "Townsend Property."

(iii) From the Moose-Wilson Road to the land commonly referred to as the "Barker Property."

(iv) From the Moose-Wilson Road to the property commonly referred to as the "Halpin Property."

(v) From Highway 26/89/191 to those lands commonly referred to as the "Meadows", the "Circle EW Ranch", the "Moulton Property", the "Levinson Property" and the "Macmahon Property."

(vi) From Cunningham Cabin pullout on U.S. 26/89/191 near Triangle X to the piece of land commonly referred to as the "Lost Creek Ranch."

(vii) The Superintendent may designate additional routes if necessary to provide reasonable access to inholdings or adjacent private property.

(viii) Maps detailing designated routes will be available from Park Headquarters.

(19) *For what purpose may I use the routes designated in paragraph (g)(18) of this section?* The routes designated in paragraph (g)(18) of this section are only to access private property within or directly adjacent to the park boundary. Use of these roads via snowmobile is authorized only for the landowners and their representatives or guests. Use of these roads by anyone else or for any other purpose is prohibited.

(20) *Is violating any of the provisions of this section prohibited?* (i) Violating any of the terms, conditions or requirements of paragraphs (g)(3) through (g)(19) of this section is prohibited.

(ii) Anyone who violates any of the terms, conditions or requirements of this regulation will be considered to have committed one separate offense for each term, condition or requirement that they violate.

Dated: November 16, 2009.

Thomas L Strickland,

Assistant Secretary for Fish and Wildlife and Parks.

[FR Doc. E9-27894 Filed 11-17-09; 4:15 pm]

BILLING CODE 4310-CX-P

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R03-OAR-2009-0674; FRL-8983-1]

Approval and Promulgation of Air Quality Implementation Plans; Virginia; Transportation Conformity Regulations

AGENCY: Environmental Protection Agency (EPA).

ACTION: Direct final rule.

SUMMARY: EPA is taking direct final action to approve revisions to the State Implementation Plan (SIP) submitted by the Commonwealth of Virginia. This revision establishes Virginia's transportation conformity requirements.

After they have been approved, the Commonwealth's regulations will govern transportation conformity determinations in the Commonwealth of Virginia. EPA is approving these revisions in accordance with the requirements of the Clean Air Act (CAA).

DATES: This rule is effective on January 19, 2010 without further notice, unless EPA receives adverse written comment by December 21, 2009. If EPA receives such comments, it will publish a timely withdrawal of the direct final rule in the **Federal Register** and inform the public that the rule will not take effect.

ADDRESSES: Submit your comments, identified by Docket ID Number EPA-R03-OAR-2009-0674 by one of the following methods:

A. *http://www.regulations.gov.* Follow the on-line instructions for submitting comments.

B. *E-mail:*

fernandez.cristina@epa.gov.

C. *Mail:* EPA-R03-OAR-2009-0674, Cristina Fernandez, Associate Director, Office of Air Program Planning, Mailcode 3AP30, U.S. Environmental Protection Agency, Region III, 1650 Arch Street, Philadelphia, Pennsylvania 19103.

D. *Hand Delivery:* At the previously-listed EPA Region III address. Such deliveries are only accepted during the Docket's normal hours of operation, and special arrangements should be made for deliveries of boxed information.

Instructions: Direct your comments to Docket ID No. EPA-R03-OAR-2009-0674. EPA's policy is that all comments received will be included in the public docket without change, and may be made available online at *http://www.regulations.gov*, including any personal information provided, unless the comment includes information claimed to be Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. Do not submit information that you consider to be CBI or otherwise protected through *http://www.regulations.gov* or e-mail. The *http://www.regulations.gov* Web site is an anonymous access system, which means EPA will not know your identity or contact information unless you provide it in the body of your comment. If you send an e-mail comment directly to EPA without going through *http://www.regulations.gov*, your e-mail address will be automatically captured and included as part of the comment that is placed in the public docket and made available on the Internet. If you submit an electronic comment, EPA recommends that you include your

name and other contact information in the body of your comment and with any disk or CD-ROM you submit. If EPA cannot read your comment due to technical difficulties and cannot contact you for clarification, EPA may not be able to consider your comment. Electronic files should avoid the use of special characters, any form of encryption, and be free of any defects or viruses.

Docket: All documents in the electronic docket are listed in the <http://www.regulations.gov> index. Although listed in the index, some information is not publicly available, i.e., CBI or other information whose disclosure is restricted by statute. Certain other material, such as copyrighted material, is not placed on the Internet and will be publicly available only in hard copy form. Publicly available docket materials are available either electronically in <http://www.regulations.gov> or in hard copy during normal business hours at the Air Protection Division, U.S. Environmental Protection Agency, Region III, 1650 Arch Street, Philadelphia, Pennsylvania 19103. Copies of the State submittal are available at the Virginia Department of Environmental Quality, 629 East Main Street, Richmond, Virginia 23219.

FOR FURTHER INFORMATION CONTACT: Martin Kotsch, (215) 814-3335 or by e-mail at kotsch.martin@epa.gov.

SUPPLEMENTARY INFORMATION: Throughout this document whenever "we," "us," or "our" is used, we mean EPA.

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I. What Is Transportation Conformity?

Transportation conformity is required under section 176(c) of the Clean Air Act to ensure that Federally supported highway, transit projects, and other activities are consistent with (conform to) the purpose of the SIP. Conformity currently applies to areas that are designated nonattainment, and those redesignated to attainment after 1990 (maintenance areas), with plans developed under section 175A of the Clean Air Act for the following transportation related criteria pollutants: Ozone, particulate matter (PM_{2.5} and PM₁₀), carbon monoxide (CO), and nitrogen dioxide (NO₂).

Conformity to the purpose of the SIP means that transportation activities will not cause new air quality violations, worsen existing violations, or delay timely attainment of the relevant national ambient air quality standards (NAAQS). The transportation conformity regulation is found in 40 CFR part 93 and provisions related to conformity SIPs are found in 40 CFR 51.390.

II. What Is the Background for This Action?

On August 10, 2005, the Safe, Accountable, Flexible, Efficient, Transportation Equity Act: A Legacy for Users (SAFETEA-LU) was signed into law. SAFETEA-LU revised certain provisions of section 176(c) of the Clean Air Act, related to transportation conformity. Prior to SAFETEA-LU, states were required to address all of the Federal conformity rule's provisions in their conformity SIPs. After SAFETEA-LU, state SIPs were required to contain all or portions of only the following three sections of the Federal rule, modified as appropriate to each state's circumstances: 40 CFR 93.105 (consultation procedures); 40 CFR 93.122(a)(4)(ii) (written commitments to implement certain kinds of control measures); and 40 CFR 93.125(c) (written commitments to implement certain kinds of mitigation measures). States are no longer required to submit conformity SIP revisions that address the other sections of the Federal conformity rule.

III. What Did the State Submit and How Did We Evaluate It?

On July 9, 2007, the Virginia Department of Environmental Quality (VADEQ) submitted a revision to its SIP for Transportation Conformity purposes. The SIP revision consists of the State Regulation for Transportation Conformity (9 VAC 5 Chapter 151). This SIP revision addresses the three provisions of the EPA Conformity Rule required under SAFETEA-LU: 40 CFR 93.105 (consultation procedures); 40 CFR 93.122(a)(4)(ii) (control measures) and 40 CFR 93.125(c) (mitigation measures).

We reviewed the submittal to assure consistency with the February 14, 2006 "Interim Guidance for Implementing the Transportation Conformity provisions in the Safe, Accountable, Flexible, Efficient, Transportation Equity Act: A Legacy for Users (SAFETEA-LU)." The guidance document can be found at <http://epa.gov/otaq/stateresources/transconf/policy.htm>. The guidance document states that each state is only required to address and tailor the afore-

mentioned three sections of the Federal Conformity Rule to be included in their state conformity SIPs.

EPA's review of Virginia's proposed SIP indicates that it is consistent with EPA's guidance in that it includes the three elements specified by SAFETEA-LU. Consistent with the EPA Conformity Rule at 40 CFR 93.105 (consultation procedures), Regulation 9 VAC 5 Chapter 151-70 identifies the appropriate agencies, procedures and allocation of responsibilities as required under 40 CFR 93.105 for consultation procedures. In addition, Regulation 9 VAC 5 Chapter 151-50 and Regulation 9 VAC 5 Chapter 151-60 provide for appropriate public consultation/public involvement consistent with 40 CFR 93.105. With respect to the requirements of 40 CFR 93.122(a)(4)(ii) and 40 CFR 93.125(c), Regulation 9 VAC 5 Chapter 151-50 also specifies that written commitments for control measures and mitigation measures for meeting these requirements will be provided as needed.

IV. General Information Pertaining to SIP Submittals From the Commonwealth of Virginia

In 1995, Virginia adopted legislation that provides, subject to certain conditions, for an environmental assessment (audit) "privilege" for voluntary compliance evaluations performed by a regulated entity. The legislation further addresses the relative burden of proof for parties either asserting the privilege or seeking disclosure of documents for which the privilege is claimed. Virginia's legislation also provides, subject to certain conditions, for a penalty waiver for violations of environmental laws when a regulated entity discovers such violations pursuant to a voluntary compliance evaluation and voluntarily discloses such violations to the Commonwealth and takes prompt and appropriate measures to remedy the violations. Virginia's Voluntary Environmental Assessment Privilege Law, Va. Code Section 10.1-1198, provides a privilege that protects from disclosure documents and information about the content of those documents that are the product of a voluntary environmental assessment. The Privilege Law does not extend to documents or information (1) That are generated or developed before the commencement of a voluntary environmental assessment; (2) that are prepared independently of the assessment process; (3) that demonstrate a clear, imminent and substantial danger to the public health or

environment; or (4) that are required by law.

On January 12, 1998, the Commonwealth of Virginia Office of the Attorney General provided a legal opinion that states that the Privilege Law, Va. Code Section 10.1-1198, precludes granting a privilege to documents and information "required by law," including documents and information "required by Federal law to maintain program delegation, authorization or approval," since Virginia must "enforce Federally authorized environmental programs in a manner that is no less stringent than their Federal counterparts * * *." The opinion concludes that "[r]egarding 10.1-1198, therefore, documents or other information needed for civil or criminal enforcement under one of these programs could not be privileged because such documents and information are essential to pursuing enforcement in a manner required by Federal law to maintain program delegation, authorization or approval."

Virginia's Immunity law, Va. Code Section 10.1-1199, provides that "[t]o the extent consistent with requirements imposed by Federal law," any person making a voluntary disclosure of information to a state agency regarding a violation of an environmental statute, regulation, permit, or administrative order is granted immunity from administrative or civil penalty. The Attorney General's January 12, 1998 opinion states that the quoted language renders this statute inapplicable to enforcement of any Federally authorized programs, since "no immunity could be afforded from administrative, civil, or criminal penalties because granting such immunity would not be consistent with Federal law, which is one of the criteria for immunity."

Therefore, EPA has determined that Virginia's Privilege and Immunity statutes will not preclude the Commonwealth from enforcing its program consistent with the Federal requirements. In any event, because EPA has also determined that a state audit privilege and immunity law can affect only state enforcement and cannot have any impact on Federal enforcement authorities, EPA may at any time invoke its authority under the CAA, including, for example, sections 113, 167, 205, 211 or 213, to enforce the requirements or prohibitions of the state plan, independently of any state enforcement effort. In addition, citizen enforcement under section 304 of the CAA is likewise unaffected by this, or any, state audit privilege or immunity law.

V. What Action Is EPA Taking Today?

EPA is approving the Virginia SIP revision for Transportation Conformity, which was submitted on July 9, 2007. EPA is publishing this rule without prior proposal because the Agency views this as a noncontroversial amendment and anticipates no adverse comment. However, in the Proposed Rules section of today's **Federal Register**, EPA is publishing a separate document that will serve as the proposal to approve the SIP revision if adverse comments are filed. This rule will be effective on January 19, 2010 without further notice unless EPA receives adverse comment by December 21, 2009. If EPA receives adverse comment, EPA will publish a timely withdrawal in the **Federal Register** informing the public that the rule will not take effect. EPA will address all public comments in a subsequent final rule based on the proposed rule. EPA will not institute a second comment period on this action. Any parties interested in commenting must do so at this time. Please note that if EPA receives adverse comment on an amendment, paragraph, or section of this rule and if that provision may be severed from the remainder of the rule, EPA may adopt as final those provisions of the rule that are not the subject of an adverse comment.

VI. Statutory and Executive Order Reviews

A. General Requirements

Under the Clean Air Act, the Administrator is required to approve a SIP submission that complies with the provisions of the Act and applicable Federal regulations 42 U.S.C. 7410(k); 40 CFR 52.02(a). Thus, in reviewing SIP submissions, EPA's role is to approve state choices, provided that they meet the criteria of the Clean Air Act. Accordingly, this action merely approves state law as meeting Federal requirements and does not impose additional requirements beyond those imposed by state law. For that reason, this action:

- Is not a "significant regulatory action" subject to review by the Office of Management and Budget under Executive Order 12866 (58 FR 51735, October 4, 1993);
- Does not impose an information collection burden under the provisions of the Paperwork Reduction Act (44 U.S.C. 3501 *et seq.*);
- Is certified as not having a significant economic impact on a substantial number of small entities under the Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*);

- Does not contain any unfunded mandate or significantly or uniquely affect small governments, as described in the Unfunded Mandates Reform Act of 1995 (Pub. L. 104-4);

- Does not have Federalism implications as specified in Executive Order 13132 (64 FR 43255, August 10, 1999);

- Is not an economically significant regulatory action based on health or safety risks subject to Executive Order 13045 (62 FR 19885, April 23, 1997);

- Is not a significant regulatory action subject to Executive Order 13211 (66 FR 28355, May 22, 2001);

- Is not subject to requirements of section 12(d) of the National Technology Transfer and Advancement Act of 1995 (15 U.S.C. 272 note) because application of those requirements would be inconsistent with the Clean Air Act; and

- Does not provide EPA with the discretionary authority to address, as appropriate, disproportionate human health or environmental effects, using practicable and legally permissible methods, under Executive Order 12898 (59 FR 7629, February 16, 1994).

In addition, this rule does not have tribal implications as specified by Executive Order 13175 (65 FR 67249, November 9, 2000), because the SIP is not approved to apply in Indian country located in the state, and EPA notes that it will not impose substantial direct costs on tribal governments or preempt tribal law.

B. Submission to Congress and the Comptroller General

The Congressional Review Act, 5 U.S.C. 801 *et seq.*, as added by the Small Business Regulatory Enforcement Fairness Act of 1996, generally provides that before a rule may take effect, the agency promulgating the rule must submit a rule report, which includes a copy of the rule, to each House of the Congress and to the Comptroller General of the United States. EPA will submit a report containing this action and other required information to the U.S. Senate, the U.S. House of Representatives, and the Comptroller General of the United States prior to publication of the rule in the **Federal Register**. A major rule cannot take effect until 60 days after it is published in the **Federal Register**. This action is not a "major rule" as defined by 5 U.S.C. 804(2).

C. Petitions for Judicial Review

Under section 307(b)(1) of the CAA, petitions for judicial review of this action must be filed in the United States Court of Appeals for the appropriate circuit by *January 19, 2010*. Filing a

petition for reconsideration by the Administrator of this final rule does not affect the finality of this rule for the purposes of judicial review nor does it extend the time within which a petition for judicial review may be filed, and shall not postpone the effectiveness of such rule or action. This action to approve the Virginia Transportation Conformity Regulation may not be challenged later in proceedings to enforce its requirements. (See, section 307(b)(2)).

List of Subjects in 40 CFR Part 52

Environmental protection, Air pollution control, Carbon monoxide, Incorporation by reference, Intergovernmental relations, Nitrogen dioxide, Ozone, Particulate matter, Reporting and recordkeeping requirements, Volatile organic compounds.

Dated: November 5, 2009.

William C. Early,
Acting Regional Administrator, Region III.

■ 40 CFR part 52 is amended as follows:

PART 52—[AMENDED]

■ 1. The authority citation for 40 CFR part 52 continues to read as follows:
Authority: 42 U.S.C. 7401 *et seq.*

Subpart VV—Virginia

■ 2. In § 52.2420, the table in paragraph (c) is amended by adding an entry for Chapter 151 after the existing Chapter 140 to read as follows:

52.2420 Identification of plan.

* * * * *
(c) * * *

EPA-APPROVED VIRGINIA REGULATIONS AND STATUTES

State citation (9 VAC 5)	Title/subject	State effective date	EPA approval date	Explanation [former SIP citation]
*	*	*	*	*
Chapter 151 Transportation Conformity				
Part I General Definitions				
5-151-10	Definitions	12/31/08	11/20/09	[Insert page number where the document begins].
Part II General Provisions				
5-151-20	Applicability	12/31/08	11/20/09	[Insert page number where the document begins].
5-151-30	Authority of Board and DEQ.	12/31/08	11/20/09	[Insert page number where the document begins].
Part III Criteria and Procedures for Making Conformity Determinations				
5-151-40	General	12/31/08	11/20/09	[Insert page number where the document begins].
5-151-50	Designated provisions	12/31/08	11/20/09	[Insert page number where the document begins].
5-151-60	Word or phrase substitutions.	12/31/08	11/20/09	[Insert page number where the document begins].
5-151-70	Consultation	12/31/08	11/20/09	[Insert page number where the document begins].
*	*	*	*	*

* * * * *
[FR Doc. E9-27814 Filed 11-19-09; 8:45 am]
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ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R05-OAR-2009-0771; FRL-8980-4]

Approval and Promulgation of Air Quality Implementation Plans; Indiana

AGENCY: Environmental Protection Agency (EPA).

ACTION: Direct final rule.

SUMMARY: EPA is approving a request submitted by the Indiana Department of

Environmental Management (IDEM) on September 25, 2009, to revise the Indiana State Implementation Plan (SIP). The submission revises the Indiana Administrative Code (IAC) by amending and updating the definition of "References to the Code of Federal Regulations," to refer to the 2008 edition.

DATES: This rule is effective on January 19, 2010, unless EPA receives adverse written comments by December 21, 2009. If EPA receives adverse comments, EPA will publish a timely withdrawal of the rule in the **Federal Register** and inform the public that the rule will not take effect.

ADDRESSES: Submit your comments, identified by Docket ID No. EPA-R05-

OAR-2009-0771 by one of the following methods:

- <http://www.regulations.gov>: Follow the on-line instructions for submitting comments.
- *E-mail:* mooney.john@epa.gov.
- *Fax:* (312) 692-2551.
- *Mail:* John Mooney, Chief, Criteria Pollutant Section, Air Programs Branch (AR-18J), U.S. Environmental Protection Agency, 77 West Jackson Boulevard, Chicago, Illinois 60604.
- *Hand Delivery:* John Mooney, Chief, Criteria Pollutant Section, Air Programs Branch (AR-18J), U.S. Environmental Protection Agency, 77 West Jackson Boulevard, Chicago, Illinois 60604. Such deliveries are only accepted during the Regional Office normal hours of operation, and special arrangements

should be made for deliveries of boxed information. The Regional Office official hours of business are Monday through Friday, 8:30 a.m. to 4:30 p.m. excluding Federal holidays.

Instructions: Direct your comments to Docket ID No. EPA-R05-OAR-2009-0771. EPA's policy is that all comments received will be included in the public docket without change and may be made available online at <http://www.regulations.gov>, including any personal information provided, unless the comment includes information claimed to be Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. Do not submit information that you consider to be CBI or otherwise protected through <http://www.regulations.gov> or e-mail. The <http://www.regulations.gov> Web site is an "anonymous access" system, which means EPA will not know your identity or contact information unless you provide it in the body of your comment. If you send an e-mail comment directly to EPA without going through <http://www.regulations.gov> your e-mail address will be automatically captured and included as part of the comment that is placed in the public docket and made available on the Internet. If you submit an electronic comment, EPA recommends that you include your name and other contact information in the body of your comment and with any disk or CD-ROM you submit. If EPA cannot read your comment due to technical difficulties and cannot contact you for clarification, EPA may not be able to consider your comment. Electronic files should avoid the use of special characters, any form of encryption, and be free of any defects or viruses.

Docket: All documents in the docket are listed in the <http://www.regulations.gov> index. Although listed in the index, some information is not publicly available, e.g., CBI or other information whose disclosure is restricted by statute. Certain other material, such as copyrighted material, will be publicly available only in hard copy. Publicly available docket materials are available either electronically in <http://www.regulations.gov> or in hard copy at the Environmental Protection Agency, Region 5, Air and Radiation Division, 77 West Jackson Boulevard, Chicago, Illinois 60604. This Facility is open from 8:30 a.m. to 4:30 p.m., Monday through Friday, excluding legal holidays. We recommend that you telephone Charles Hatten, Environmental Engineer, at (312) 886-6031 before visiting the Region 5 office.

FOR FURTHER INFORMATION CONTACT:

Charles Hatten, Environmental Engineer, Criteria Pollutant Section, Air Programs Branch (AR-18J), Environmental Protection Agency, Region 5, 77 West Jackson Boulevard, Chicago, Illinois 60604, (312) 886-6031, hatten.charles@epa.gov.

SUPPLEMENTARY INFORMATION:

Throughout this document whenever "we," "us," or "our" is used, we mean EPA. This supplementary information section is arranged as follows:

- I. What is the background for this action?
 - A. When did the State submit the requested SIP revision to EPA?
 - B. Did Indiana hold public hearings on this SIP revision?
- II. What revision did the State request be incorporated into the SIP?
- III. What action is EPA taking today?
- IV. Statutory and Executive Order Reviews

I. What is the background for this action?

A. When did the State submit the requested SIP revision to EPA?

IDEM submitted the requested SIP revision on September 25, 2009.

B. Did Indiana hold public hearings on this SIP revision?

IDEM held public hearings on March 4, 2009. IDEM did not receive any public comments concerning the SIP revision.

II. What revision did the State request be incorporated into the SIP?

The State has requested that EPA approve revisions to 326 IAC 1-1-3 to update references to the Code of Federal Regulations (CFR) at 326 IAC 1-1-3.

Rule 326 IAC 1-1-3, definition of "References to Code of Federal Regulations." IDEM updated the reference to the CFR in 326 IAC 1-1-3 from the 2007 edition to the 2008 edition. This is solely an administrative change that allows Indiana to reference a more current version of the CFR.

III. What action is EPA taking today?

We are approving a revision to the Indiana SIP to update the definition at 326 IAC 1-1-3, "References to the CFR", to refer to the 2008 edition.

We are publishing this action without prior proposal because we view this as a noncontroversial amendment and anticipate no adverse comments.

However, in the proposed rules section of this **Federal Register** publication, we are publishing a separate document that will serve as the proposal to approve the state plan if relevant adverse written comments are filed. This rule will be effective January 19, 2010 without further notice unless we receive relevant

adverse written comments by *December 21, 2009*. If we receive such comments, we will withdraw this action before the effective date by publishing a subsequent document that will withdraw the final action. All public comments received will then be addressed in a subsequent final rule based on the proposed action. The EPA will not institute a second comment period. Any parties interested in commenting on this action should do so at this time. If we do not receive any comments, this action will be effective January 19, 2010.

IV. Statutory and Executive Order Reviews

Under the Clean Air Act, the Administrator is required to approve a SIP submission that complies with the provisions of the Clean Air Act and applicable Federal regulations. 42 U.S.C. 7410(k); 40 CFR 52.02(a). Thus, in reviewing SIP submissions, EPA's role is to approve state choices, provided that they meet the criteria of the Clean Air Act. Accordingly, this action merely approves state law as meeting Federal requirements and does not impose additional requirements beyond those imposed by state law. For that reason, this action:

- Is not a "significant regulatory action" subject to review by the Office of Management and Budget under Executive Order 12866 (58 FR 51735, October 4, 1993);

- Does not impose an information collection burden under the provisions of the Paperwork Reduction Act (44 U.S.C. 3501 *et seq.*);

- Is certified as not having a significant economic impact on a substantial number of small entities under the Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*);

- Does not contain any unfunded mandate or significantly or uniquely affect small governments, as described in the Unfunded Mandates Reform Act of 1995 (Pub. L. 104-4);

- Does not have Federalism implications as specified in Executive Order 13132 (64 FR 43255, August 10, 1999);

- Is not an economically significant regulatory action based on health or safety risks subject to Executive Order 13045 (62 FR 19885, April 23, 1997);

- Is not a significant regulatory action subject to Executive Order 13211 (66 FR 28355, May 22, 2001);

- Is not subject to requirements of Section 12(d) of the National Technology Transfer and Advancement Act of 1995 (15 U.S.C. 272 note) because application of those requirements would

be inconsistent with the Clean Air Act; and

- Does not provide EPA with the discretionary authority to address, as appropriate, disproportionate human health or environmental effects, using practicable and legally permissible methods, under Executive Order 12898 (59 FR 7629, February 16, 1994).

In addition, this rule does not have tribal implications as specified by Executive Order 13175 (65 FR 67249, November 9, 2000), because the SIP is not approved to apply in Indian country located in the state, and EPA notes that it will not impose substantial direct costs on tribal governments or preempt tribal law.

The Congressional Review Act, 5 U.S.C. 801 *et seq.*, as added by the Small Business Regulatory Enforcement Fairness Act of 1996, generally provides that before a rule may take effect, the agency promulgating the rule must submit a rule report, which includes a copy of the rule, to each House of the Congress and to the Comptroller General of the United States. EPA will submit a report containing this action and other required information to the U.S. Senate, the U.S. House of Representatives, and the Comptroller General of the United States prior to publication of the rule in the **Federal Register**. A major rule cannot take effect until 60 days after it is published in the **Federal Register**. This action is not a "major rule" as defined by 5 U.S.C. 804(2).

Under section 307(b)(1) of the Clean Air Act, petitions for judicial review of this action must be filed in the United States Court of Appeals for the appropriate circuit by January 19, 2010. Filing a petition for reconsideration by the Administrator of this final rule does not affect the finality of this action for the purposes of judicial review nor does it extend the time within which a petition for judicial review may be filed, and shall not postpone the effectiveness of such rule or action. This action may not be challenged later in proceedings to enforce its requirements. (See section 307(b)(2).)

List of Subjects in 40 CFR Part 52

Environmental protection, Air pollution control, Carbon monoxide, Incorporation by reference, Intergovernmental relations, Lead, Nitrogen dioxide, Ozone, Particulate matter, Reporting and recordkeeping requirements, Sulfur oxides, Volatile organic compounds.

Dated: November 3, 2009.

Bharat Mathur,

Acting Regional Administrator, Region 5.

■ For the reasons stated in the preamble, part 52, chapter I, of title 40 of the Code of Federal Regulations is amended as follows:

PART 52—[AMENDED]

■ 1. The authority citation for part 52 continues to read as follows:

Authority: 42 U.S.C. 7401 *et seq.*

Subpart P—Indiana

■ 2. Section 52.770 is amended by adding paragraph (c)(192) to read as follows:

§ 52.770 Identification of plan.

* * * * *

(c) * * *

(192) The Indiana Department of Environmental Management submitted a revision to Indiana's State Implementation plan on September 25, 2009, to amend 326 IAC 1-1-3, "References to the Code of Federal Regulations". The revision to 326 IAC 1-1-3 updates the references to CFR from the 2007 edition to the 2008 edition.

(i) *Incorporation by reference.* Title 326 of the Indiana Administrative Code (IAC), section 1-1-3, "References to the Code of Federal Regulations" is incorporated by reference. The rule was filed with the Publisher of the Indiana Register on July 1, 2009, and became effective on July 31, 2009. Published in the Indiana Register, on July 29, 2009 (DIN: 20090729-IR-326080901FRA).

[FR Doc. E9-27817 Filed 11-19-09; 8:45 am]

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ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R03-OAR-2009-0199; EPA-R03-OAR-2009-0547; FRL-8982-6]

Approval and Promulgation of Air Quality Implementation Plans; West Virginia; Maryland; Ohio; Determinations of Attainment for the 1997 Fine Particulate Matter Standard

AGENCY: Environmental Protection Agency (EPA).

ACTION: Final rule.

SUMMARY: EPA is making determinations that three areas designated nonattainment for the 1997 fine particulate (PM_{2.5}) National Ambient Air Quality Standard (NAAQS) have

attained the 1997 PM_{2.5} NAAQS. These are the Martinsburg-Hagerstown, WV-MD nonattainment area; the Parkersburg-Marietta, WV-OH nonattainment area; and the Wheeling, WV-OH nonattainment area. These determinations are based upon complete, quality assured, quality controlled, and certified ambient air monitoring data that show that these areas have monitored attainment of the 1997 PM_{2.5} NAAQS during the 2006-2008 monitoring period. Currently available monitoring data for 2009 are consistent with continued attainment of the standard. The intended effect of these actions is to finalize these attainment determinations for these areas. With these final determinations, the requirements for States to submit for these areas an attainment demonstration, associated reasonably available control measures, a reasonable further progress plan, contingency measures, and other planning State Implementation Plans (SIPs) related to attainment of the standard are suspended for so long as the areas continue to meet the 1997 PM_{2.5} NAAQS. EPA's determinations that these areas have attained the 1997 PM_{2.5} NAAQS are not equivalent to the redesignation of the areas to attainment. These actions do not constitute redesignations to attainment under section 107(d)(3) of the Clean Air Act (CAA), because we do not yet have an approved maintenance plan for these areas as required under that section and section 175A of the CAA, nor a determination that these areas have met the other requirements for redesignation. The designation status of these areas remains nonattainment for the 1997 PM_{2.5} NAAQS until such time as EPA determines that these areas meet the CAA requirements for redesignation to attainment.

DATES: *Effective Date:* These final rules are effective on November 20, 2009.

ADDRESSES: EPA has established dockets for this action under Docket ID Numbers EPA-R03-OAR-2009-0199 and EPA-R03-OAR-2009-0547. All documents in the dockets are listed in the <http://www.regulations.gov> Web site. Although listed in the electronic docket, some information is not publicly available, i.e., confidential business information (CBI) or other information whose disclosure is restricted by statute. Certain other material, such as copyrighted material, is not placed on the Internet and will be publicly available only in hard copy form. Publicly available docket materials are available either electronically through <http://www.regulations.gov> or in hard

copy for public inspection during normal business hours at the Air Protection Division, U.S. Environmental Protection Agency, Region III, 1650 Arch Street, Philadelphia, Pennsylvania 19103.

FOR FURTHER INFORMATION CONTACT: Marilyn Powers, (215) 814-2308, or by e-mail at powers.marilyn@epa.gov.

SUPPLEMENTARY INFORMATION: Throughout this document whenever “we,” “us,” or “our” is used, we mean EPA.

Organization of this document. The following outline is provided to aid in locating information in this preamble.

- I. What Actions is EPA Taking?
- II. Responses to Comments
- III. What is the Effect of these Actions?
- IV. When are these Actions Effective?
- V. What are EPA’s Final Actions?
- VI. What are the Statutory and Executive Order Reviews?

I. What Actions Is EPA Taking?

EPA is determining that the Martinsburg-Hagerstown, WV-MD, Parkersburg-Marietta, WV-OH, and Wheeling, WV-OH PM_{2.5} nonattainment areas have attained the 1997 PM_{2.5} NAAQS. These determinations are based upon three years of complete, quality assured, quality controlled, and certified ambient air monitoring data that show the areas have monitored attainment of the 1997 PM_{2.5} NAAQS during the 2006–2008 monitoring period. Currently available monitoring data for 2009 are consistent with continued attainment. Other specific details of the determinations and the rationale for EPA’s proposed actions are explained in the notices of proposed rulemaking (NPRs) published on July 31, 2009 (74 FR 38154 and 74 FR 38161) and the subsequent NPRs reopening the comment period, published on September 29, 2009 (74 FR 49833 and 74 FR 49834), and will not be restated here. Although on July 31, 2009 EPA issued separate notices of proposed rulemaking for the Maryland portion of the Martinsburg-Hagerstown nonattainment area (74 FR 38161) and the West Virginia portion (74 FR 38154), in this notice of final rulemaking EPA is addressing in combination both State portions of this nonattainment area. EPA considers its determination of attainment for the Hagerstown-Martinsburg area, and the comments and responses relating to it, as applicable to the entire nonattainment area.¹ Similarly, although EPA

articulated its proposals for determinations of attainment for the Parkersburg-Marietta, WV-OH and Wheeling, WV-OH nonattainment areas (74 FR 38154) in terms of the West Virginia and Ohio portions of the nonattainment area, in this final rulemaking EPA addresses in combination both State portions of the nonattainment area. EPA considers its determinations of attainment for those areas, and any comments and responses relating to them, as applicable to each entire nonattainment area.

II. Responses to Comments

EPA received comments in response to the NPRs published on July 31, 2009 (74 FR 38154 and 74 FR 38161) and the subsequent NPRs reopening the comment period, published on September 29, 2009 (74 FR 49833 and 74 FR 49834). EPA received both supporting and adverse comments. By this notice EPA is responding to adverse public comments received in response to these NPRs. EarthJustice, on behalf of the Sierra Club, submitted comments by letters dated August 31, 2009 and October 29, 2009. In its comment letter dated August 31, 2009, EarthJustice included a request that the NPRs either be revised or the comment period be reopened to allow the public to consider whether the 2009 data supports the proposed attainment determinations. EPA agreed to reopen the comment period to provide the requested data. In the technical support documents (TSDs) for the NPRs that reopened the comment period, the 2009 data were included, as was additional information that clarified the 2006–2008 data that formed the foundation for the proposed clean data determinations. EPA explained in the TSDs that its determinations were based on complete, quality assured 2006–2008 data, and that the available 2009 data, while not yet complete or quality assured, were consistent with continued attainment. On October 29, 2009, EarthJustice, on behalf of the Sierra Club, submitted additional comments on the supplemental proposals and TSDs.

The only adverse comments received were submitted by the Sierra Club and may be categorized as follows: (a) Comments specifically aimed at the 2006–2008 data that form the

PM_{2.5} nonattainment area. EarthJustice, on behalf of the Sierra Club, also submitted comments on this proposal in conjunction with its comments on the other proposed rulemakings discussed in this final notice. EPA is not finalizing its proposed attainment determination for the Baltimore, MD PM_{2.5} nonattainment area in this notice of final rulemaking, and is therefore not addressing the comments relating to Baltimore that were submitted by the commenter with respect to that proposal.

foundation of EPA’s attainment determination for the Martinsburg-Hagerstown nonattainment area and (b) comments relating generally to monitoring data for 2009. Through this notice, EPA first addresses the adverse comments specifically directed at the 2006–2008 data for the Martinsburg-Hagerstown nonattainment area, and then addresses the comments relating to monitoring data for 2009.

Martinsburg-Hagerstown Nonattainment Area

Comment

Through its comment letters dated August 31, 2009 and October 30, 2009, the commenter disagrees with EPA’s proposed attainment determination for the Martinsburg-Hagerstown nonattainment area, based on information relating to a Martinsburg, WV monitor that the commenter obtained from EPA’s AirData database (<http://www.epa.gov/air/data/index.html>). As part of its August 31, 2009 comments, the commenter submitted copies of three Monitor Values Reports obtained from EPA’s AirData Web site on August 18, 2009. Each of these reports provides data showing that the Martinsburg, WV monitor (monitor ID #540030003) located in Berkeley County, West Virginia monitored annual PM_{2.5} mean concentrations of 14.93 µg/m³ in 2006, 15.61 µg/m³ in 2007 and 15.36 µg/m³ in 2008. Based upon the data set forth in these three reports, the commenter asserts that the design value calculated from these annual means results in nonattainment of the annual NAAQS over the 2006–2008 period. The commenter believes this is contrary to the data provided in the July 31, 2009 NPRs, and that it renders EPA’s proposed determination of attainment for the Martinsburg-Hagerstown area inaccurate; the commenter contends that finalizing the determination would be arbitrary and unlawful. Through its October 30, 2009 comments, the commenter also noted that, in accordance with a September 4, 1992 memo entitled “Procedures for Processing Requests to Redesignate Areas to Attainment” from John Calcagni, Director, Air Quality Management Division, EPA’s Office of Air Quality Planning and Standards, data used to demonstrate attainment should be recorded in the Aerometric Information Retrieval System (AIRS) in order for it to be available to the public for review. The commenter asserted that EPA cannot rely on data that is not published in AIRS to support a finding of attainment.

¹ EPA’s proposed rulemaking for the Maryland portion of the Martinsburg-Hagerstown area also contained a separate and independent proposed determination of attainment for the Baltimore, MD

Response

EPA has reviewed the AirData Monitor Values Reports that were submitted by the commenter. The "airdata files" submitted by the commenter were obtained using EPA's "AirData" Web site which accesses an extracted static subset of the Aerometric Information Retrieval System Air Quality System (AQS) database. This distinction is relevant because, although, prior to February 2009, the "airdata files" were extracted from AQS on a monthly basis,² some values may be absent due to incomplete reporting and some values may be subsequently changed after being subjected to quality assurance activities. EPA does not rely on incomplete, non-quality assured data to make determinations of attainment with the NAAQS. Instead, EPA uses certified air monitoring data generally from AQS that meet requirements found in Appendices A, C, D, and E of 40 CFR Part 58, to determine compliance with the NAAQS. Each of the AirData Monitor Values Reports that was submitted by the commenter includes a disclaimer which states in part: "AirData reports are produced from a monthly extract of EPA's air pollution database, AQS. Data for this report were extracted on January 10, 2009. They represent the best information available to EPA from state agencies on that date. However, some values may be absent due to incomplete reporting, and some values subsequently may be changed due to quality assurance activities." This disclaimer is particularly relevant to the EPA's review of the data presented in the Monitor Values Report for the Martinsburg, WV monitor (monitor ID #540030003) located in Berkeley County, West Virginia ("the Martinsburg monitor") for the 2008 calendar year. The monitored data set forth in this 2008 Monitor Values Report were incomplete because this report did not include monitoring data for the fourth quarter of 2008. The submitted report included data for only the first three quarters of 2008. Due to the absence of the fourth quarter 2008 monitoring data, the 2008 design value

² EPA no longer supports the monthly updating of the AirData Web site from official data contained within AQS. Ambient data reported to AQS after January 2009 would not be available through this tool. Due to resources constraints and EPA's plan to replace AirData with a new method allowing up-to-date access by the public to data in AQS, as of the date of this response EPA has not updated the AirData summary data to reflect all 2008 data submitted to AQS. At the present time, the preferred method for the general public to access the publicly available PM_{2.5} AQS data is to use the data and information available as part of EPA's AirTrends Site at: <http://www.epa.gov/airtrends/values.html>.

for the Martinsburg monitor cannot be accurately calculated using the data set forth in the submitted 2008 Monitor Values Report. The fourth quarter monitored PM_{2.5} mean concentration level at the Martinsburg monitor, which became available after January 2009, was 10.68 µg/m³. The annual PM_{2.5} mean concentration for 2008 is derived from the average of the quarterly means for each of the four quarters of 2008: 15.70 µg/m³, 13.80 µg/m³, 16.57 µg/m³, and 10.68 µg/m³; and such calculation results in a finding that the annual PM_{2.5} mean concentration for 2008 is 14.19 µg/m³. The design value for the Martinsburg monitor for the 2006–2008 period is 14.9 µg/m³; this is the average of the monitored annual PM_{2.5} mean concentrations of: 14.93 µg/m³ in 2006, 15.61 µg/m³ in 2007, and 14.19 µg/m³ in 2008. The design value shown in the initial NPRs for the Martinsburg-Hagerstown, WV-MD nonattainment area published on July 31, 2009 (74 FR 38154 and 74 FR 38161) was 14.9 µg/m³. The annual PM_{2.5} mean concentrations data and additional 2008 data, that form the foundation for EPA's attainment determination were supplied and made available for public comment when EPA re-opened its public comment period and issued its supplemental NPRs and TSDs. For further information, see paragraph C(1) on page 3 of the TSDs for the September 29, 2009 NPR reopening of the comment period (74 FR 49833 and 74 FR 49834).

Comment

Based on the commenter's contention, set forth above in the preceding comment, that in 2006–2008 a monitor in Martinsburg, WV recorded a violation of the PM_{2.5} NAAQS, the commenter argues that EPA cannot make a determination of attainment solely for the Maryland portion of the Martinsburg-Hagerstown area.

Response

As set forth in the response to comment above, EPA disagrees that the Martinsburg monitor in the Martinsburg-Hagerstown nonattainment area shows a violation of the annual PM_{2.5} standard for the period 2006–2008. EPA agrees, however, that if, hypothetically, the commenter were correct and EPA determines that complete, quality assured data from any monitor in the area eligible for comparison to the NAAQS shows nonattainment, the violation would affect the attainment status of the entire area that had been designated nonattainment. As set forth in detail above, however, the complete, quality-assured data for both monitors in the

Martinsburg-Hagerstown area show that the entire area is in attainment of the NAAQS. EPA agrees that 40 CFR 51.1004(c) applies "upon a determination by EPA that an area designated nonattainment for the PM_{2.5} NAAQS has attained the standard." EPA is making the determination here that the entire Martinsburg-Hagerstown area has attained the PM_{2.5} NAAQS, based on monitors in both portions of the bi-state area.

2009 Data

Comment

In its August 31, 2009 comments, the commenter requested that the 2009 data referred to in the NPRs be made available to the public via either revised NPRs or a reopening of the comment period to enable the public to comment on whether the 2009 data supports a finding of attainment. In its October 29, 2009 comments, the commenter agreed that "EPA cannot rely on the 2009 data to make a finding of attainment as it is not complete and does not meet EPA requirements for demonstration of attainment."

Response

EPA granted the commenter's August 31, 2009 request, presented available 2009 data, and reopened the comment period. As requested by the commenter, the portion of the 2009 data that was available at the time of EPA's July 31, 2009 proposed rulemaking was included as part of the docket, and the public comment period was reopened on September 29, 2009 (74 FR 49833 and 74 FR 49834). In addition to the 2009 data, additional information was provided in the September 29, 2009 NPRs that clarifies how the design value was calculated for the 2006–2008 period. Thus the commenter and the public have been provided an opportunity to review data and analyses relating to EPA's determinations of attainment that are the subject of today's rulemakings.

At the time of EPA's proposed determinations EPA did not have complete, quality-assured, State-certified air quality data for the entire 2009 calendar year. Nor does EPA have those data at the time of this final rulemaking. The complete, quality assured, State certified air quality data for the entire 2009 calendar year will not be available until well into calendar year 2010, and, therefore, cannot be used at this time for purposes of design value calculations during calendar year 2009. In accordance with 40 CFR Part 50 Appendix N and standard EPA practice, EPA's determinations of attainment are

based on the three most recent years of complete, quality-assured data, from 2006 to 2008. Appendix N does not provide for examining partial years of data, because various seasons of the year reflect various influences on PM_{2.5} concentrations, and a partial year's data may not be representative of values that would be determined from a full year's data set. Nevertheless, EPA also examined currently available data from 2009 for the limited purpose of determining whether they are consistent with its determination of attainment. The available data for 2009, though not the basis of EPA determinations of attainment, indicate a continuing trend that is consistent with EPA's determination of attainment, based on 2006 to 2008 data, that the Martinsburg-Hagerstown, Parkersburg-Marietta, and Wheeling 1997 PM_{2.5} nonattainment areas are attaining the 1997 PM_{2.5} standards.

In its October 29, 2009 comments, which the Sierra Club submitted after having an opportunity to review the 2009 data it requested, the Sierra Club did not raise any additional concerns about the data apart from their incomplete and preliminary nature. EPA has addressed these concerns above and in its TSDs. EPA has thus fully explained the scope of its review of these data, and the basis for its determinations of attainment based on 2006–2008 data that are complete and quality assured.

III. What Is the Effect of These Actions?

These final actions, in accordance with 40 CFR 51.1004(c), suspend the requirements for the Martinsburg-Hagerstown, WV-MD nonattainment area, the Parkersburg-Marietta, WV-OH nonattainment area, and the Wheeling, WV-OH nonattainment area to submit attainment demonstrations, associated reasonably available control measures, reasonable further progress plans, contingency measures, and other planning SIPs related to attainment of the 1997 PM_{2.5} NAAQS for so long as that area continues to attain the 1997 PM_{2.5} NAAQS.

IV. When Are These Actions Effective?

EPA finds that there is good cause for these determinations to become effective on the date of publication of this action in the **Federal Register**, because a delayed effective date is unnecessary due to the nature of the approval. The expedited effective date for these actions is authorized under both 5 U.S.C. 553(d)(1), which provides that rule actions may become effective less than 30 days after publication if the rule “grants or recognizes an exemption

or relieves a restriction” and 5 U.S.C. 553(d)(3), which allows an effective date less than 30 days after publication “as otherwise provided by the agency for good cause found and published with the rule.” As noted above, these determinations of attainment suspend the requirements for the Martinsburg-Hagerstown, Parkersburg-Marietta, and Wheeling PM_{2.5} nonattainment areas to submit an attainment demonstration, associated reasonably available control measures, a reasonable further progress plan, contingency measures, and any other planning SIPs related to attainment of the standard for so long as these areas continue to meet the 1997 PM_{2.5} NAAQS. The suspension of these requirements is sufficient reason to allow an expedited effective date of this rule under 5 U.S.C. 553(d)(1). In addition, the suspension of these requirements provide good cause to make this rule effective on the date of publication of this action in the **Federal Register**, pursuant to 5 U.S.C. 553(d)(3). The purpose of the 30-day waiting period prescribed in 5 U.S.C. 553(d) is to give affected parties a reasonable time to adjust their behavior and prepare before the final rule takes effect. Where, as here, the final rules suspend requirements rather than imposing obligations, affected parties do not need time to adjust and prepare before the rule takes effect.

V. What Are EPA's Final Actions?

EPA is determining that the Martinsburg-Hagerstown, Parkersburg-Marietta, and Wheeling nonattainment areas have attained the standards for the 1997 PM_{2.5} NAAQS. These determinations are based upon complete, quality assured, quality controlled, and certified ambient air monitoring data showing that these areas have monitored attainment of the 1997 PM_{2.5} NAAQS based on the 2006–2008 data. In addition, preliminary air quality data available for 2009 are consistent with continuing attainment. These final actions, in accordance with 40 CFR 51.1004(c), will suspend the requirements for States to submit attainment demonstrations, associated reasonably available control measures, reasonable further progress plans, contingency measures, and other planning SIPs related to attainment of the 1997 PM_{2.5} NAAQS for each area, for so long as that area continues to meet the 1997 PM_{2.5} NAAQS. EPA's determination that these areas have attained the 1997 PM_{2.5} NAAQS are not equivalent to the redesignation of these areas to attainment. These actions do not constitute redesignations to attainment under section 107(d)(3) of

the CAA, because we do not yet have an approved maintenance plan for these areas as required under section 175A of the CAA, nor a determination that these areas have met the other requirements for redesignation. The designation status of these areas remains nonattainment for the 1997 PM_{2.5} NAAQS until such time as EPA determines that these areas meet the CAA requirements for redesignation to attainment.

VI. What Are the Statutory and Executive Order Reviews?

A. General Requirements

Under Executive Order 12866 (58 FR 51735, October 4, 1993), these actions are not “significant regulatory action” and therefore is not subject to review by the Office of Management and Budget. For this reason, these actions are not subject to Executive Order 13211, “Actions Concerning Regulations That Significantly Affect Energy Supply, Distribution, or Use” (66 FR 28355, May 22, 2001). These actions make determinations based on air quality data and result in the suspension of certain Federal requirements. Accordingly, the Administrator certifies that these rules will not have a significant economic impact on a substantial number of small entities under the Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*). Because these rules make determinations based on air quality data, and result in the suspension of certain Federal requirements, they do not contain any unfunded mandate or significantly or uniquely affect small governments, as described in the Unfunded Mandates Reform Act of 1995 (Pub. L. 104–4).

These rules also do not have tribal applications because they will not have a substantial direct effect on one or more Indian tribes, on the relationship between the Federal Government and Indian tribes, or on the distribution of power and responsibilities between the Federal Government and Indian tribes, as specified by Executive Order 13175 (65 FR 67249, November 9, 2000). These actions also do not have Federalism implications because they do not have substantial direct effects on the States, on the relationship between the national government and the States, or on the distribution of power and responsibilities among the various levels of government, as specified in Executive Order 13132 (64 FR 43255, August 10, 1999), because they merely make determinations based on air quality data and result in the suspension of certain Federal requirements, and do not alter the relationship or the distribution of power and responsibilities established in the

CAA. These rules also are not subject to Executive Order 13045 "Protection of Children from Environmental Health Risks" (62 FR 19885, April 23, 1997) because they determine that air quality in the affected areas are meeting Federal standards.

The requirements of section 12(d) of the National Technology Transfer and Advancement Act of 1995 (15 U.S.C. 272 note) do not apply because it would be inconsistent with applicable law for EPA, when determining the attainment status of an area, to use voluntary consensus standards in place of promulgated air quality standards and monitoring procedures to otherwise satisfy the provisions of the CAA.

These rules do not impose an information collection burden under the provisions of the Paper Reduction Act of 1995 (44 U.S.C. 3501 *et seq.*)

Under Executive Order 12898, EPA finds that these rules involve determinations of attainment based on air quality data and will not have disproportionately high and adverse human health or environmental effects on any communities in these areas, including minority and low-income communities.

B. Submission to Congress and the Comptroller General

The Congressional Review Act, 5 U.S.C. 801 *et seq.*, as added by the Small Business Regulatory Enforcement Fairness Act of 1996, generally provides that before a rule may take effect, the agency promulgating the rule must submit a rule report, which includes a copy of the rule, to each House of the Congress and to the Comptroller General of the United States. EPA will submit a report containing these actions and other required information to the U.S. Senate, the U.S. House of Representatives, and the Comptroller General of the United States prior to publication of the rule in the **Federal Register**. A major rule cannot take effect until 60 days after it is published in the **Federal Register**. These actions are not "major rules" as defined by 5 U.S.C. 804(2).

C. Petitions for Judicial Review

Under section 307(b)(1) of the Clean Air Act, petitions for judicial review of these actions must be filed in the United States Court of Appeals for the appropriate circuit by January 19, 2010. Filing a petition for reconsideration by the Administrator of this final rule does not affect the finality of these actions for the purposes of judicial review nor does it extend the time within which a petition for judicial review may be filed,

and shall not postpone the effectiveness of such rule or actions.

These actions, pertaining to the determinations of attainment for the 1997 fine particulate matter standard for the Martinsburg-Hagerstown, Parkersburg-Marietta, and Wheeling PM_{2.5} nonattainment areas, may not be challenged later in proceedings to enforce requirements. (See section 307(b)(2).)

List of Subjects in 40 CFR Part 52

Environmental protection, Air pollution control, Incorporation by reference, Particulate matter.

Dated: November 10, 2009.

William C. Early,

Acting Regional Administrator, Region III.

■ 40 CFR part 52 is amended as follows:

PART 52—[AMENDED]

■ 1. The authority citation for part 52 continues to read as follows:

Authority: 42 U.S.C. 7401 *et seq.*

Subpart V—Maryland

■ 2. Section 52.1081 is amended by designating the existing paragraph as paragraph (a) and adding paragraph (b) to read as follows:

§ 52.1081 Control strategy: Particulate matter.

* * * * *

(b) *Determination of Attainment.* EPA has determined, as of November 20, 2009, the Martinsburg-Hagerstown, WV-MD PM_{2.5} nonattainment area has attained the 1997 PM_{2.5} NAAQS. This determination, in accordance with 40 CFR 52.1004(c), suspend the requirements for this area to submit an attainment demonstration, associated reasonably available control measures, a reasonable further progress plan, contingency measures, and other planning SIPs related to attainment of the standard for as long as this area continues to meet the 1997 PM_{2.5} NAAQS.

Subpart KK—Ohio

■ 3. Section 52.1880 is amended by adding paragraph (k) to read as follows:

§ 52.1880 Control strategy: Particulate matter.

* * * * *

(k) *Determinations of Attainment.* EPA has determined, as of November 20, 2009, the Parkersburg-Marietta, WV-OH and the Wheeling, WV-OH PM_{2.5} nonattainment areas have attained the 1997 PM_{2.5} NAAQS. These determinations, in accordance with 40 CFR 52.1004(c), suspend the

requirements for these areas to submit an attainment demonstration, associated reasonably available control measures, a reasonable further progress plan, contingency measures, and other planning SIPs related to attainment of the standard for as long as these areas continue to meet the 1997 PM_{2.5} NAAQS.

Subpart XX—West Virginia

■ 4. Section 52.2526 is amended by designating the existing paragraph as paragraph (a) and by adding paragraph (b) to read as follows:

§ 52.2526 Control strategy: Particulate matter.

* * * * *

(b) *Determinations of Attainment.* EPA has determined, as of November 20, 2009, the Martinsburg-Hagerstown, WV-MD, the Parkersburg-Marietta, WV-OH and the Wheeling, WV-OH PM_{2.5} nonattainment areas have attained the 1997 PM_{2.5} NAAQS. These determinations, in accordance with 40 CFR 52.1004(c), suspend the requirements for these areas to submit an attainment demonstration, associated reasonably available control measures, a reasonable further progress plan, contingency measures, and other planning SIPs related to attainment of the standard for as long as these areas continue to meet the 1997 PM_{2.5} NAAQS.

[FR Doc. E9-27824 Filed 11-19-09; 8:45 am]

BILLING CODE 6560-50-P

DEPARTMENT OF HOMELAND SECURITY

Federal Emergency Management Agency

44 CFR Part 206

[Docket ID FEMA-2006-0028]

RIN 1660-AA45

Public Assistance Eligibility

AGENCY: Federal Emergency Management Agency, DHS.

ACTION: Final rule.

SUMMARY: The Federal Emergency Management Agency (FEMA) provides financial assistance to State, local, and Tribal governments, as well as certain private non-profit organizations, for response and recovery activities required as a result of a presidentially-declared major disaster or emergency. Assistance may include reimbursement for sheltering and evacuation costs

incurred to assist individuals displaced by a declared major disaster or emergency. This rule finalizes the July 2006 interim rule which amended FEMA's Public Assistance eligibility regulations to allow grantees to seek reimbursement for sheltering and evacuation costs incurred outside of the area designated under a Presidential emergency or major disaster declaration, if such costs are otherwise eligible for FEMA Public Assistance. This rule further clarifies those regulations to specify which entities may be eligible for reimbursement for costs incurred from providing evacuation and sheltering services outside the area of the declared emergency or major disaster, and the procedures FEMA will use to reimburse those applicants. The rule also establishes the terms "impact-State" and "host-State" to differentiate between the State for which the President has issued a declaration and that requests evacuation and/or sheltering assistance, and the State (or Tribe) that provides the sheltering and/or evacuation assistance, respectively. Finally, the rule makes a procedural change to the way in which a host-State receives reimbursement for the regular salary or hourly wages and benefits paid to its permanent employees.

DATES: *Effective Date:* December 21, 2009.

ADDRESSES: The electronic docket for this rulemaking is available on the Federal eRulemaking Portal at www.regulations.gov, in Docket ID "FEMA-2006-0028." A hard copy of the docket may also be viewed at FEMA, Office of Chief Counsel, Room 835, 500 C Street, SW., Washington, DC 20472-3100.

FOR FURTHER INFORMATION CONTACT: Tod Wells, Acting Director, Public Assistance Division, Federal Emergency Management Agency, 500 C Street, SW., Washington, DC 20472-3100, (phone) 202-646-3936, or (e-mail) tod.wells@dhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

FEMA, through its Public Assistance program, provides financial assistance to State, Tribal, and local governments, as well as certain private non-profit organizations to quickly respond to and assist communities to recover from major disasters or emergencies declared by the President. In providing assistance through this program, FEMA provides a grant to a "grantee," which is typically a State, but may also be an Indian Tribal government. 44 CFR 206.201(e) and 206.202(f). The grantee administers the program and provides funding directly

to "subgrantees," which may be local governments, eligible private non-profit organizations, and Indian Tribal governments. An Indian Tribal government may choose to be either a grantee or a subgrantee. The grantee submits eligible costs incurred by it and/or its subgrantees to FEMA for reimbursement.

Traditionally, the grantee is the State that requests and receives a major disaster or emergency declaration from the President, and the costs eligible for reimbursement from FEMA are costs incurred in the area designated in the major disaster or emergency declaration. Costs incurred outside the declared area were not reimbursable. When a State, Indian Tribal, or local government, or an eligible private non-profit incurred costs as a result of a disaster that occurred elsewhere, the State¹ was required to seek its own emergency declaration to have those costs reimbursed.

In response to Hurricanes Katrina and Rita in 2005, 45 States requested and received emergency declarations to recover sheltering costs for tens of thousands of evacuees from the Gulf Coast States. Declaring an emergency in each of these States² was an imperfect method of responding because each State incurred administrative costs to request an emergency declaration, and requesting States were subject to the cost share requirements of The Robert T. Stafford Disaster Relief and Emergency Assistance Act³ (Stafford Act). FEMA concluded that "host" States, Tribes, and local governments could better assist "impact" States needing assistance evacuating and sheltering their residents as a result of declared major disasters or emergencies if the host-State could obtain reimbursement directly from either the impact State, Tribe or local government or directly from FEMA without first obtaining an individual emergency declaration.

On July 14, 2006, FEMA published an interim rule amending its Public

Assistance eligibility regulations to allow grantees to seek reimbursement for sheltering and evacuation costs incurred outside of the area designated under a Presidential emergency or major disaster declaration, if such costs are otherwise eligible for FEMA Public Assistance funding. 71 FR 40025. FEMA, in promulgating the 2006 interim rule, recognized the benefit in reimbursing grantees outside of a designated area when they are requested to provide, and consequently incur, costs for sheltering and evacuation support to evacuees from another State, Tribe, or local government.

In making this change, the 2006 interim rule expanded eligible costs to include sheltering and evacuation costs that occur outside of a declared area, and allowed States to recover these costs through direct Federal reimbursement when such assistance was requested by the State with a declaration.

The expansion of eligible costs to include sheltering and evacuation activities that occur outside a declared area affected costs both within and outside a declared State. The evacuation and sheltering costs provided between local governments within a State may be covered by mutual aid agreements, whereas assistance provided outside a state may be covered either by a mutual aid agreement between States or direct Federal reimbursement to a State from FEMA pursuant to a disaster declaration.

Many local governments have pre-existing mutual aid agreements and share materials and services with one another in times of need. Under mutual aid agreements, a requesting jurisdiction within a designated area may request another jurisdiction outside of the designated area to provide evacuation and sheltering services for the requesting jurisdiction's residents. Additionally, many States participate in the Emergency Management Assistance Compact (EMAC), or have other similar agreements, through which the States agree to provide assistance to one another when requested. Under a mutual aid arrangement, the jurisdiction that requested the assistance reimburses the providing jurisdiction for its costs, and then in turn, seeks reimbursement from FEMA (through the State, if the requesting entity is an eligible subgrantee).

The 2006 interim rule also allowed other States that provide evacuation and sheltering services to recover their costs from FEMA after a declared State requests direct Federal assistance pursuant to 44 CFR 206.208. FEMA provides direct Federal assistance to the

¹ As defined in the Stafford Act, only States may receive major disaster or emergency declarations. Although Indian Tribal governments may be grantees under a State's declaration, the President does not have the authority to issue a declaration for a Tribe. See 42 U.S.C. 5122, 5170, and 5191.

² The standard Federal/State cost share rate is 75/25, although it may be raised to 90/10 or 100% Federal. See 42 U.S.C. 5170(b), 5193(a), and 44 CFR 206.47. The declarations issued to States that provided sheltering and/or evacuation services as a result of Hurricanes Katrina and Rita in 2005 and Hurricane Gustav in 2008 were set by the President at a 100 percent Federal rate. There is no guarantee, however, that future disasters will receive a 100 percent rate as the rate is set on a case by case basis at the President's discretion.

³ Disaster Relief Act of 1974, Public Law 93-288, 88 Stat. 143 (May 22, 1974), as amended 42 U.S.C. 5121 *et seq.*

declared State when a State or Indian Tribal government outside the declared area, at FEMA's request, provides sheltering and evacuation services to that declared State. States, Tribes, and local governments that provide sheltering and/or evacuation assistance do not seek direct Federal assistance; the State with the declaration makes the request, and then FEMA identifies States and Tribes that are willing and able to help. This change, therefore, only affects costs outside a State that was granted an emergency or major disaster declaration; it will not affect local governments or private non-profit entities within the declared State. Only when an impact-State is overwhelmed and lacks the capability to perform or contract for emergency work would it turn to FEMA for direct Federal assistance. Since a State must exhaust its resources before receiving direct Federal assistance from FEMA, there would be no resources available within the State to provide evacuation and sheltering services.

FEMA evaluated the effectiveness of the 2006 interim rule following mass evacuations from Louisiana, including the City of New Orleans, in advance of Hurricane Gustav in August 2008. Although FEMA has found that the changes made by the 2006 Public Assistance Eligibility interim rule significantly improved the reimbursement process during the 2008 hurricane season, FEMA identified several areas to further improve the procedures for reimbursing evacuation and sheltering assistance. For example, although some States preferred to be directly reimbursed by FEMA, they requested clarification regarding the reimbursement process and which entities would be eligible for direct reimbursement. In the Public Assistance program, typically the State for which the major disaster or emergency is declared (the "impact-State") is the grantee, but, in this case, a State without a major disaster or emergency declaration providing the evacuation services may receive a grant. This new situation raised questions as to whether the State without a declaration has the responsibilities of a grantee with respect to its grant, or whether it is a subgrantee of the impact-State. States did not understand who was responsible for the non-Federal cost share under the 2006 interim rule. The 2006 interim rule did not answer whether an undeclared State that provided sheltering and evacuation services stood as a grantee or as a subgrantee to the declared State, and there was no clear application process in place. Grantees are typically

responsible for paying the non-Federal cost share and for oversight under 44 CFR part 13.

This confusion led to delays and duplicative application requirements for those seeking to recover regular salary or hourly wages and benefits paid to an applicant's permanent employees, referred to as "straight-time force account labor costs." For example, because straight-time force account labor costs were eligible only when incurred through mutual aid agreements between States, States sought reimbursement for these costs through mutual aid agreements and would apply for direct funding from FEMA for the remaining costs. Thus, a clear understanding of the procedures for addressing out-of-state evacuation and sheltering is essential to FEMA's effective management of the Public Assistance program. This final rule clarifies the process for FEMA reimbursement of those entities outside a declared area that provide sheltering and/or evacuation assistance. Further, it will provide a more efficient grant process that is likely to result in more States being willing to provide their resources to protect residents of another State impacted by a major disaster or emergency.

FEMA recognized, in addition, the need to reimburse straight-time force account labor costs through the direct Federal assistance process. Public Assistance grants are generally not available to reimburse force account straight-time for emergency work. 44 CFR 206.228(a)(2). Since an applicant's costs for permanently employed personnel are pre-disaster existing resource costs the employer would incur in addressing its responsibilities regardless of whether the event occurred, these costs are not eligible. Overtime wages are reimbursable, however, for permanent employees working extra hours in performing eligible emergency work as a result of the declared emergency or major disaster. Labor costs, including overtime wages, moreover, to backfill employees assigned to perform eligible emergency work in support of the declared emergency or major disaster are also reimbursable.

FEMA currently reimburses straight-time force account labor costs when States use the mutual aid process (such as EMAC). These costs are eligible under mutual aid because the jurisdiction providing the assistance under the agreement is considered a contractor hired as a result of the declared event to address the needs of another jurisdiction. Contrary to the typical disaster assistance subgrantee,

States that host another State's residents are not expending pre-budgeted costs to address their own governmental responsibilities. FEMA has been repeatedly advised that States assisting other States' residents are unable and unwilling to assume this added expense should a future disaster occur. This final rule, therefore, makes a change in procedure that allows for the reimbursement of straight-time force account labor to host-States directly from FEMA, rather than solely through the mutual aid process. This change is strictly procedural, and does not otherwise affect the eligibility of those costs, or the amount reimbursed. This change is expected to result in more States being willing to provide host-State sheltering assistance.

FEMA, in this rule, addresses public comments received on the interim rule, finalizes the regulations, and implements these procedural improvements. This rule establishes definitions for "impact-State" and "host-State" to clearly differentiate between the State that is being directly impacted by the event resulting in a Presidential emergency or disaster declaration and has requested direct Federal assistance to address its evacuation and sheltering needs out of state, and the State that is, at FEMA's request, providing the evacuation and sheltering to residents from the designated areas. The rule more clearly articulates the entities that may be eligible to act as a host-State, and establishes application procedures for host-States seeking reimbursement for evacuation and sheltering activities directly from FEMA. Finally, the rule revises the procedure by which host-States receive reimbursement of straight-time force account labor costs. As with the 2006 interim rule, this rule allows for both mechanisms for reimbursement—a host-State may receive reimbursement either through a mutual aid agreement or by direct reimbursement from FEMA.

II. Discussion

A. Amendments to FEMA's Public Assistance Regulations Under This Final Rule

1. Designation of Affected Areas—Clarification of Terminology

FEMA's regulations occasionally refer to "disaster-affected" areas or "designated disaster" areas in sections that apply to both emergencies and major disasters. To remove the potential that one could misconstrue the use of the term "disaster" as FEMA's intent to exclude application during declared emergencies, FEMA has revised the

language to be more precise. The term “disaster-affected” has been replaced with the term “affected” in 44 CFR 206.40(b). The word “disaster” has been removed before “affected” in 44 CFR 206.2(a)(6). The term “designated disaster area” has been replaced with “designated area” in 44 CFR 206.223(a)(2), and the words “emergency or,” have been added before the phrase “major disaster event” in 44 CFR 206.223(a)(1). Similarly, to clarify that 44 CFR 206.208(a) applies to emergency assistance under an emergency declaration as well as a major disaster declaration, this rule adds Stafford Act citations to that section.

2. Direct Reimbursement for Host-State Evacuation and/or Sheltering—Clarification of Procedure and General Eligibility

As a result of the 2006 interim rule, a State or Tribe may be reimbursed for costs incurred from evacuation and sheltering activities performed outside the designated area. This rule amends FEMA regulations to align with the preamble of the 2006 interim rule and clarify that a State with a Stafford Act declaration may request direct Federal assistance from FEMA for evacuation and/or sheltering activities that occur outside the State. In doing so, the rule points applicants to the eligibility requirements for those who may provide evacuation and/or sheltering assistance when requested, what costs are eligible for reimbursement, and establishes the procedures the providing entity must follow to seek reimbursement. The State with a Presidential declaration may also request assistance from another State on its own, through a mutual aid agreement (such as EMAC), but this rule does not specifically address that option since mutual aid costs are already reimbursed by FEMA.

Direct Federal assistance under 44 CFR 206.208 applies when a State lacks the capability to perform or contract for emergency work. When this occurs, the State asks FEMA for assistance. In this rule the requesting State is referred to as the “impact-State.” When such a request is made, FEMA will ask another State or an Indian Tribal government if it is capable and willing to provide sheltering and/or evacuation assistance to the impact-State. If such State or Indian Tribal Government is capable and willing, FEMA will then provide direct reimbursement through a grant to the State or Indian Tribal government that provides evacuation and/or sheltering activities. The providing State or Indian Tribal government is referred to in this rule as the “host-State” or

“host-Tribe,” respectively, or collectively as the “host-State.” Through the direct Federal assistance process, the host-State is a grantee. Although it is obtaining assistance as a result of the impact-State’s declaration, it is not a subgrantee of the impact-State. This means that although the impact-State will continue to incur the Federal cost-share for the assistance, the impact-State is not responsible for the oversight of the host-State’s grant under the requirements of 44 CFR part 13 as it would for subgrantees. The cost-share requirements for impact-States are discussed more fully elsewhere in this preamble.

FEMA’s regulations set out the criteria that routinely apply when direct Federal assistance is requested by and provided to a State that has received a Stafford Act declaration. 44 CFR 206.208. This rule clarifies that the criteria also apply to a host-State and an impact-State. For example, the impact-State is responsible for the non-Federal cost share under 44 CFR 206.208(b)(iii) and as required in 44 CFR 206.208(c), the requested work must be eligible under the Public Assistance eligibility criteria contained in Subpart H, *Public Assistance Eligibility*. Since the criteria set out in 44 CFR 206.208 apply to an impact-State’s request for direct Federal assistance, as well as to how FEMA can provide such assistance, a provision has been added to 44 CFR 206.208 that specifically addresses host-State reimbursement.

The 2006 interim rule was also silent with respect to when and to which entity FEMA would award a grant for direct Federal reimbursement. FEMA must take into consideration the host-State’s evacuation and sheltering capabilities before it can award a grant to the host-State to protect against the possibility of individuals being sent to States that are unable to appropriately shelter them. Neither FEMA nor the impact-State should send people to a host-State, as a matter of policy, if that host-State is unable to meet the needs of the evacuees. A grant to a State that cannot host evacuees would not serve the purpose of aiding the impact-State. The determination of a host-State’s capability will be made on a case-by-case basis and the criteria will vary depending upon the specific needs of the impact-State, but will generally focus on the availability of short or mid-term housing, and equipment for evacuation activities. This rule adds a provision to 44 CFR 206.208 to address this need, providing that a grant to a host-State is available when FEMA determines that a host-State has sufficient capability to meet some or all

of the sheltering and/or evacuation needs of the impact-State.

To establish a record of the agreement and reduce confusion and miscommunication, this rule adds a requirement to 44 CFR 206.208 that the host-State must agree in writing to provide evacuation and/or sheltering assistance to individuals from the impact-State. This agreement is referred to as the commitment letter, and is provided to FEMA before the execution of the FEMA/Host-State Agreement.

The 2006 interim rule also lacked sufficient clarity with respect to the host-State’s obligation to enter into a written agreement with FEMA. This rule clarifies that a host-State must enter into a FEMA/Host-State Agreement (similar to a FEMA/State Agreement) before grant funds will be awarded. This FEMA/Host-State Agreement, which covers the conditions of the grant award, is consistent with that required of a declared State grantee pursuant to 44 CFR 206.44. The FEMA/Host-State Agreement also includes a provision on the cost share.

Grantees are required by the Stafford Act and FEMA’s implementing regulations to pay a percentage share of the costs of the Federal assistance, known as the non-Federal cost share. See 42 U.S.C. 5170(b), 5193(a), and 44 CFR 206.47. Such costs would include those for evacuation and sheltering activities. This cost share requirement applies whether the cost is incurred through mutual aid or through direct Federal assistance. The Federal/non-Federal cost share for a grant to a host-State to evacuate and/or shelter individuals from the impact-State is the same as the cost share established for all other Category B, Emergency Protective Measures, for the declared major disaster or emergency. As with all other assistance under the declaration, the non-Federal cost share for host-State sheltering is the responsibility of the impact-State under its declaration. This means that the host-State will be reimbursed for 100 percent of its eligible costs and the impact-State will continue to be responsible for the non-Federal cost share as agreed to in its FEMA/State Agreement. FEMA finds that the host-State should be reimbursed for 100 percent of its eligible costs because it is using its State resources to aid individuals from another State. Such costs are not part of a host-State’s annual budget. For impact-States, these costs would have been borne by the impact-State had they sheltered their residents in-State, or requested assistance from the host-State themselves through mutual aid. This clarification, therefore, adds no new

costs for the impact-State. An impact-State must agree, when requesting direct Federal assistance for evacuation and sheltering, to provide the non-Federal cost share for all eligible costs incurred by any host-State to ensure that no improper Federalism implications occur and that impact-States knowingly and willingly agree to incur these costs.

States have expressed some confusion as to whether host-State grantees are required to submit the same information, and undertake the same obligations as other grantees. The requirements for host-State direct reimbursement under 44 CFR 206.202(f)(1) and 206.208 should be read together. A host-State's responsibilities, including the requirement to assume the responsibilities of a Public Assistance grantee with respect to its grant award, are set out in 44 CFR 206.202(f). The host-State assumes these responsibilities because the host-State is receiving a direct grant from FEMA and is therefore acting as a grantee. For clarity, in 44 CFR 206.208, this rule specifically adds a reference to 44 CFR 206.202(f)(1), Host-State Evacuation and/or Sheltering.

This rule clearly states that, as a grantee, the host-State must submit a Standard Form 424, *Application for Federal Assistance*, to apply for reimbursement from FEMA. SF-424 is not a new requirement, as FEMA requires this form from all grantees under 44 CFR 206.202(e). The host-State is responsible for this and other grants management provisions in the regulation only with respect to its evacuation/sheltering grant. FEMA also requires all grantees to develop a State administrative plan. See 44 CFR 206.207. The State administrative plan includes the designation of State agencies responsible for program administration, identifies Public Assistance staffing functions, and includes procedures for conducting briefings, notifying potential applicants, processing appeal requests, and other procedures for administering the Public Assistance program. Grantees are required to update their administrative plans under 44 CFR 206.207. This rule clearly states that this requirement also applies to host-States under 44 CFR 206.202(f).

3. Straight-Time Force Account Labor

As discussed above, FEMA currently reimburses applicants for the overtime costs of their permanently employed personnel who perform emergency work as a result of a declared event when direct Federal assistance is provided. FEMA does not, however, reimburse the

straight-time wages for these employees. When a host-State provides evacuation and/or sheltering assistance under a mutual aid agreement, however, FEMA does reimburse host-State force account labor for both straight-time and overtime. FEMA treats the costs incurred by the host-State (referred to as a "providing entity") under a mutual aid agreement as contract labor, with regular time and overtime wages and certain benefits eligible, provided the labor rates are reasonable.

FEMA's reimbursement of regular- or "straight-time" salaries is generally governed by 44 CFR part 13 (*Uniform Administrative Requirements for Grants and Cooperative Agreements to State and Local Governments*); 2 CFR Part 225 (*Cost Principles for State, Local, and Indian Tribal Governments*); OMB Circular A-102, *Grants and Cooperative Agreements with State and Local Governments*; and OMB Circular A-87, *Principles for Determining Costs Applicable to Grants and Contracts with State, Local, and Federally Recognized Indian Tribal Governments*. FEMA has determined that it is appropriate to reimburse the regular- or straight-time salaries of a host-State's permanent employees' eligible evacuation and sheltering activities on behalf of an impact-State because a host-State is providing assistance to an impact-State's residents. The host-State is using its own resources for another State's residents, and, therefore, should be wholly compensated for the assistance that it has not budgeted. This assistance is not being provided for the benefit of the host-State's taxpayers.

As a result of Hurricanes Katrina and Rita, FEMA recognized the importance of host-State evacuation and sheltering activities in response to a large-scale event. Host-States should be encouraged to provide such assistance for future large scale events, as necessary, and delay in reimbursement through the impact-State discourages such assistance. Allowing reimbursement of straight-time force account labor under both the mutual aid and direct grant mechanisms ensures consistency and fairness in reimbursement of these eligible costs. Allowing this reimbursement also avoids any potential administrative burden of the States being required to consider differences in eligible costs when considering which reimbursement mechanism is most suitable. This rule, therefore, establishes a process for FEMA to provide direct Federal reimbursement to a host-State for straight-time salaries and benefits of a host-State's permanently employed personnel who perform evacuation and/or sheltering activities. A host-State's

straight and overtime costs may be directly reimbursed through the host-State's grant from FEMA.

4. Definitions of Host-State and Impact-State

FEMA makes frequent reference to entities within the designated area of a Presidential emergency or major disaster declaration and entities that provide evacuation and/or sheltering assistance outside the State receiving the emergency or major disaster declaration. FEMA has recognized the need to assign shorter, uniform terms to identify these entities. A uniform definition ensures consistency and clarity in implementation of this regulation. This final rule therefore adds definitions for "host-State" and "impact-State" to 44 CFR 206.201, which is the definitions section for the Public Assistance project administration regulations.

A "host-State" is a State or Indian Tribal government that by agreement with FEMA is providing sheltering and/or evacuation support to evacuees from an impact-State. An "impact-State" is the State for which the President has declared an emergency or major disaster and that, due to a need to protect its affected residents, requests assistance from FEMA pursuant to 44 CFR 206.208 to evacuate and/or shelter such individuals outside the State.

5. Definitions of Grantee and Indian Tribal Government

Since host-States are grantees, as described in this rulemaking, FEMA is updating the definition of "grantee." Typically, the declared State is the grantee eligible to receive assistance under the emergency or major disaster declaration, and is responsible for the administration and use of assistance provided under the Public Assistance program for that declaration. The revised definition of grantee states that for purposes of the Public Assistance regulations, the declared State is the grantee, except as noted in 44 CFR 206.202(f). The exception under paragraph (f) allows a host-State to apply for a grant for the specific purpose of providing sheltering and evacuation activities to the impact-State that requested direct Federal assistance from FEMA. Under this exception, a host-State reimbursed by FEMA pursuant to 44 CFR 206.208 for sheltering and/or evacuation activities has all of the responsibilities of the declared State in administering its Public Assistance grant. FEMA makes a similar clarifying amendment to the definition of "grantee" in 44 CFR 206.431 to note that the grantee is generally the declared State. FEMA has added this clarification

because before this rulemaking the “grantee” has always been presumed to be the declared State for both the Public Assistance and Hazard Mitigation Assistance programs. Without this change, FEMA was concerned that in the absence of express language to the contrary, the definition would leave the impression that a State is not required to be the declared one to receive assistance under the Hazard Mitigation Grant Program.

Before this final rule, the definition of grantee referred to a State, and 44 CFR 206.202(f) created an exception that allowed Indian Tribal governments affected by an emergency or major disaster to apply directly to FEMA for a grant when State law prohibits a State to act as grantee for an Indian Tribal government. This rule merges the exception for Indian Tribal governments that appeared in paragraph (f) into the definition of grantee to clarify that an Indian Tribal government in the affected area may choose to be a grantee, or it may act as a subgrantee under the State receiving the declaration. This merger gives Indian Tribal governments the level of recognition commensurate with the declared States because both can apply directly to FEMA for disaster assistance and is consistent with the other program definitions of the term “grantee” throughout FEMA’s regulations. This merger into the definition is consistent with FEMA’s established practice, recognition of, and commitment to, a government-to-government relationship with Indian Tribal governments. FEMA recognizes the tribal right of self-government that flows from the inherent sovereignty of Tribes as nations, and that Federally-recognized Tribes have a unique and direct relationship with the Federal government. This sovereign status also permits a qualified Indian Tribal government to deal directly with FEMA with respect to Public Assistance funding for which it is eligible under a Presidentially-declared emergency or major disaster declaration. In choosing to act as grantee, the Indian Tribal government assumes the responsibilities of grantees, including the reporting, recordkeeping, and other requirements contained in the program regulations. This choice and assumption also comports with the intent of FEMA’s policy, *Final Agency Policy for Government-to-Government Relations with American Indian and Alaska Native Tribal Governments*, 64 FR 2096 (Jan. 12, 1999), as available at <http://www.fema.gov/government/tribal/natamerpolcy.shtm>, which permits a qualified Tribal government to interact

directly with FEMA and act as its own grantee.

Finally, unlike many of FEMA’s other regulatory parts, Subpart G lacked a definition of the term “Indian Tribal government.” The definition added to 44 CFR 206.201 matches the definition of “Indian Tribal government” in other sections of FEMA regulations, such as at 44 CFR 201.2 (Mitigation Planning), 206.430 (Hazard Mitigation Grant Program), and 207.2 (Management Costs).

FEMA will be updating its guidance to the States to reflect the changes made in this rule. When these documents are available, they will be posted to FEMA’s Web site at <http://www.fema.gov>, as well as to the docket for this rulemaking at <http://www.regulations.gov>, Docket ID: FEMA–2006–0028.

B. Discussion of Public Comments on the 2006 Interim Rule

FEMA received four comments on the 2006 interim rule. The commenters included one emergency management organization, one State, and two local governments.

1. General Comments

The International Association of Emergency Managers (IAEM) stated that it received responses from 10 of its members, all in favor of the rule. The Georgia Emergency Management Agency (GEMA) stated that it was in favor of the rule because it facilitates a reasonable means of providing sheltering and evacuation support outside the impacted areas without imposing all of the other requirements associated with providing access to Public Assistance funding. The Onslow County North Carolina Emergency Services & Homeland Security Department (Onslow County) stated that the rule allowed non-affected counties to better support affected areas without absorbing the costs directly in their smaller budgets.

These comments reflect one of the main reasons FEMA promulgated the interim rule: to reduce the costs and administrative burden placed on the host-State. By eliminating the requirement that a host-State request and receive an emergency declaration from the President before recouping eligible costs for evacuation and sheltering activities, the host-State is not required to activate the same level of emergency management plans, staff, and resources that are normally required to manage and coordinate operations with FEMA.

2. Self-Evacuees

GEMA expressed concern that communities outside the designated areas that provide sheltering and evacuation for self-evacuees would not be eligible for direct reimbursement from FEMA. GEMA was concerned that the volume of individuals that may choose to evacuate and seek shelter without government support, could overwhelm existing resources and necessitate the opening of mass sheltering operations to provide basic services to the evacuees. For example, displaced individuals may choose to evacuate without government support temporarily to a more distant region due to family connections or other perceived advantages, even if FEMA-funded shelter operations were available in other jurisdictions nearer the impacted area. If a sufficient number of evacuees requiring shelter chose to relocate to areas other than those designated by the impacted communities or by FEMA, then, GEMA asserted, the receiving community should have some recourse to seek financial reimbursement in the event that the number of displaced, non-housed persons necessitates the opening of sheltering services.

FEMA generally does not have the authority to provide grant funds under the Stafford Act outside of the designated areas of the Presidential declaration. If a local government finds itself overwhelmed by self evacuees, but is not included in the State’s designated areas, the State may find it is appropriate to request that FEMA include the county among the designated areas under the declaration. As discussed elsewhere in this preamble, FEMA has been delegated the authority to amend emergency and major disaster declarations to add counties when appropriate, and frequently exercises this authority.

If a State without a Stafford Act declaration is burdened with providing sheltering support to self-evacuees, the State may ask the declared State to seek direct Federal assistance from FEMA under the provisions of this rule, or seek reimbursement through a mutual aid agreement with the declared State. Although FEMA recommends mutual aid, in some cases where there is a large-scale event, direct reimbursement from FEMA may be available in accordance with this rule.

3. Mutual Aid Agreements—Burden on Local Governments

The City of Plano was concerned that mutual aid agreements would burden local governments. The commenter stated that it would not be practical for

local governments to administer mutual aid agreements because it would be inefficient and a complex task for local governments to predetermine the host entities across the United States with which it should enter into a mutual aid agreement. The commenter also expressed concern that interstate agreements would be difficult to enforce and that local governments would not have sufficient funds to reimburse the host entities. Further, the City of Plano asserts that cities would be less likely to participate if, as stated in the 2006 interim rule, the eligible applicant will reimburse the providing entity and then be reimbursed by FEMA.

FEMA encourages the use of mutual aid agreements, including the Emergency Management Assistance Compact (EMAC). A mutual aid agreement is an efficient mechanism for providing evacuation and sheltering services when there are a relatively small number of disaster victims. States may enter into post-event mutual aid agreements, which would negate the difficulty local governments may have in determining host entities in advance. A providing entity's costs for evacuation and sheltering services under a mutual aid agreement are eligible for reimbursement by FEMA through the declared State, just as those costs are eligible if the declared State seeks direct assistance from FEMA.

The 2006 interim rule provides, and this rule clarifies, that a State (or Tribal government) may become a host-State when an impact-State requests direct Federal assistance from FEMA, FEMA approves the request, and requests the host-State to provide evacuation and sheltering services outside of the designated area. In this situation, the host-State would receive direct reimbursement from FEMA. This provides an alternate method to mutual aid agreements and may be more appropriate for large-scale events, such as Hurricanes Katrina and Rita, where the impact-State is overwhelmed and lacks the capability to respond to the need.

IV. Regulatory Requirements

A. Administrative Procedure Act

The Administrative Procedure Act requires FEMA to publish notice and consider public comments before promulgating substantive amendments to regulations, 5 U.S.C. 553(b), except when the amendment is a "rule[] of agency organization, procedure, or practice * * *." 5 U.S.C. 553(b)(3)(B). This rule makes one change that was not contemplated in the 2006 interim rule to the manner in which FEMA reimburses

a host-State for straight-time force account labor costs incurred in support of evacuation from an impacted State. As discussed throughout this preamble, straight-time force account labor costs are fully reimbursable by FEMA if the service is provided through a mutual aid agreement with the impact-State. This rule amends the regulations to permit these costs to be directly reimbursable by FEMA. The rule does not increase or decrease those costs, but merely changes the method by which host-States obtain the funds. This amendment is a rule of agency organization, procedure, or practice that is exempt from the notice and comments requirements under 5 U.S.C. 553(b)(A).

B. Executive Order 12866, Regulatory Planning and Review

FEMA has prepared and reviewed this rule consistent with Executive Order 12866, Regulatory Planning and Review. This rule has been deemed a significant, but not economically significant regulatory action by the Office of Management and Budget (OMB), and has, therefore, been reviewed by OMB.

This rule results in \$51,681 in cost savings for each large scale disaster that requires evacuation and sheltering activities to occur outside the area designated by the major disaster or emergency declaration. These savings are due to administrative savings resulting from States not being required to request Presidential declarations of their own for the event, but being able to act as "host-States" under another State's declaration. As a result, States do not need to prepare, and FEMA is not required to review and analyze, those declaration requests to make a recommendation to the President, thereby avoiding the administrative cost associated with such a review. This rule does not change the amount of assistance provided by FEMA for evacuation and sheltering activities, only the procedures by which States seek and receive reimbursement from FEMA for those costs.

Host-State evacuation and sheltering assistance is needed in only rare occurrences, and to date has only occurred twice—for Hurricanes Katrina and Rita in 2005 and Hurricane Gustav in 2008. In 2005, States not directly impacted by Hurricanes Katrina or Rita received a large number of evacuees from the impacted States of Louisiana, Mississippi, and Alabama. Although they were not actually struck by the storm, these States that provided evacuation and sheltering services to evacuees from the impacted States incurred costs. To reimburse these costs, the President declared emergencies in

many of these States, thereby making Federal assistance available for the eligible costs they incurred in providing evacuation and sheltering assistance to evacuees from the impacted States. Without obtaining a declaration, costs incurred by these States were not eligible for Federal reimbursement because the evacuation and sheltering assistance was provided outside the designated areas of the impacted States.

At the time Hurricanes Katrina and Rita struck, costs eligible for reimbursement were limited to those incurred within a designated area. Therefore, if a State incurred costs to evacuate and/or shelter residents from another State, that "host-State" was required to request and obtain its own emergency declaration to recoup eligible costs. Forty-five of the fifty States received Presidentially-declared emergencies so that they could receive Federal assistance for costs incurred after Hurricanes Katrina and Rita for evacuation and sheltering activities.⁴ FEMA provided approximately \$752.62 million in Public Assistance funding for reimbursement for host-State evacuation and sheltering activities for Hurricanes Katrina and Rita. The Federal cost share (which was 100 percent) for some States, such as Texas, Arkansas, and Tennessee, where costs totaled \$558.28 million, \$44.28 million and \$33.66 million, respectively, was substantial. Even States geographically distant from States directly struck by Katrina received Federal reimbursement for their costs. For example, Massachusetts received \$5.72 million. It became apparent that an emergency declaration was not the appropriate vehicle by which FEMA should reimburse a host-State for sheltering and evacuation activities. Sheltering and evacuation are a limited set of activities that normally, by themselves, would not warrant a Presidentially-declared emergency. FEMA needed a mechanism other than a host-State declaration to allow reimbursement for sheltering and evacuation activities outside of the areas contained in a Presidential declaration.

FEMA published the interim rule to address this need and to allow FEMA to reimburse sheltering and evacuation costs incurred by State, local, and Tribal governments that were located outside of a Presidentially-declared emergency or major disaster area, if the costs were otherwise eligible for Public Assistance funding.

Two mechanisms are provided for reimbursement. Under one mechanism,

⁴Data Source: National Emergency Management Information System (NEMIS), FEMA 2009; Enterprise Data Warehouse, FEMA 2009.

an impacted State may request an entity outside of the designated area to provide evacuation and sheltering services for the impacted State's citizens. The entity that provides the evacuation or sheltering services may seek reimbursement under a mutual aid or similar agreement with the impacted State. Under the other mechanism, the impacted State may seek direct Federal assistance from FEMA, and FEMA may, in turn, request an entity outside of the designated area to provide evacuation and sheltering services for the impacted State. This mechanism would allow the providing entity to directly receive reimbursement of its eligible costs from FEMA.

States that provide evacuation and sheltering services outside of the designated area(s) are no longer required to request and receive an emergency declaration from the President to recoup eligible Public Assistance costs for those services under the 2006 interim rule. States avoid the administrative requirements associated with requesting an emergency declaration or requesting

additional designated areas to an existing emergency or major disaster declaration. As a result, FEMA is not required to review and analyze those declaration requests to make a recommendation to the President, thereby avoiding the administrative cost associated with such a review.

The Governor of the State requesting an emergency declaration from the President submits:

- Confirmation that the Governor has executed the State Emergency Plan;
- Preliminary damage assessment;
- State resources committed (a description of State and local resources that have already been committed) and an estimate of Federal assistance needed; and
- Certification that the State will comply with the cost-sharing requirements of the Stafford Act.

States incur costs to gather and submit this information to FEMA. FEMA estimates 33 burden hours for a State to prepare and submit a major disaster or emergency declaration.⁴ To determine that figure, FEMA assumes that the 33 burden hours include 9

hours of work spent by management staff and 24 hours by technical staff per major disaster or emergency declaration.

FEMA obtained the national average hourly wages for managerial (\$46.91) and technical (\$24.03) positions in State government from the Bureau of Labor Statistics.⁵ The managerial wage rate was for the "Chief Executive" position (standard occupational classification (SOC) code #: 11-1021). The technical wage rate was for the "First-Line Supervisors/Managers of Office and Administrative Support Workers" position (SOC code #43-1011) in State government. The hourly wage reflects only the direct cost of employment. FEMA multiplied the wage rates by 1.4 to derive the full employment costs for managerial (\$65.67) and technical (\$33.64) positions in State governments. Using these figures, FEMA estimates the cost savings experienced by States for not having to request a major disaster or emergency declaration is \$1,398. Table 1 details the cost to a State for submitting a major disaster or emergency declaration.

Activities	Managerial (\$65.67)	Technical (\$33.64)	Hours by activities
Data gathering for Governor's request	0	24	24
Preparing and submitting Governor's request	9	0	9
Total burden hours	9	24	33
Estimated cost savings	\$591	\$807	\$1,398

As part of FEMA's review of a declaration request, FEMA regional staff analyzes the information obtained by joint Federal, State, and local preliminary damage assessments. FEMA's regional summary, regional analysis, and recommendation includes a discussion of State and local resources and capabilities, and other assistance available to meet disaster-related needs. The Administrator of FEMA then submits a recommendation to the President and provides a copy of the Governor's request. FEMA takes the following steps in reviewing a major disaster or emergency declaration request:

- Federal officials, with the assistance of State, local, and Tribal officials, prepare a preliminary damage assessment.
- The FEMA Regional Administrator evaluates the damage and requirements

for Federal assistance and makes a recommendation to the FEMA Administrator.

- The FEMA Administrator reviews the Governor's request and the regional analysis and then makes a recommendation to the President.

FEMA estimates that it expends 48 burden hours in reviewing a major disaster or emergency declaration request. The 48 burden hours represent 9.6 hours spent by 5 management-level employees. This time is not consecutive, as FEMA often submits recommendations to the President on declaration requests within the span of a single day. These individuals represent program specialists, attorneys, and other senior officials, and the time includes work to review the Governor's request, generate FEMA's recommendation to the President, and activities that occur after the President

grants or denies the Governor's request (such as publishing a **Federal Register** Notice).

FEMA obtained the hourly wages for a managerial (GS 15, Step 5, \$65.62), position in the Federal government from the U.S. Office of Personnel Management.⁶ This hourly wage includes the locality pay for the area of Washington, DC and reflects only the direct cost of employment. The full employment cost is \$91.87. FEMA used the same factor of 1.4 to derive the full cost wage for Federal employees as it used for State employees.

FEMA estimates that the cost to FEMA to review a request for a major disaster or emergency declaration and to make a recommendation to the President is \$4,410 (= \$91.87 × 48). Therefore, the total administrative cost savings both to FEMA and State governments per major disaster or

⁴ 74 FR 36498 (2009), Collection of Information Notice, The Declaration Process. On an annual basis, FEMA estimates 56 respondents average 6 responses per year at 33 hours per response, totaling an estimated 11,088 burden hours per year for submission of a declaration request.

⁵ The Bureau of Labor Statistics (2009). "May 2007 National Industry-specific Occupational Employment and Wage Estimates, NAICS 999200—State Government (OES Designation)." http://www.bls.gov/oes/current/naics4_999200.htm#b43-0000.

⁶ U.S. Office of Personnel Management (2009). Salary Table 2009—Washington, DC Area, http://www.opm.gov/flsa/oca/09tables/html/dcb_h.asp.

emergency declaration is \$5,808 (= \$1,398 + \$4,410).

Hurricane Gustav in August 2008 has been the only disaster event since the 2006 interim rule was published that required assistance⁷ from host-States for sheltering and evacuation. As a result of Hurricane Gustav, FEMA provided approximately \$42 million to the nine host-States: Alabama, Arkansas, Indiana, Kentucky, Missouri, New Mexico, Oklahoma, Tennessee, and Texas.⁸ After this disaster event, which was FEMA's first opportunity to implement the 2006 interim rule, FEMA realized a need to clarify the eligibility of host-States and the reimbursement process. As a result, this final rule clarifies the eligibility of host-States by adding definitions for the terms "host-State" and "impact-State," and by revising the definition of "grantee." The final rule also provides additional information to clarify how a host-State receives a direct Federal grant from FEMA. The final rule clarifies that the host-State must submit a Standard Form SF-424 (Application for Federal Assistance) directly to FEMA to apply for reimbursement, that a host-State must enter into a FEMA/Host-State agreement (similar to a FEMA/State Agreement), and that a host-State is required to prepare any amendments to the State administrative plan to meet current policy guidance. However, these changes are not new requirements for grantees and this rule simply clarifies that these requirements apply to host-States. FEMA does not expect that these changes will result in any additional costs to the States.

FEMA also requires that a host-State must agree in writing to provide evacuation and/or sheltering support to the impact-State. FEMA refers to this agreement as the commitment letter, which the host-State submits to FEMA before the execution of the FEMA/Host-State Agreement. FEMA estimates that it will take one managerial employee one hour to draft and submit this letter. FEMA does not expect to use the host-

State sheltering provisions regularly. Federal host-State sheltering assistance has only been needed twice—for Hurricanes Katrina and Rita in 2005, and Hurricane Gustav in 2008. However, for the purpose of this economic analysis, FEMA conservatively estimates that it will be implemented once a year. Using Hurricane Gustav as a "typical" example, FEMA expects nine states to submit this letter when FEMA uses host-State sheltering. Therefore, the cost to comply with this new requirement will be \$591 (= 1 × 9 × \$65.67) per year.

FEMA also added a provision in this final rule that allows the agency to directly reimburse the regular-time salaries and benefits of a host-State's permanently employed personnel that perform evacuation and/or sheltering activities. These costs assist individuals who are not taxpayers in the host-State. In providing these services, a host-State incurs costs for a task that is not otherwise its responsibility, and therefore the Federal government should wholly compensate host-States for those services provided. Currently, a host-State can seek reimbursement for force account labor costs from the impact-State under a mutual aid agreement,⁹ but these costs are not reimbursable via a direct grant from FEMA pursuant to 44 CFR 206.228.

Under a mutual aid agreement, the State requesting assistance would reimburse the State providing assistance for eligible regular-time and overtime force account labor costs it incurred. The State requesting assistance would then seek reimbursement of those eligible costs from FEMA, subject to a cost share. Regular-time force account labor is reimbursable under a mutual aid agreement because FEMA considers the eligible costs incurred as contract labor. The new provision in this final rule would allow a host-State to be reimbursed regular-time force account labor costs when it provides assistance under a direct grant with FEMA. This is consistent with the eligible costs that

can be reimbursed for services provided under a mutual aid agreement. In addition, it avoids the administrative burden of a host-State seeking reimbursement for these costs through mutual aid from an impact-State when it is otherwise being reimbursed through a direct grant from FEMA. For Hurricane Gustav, FEMA has reimbursed approximately \$1 million for regular-time force account costs incurred by three (Alabama, New Mexico, and Oklahoma) of the nine host-States as of April 8, 2009. The other six states have not submitted costs, if any, to FEMA for reimbursement. The total amount obligated will likely increase once FEMA takes into account the regular-time force account costs for the other six host-States. However, this change in the regulation, which allows for straight-time reimbursement via direct grant, will not affect the amount of eligible Public Assistance funding; it merely streamlines the process by which funds reach the host-State. The cost implications of this rule are solely administrative in nature.

Although we have only experienced three disasters to date that have required the type of mass evacuation that called for host-State sheltering and evacuation assistance, to produce conservative estimates of the impact of the rule, FEMA assumes that there will be one large-scale disaster on an annual basis that will require host-States to provide sheltering and evacuation. If there is one large-scale disaster (and nine host-States per large-scale disaster), this rule will result in a reduction in administrative costs of \$39,690 to FEMA and \$11,991 to the States. Therefore, the annual impact of this rule is estimated at \$51,681 per year (= \$39,690 + \$11,991). Table 2 details the annual impact of the interim final rule. FEMA has determined that this rule will not have a significant economic impact of \$100 million or more per year.

TABLE 2—ANNUAL IMPACT OF THE RULE

Assumptions	<ul style="list-style-type: none"> • Number of large-scale disaster events that will require host-States to provide sheltering and evacuation support per year: 1. • Number of host-States per large-scale disaster (based on Hurricane Gustav): 9. 	
Administrative Cost per Major Disaster/Emergency Declaration.	FEMA -\$4,410	State, Local, and Tribal Governments -\$1,398

⁷ As noted above, evacuation and sheltering activities also occurred as a result of Hurricane Ike, but no financial assistance was required from FEMA for those purposes for that event.

⁸ This is an estimate as of April 8, 2009. FEMA continues to process reimbursement for the nine host-States for Hurricane Gustav.

⁹ Mutual aid agreements where one State or local government reimburses another State or local

government for services provided take many forms, including the Emergency Management Assistance Compact. Granting the consent of Congress to the Emergency Management Assistance Compact, Public Law 104-321, 110 Stat. 3877 (Oct. 19, 1996).

TABLE 2—ANNUAL IMPACT OF THE RULE—Continued

The Commitment Letter		\$65.67
Number of Large-Scale Disaster Events per Year	1	1
Number of host-States per Large-Scale Disaster	9	9
Administrative Cost per Year	-\$39,690 (= -\$4,410 × 1 × 9)	-\$11,991 [= (-\$1,398 + \$65.67) × 1 × 9]
Total		= -\$51,681 (- \$39,690 + - \$11,991)

C. Regulatory Flexibility Act (RFA)

The Regulatory Flexibility Act (RFA) (5 U.S.C. 601–612), and section 213(a) of the Small Business Regulatory Enforcement Fairness Act of 1996 (Pub. L. 104–121, 110 Stat. 847, 858–9 (March 29, 1996) (5 U.S.C. 601 note)) require that special consideration be given to the effects of proposed regulations on small entities. The RFA mandates that an agency conduct a RFA analysis when an agency is “required by section 553 * * * to publish general notice of proposed rulemaking for any proposed rule * * * 5 U.S.C. 603(a).” This rule finalizes an interim final rule and no initial or final regulatory flexibility analysis is required by the RFA.

D. National Environmental Policy Act (NEPA)

The National Environmental Policy Act, Public Law 91–190, 83 Stat. 852 (Jan. 1, 1970) (42 U.S.C. 4321 *et seq.*) (NEPA), as amended, requires the development of environmental impact statements in Federal actions “significantly affecting the quality of the human environment.” FEMA has adopted categorical exclusions from the preparation of an environmental assessment or environmental impact statement for essential assistance or emergency assistance. 44 CFR 10.8(d)(2)(xix)(B), (O); 44 CFR 10.8(d)(2)(ii). Actions taken or assistance provided under sections 403 and 502 of the Stafford Act are also statutorily excluded from NEPA review. 42 U.S.C. 5170b and 5192; 44 CFR 10.8(c)(1). The promulgation of this rule, accordingly, does not require the preparation of either an environmental assessment or an environmental impact statement as defined by NEPA.

E. Executive Order 12898, Environmental Justice

Executive Order 12898, Federal Actions to Address Environmental Justice in Minority Populations and Low-Income Populations, 59 FR 7629, February 16, 1994, requires agencies to incorporate environmental justice into policies and programs, and to conduct programs, policies, and activities that substantially affect human health or the

environment in a manner that ensures that those programs, policies, and activities do not have the effect of excluding persons from participation in those programs, denying persons the benefits of those programs, or subjecting persons to discrimination because of their race, color, or national origin. FEMA does not anticipate any action under this rule would have a disproportionately high or adverse human health and environmental effect on any segment of the population.

F. Congressional Review of Agency Rulemaking

FEMA will send this rule to the Congress and to the General Accounting Office under the Congressional Review of Agency Rulemaking Act, (Congressional Review Act), Public Law 104–121, 110 Stat. 873 (March 29, 1996) (5 U.S.C. 804). This rule is not a “major rule” within the meaning of the Congressional Review Act.

G. Unfunded Mandates Reform Act of 1995

The Unfunded Mandates Reform Act of 1995 (UMRA), Public Law 104–4, 109 Stat. 48 (March 22, 1995) (2 U.S.C. 1501 *et seq.*), requires agencies to prepare a written assessment of the costs, benefits, and other effects of proposed or final rules that include a Federal mandate likely to result in the expenditure by State, local, or tribal governments, in the aggregate, or by the private sector, of \$100 million or more, adjusted for inflation, in any one year. 2 U.S.C. 1532(a). FEMA has determined that this rule will not result in the expenditure by State, local, and Tribal governments, in the aggregate, nor by the private sector, of \$100 million or more in any one year as a result of a Federal mandate, and it will not significantly or uniquely affect small governments.

H. Executive Order 13132, Federalism

Executive Order 13132, Federalism, 64 FR 43255, August 4, 1999, sets forth principles and criteria that agencies must adhere to in formulating and implementing policies that have federalism implications, that is, regulations that have substantial direct effects on the States, or on the

distribution of power and responsibilities among the various levels of government. Federal agencies must closely examine the statutory authority supporting any action that would limit the policymaking discretion of the States, and to the extent practicable, must consult with State and local officials before implementing any such action. This rule involves only principles and criteria that affect the eligibility for and manner in which FEMA reimburses States, Tribes and political subdivisions for costs incurred in support of disaster recovery and does not have federalism implications under Executive Order 13132.

I. Paperwork Reduction Act of 1995

The Paperwork Reduction Act (PRA), 44 U.S.C. 3501 *et seq.*, requires government agencies to acquire approval from the Office of Management and Budget (OMB) for uses of forms and collections of information from the public. This final rule addresses the collection of four documents: The SF–424 Application for Federal Assistance, which is approved under OMB control number 1660–0025 until August 31, 2011, a State Administrative Plan, a FEMA/Host-State Agreement (similar to the FEMA/State Agreement), and the Commitment Letter. Collections of the State Administrative Plan, FEMA/Host-State Agreement, and Commitment Letter have not been approved by OMB under the Paperwork Reduction Act.

The PRA applies when a request for information is addressed to 10 or more persons. OMB has clarified that, “‘ten or more persons’ refers to the persons to whom a collection of information is addressed by the agency within any 12-month period.” 5 CFR 1320.3(c)(4). FEMA has determined, based on assessments of past disasters, that the likely number of respondents for host-State applications from non disaster-declared States in a 12-month period will not reach the threshold. FEMA estimates that there will be nine host-State applications and collections that would transpire in a 12-month period using Hurricane Gustav as an “average” disaster in which host-State sheltering is needed.

Collection of information from host-States is not expected to trigger the PRA because the number of host-State applicants is not likely to exceed nine. Therefore, FEMA has not sought approval from OMB for the collection of the State Administrative Plan, the FEMA/Host-State Agreement, or the Commitment Letter.

J. Executive Order 13175, Consultation and Coordination With Indian Tribal Governments

Executive Order 13175, Consultation and Coordination with Indian Tribal Governments, 65 FR 67249, November 9, 2000, applies to agency regulations that have Tribal implications, that is, regulations that have substantial direct effects on one or more Indian Tribes, on the relationship between the Federal Government and Indian Tribes, or on the distribution of power and responsibilities between the Federal Government and Indian Tribes. Under this Executive Order, to the extent practicable and permitted by law, no agency may promulgate any regulation that has Tribal implications, that imposes substantial direct compliance costs on Indian Tribal governments, and that is not required by statute, unless funds necessary to pay the direct costs incurred by the Indian Tribal government or the Tribe in complying with the regulation are provided by the Federal Government, or the agency consults with Tribal officials.

There is no substantial direct compliance cost associated with this rule; the Public Assistance Program provides funding to impact-States and host-States, including Tribal governments, for sheltering and evacuation activities. This rule would not affect the distribution of power or responsibilities of Tribal governments.

K. Executive Order 12630, Governmental Actions and Interference With Constitutionally Protected Property Rights

FEMA has reviewed this rule under Executive Order 12630, "Governmental Actions and Interference with Constitutionally Protected Property Rights" (53 FR 8859, Mar. 18, 1988) as supplemented by Executive Order 13406, "Protecting the Property Rights of the American People" (71 FR 36973, June 28, 2006). This rule will not affect the taking of private property or otherwise have taking implications under Executive Order 12630.

L. Executive Order 12988, Civil Justice Reform

FEMA has reviewed this rule under Executive Order 12988, "Civil Justice

Reform" (61 FR 4729, Feb. 7, 1996). This rule meets applicable standards to minimize litigation, eliminate ambiguity, and reduce burden.

List of Subjects in 44 CFR Part 206

Administrative practice and procedure, Coastal zone, Community facilities, Disaster assistance, Fire prevention, Grant programs—housing and community development, Housing, Insurance, Intergovernmental relations, Loan programs—housing and community development, Natural resources, Penalties, Reporting and recordkeeping requirements.

■ For the reasons discussed in the preamble, the Federal Emergency Management Agency amends 44 CFR part 206, subparts B and G, as follows:

PART 206—FEDERAL DISASTER ASSISTANCE

■ 1. The authority citation for part 206 continues to read as follows:

Authority: Robert T. Stafford Disaster Relief and Emergency Assistance Act, 42 U.S.C. 5121 through 5207; Reorganization Plan No. 3 of 1978, 43 FR 41943, 3 CFR, 1978 Comp., p. 329; Homeland Security Act of 2002, 6 U.S.C. 101; E.O. 12127, 44 FR 19367, 3 CFR, 1979 Comp., p. 376; E.O. 12148, 44 FR 43239, 3 CFR, 1979 Comp., p. 412; and E.O. 13286, 68 FR 10619, 3 CFR, 2003 Comp., p. 166.

§ 206.40 [Amended]

■ 2. In § 206.40, amend paragraph (b) by removing "disaster-affected" and adding "affected" in its place in the first sentence and by removing "A disaster-affected" and adding "An affected" in its place in the third sentence.

■ 3. In § 206.201—

■ a. Revise paragraph (e) to read as set forth below;

■ b. Redesignate paragraphs (g) through (l) as paragraphs (j) through (o); and

■ c. Add new paragraphs (g) through (i).

§ 206.201 Definitions

* * * * *

(e) *Grantee.* Grantee means the government to which a grant is awarded, and which is accountable for the use of the funds provided. The grantee is the entire legal entity even if only a particular component of the entity is designated in the grant award document. Generally, except as provided in § 206.202(f), the State for which the emergency or major disaster is declared is the grantee. However, an Indian Tribal government may choose to be a grantee, or it may act as a subgrantee under the State. If an Indian Tribal government is the grantee, it will assume the responsibilities of the "grantee" or "State" as described in this

part with respect to administration of the Public Assistance program.

* * * * *

(g) *Host-State.* A State or Indian Tribal government that by agreement with FEMA provides sheltering and/or evacuation support to evacuees from an impact-State. An Indian Tribal government may also be referred to as a "Host-Tribe."

(h) *Impact-State.* The State for which the President has declared an emergency or major disaster and that, due to a need to evacuate and/or shelter affected individuals outside the State, requests such assistance from FEMA pursuant to § 206.208.

(i) *Indian Tribal government* means any federally recognized governing body of an Indian or Alaska Native Tribe, band, nation, pueblo, village, or community that the Secretary of the Interior acknowledges to exist as an Indian Tribe under the Federally Recognized Tribe List Act of 1994, 25 U.S.C. 479a. This does not include Alaska Native corporations, the ownership of which is vested in private individuals.

* * * * *

■ 4. In § 206.202, revise paragraph (f) introductory text and paragraph (f)(1) to read as follows:

§ 206.202 Application procedures.

* * * * *

(f) *Exceptions.* The following are exceptions to the procedures and time limitations outlined in this section.

(1) *Host-State Evacuation and/or Sheltering.* (i) *General.* A grant to a host-State for sheltering and/or evacuation support is available under this section when an impact-State requests direct Federal assistance for sheltering and/or evacuation support pursuant to § 206.208. To receive this grant, a host-State must enter into a FEMA-Host-State Agreement, amend its State Administrative Plan pursuant to § 206.207, and submit a Standard Form SF424 *Application for Federal Assistance* directly to FEMA to apply for reimbursement of eligible costs for evacuating and/or sheltering individuals from an impact-State. Upon award, the host-State assumes the responsibilities of the "grantee" or "State" under this part with respect to its grant award.

(ii) *Force Account Labor Costs.* For the performance of eligible evacuation and sheltering support under sections 403 or 502 of the Stafford Act, the straight-time salaries and benefits of a host-State's permanently employed personnel are eligible for

reimbursement. This is an exception to § 206.228(a)(2).

* * * * *

■ 5. In § 206.208-

■ a. In the first complete sentence of paragraph (a), remove the phrase “sections 402(4), 403 or 407” and add the phrase “sections 402(1) and (4), 403, 407, 502(a)(1), (5) and (7)” in its place; and

■ b. Add paragraph (c)(3) to read as follows:

§ 206.208 Direct Federal assistance.

* * * * *

(c) * * *

(3) If an impact-State requests assistance in providing evacuation and sheltering support outside an impact-State, FEMA may directly reimburse a host-State for such eligible costs through a grant to a host-State under an impact-State’s declaration, consistent with § 206.202(f)(1). FEMA may award a grant to a host-State when FEMA

determines that a host-State has sufficient capability to meet some or all of the sheltering and/or evacuation needs of an impact-State, and a host-State agrees in writing to provide such support to an impact-State.

* * * * *

■ 6. In § 206.223, revise paragraphs (a)(1) and (a)(2) to read as follows:

§ 206.223 General work eligibility.

(a) * * *

(1) Be required as the result of the emergency or major disaster event;

(2) Be located within the designated area of a major disaster or emergency declaration, except that sheltering and evacuation activities may be located outside the designated area; and

* * * * *

■ 7. In § 206.431, revise the definition of “grantee” to read as follows:

§ 206.431 Definitions.

* * * * *

Grantee means the government to which a grant is awarded and which is accountable for the use of the funds provided. The grantee is the entire legal entity even if only a particular component of the entity is designated in the grant award document. Generally, the State for which the major disaster is declared is the grantee. However, an Indian tribal government may choose to be a grantee, or it may act as a subgrantee under the State. An Indian tribal government acting as a grantee will assume the responsibilities of a “state”, under this subpart, for the purposes of administering the grant.

* * * * *

Dated: November 9, 2009.

W. Craig Fugate,

Administrator, Federal Emergency Management Agency.

[FR Doc. E9-27883 Filed 11-19-09; 8:45 am]

BILLING CODE 9111-23-P

Proposed Rules

Federal Register

Vol. 74, No. 223

Friday, November 20, 2009

This section of the FEDERAL REGISTER contains notices to the public of the proposed issuance of rules and regulations. The purpose of these notices is to give interested persons an opportunity to participate in the rule making prior to the adoption of the final rules.

DEPARTMENT OF TRANSPORTATION

Federal Aviation Administration

14 CFR Part 39

[Docket No. FAA-2009-1069; Directorate Identifier 2009-NM-036-AD]

RIN 2120-AA64

Airworthiness Directives; Boeing Model 747-100, 747-100B, 747-100B SUD, 747-200B, 747-200C, 747-200F, 747-300, 747-400, 747-400D, 747-400F, and 747SR Series Airplanes

AGENCY: Federal Aviation Administration (FAA), Department of Transportation (DOT).

ACTION: Notice of proposed rulemaking (NPRM).

SUMMARY: The FAA proposes to supersede an existing airworthiness directive (AD) that applies to all Boeing Model 747-100, 747-100B, 747-100B SUD, 747-200B, 747-200C, 747-200F, 747-300, 747-400, 747-400D, 747-400F, and 747SR series airplanes. The existing AD currently requires repetitive inspections to find cracking of the web, strap, inner chords, and inner chord angle of the forward edge frame of the number 5 main entry door cutouts, and repair, if necessary. This proposed AD would expand the inspection areas to include the frame segment between stringers 16 and 23. This proposed AD would reinstate the repetitive inspections specified above for certain airplanes. This proposed AD would also require repetitive inspections for cracking of repairs. This proposed AD results from additional reports of cracks that have been found in the strap and inner chord of the forward edge frame of the number 5 main entry door cutouts, between stringers 16 and 23. We are proposing this AD to detect and correct such cracks. This condition, if not corrected, could cause damage to the adjacent body structure, which could result in depressurization of the airplane in flight.

DATES: We must receive comments on this proposed AD by January 4, 2010.

ADDRESSES: You may send comments by any of the following methods:

- **Federal eRulemaking Portal:** Go to <http://www.regulations.gov>. Follow the instructions for submitting comments.
- **Fax:** 202-493-2251.
- **Mail:** U.S. Department of Transportation, Docket Operations, M-30, West Building Ground Floor, Room W12-140, 1200 New Jersey Avenue, SE., Washington, DC 20590.
- **Hand Delivery:** U.S. Department of Transportation, Docket Operations, M-30, West Building Ground Floor, Room W12-140, 1200 New Jersey Avenue, SE., Washington, DC 20590, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

For service information identified in this proposed AD, contact Boeing Commercial Airplanes, Attention: Data & Services Management, P.O. Box 3707, MC 2H-65, Seattle, Washington 98124-2207; telephone 206-544-5000, extension 1; fax 206-766-5680; e-mail me.boecom@boeing.com; Internet <https://www.myboeingfleet.com>. You may review copies of the referenced service information at the FAA, Transport Airplane Directorate, 1601 Lind Avenue, SW., Renton, Washington. For information on the availability of this material at the FAA, call 425-227-1221 or 425-227-1152.

Examining the AD Docket

You may examine the AD docket on the Internet at <http://www.regulations.gov>; or in person at the Docket Management Facility between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays. The AD docket contains this proposed AD, the regulatory evaluation, any comments received, and other information. The street address for the Docket Office (telephone 800-647-5527) is in the **ADDRESSES** section. Comments will be available in the AD docket shortly after receipt.

FOR FURTHER INFORMATION CONTACT: Ivan Li, Aerospace Engineer, Airframe Branch, ANM-120S, FAA, Seattle Aircraft Certification Office, 1601 Lind Avenue, SW., Renton, Washington 98057-3356; telephone (425) 917-6437; fax (425) 917-6590.

SUPPLEMENTARY INFORMATION:

Comments Invited

We invite you to send any written relevant data, views, or arguments about this proposed AD. Send your comments to an address listed under the **ADDRESSES** section. Include "Docket No. FAA-2009-1069; Directorate Identifier 2009-NM-036-AD" at the beginning of your comments. We specifically invite comments on the overall regulatory, economic, environmental, and energy aspects of this proposed AD. We will consider all comments received by the closing date and may amend this proposed AD because of those comments.

We will post all comments we receive, without change, to <http://www.regulations.gov>, including any personal information you provide. We will also post a report summarizing each substantive verbal contact we receive about this proposed AD.

Discussion

On July 26, 2001, we issued AD 2001-16-02, amendment 39-12370 (66 FR 41440, August 8, 2001), for certain Boeing Model 747 series airplanes. That AD requires repetitive inspections to find cracking of the web, strap, inner chords, and inner chord angle of the forward edge frame of the number 5 main entry door cutouts, and repair, if necessary. That AD resulted from reports of cracks in the web, strap, inner chords, and inner chord angle of the forward edge frame of the number 5 main entry door cutouts. We issued that AD to detect and correct such cracks, which could result in severing of the frame, inability of the frame to react loads from the door stops, and consequent rapid depressurization of the airplane in flight.

Actions Since Existing AD Was Issued

Since we issued AD 2001-16-02, Boeing stated that production line numbers 1305 and on have an improved frame design and issued Boeing Service Bulletin 747-53A2450, Revision 3, dated July 24, 2003, which removed those line numbers from the effectivity. We referred to Boeing Alert Service Bulletin 747-53A2450, Revision 2, including Appendix A, dated January 4, 2001, as the appropriate source of service information for accomplishing the required actions of AD 2001-16-02. Based on Revision 3 of the service bulletin, we approved an alternative

method of compliance (AMOC), dated September 22, 2003, which allowed an alternative applicability to AD 2001-16-02. AD 2001-16-02 is applicable to all Model 747 airplanes except Model 747SP series airplanes; the AMOC allowed an alternative applicability of Model 747 airplanes, production line numbers 1 through 1304, excluding Model 747SP airplanes.

After approving the AMOC, we have since received reports of cracks in the left and right Station 2231 frame inner chord and strap between stringers 16 and 23. Subsequently, we have determined that line numbers 1305 and on are again subject to the unsafe condition.

Relevant Service Information

We have reviewed Boeing Alert Service Bulletin 747-53A2450, Revision 5, dated January 29, 2009. Revision 5 of the service bulletin also adds airplanes having production line numbers 1305 and on to the effectivity. Revision 5 describes the same repetitive inspections as those specified in Revision 2 of the service bulletin but it also identifies expanded inspection areas that include the frame segment between stringers 16 and 23.

Revision 5 also specifies repetitive post-repair inspections of the repaired frame segments for cracks and repair if necessary. The post-repair inspections include a detailed inspection, an open hole high frequency eddy current (HFEC) inspection, a surface HFEC

inspection, and a subsurface low frequency eddy current inspection, and corrective actions if necessary.

FAA’s Determination and Requirements of the Proposed AD

We have evaluated all pertinent information and identified an unsafe condition that is likely to develop on other airplanes of the same type design. For this reason, we are proposing this AD, which would supersede AD 2001-16-02 and would retain the requirements of the existing AD. This proposed AD would also require accomplishing the actions specified in Relevant Service Information described previously, except as discussed under “Differences Between the Proposed AD and Relevant Service Information.”

Difference Between the Proposed AD and Relevant Service Information

The service bulletin specifies to contact the manufacturer for instructions on how to repair certain conditions, but this proposed AD would require repairing those conditions in one of the following ways:

- Using a method that we approve; or
- Using data that meet the certification basis of the airplane, and that have been approved by an

Authorized Representative for the Boeing Commercial Airplanes Delegation Option Authorization Organization whom we have authorized to make those findings.

Change to Existing AD

This proposed AD would retain all requirements of AD 2001-16-02. Since AD 2001-16-02 was issued, the AD format has been revised, and certain paragraphs have been rearranged. As a result, the corresponding paragraph identifiers have changed in this proposed AD, as listed in the following table.

REVISED PARAGRAPH IDENTIFIERS

Requirement in AD 2001-16-02	Corresponding requirement in this proposed AD
paragraph (a)	paragraph (g)
paragraph (b)	paragraph (h)
paragraph (c)	paragraph (i)

Boeing Commercial Airplanes has received a Delegation Option Authorization (DOA). In paragraph (l) of this AD, we have referred to paragraph (m) of this AD to delegate the authority to approve an alternative method of compliance for any repair required by this AD to an Authorized Representative for the Boeing Commercial Airplanes DOA rather than a Designated Engineering Representative (DER).

Costs of Compliance

There are about 163 airplanes of the affected design in the worldwide fleet. The following table provides the estimated costs for U.S. operators to comply with this proposed AD.

ESTIMATED COSTS

Action	Work hours	Average labor rate per hour	Parts	Cost per airplane	Number of U.S.-registered airplanes	Fleet cost
Inspections (required by AD 2001-16-02).	16	\$80	None	\$1,280, per inspection cycle.	163	\$208,640, per inspection cycle.
Inspections (new proposed action).	28 depending on airplane configuration.	80	None	Up to \$2,240, per inspection cycle; depending on airplane configuration.	163	Up to \$365,120, per inspection cycle; depending on airplane configuration.

Authority for This Rulemaking

Title 49 of the United States Code specifies the FAA’s authority to issue rules on aviation safety. Subtitle I, Section 106, describes the authority of the FAA Administrator. Subtitle VII, Aviation Programs, describes in more detail the scope of the Agency’s authority.

We are issuing this rulemaking under the authority described in Subtitle VII, Part A, Subpart III, Section 44701, “General requirements.” Under that section, Congress charges the FAA with promoting safe flight of civil aircraft in

air commerce by prescribing regulations for practices, methods, and procedures the Administrator finds necessary for safety in air commerce. This regulation is within the scope of that authority because it addresses an unsafe condition that is likely to exist or develop on products identified in this rulemaking action.

Regulatory Findings

We have determined that this proposed AD would not have federalism implications under Executive Order 13132. This proposed AD would not have a substantial direct effect on the

States, on the relationship between the national Government and the States, or on the distribution of power and responsibilities among the various levels of government.

For the reasons discussed above, I certify that the proposed regulation:

1. Is not a “significant regulatory action” under Executive Order 12866;
2. Is not a “significant rule” under the DOT Regulatory Policies and Procedures (44 FR 11034, February 26, 1979); and
3. Will not have a significant economic impact, positive or negative, on a substantial number of small entities

under the criteria of the Regulatory Flexibility Act.

We prepared a regulatory evaluation of the estimated costs to comply with this proposed AD and placed it in the AD docket. See the **ADDRESSES** section for a location to examine the regulatory evaluation.

List of Subjects in 14 CFR Part 39

Air transportation, Aircraft, Aviation safety, Incorporation by reference, Safety.

The Proposed Amendment

Accordingly, under the authority delegated to me by the Administrator, the FAA proposes to amend 14 CFR part 39 as follows:

PART 39—AIRWORTHINESS DIRECTIVES

1. The authority citation for part 39 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701.

§ 39.13 [Amended]

2. The FAA amends § 39.13 by removing amendment 39–12370 (66 FR

41440, August 8, 2001) and adding the following new AD:

Boeing: Docket No. FAA–2009–1069; Directorate Identifier 2009–NM–036–AD.

Comments Due Date

(a) The FAA must receive comments on this AD action by January 4, 2010.

Affected ADs

(b) This AD supersedes AD 2001–16–02.

Applicability

(c) This AD applies to Boeing Model 747–100, 747–100B, 747–100B SUD, 747–200B, 747–200C, 747–200F, 747–300, 747–400, 747–400D, 747–400F, and 747SR series airplanes, certificated in any category, as identified in Boeing Alert Service Bulletin 747–53A2450, Revision 5, dated January 29, 2009.

Subject

(d) Air Transport Association (ATA) of America Code 53: Fuselage.

Unsafe Condition

(e) This AD results from additional reports of cracks that have been found in the strap and inner chord of the forward edge frame of the number 5 main entry door cutouts, between stringers 16 and 23. Based on these reports, we have determined that the frame

segment between stringers 16 and 23 is also susceptible to the unsafe condition. The Federal Aviation Administration is issuing this AD to detect and correct such cracks. This condition, if not corrected, could cause damage to the adjacent body structure, which could result in depressurization of the airplane in flight.

Compliance

(f) You are responsible for having the actions required by this AD performed within the compliance times specified, unless the actions have already been done.

Restatement of AD 2001–16–02, With New Service Information

Repetitive Inspections for Frame Segment Between Stringers 23 and 31 (No Terminating Action)

(g) For airplanes having line numbers 1 through 1304 inclusive: Inspect the airplane for cracks between stringers 23 and 31 per Boeing Alert Service Bulletin 747–53A2450, Revision 2, including Appendix A, dated January 4, 2001; or Boeing Alert Service Bulletin 747–53A2450, Revision 5, dated January 29, 2009; at the later of the times specified in either paragraph (h) or (i) of this AD, per Table 1, as follows. After the effective date of this AD, use only Revision 5 of Boeing Service Bulletin 747–53A2450.

TABLE 1—INSPECTION REQUIREMENTS

Type of inspection	Area to inspect
(1) Detailed Visual	Strap inner chords forward and aft of the web, and exposed web adjacent to the inner chords on station 2231 frame from stringer 23 through 31 per Figure 5 or Figure 6 of the service bulletin, as applicable.
(2) Surface High Frequency Eddy Current (HFEC)	Station 2231 inner chord angles at lower main sill interface per Figure 5 or Figure 6 of the service bulletin, as applicable.
(3) Open Hole HFEC	Station 2231 frame fastener locations per Figures 4 and 7, and either Figure 5 or 6 of the service bulletin, as applicable.
(4) Surface HFEC	Around fastener locations on station 2231 inner chords from stringer 23 through 31 per Figure 5 or Figure 6 of the service bulletin, as applicable.
(5) Low Frequency Eddy Current	Station 2231 frame strap in areas covered by the reveal per Figure 5 or Figure 6 of the service bulletin, as applicable.

(h) Do the inspections specified in paragraph (g) of this AD at the applicable times specified in paragraph (h)(1) or (h)(2) of this AD. Repeat the inspections at intervals not to exceed 3,000 flight cycles.

(1) Do the inspections per Table 1 of this AD at the applicable time specified in the logic diagram in Figure 1 of Boeing Alert Service Bulletin 747–53A2450, Revision 2, including Appendix A, dated January 4, 2001. Where the compliance time in the logic diagram specifies a compliance time beginning, “from receipt of this service bulletin,” this AD requires that the compliance time begin “after September 12, 2001 (the effective date of AD 2001–16–02).”

(2) After the effective date of this AD, do the inspections per Table 1 of this AD at the applicable compliance time specified in paragraph 1.E., “Compliance” of the Boeing Alert Service Bulletin 747–53A2450, Revision 5, dated January 29, 2009. Where the compliance time in the service bulletin specifies a compliance time beginning, “after the date on Revision 2 of this service

bulletin,” this AD requires that the compliance time begin “after September 12, 2001 (the effective date of AD 2001–16–02).”

(i) Within 3,000 flight cycles after accomplishment of the inspections specified in Figure 1 of Boeing Alert Service Bulletin 747–53A2450, dated May 4, 2000, or Revision 1, dated July 6, 2000, repeat the inspections at intervals not to exceed 3,000 flight cycles.

Note 1: There is no terminating action currently available for the inspections required by paragraph (g) of this AD.

Note 2: Where there are differences between the AD and the alert service bulletin, the AD prevails.

New Requirements of This AD

Additional Repetitive Inspections (For Frame Segment Between Stringers 16 and 23)

(j) For all airplanes: Before the accumulation of 16,000 total flight cycles, or

within 1,500 flight cycles after the effective date of this AD, whichever occurs later, do a detailed inspection, an open hole high frequency eddy current (HFEC) inspection, a surface HFEC inspection, and a subsurface low frequency eddy current (LFEC) inspection for cracking of the forward edge frame of the number 5 main entry door cutouts, at station 2231, between stringers 16 and 23; in accordance with the Accomplishment Instructions of Boeing Alert Service Bulletin 747–53A2450, Revision 5, dated January 29, 2009. Repeat the inspections thereafter at intervals not to exceed 3,000 flight cycles.

Repetitive Inspections for Line Numbers 1305 and On (For Frame Segment Between Stringers 23 and 31)

(k) For airplanes having line numbers 1305 and on: Before 16,000 total flight cycles or within 1,500 flight cycles after the effective date of this AD, whichever occurs later, do a detailed inspection, an open hole HFEC inspection, a surface HFEC inspection, and a

subsurface LFEC inspection for cracking of the forward edge frame of the number 5 main entry door cutouts, at station 2231, between stringers 23 and 31; in accordance with the Accomplishment Instructions of Boeing Alert Service Bulletin 747-53A2450, Revision 5, dated January 29, 2009. Repeat the inspections thereafter at intervals not to exceed 3,000 flight cycles.

Corrective Action

(l) If any crack is found during any inspection required this AD, before further flight repair the crack per a method approved by the Manager, Seattle Aircraft Certification Office (SACO), FAA; Per data meeting the type certification basis of the airplane approved by a Boeing Company Designated Engineering Representative who has been authorized by the Manager, Seattle ACO, to make such findings; or in accordance with Boeing Alert Service Bulletin 747-53A2450, Revision 5, dated January 29, 2009. For a repair method to be approved by the Manager, Seattle ACO, as required by this paragraph, the approval letter must specifically reference this AD. As of the effective date of this AD, repair the crack using a method approved in accordance with the procedures specified in paragraph (o) of this AD.

Post-Repair Inspections

(m) Except as required by paragraph (n) of this AD, for airplanes on which the forward edge frame of the number 5 main entry door cutouts, at station 2231, between stringers 16 and 31, is repaired in accordance with Boeing Alert Service Bulletin 747-53A2450: Within 3,000 flight cycles after doing the repair or within 1,500 flight cycles after the effective date of this AD, whichever occurs later, do the detailed, LFEC, and HFEC inspections of the repaired area for cracks in accordance with the Accomplishment Instructions of Boeing Alert Service Bulletin 747-53A2450, Revision 5, dated January 29, 2009. If no cracking is found, repeat the inspections thereafter at intervals not to exceed 3,000 flight cycles. If any crack is found, before further flight, repair using a method approved in accordance with the procedures specified in paragraph (o) of this AD. Doing the inspections specified in paragraph (m) of this AD terminates the repetitive inspections required by paragraphs (g), (h), (i), (j), and (k) of this AD for the repaired area.

(n) For any frame that is repaired in accordance with a method other than the Accomplishment Instructions of Boeing Alert Service Bulletin 747-53A2450, Revision 5, dated January 29, 2009, do the inspection in accordance with a method approved in accordance with the procedures specified in paragraph (o) of this AD.

Alternative Methods of Compliance (AMOCs)

(o)(1) The Manager, Seattle Aircraft Certification Office (SACO), FAA, has the authority to approve AMOCs for this AD, if requested using the procedures found in 14 CFR 39.19. Send information to ATTN: Ivan Li, Aerospace Engineer, Airframe Branch, ANM-120S, FAA, Seattle Aircraft Certification Office, 1601 Lind Avenue, SW.,

Renton, Washington 98057-3356; telephone (425) 917-6437; fax (425) 917-6590; Or, e-mail information to 9-ANM-Seattle-ACO-AMOC-Requests@faa.gov.

(2) To request a different method of compliance or a different compliance time for this AD, follow the procedures in 14 CFR 39.19. Before using any approved AMOC on any airplane to which the AMOC applies, notify your principal maintenance inspector (PMI) or principal avionics inspector (PAI), as appropriate, or lacking a principal inspector, your local Flight Standards District Office. The AMOC approval letter must specifically reference this AD.

(3) An AMOC that provides an acceptable level of safety may be used for any repair required by this AD, if it is approved by an Authorized Representative for the Boeing Commercial Airplanes Delegation Option Authorization Organization who has been authorized by the Manager, Seattle ACO, to make those findings. For a repair method to be approved, the repair must meet the certification basis of the airplane, and the approval must specifically refer to this AD.

(4) AMOCs approved previously in accordance with AD 2001-16-02, amendment 39-12370, are approved as AMOCs for the corresponding provisions of paragraphs (g), (h), (i), and (l) of this AD.

Issued in Renton, Washington, on November 6, 2009.

Ali Bahrami,

Manager, Transport Airplane Directorate, Aircraft Certification Service.

[FR Doc. E9-27963 Filed 11-19-09; 8:45 am]

BILLING CODE 4910-13-P

DEPARTMENT OF TRANSPORTATION

Federal Aviation Administration

14 CFR Parts 91, 119, 125, 133, 137, 141, 142, 145 and 147

[Docket No. FAA-2008-1154; Notice No. 09-13]

RIN 2120-AJ36

Restrictions on Operators Employing Former Flight Standards Service Aviation Safety Inspectors

AGENCY: Federal Aviation Administration (FAA), DOT.

ACTION: Notice of proposed rulemaking (NPRM).

SUMMARY: This proposed rule would prohibit any person holding a certificate to conduct certain operations from knowingly employing, or making a contractual arrangement with, certain individuals to act as an agent or a representative of the certificate holder in any matter before the FAA under certain conditions. These restrictions would apply if the individual, in the preceding 2-year period: Directly served as, or was directly responsible for the

oversight of, a Flight Standards Service Aviation Safety Inspector; and had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder. This proposed rule would also apply to persons who own or manage fractional ownership program aircraft that are used to conduct operations under specific regulations described in this document. This proposed rule would establish these restrictions to prevent potential organizational conflicts of interests which could adversely affect aviation safety.

DATES: Send your comments to reach us on or before February 18, 2010.

ADDRESSES: You may send comments identified by Docket Number FAA-2008-1154 using any of the following methods:

- *Federal eRulemaking Portal:* Go to <http://www.regulations.gov> and follow the online instructions for sending your comments electronically.

- *Mail:* Send comments to Docket Operations, M-30; U.S. Department of Transportation, 1200 New Jersey Avenue, SE., Room W12-140, West Building Ground Floor, Washington, DC 20590-0001.

- *Hand Delivery or Courier:* Take comments to Docket Operations in Room W12-140 of the West Building Ground Floor at 1200 New Jersey Avenue, SE., Washington, DC, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

- *Fax:* Fax comments to Docket Operations at 202-493-2251.

For more information on the rulemaking process, see the **SUPPLEMENTARY INFORMATION** section of this document.

Privacy: We will post all comments we receive, without change, to <http://www.regulations.gov>, including any personal information you provide. Using the search function of our docket Web site, anyone can find and read the electronic form of all comments received into any of our dockets, including the name of the individual sending the comment (or signing the comment for an association, business, labor union, etc.). You may review DOT's complete Privacy Act Statement in the **Federal Register** published on April 11, 2000 (65 FR 19477-78) or you may visit <http://DocketsInfo.dot.gov>.

Docket: To read background documents or comments received, go to <http://www.regulations.gov> at any time and follow the online instructions for accessing the docket, or, go to the Docket Operations in Room W12-140 of the West Building Ground Floor at 1200 New Jersey Avenue, SE., Washington,

DC, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

FOR FURTHER INFORMATION CONTACT: For technical questions concerning this proposed rule, contact Nancy Lauck Claussen, Air Transportation Division, AFS-200, Federal Aviation Administration, 800 Independence Avenue, SW., Washington, DC 20591; telephone (202) 267-8166, e-mail Nancy.L.Claussen@faa.gov. For legal questions concerning this proposed rule, contact Paul G. Greer, Federal Aviation Administration, Office of the Chief Counsel, 800 Independence Avenue, SW., Washington, DC 20591; Telephone: 202-267-3073, e-mail: Paul.G.Greer@faa.gov.

SUPPLEMENTARY INFORMATION: Later in this preamble under the Additional Information section, we discuss how you can comment on this proposal and how we will handle your comments. Included in this discussion is related information about the docket, privacy, and the handling of proprietary or confidential business information. We also discuss how you can get a copy of related rulemaking documents.

Authority for This Rulemaking

The FAA's authority to issue rules on aviation safety is found in Title 49 of the United States Code. Subtitle I, Section 106 describes the authority of the FAA Administrator, to include the authority to issue, rescind, and revise regulations. Subtitle VII, Aviation Programs, describes in more detail the scope of the agency's authority. This rulemaking is promulgated under the authority described in Subtitle VII, Part A, Chapter 447, Safety Regulation. Under Section 44701(a) the FAA is charged with promoting the safe flight of civil aircraft in air commerce by prescribing regulations and minimum standards for other practices, methods, and procedures necessary for safety in air commerce and national security.

I. Background

On March 5, 2008, the FAA proposed a \$10.2 million civil penalty against a major airline for operating 46 airplanes without performing mandatory inspections for fuselage fatigue cracking. The FAA alleged that the airline operated 46 Boeing 737 airplanes on almost 60,000 flights from June 2006 to March 2007 while failing to comply with an existing FAA Airworthiness Directive (AD) that required repetitive inspections of certain fuselage areas to detect fatigue cracking.

After investigating these events, the FAA took steps to improve its safety systems and strengthen regulations to

minimize the risk of recurrence of these or similar events. One such step was to toughen Aviation Safety Inspector (ASI) post employment restrictions to prevent conflicts of interest. This proposed rulemaking would establish restrictions on persons employing former Flight Standards Service (AFS) ASIs and those responsible for their oversight.

Review of FAA's Safety Oversight of Airlines and Use of Regulatory Partnership Programs

On June 30, 2008, the Department of Transportation (DOT) Office of Inspector General issued its review of the FAA's oversight of airlines and use of regulatory partnership programs. The report concluded that the FAA Certificate Management Office overseeing the airline that failed to perform the required inspections had developed an overly collaborative relationship with the airline. That relationship allowed repeated self-disclosures of AD violations without ensuring that the airline had developed a comprehensive solution for those reported safety problems.

The report noted that the Regulatory Compliance Manager for the airline was a former FAA ASI who reported directly to the FAA Principal Maintenance Inspector assigned to the airline when the former ASI worked for the FAA. The former employee had become a manager at the airline two weeks after leaving the FAA. In his new position at the airline, the former ASI served as the liaison between the carrier and the FAA and managed both the airline's AD Compliance Program and its Voluntary Disclosure Reporting Program.

The report also concluded that the overly collaborative relationship with the air carrier occurred because the FAA lacked effective management controls over its partnership program. The report stated that effective management controls would address: (1) Adequate segregation of duties; (2) the avoidance of potential conflicts of interests among its employees dealing with the carrier; and (3) verification of the propriety and integrity of corrective actions taken.

The report recommended that the FAA should enhance management controls by implementing post-employment guidance that includes a "cooling-off" period to prohibit an air carrier from hiring an FAA ASI who previously inspected the air carrier from acting in any type of liaison capacity between that air carrier and the FAA. A full copy of the report is contained in the docket for this rulemaking.

Proposed Legislation

On July 15, 2008, Congressman James L. Oberstar introduced the Aviation Safety Enhancement Act of 2008 (H.R. 6493). Section 4 of the proposed legislation included post employment restrictions for AFS ASIs. The proposed legislation would prohibit certificate holders from employing or contracting with a former AFS ASI or other person with certificate holder oversight responsibilities to represent that certificate holder in any matter before the FAA for a 2-year period after leaving the FAA. The proposed legislation was passed unanimously by the House of Representatives on July 22, 2008. However, it was not subsequently passed by the Senate prior to adjournment of the 110th Congress.

On May 21, 2009, the House of Representatives passed the FAA Reauthorization Act of 2009 (H.R. 915). Section 333 of the proposed legislation contains language identical to that proposed earlier in section 4 of the Aviation Safety Enhancement Act of 2008. Similar provisions are also found in Section 513 of the FAA Air Transportation Modernization and Safety Improvement Act which was introduced in the Senate on July 14, 2009 (S. 1451).

Managing Risks in Civil Aviation: A Review of the FAA's Approach to Safety

On May 1, 2008, former Secretary of Transportation, Mary E. Peters, appointed an independent review team to examine the FAA's safety culture and its implementation of safety management systems. She asked the team to prepare recommendations that would optimize the FAA's regulatory effectiveness. On September 2, 2008, the independent review team issued its report titled, "Managing Risks in Civil Aviation: A Review of the FAA's Approach to Safety." A full copy of the report may be found in the docket for this rulemaking.

The report stated that "[t]he FAA, like all other regulators, faces the danger of regulatory capture. Capture occurs when a regulatory agency draws so close to those with whom it deals on a daily basis (*i.e.* the regulated) that the agency ends up elevating their concerns at the expense of the agency's core mission. One feature of the FAA's current structure has the potential to increase this risk: the inspection teams are mostly organized around airlines, rather than cutting across multiple airlines and organizing around some other dimension, like geography, or type of plane. Most regulatory agencies organize by broad functional areas (like

enforcement, education, etc.) and also by geography; as a result, any one inspector normally deals with multiple corporations on a daily basis. By contrast, the majority of FAA airline inspectors are assigned to a specific Certificate Management Office, and deal with one airline, full time, and for many years at a stretch * * *

Further, the report stated that the panel does “understand the enhanced risk of regulatory capture that long-standing relationships between regulators and regulated entities might produce. We understand also the countervailing value in accumulating a detailed knowledge of a specific airline’s operations. We believe that any enhanced risk of capture can be properly mitigated * * *” This proposal would serve to mitigate the risks associated with regulatory capture by establishing a “cooling off” period for former AFS ASIs, while allowing AFS ASIs assigned to a specific operator to acquire the level of knowledge necessary to conduct effective oversight.

Current Post Employment Restrictions of Former Employees

Section 207(a)(1) of Title 18, United States Code (18 U.S.C.) generally places a permanent restriction on former executive branch employees (including FAA employees) regarding their ability to represent any other person in connection with a particular matter in which the United States government has a direct and substantial interest and in which that person participated personally and substantially.

In addition, it also places a 2-year restriction on those same former employees concerning their ability to represent any other person in connection with a particular matter in which the U.S. government has a direct and substantial interest and which that person knew, or reasonably should have known, was pending under his or her official responsibility within 1 year of their separation. Section 207(a)(2) basically restricts a person’s ability to represent an entity before the FAA on particular matters in which they were involved. It does not limit a former FAA employee’s ability to obtain employment with any entity.

Current FAA Flight Standards Service Policy

In order to minimize the influence of a particular carrier on the FAA, AFS policy provides for a 2-year “cooling off” period for newly employed ASIs, which prohibits them from having certificate management responsibilities for their former aviation employer. The proposed rule would not change this

longstanding FAA policy. It would, however, create a corresponding requirement applicable to operators who seek to employ certain former FAA ASIs and those responsible for their oversight. Current AFS policy was first set forth in a memorandum, dated May 10, 1990 from the Director, Flight Standards Service (AFS–1) to all AFS staff. It was reiterated in two subsequent AFS–1 memoranda dated July 18, 1996 and April 9, 2008.

II. Discussion of the Proposal

The FAA has considered the proposed legislation, the current ethics regulations, and the recommendations raised in the previously discussed reports. Although 18 U.S.C. 207 establishes some general restrictions for Federal employees after they leave government service, the FAA proposes additional safety-based restrictions on certificate holders conducting operations under parts 121, 125, 133, 135, 137, 141, 142, 145 or 147. (Parts 121, 125, 133, 135, 137, 141, 142, 145 and 147 apply to: Air carriers conducting domestic, flag, or supplemental operations; operators of airplanes having a seating capacity of 20 or more passengers or a maximum payload capacity of 6,000 pounds or more; rotorcraft external-load operations; commuter and on-demand operations; agricultural aircraft operations; pilot schools; training centers; repair stations; and aviation maintenance technician schools, respectively). The proposed restrictions would apply if the certificate holder employs (or makes a contractual arrangement with) a former AFS ASI or a person directly responsible for the oversight of the ASI and either person had direct responsibility to inspect, or oversee the inspection of, the certificate holder. The proposed restrictions would also apply to persons who own or manage fractional ownership program aircraft that are used to conduct operations using fractional ownership program aircraft under subpart K of part 91.

The proposed rule would address a significant concern highlighted in the report issued by the independent review team—the need to address “regulatory capture” to mitigate risk. Although the report did not specifically recommend a “cooling off period” for former AFS ASIs after they leave the FAA, this proposed rule is consistent with the FAA’s commitment to take steps to mitigate the risk that a current FAA employee may engage inappropriately with a regulated party. This proposed rule would establish restrictions on these operators that exceed current

restrictions applicable to most businesses who hire former Federal employees.

The proposed rule would specifically apply to AFS ASIs and those persons directly responsible for their oversight. The FAA considers an AFS ASI to be a properly credentialed individual who holds FAA Form 110A and is authorized under the provisions of 49 U.S.C. 40113 to perform inspections and investigations.

This proposal would prohibit any person conducting operations under parts 121, 125, 133, 135, 137, 141, 142, 145, 147, or subpart K of part 91 from knowingly employing or contracting with a former AFS ASI (Avionics, Cabin Safety, Dispatch, Maintenance, or Operations), or other person with oversight responsibilities for that operator, to represent that operator in any matter before the FAA. These restrictions would apply if the person, in the preceding 2-year period has served as, or was directly responsible for the oversight of, an AFS ASI and had the direct responsibility to inspect, or oversee the inspection of, the operator. Operators, however, would only be restricted from employing or making a contractual arrangement with former AFS ASIs who had inspection or oversight responsibilities for that particular operator. The proposed rule would not apply if an operator employs or contracts with an AFS ASI who had inspection or oversight responsibilities for another operator that has (or may have had) a marketing, code share, business partnership, or similar relationship with the operator. The FAA contends that these often temporary business arrangements between separate and distinct operators do not warrant the application of the restrictions set forth in this proposed rule.

The FAA would consider the proposed restrictions to apply only to those operators employing persons who had an office location in a Flight Standards District Office or a Certificate Management Office with oversight responsibilities for the operator (e.g. Office Managers, Assistant Office Managers, Unit Supervisors, and Aviation Safety Inspectors). AFS ASIs directly engaged in certificate management typically develop extensive knowledge of an operator’s practices. They also develop close working relationships with other AFS ASIs with whom they share direct oversight responsibilities for that particular operator. The FAA believes that aviation safety could be compromised if a former AFS ASI, acting on behalf of the operator, is able to exert undue influence on current

FAA employees with whom he or she had established close working relationships while working at a Flight Standards District Office or a Certificate Management Office. This proposed rule would address these concerns.

The intent of the proposed rule is not to affect employment relationships entered into prior to the effective date of this rule. Therefore, the proposed rule would not affect any operator currently employing a former AFS ASI in any capacity. A former AFS ASI hired by an operator prior to the effective date of the rule may continue to act as a representative of that operator in any matter before the FAA. The proposal would only prohibit an operator from hiring or making a contractual arrangement with an individual to act as a representative of the operator in any matter before the FAA if the individual had direct certificate oversight responsibilities for that operator in the previous 2 years and that employment commenced on or after the effective date of the rule.

The following examples further explain the provisions of this proposed rule:

(1) A former AFS ASI who was assigned direct oversight responsibilities for air carrier X, who is currently working for air carrier X in any position which includes representing air carrier X to the FAA prior to the effective date of the rule, may continue in that position.

(2) In order to be hired by training center A for a position which includes representing the training center in any matter before the FAA, on or after the effective date of the rule, the former AFS ASI must be able to look back over the 2 years preceding his or her being hired by training center A and determine that during that preceding 2 years the former ASI was not assigned oversight responsibilities for training center A.

(3) A former AFS ASI who was assigned direct oversight responsibilities for repair station Q may immediately go to work for any repair station other than repair station Q in any position.

(4) A former AFS ASI who was assigned direct oversight responsibilities for aviation maintenance technician school Q may immediately go to work for aviation maintenance technician school Q in any position that does not require representing aviation maintenance technician school Q to the FAA.

The FAA has many employees other than AFS ASIs with direct oversight responsibilities for various regulated entities. However, after considering the

potential safety risks and in light of the findings of recent reports, the FAA proposes only to establish restrictions for operators who employ or make contractual arrangements with former AFS ASIs who previously had direct oversight responsibility for that operator. This action is necessary to address the development of overly collaborative relationships that may occur during routine AFS surveillance of certain operators. Such relationships occur when a regulatory agency draws so close to those with whom it deals on a daily basis (*i.e.* the regulated) that the agency ends up elevating their concerns at the expense of the agency's core safety mission.

The proposed rule would not prohibit an operator from employing a former AFS ASI to serve in any capacity if that former AFS ASI did not have direct oversight responsibilities for that operator within the previous 2 years. The FAA acknowledges that the skills and expertise former FAA employees bring to the aviation industry are valuable and enhance safety. The agency notes that there are many employment opportunities for former FAA employees that would not be restricted by the proposed rule. There are numerous positions that would typically not require representing an operator to the FAA, but would take advantage of the unique skill set that a former AFS ASI would possess. For example, under most circumstances, working in operations or maintenance as an aircraft dispatcher, flight attendant, maintenance technician, training instructor, or pilot would not be prohibited by the proposed rule. As long as the covered employee did not act as an agent or representative of the operator before the FAA, the employee would be able to provide highly beneficial expertise and enhance safety in areas such as safety management systems, continuous analysis programs, operational training programs, crewmember training programs, maintenance training programs, aircraft dispatcher training programs, ETOPs (Extended Range Operations), operational control systems, maintenance, accident investigation, and regulatory compliance.

Based on recent events and reviews of the FAA's safety oversight programs, the agency has determined that the proposed restrictions set forth in this notice must be placed on the employment of persons holding certain agency positions that could lead to organizational conflicts of interest. This proposed rule would enhance the FAA's ability to properly perform its safety

mission and ensure the integrity of the programs administered by the FAA.

During the development of this proposal, the FAA considered a prohibition on operators employing a former AFS ASI to serve in any capacity if that former AFS ASI had direct oversight responsibilities for that operator within the previous 2 years. The FAA determined that as long as the former AFS ASI did not act as an agent or a representative of the operator in any matter before the FAA, serving in other positions with the operator (e.g. aircraft dispatcher, flight attendant, maintenance technician, pilot, or training instructor) would not be prohibited by the proposed rule. The FAA also consulted with representatives of the Professional Aviation Safety Specialists (PASS) to determine their views on the scope of the restrictions; a record of that meeting is available in the docket. The FAA is seeking specific comments on whether the prohibition on operators should be more restrictive than as proposed.

In addition, the agency is proposing the period of restriction as a sliding timeline, with the 2-year clock starting on the last day the AFS ASI or supervisor had direct responsibility for oversight of the operator. The FAA is also seeking specific comments on whether the prohibition should instead begin on the date the individual's employment by the FAA is terminated.

Paperwork Reduction Act

The Paperwork Reduction Act of 1995 (44 U.S.C. 3507(d)) requires that the FAA consider the impact of paperwork and other information collection burdens imposed on the public. We have determined that there is no new information collection requirement associated with this proposed rule.

International Compatibility

In keeping with U.S. obligations under the Convention on International Civil Aviation, it is FAA policy to comply with International Civil Aviation Organization (ICAO) Standards and Recommended Practices to the maximum extent practicable. The FAA has determined that there are no ICAO Standards and Recommended Practices that correspond to these proposed regulations.

III. Regulatory Evaluation, Regulatory Flexibility Determination, International Trade Impact Assessment, and Unfunded Mandates Assessment

Changes to Federal regulations must undergo several economic analyses. First, Executive Order 12866 directs that each Federal agency shall propose or

adopt a regulation only upon a reasoned determination that the benefits of the intended regulation justify its costs. Second, the Regulatory Flexibility Act of 1980 (Pub. L. 96–354) requires agencies to analyze the economic impact of regulatory changes on small entities. Third, the Trade Agreements Act (Pub. L. 96–39) prohibits agencies from setting standards that create unnecessary obstacles to the foreign commerce of the United States. In developing U.S. standards, this Trade Act requires agencies to consider international standards and, where appropriate, that they be the basis of U.S. standards. Fourth, the Unfunded Mandates Reform Act of 1995 (Pub. L. 104–4) requires agencies to prepare a written assessment of the costs, benefits, and other effects of proposed or final rules that include a Federal mandate likely to result in the expenditure by State, local, or tribal governments, in the aggregate, or by the private sector, of \$100 million or more annually (adjusted for inflation with base year of 1995). This portion of the preamble summarizes the FAA’s analysis of the economic impacts of this proposed rule.

Department of Transportation Order DOT 2100.5 prescribes policies and procedures for simplification, analysis, and review of regulations. If the expected cost impact is so minimal that a proposed or final rule does not warrant a full evaluation, this order permits that a statement to that effect and the basis for it to be included in the preamble if a full regulatory evaluation of the cost and benefits is not prepared. Such a determination has been made for this proposed rule. The reasoning for this determination follows:

The proposed rule would prohibit any of the previously mentioned certificate holders from employing or making a contractual agreement with an individual who was responsible for the direct oversight of an operator as an FAA AFS ASI or who had responsibility to inspect or oversee the inspections of the operator during the preceding 2 years. This proposed rule would also apply to fractional owners or fractional ownership program managers who conduct operations under subpart K of part 91. These proposed restrictions would prevent potential organizational conflicts of interest that could adversely affect aviation safety or create a perception of such conflicts of interest. The proposed rule would have minimal economic impact. The affected former FAA employees would be allowed to work for other operators for which they did not have direct oversight responsibilities. In addition, they would be able to work for operators for which

they did have direct oversight responsibilities provided that they do not represent the operator in any matter before the FAA.

Who Would Be Potentially Affected by This Proposed Rule

This proposal would affect current and future AFS ASIs and persons responsible for their oversight who would perform work after the effective date of the rule for an operator for which they had direct oversight responsibilities when employed by the FAA. In addition, this proposal would affect operators that would have hired former FAA employees who had oversight responsibilities for those operators.

Potential Benefits and Costs

The benefits associated with this proposal would arise from preventing potential organizational conflicts of interest. There would also be benefits from reducing the potential public perception that: (1) A current AFS ASI who was offered post-FAA employment with an operator he or she regulates could compromise current aviation safety; and (2) future aviation safety could be compromised if a former FAA employee working for an operator would be able to exert undue influence on current FAA employees with whom he or she had established close working relationships. This prohibition would also apply to the more likely case of former AFS ASIs who would become consultants to the operator. By prohibiting such a close relationship between a former AFS ASI and the operator for which he or she had direct oversight responsibilities, the potential for an overly collaborative relationship leading to a possible lapse in safety standards would be avoided, increasing the public’s confidence in the safety and integrity of the FAA inspection system. Such benefits cannot be quantified.

The proposed rule would also create some minor inefficiencies. In general, an operator can benefit from employing a former AFS ASI because that ASI knows more about FAA processes than someone who had not worked for the FAA. In addition, that ASI would know more about the operator than some other former AFS ASI. Further, a former AFS ASI from a specific Flight Standards District Office or Certificate Management Office will have greater knowledge about that office (as well as be better acquainted with the people in that office) than would a former AFS ASI from a different office.

For example, some operators may believe that employing a former AFS ASI who recently had direct oversight

responsibilities for their operations would reduce the time to obtain FAA approval for manual revisions partially due to the personal relationships between the former ASI and current FAA employees. Due to the general similarities among the groups of operators, the potential inefficiencies from employing a former ASI who had not had direct oversight responsibilities for that operator would not be significant. Thus, from the societal point of view, the overall losses to some individual former FAA inspectors would be largely offset by gains to other former FAA inspectors or qualified personnel. Although the proposed rule would create income transfers among individuals, at this time, we cannot quantify this overall loss on an individual basis. From a societal basis, the safety differential paid for the incremental loss in knowledge will be very small.

The number of former AFS ASIs who leave the FAA varies from year to year. We took the time period of October 1, 2007 to October 2, 2008 as a representative year-long period. As shown in Table 1, of the 208 AFS ASIs who left FAA employment, 138 voluntarily retired, 8 retired due to disability, 27 resigned, 10 were removed, 10 were terminated during their probation period, 4 had their appointments terminated, and 11 died. Of the voluntary retirements, 13 personnel were from FAA headquarters and were not specifically assigned to an operator. They would not be affected by the proposed rule. The maximum number of AFS ASIs who would have been affected had the proposed rule been in effect are the 160 non-headquarters personnel who retired, resigned, or became disabled. (We assumed that ASIs terminated or removed from their FAA position would be unlikely to be hired by an operator to work with their former FAA office in the absence of this proposed rule, and therefore would not be part of the potential economically affected population.)

TABLE 1—THE NUMBER OF AFS ASIS WHO LEFT FAA EMPLOYMENT BETWEEN 10/1/07 AND 10/2/08

Reason for separation	Number of inspectors
Voluntary Retirement	138
Disability Retirement	8
Resignation	27
Removal	10
Termination During Probation Period	10
Termination of Appointment	4

TABLE 1—THE NUMBER OF AFS ASIS WHO LEFT FAA EMPLOYMENT BETWEEN 10/1/07 AND 10/2/08—Continued

Reason for separation	Number of inspectors
Death	11
Total	208

Currently, the FAA does not officially track the status of former AFS ASIs. We believe that few of these former AFS ASIs would become involved in post-FAA employment that would be subject to the restrictions of the proposed rule. Although the proposal may affect only a small number of former AFS ASIs, inappropriate action by a single ASI could potentially lead to significant safety issues. We further believe that this overall economic impact would be minimal, with the potential benefits exceeding the costs. We request comments on this analysis.

The FAA has, therefore, determined that this proposed rule would impose minimal cost, and under DOT 2100.5 we did not prepare a full regulatory evaluation.

Regulatory Flexibility Determination

The Regulatory Flexibility Act of 1980 (Pub. L. 96–354) (RFA) establishes “as a principle of regulatory issuance that agencies shall endeavor, consistent with the objectives of the rule and of applicable statutes, to fit regulatory and informational requirements to the scale of the businesses, organizations, and governmental jurisdictions subject to regulation. To achieve this principle, agencies are required to solicit and consider flexible regulatory proposals and to explain the rationale for their actions to assure that such proposals are given serious consideration.” The RFA covers a wide range of small entities, including small businesses, not-for-profit organizations, and small governmental jurisdictions.

Agencies must perform a review to determine whether a rule will have a significant economic impact on a substantial number of small entities. If the agency determines that it will, the agency must prepare a regulatory flexibility analysis as described in the RFA.

However, if an agency determines that a rule is not expected to have a significant economic impact on a substantial number of small entities, section 605(b) of the RFA provides that the head of the agency may so certify and a regulatory flexibility analysis is not required. The certification must

include a statement providing the factual basis for this determination, and the reasoning should be clear.

The proposed rule would only prevent an AFS ASI and persons responsible for their oversight from being employed by the operator for which he or she had direct oversight responsibilities. The cost to an operator of being unable to employ a specific individual would be minimal because other individuals with similar professional qualifications as those possessed by the former AFS ASI would be available.

Therefore the FAA certifies that this proposed rule would not have a significant economic impact on a substantial number of small entities. The FAA requests comments on this certification.

International Trade Impact Analysis

The Trade Agreements Act of 1979 (Pub. L. 96–39), as amended by the Uruguay Round Agreements Act (Pub. L. 103–465), prohibits Federal agencies from establishing any standards or engaging in related activities that create unnecessary obstacles to the foreign commerce of the United States. Pursuant to these Acts, the establishment of standards is not considered an unnecessary obstacle to the foreign commerce of the United States, so long as the standards have a legitimate domestic objective, such as the protection of safety, and do not operate in a manner that excludes imports that meet this objective. The statute also requires consideration of international standards and, where appropriate, that they be the basis for U.S. standards. The FAA notes the purpose is to ensure the safety of the American public, and has assessed the effects of this rule to ensure that it does not exclude imports that meet this objective. As a result, this rule is not considered as creating an unnecessary obstacle to foreign commerce.

Unfunded Mandates Assessment

Title II of the Unfunded Mandates Reform Act of 1995 (Pub. L. 104–4) requires each Federal agency to prepare a written statement assessing the effects of any Federal mandate in a proposed or final agency rule that may result in an expenditure of \$100 million or more (in 1995 dollars) in any one year by State, local, and tribal governments, in the aggregate, or by the private sector; such a mandate is deemed to be a “significant regulatory action.” The FAA currently uses an inflation-adjusted value of \$136.1 million in lieu of \$100 million. This proposed rule does not contain such a mandate; therefore, the

requirements of Title II of the Act do not apply.

Executive Order 13132, Federalism

The FAA has analyzed this proposed rule under the principles and criteria of Executive Order 13132, Federalism. We determined that this action would not have a substantial direct effect on the States, on the relationship between the national Government and the States, or on the distribution of power and responsibilities among the various levels of government, and, therefore, would not have federalism implications.

Environmental Analysis

FAA Order 1050.1E identifies FAA actions that are categorically excluded from preparation of an environmental assessment or environmental impact statement under the National Environmental Policy Act in the absence of extraordinary circumstances. The FAA has determined this proposed rulemaking action qualifies for the categorical exclusion identified in paragraph 312f and involves no extraordinary circumstances.

Regulations That Significantly Affect Energy Supply, Distribution, or Use

The FAA has analyzed this NPRM under Executive Order 13211, Actions Concerning Regulations that Significantly Affect Energy Supply, Distribution, or Use (May 18, 2001). While this NPRM is a “significant regulatory action” under Executive Order 12866, we have determined that it is not a “significant energy action” under the executive order because it is not likely to have a significant adverse effect on the supply, distribution, or use of energy.

Additional Information

Comments Invited

The FAA invites interested persons to participate in this rulemaking by submitting written comments, data, or views. We also invite comments relating to the economic, environmental, energy, or federalism impacts that might result from adopting the proposals in this document. The most helpful comments reference a specific portion of the proposal, explain the reason for any recommended change, and include supporting data. To ensure the docket does not contain duplicate comments, please send only one copy of written comments, or if you are filing comments electronically, please submit your comments only one time.

We will file in the docket all comments we receive, as well as a report summarizing each substantive public contact with FAA personnel

concerning this proposed rulemaking. Before acting on this proposal, we will consider all comments we receive on or before the closing date for comments. We will consider comments filed after the comment period has closed if it is possible to do so without incurring expense or delay. We may change this proposal in light of the comments we receive.

Proprietary or Confidential Business Information

Do not file in the docket information that you consider to be proprietary or confidential business information. Send or deliver this information directly to the person identified in the **FOR FURTHER INFORMATION CONTACT** section of this document. You must mark the information that you consider proprietary or confidential. If you send the information on a disk or CD-ROM, mark the outside of the disk or CD-ROM and also identify electronically within the disk or CD-ROM the specific information that is proprietary or confidential.

Under 14 CFR 11.35(b), when we are aware of proprietary information filed with a comment, we do not place it in the docket. We hold it in a separate file to which the public does not have access, and we place a note in the docket that we have received it. If we receive a request to examine or copy this information, we treat it as any other request under the Freedom of Information Act (5 U.S.C. 552). We process such a request under the DOT procedures found in 49 CFR part 7.

Availability of Rulemaking Documents

You can get an electronic copy of rulemaking documents using the Internet by—

1. Searching the Federal eRulemaking Portal (<http://www.regulations.gov>);
2. Visiting the FAA's Regulations and Policies Web page at: http://www.faa.gov/regulations_policies/; or
3. Accessing the Government Printing Office's Web page at: <http://www.gpoaccess.gov/fr/index.html>.

You can also get a copy by sending a request to the Federal Aviation Administration, Office of Rulemaking, ARM-1, 800 Independence Avenue, SW., Washington, DC 20591, or by calling (202) 267-9680. Make sure to identify the docket or notice number of this rulemaking.

You may access all documents the FAA considered in developing this proposed rule, including economic analyses and technical reports, from the Internet through the Federal eRulemaking Portal referenced in paragraph (1).

List of Subjects

14 CFR Part 91

Aircraft, Airmen, Airports, Aviation safety.

14 CFR Part 119

Air carriers, Aircraft, Aviation safety.

14 CFR Part 125

Aircraft, Aviation safety.

14 CFR Part 133

Aircraft, Aviation safety.

14 CFR Part 137

Aircraft, Aviation safety.

14 CFR Part 141

Educational facilities, Schools.

14 CFR Part 142

Educational facilities, Schools.

14 CFR Part 145

Aircraft, Aviation safety.

14 CFR Part 147

Aircraft, Educational facilities, Schools.

The Proposed Amendment

In consideration of the foregoing, the Federal Aviation Administration proposes to amend Chapter I of Title 14, Code of Federal Regulations, as follows:

PART 91—GENERAL OPERATING AND FLIGHT RULES

1. The authority citation for part 91 continues to read as follows:

Authority: 49 U.S.C. 106(g), 1155, 40103, 40113, 40120, 44101, 44111, 44701, 44704, 44709, 44711, 44712, 44715, 44716, 44717, 44722, 46306, 46315, 46316, 46504, 46506–46507, 47122, 47508, 47528–47531, articles 12 and 29 of the Convention on International Civil Aviation (61 Stat. 1180).

2. Add § 91.1050 to read as follows:

§ 91.1050 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no fractional owner or fractional ownership program manager may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the fractional owner or fractional ownership program manager in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the

operations of the fractional owner or fractional ownership program manager.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a fractional owner or fractional ownership program manager in a matter before the agency if the individual makes any written or oral communication on behalf of the fractional owner or fractional ownership program manager to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a fractional owner or fractional ownership program manager from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the fractional owner or fractional ownership program manager in any matter before the Federal Aviation Administration if the individual was employed by the fractional owner or fractional ownership program manager before [effective date of the rule].

PART 119—CERTIFICATION: AIR CARRIERS AND COMMERCIAL OPERATORS

3. The authority citation for part 119 continues to read as follows:

Authority: 49 U.S.C. 106(g), 1153, 40101, 40102, 40103, 40113, 44105, 44106, 44111, 44701–44717, 44722, 44901, 44903, 44904, 44906, 44912, 44914, 44936, 44938, 46103, 46105.

4. Add § 119.73 to read as follows:

§ 119.73 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no certificate holder conducting operations under part 121 or 135 of this chapter may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be

acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a certificate holder from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the certificate holder before [effective date of the rule].

PART 125—CERTIFICATION AND OPERATIONS: AIRPLANES HAVING A SEATING CAPACITY OF 20 OR MORE PASSENGERS OR A MAXIMUM PAYLOAD CAPACITY OF 6,000 POUNDS OR MORE; AND RULES GOVERNING PERSONS ON BOARD SUCH AIRCRAFT

5. The authority citation for part 125 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701–44702, 44705, 44710–44711, 44713, 44716–44717, 44722.

6. Add § 125.26 to read as follows:

§ 125.26 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no certificate holder may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific

party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a certificate holder from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the certificate holder before [effective date of the rule].

PART 133—ROTORCRAFT EXTERNAL-LOAD OPERATIONS

7. The authority citation for part 133 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701–44702.

8. Add § 133.22 to read as follows:

§ 133.22 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no certificate holder may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a certificate holder from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the

certificate holder before [effective date of the rule].

PART 137—AGRICULTURAL AIRCRAFT OPERATIONS

9. The authority citation for part 137 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40103, 40113, 44701–44702.

10. Add § 137.40 to read as follows:

§ 137.40 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no certificate holder may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a certificate holder from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the certificate holder before [effective date of the rule].

PART 141—PILOT SCHOOLS

11. The authority citation for part 141 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701–44703, 44707, 44709, 44711, 45102–45103, 45301–45302.

12. Add § 141.34 to read as follows:

§ 141.34 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no holder of a pilot

school certificate or a provisional pilot school certificate may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a holder of a pilot school certificate or a provisional pilot school certificate from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the certificate holder before [effective date of the rule].

PART 142—TRAINING CENTERS

13. The authority citation for part 142 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 40119, 44101, 44701–44703, 44705, 44707, 44709–44711, 45102–45103, 45301–45302.

14. Add § 142.14 to read as follows:

§ 142.14 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no holder of a training center certificate may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a holder of a training center certificate from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the certificate holder before [effective date of the rule].

PART 145—REPAIR STATIONS

15. The authority citation for part 145 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701–44702, 44707, 44709, 44717.

16. Add § 145.160 to read as follows:

§ 145.160 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no holder of a repair station certificate may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had

responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a holder of a repair station certificate from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the certificate holder before [effective date of the rule].

PART 147—AVIATION MAINTENANCE TECHNICIAN SCHOOLS

17. The authority citation for part 147 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701–44702, 44707–44709.

18. Add § 147.8 to read as follows:

§ 147.8 Employment of former FAA employees.

(a) Except as specified in paragraph (c) of this section, no holder of an aviation maintenance technician certificate may knowingly employ or make a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual, in the preceding 2 years—

(1) Served as, or was directly responsible for the oversight of, a Flight Standards Service aviation safety inspector; and

(2) Had direct responsibility to inspect, or oversee the inspection of, the operations of the certificate holder.

(b) For the purpose of this section, an individual shall be considered to be acting as an agent or representative of a certificate holder in a matter before the agency if the individual makes any written or oral communication on behalf of the certificate holder to the agency (or any of its officers or employees) in connection with a particular matter, whether or not involving a specific party and without regard to whether the individual has participated in, or had responsibility for, the particular matter while serving as a Flight Standards Service aviation safety inspector.

(c) The provisions of this section do not prohibit a holder of an aviation maintenance technician school certificate from knowingly employing or making a contractual arrangement which permits an individual to act as an agent or representative of the certificate holder in any matter before the Federal Aviation Administration if the individual was employed by the

certificate holder before [effective date of the rule].

Issued in Washington, DC, on November 9, 2009.

John M. Allen,

Director, Flight Standards Service.

[FR Doc. E9-27852 Filed 11-19-09; 8:45 am]

BILLING CODE 4910-13-P

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R05-OAR-2009-0771; FRL-8980-5]

Approval and Promulgation of Air Quality Implementation Plans; Indiana

AGENCY: Environmental Protection Agency (EPA).

ACTION: Proposed rule.

SUMMARY: EPA is proposing to approve a request submitted by the Indiana Department of Environmental Management on September 25, 2009, to revise the Indiana State Implementation Plan (SIP). The submission revises the Indiana Administrative Code (IAC) by amending and updating the definition of "References to Code of Federal Regulations," to refer to the 2008 edition.

DATES: Comments must be received on or before December 21, 2009.

ADDRESSES: Submit your comments, identified by Docket ID No. EPA-R05-OAR-2009-0771 by one of the following methods:

- <http://www.regulations.gov>: Follow the on-line instructions for submitting comments.

- *E-mail:* mooney.john@epa.gov.

- *Fax:* (312) 692-2551.

- *Mail:* John Mooney, Chief, Criteria Pollutant Section, Air Programs Branch (AR-18J), U.S. Environmental Protection Agency, 77 West Jackson Boulevard, Chicago, Illinois 60604.

- *Hand Delivery:* John Mooney, Chief, Criteria Pollutant Section, Air Programs Branch (AR-18J), U.S. Environmental Protection Agency, 77 West Jackson Boulevard, Chicago, Illinois 60604. Such deliveries are only accepted during the Regional Office normal hours of operation, and special arrangements should be made for deliveries of boxed information. The Regional Office official hours of business are Monday through Friday, 8:30 a.m. to 4:30 p.m. excluding Federal holidays.

Please see the direct final rule which is located in the Rules section of this **Federal Register** for detailed instructions on how to submit comments.

FOR FURTHER INFORMATION CONTACT:

Charles Hatten, Environmental Engineer, Criteria Pollutant Section, Air Programs Branch (AR-18J), Environmental Protection Agency, Region 5, 77 West Jackson Boulevard, Chicago, Illinois 60604, (312) 886-6031, hatten.charles@epa.gov.

SUPPLEMENTARY INFORMATION: In the Final Rules section of this **Federal Register**, EPA is approving the State's SIP submittal as a direct final rule without prior proposal because the Agency views this as a noncontroversial submittal and anticipates no adverse comments. A detailed rationale for the approval is set forth in the direct final rule. If no adverse comments are received in response to this rule, no further activity is contemplated. If EPA receives adverse comments, the direct final rule will be withdrawn and all public comments received will be addressed in a subsequent final rule based on this proposed rule. EPA will not institute a second comment period. Any parties interested in commenting on this action should do so at this time. Please note that if EPA receives adverse comment on an amendment, paragraph, or section of this rule, and if that provision may be severed from the remainder of the rule, EPA may adopt as final those provisions of the rule that are not the subject of an adverse comment. For additional information, see the direct final rule which is located in the Rules section of this **Federal Register**.

Dated: November 3, 2009.

Bharat Mathur,

Acting Regional Administrator, Region 5.

[FR Doc. E9-27816 Filed 11-19-09; 8:45 am]

BILLING CODE 6560-50-P

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA-R03-OAR-2009-0674; FRL-8983-2]

Approval and Promulgation of Air Quality Implementation Plans; Virginia; Transportation Conformity Regulations

AGENCY: Environmental Protection Agency (EPA).

ACTION: Proposed rule.

SUMMARY: EPA proposes to approve the State Implementation Plan (SIP) revision submitted by the Commonwealth of Virginia for Transportation Conformity regulations. In the Final Rules section of this **Federal Register**, EPA is approving the State's SIP submittal as a direct final

rule without prior proposal because the Agency views this as a noncontroversial submittal and anticipates no adverse comments. A detailed rationale for the approval is set forth in the direct final rule. If no adverse comments are received in response to this action, no further activity is contemplated. If EPA receives adverse comments, the direct final rule will be withdrawn and all public comments received will be addressed in a subsequent final rule based on this proposed rule. EPA will not institute a second comment period. Any parties interested in commenting on this action should do so at this time.

DATES: Comments must be received in writing by December 21, 2009.

ADDRESSES: Submit your comments, identified by Docket ID Number EPA-R03-OAR-2009-0674 by one of the following methods:

- A. <http://www.regulations.gov>: Follow the on-line instructions for submitting comments.

- B. *E-mail:* fernandez.cristina@epa.gov.

- C. *Mail:* EPA-R03-OAR-2009-0674, Cristina Fernandez, Associate Director, Office of Air Program Planning, Mailcode 3AP30, U.S. Environmental Protection Agency, Region III, 1650 Arch Street, Philadelphia, Pennsylvania 19103.

- D. *Hand Delivery:* At the previously-listed EPA Region III address. Such deliveries are only accepted during the Docket's normal hours of operation, and special arrangements should be made for deliveries of boxed information.

Instructions: Direct your comments to Docket ID No. EPA-R03-OAR-2009-0674. EPA's policy is that all comments received will be included in the public docket without change, and may be made available online at <http://www.regulations.gov>, including any personal information provided, unless the comment includes information claimed to be Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. Do not submit information that you consider to be CBI (or otherwise protected) through <http://www.regulations.gov> or e-mail. The <http://www.regulations.gov> Web site is an anonymous access system, which means EPA will not know your identity or contact information unless you provide it in the body of your comment. If you send an e-mail comment directly to EPA without going through <http://www.regulations.gov>, your e-mail address will be automatically captured and included as part of the comment that is placed in the public docket and made available on the Internet. If you

submit an electronic comment, EPA recommends that you include your name and other contact information in the body of your comment and with any disk or CD-ROM you submit. If EPA cannot read your comment due to technical difficulties and cannot contact you for clarification, EPA may not be able to consider your comment. Electronic files should avoid the use of special characters, any form of encryption, and be free of any defects or viruses.

Docket: All documents in the electronic docket are listed in the <http://www.regulations.gov> index. Although listed in the index, some information is not publicly available, *i.e.*, CBI or other information whose disclosure is restricted by statute. Certain other material, such as copyrighted material, is not placed on the Internet and will be publicly available only in hard copy form. Publicly available docket materials are available either electronically in <http://www.regulations.gov> or in hard copy during normal business hours at the Air Protection Division, U.S. Environmental Protection Agency, Region III, 1650 Arch Street, Philadelphia, Pennsylvania 19103. Copies of the State submittal are available at the Virginia Department of the Environmental Quality, 629 East Main Street, Richmond, Virginia 23219.

FOR FURTHER INFORMATION CONTACT: Martin Kotsch, (215) 814-3335, or by e-mail at kotsch.martin@epa.gov.

SUPPLEMENTARY INFORMATION: For further information, please see the information provided in the direct final action, with the same title, that is located in the Rules and Regulations section of this **Federal Register** publication. Please note that if EPA receives adverse comment on an amendment, paragraph, or section of this rule and if that provision may be severed from the remainder of the rule, EPA may adopt as final those provisions of the rule that are not the subject of an adverse comment.

Dated: November 5, 2009.

William C. Early,

Acting Regional Administrator, Region III.
[FR Doc. E9-27813 Filed 11-19-09; 8:45 am]

BILLING CODE 6560-50-P

DEPARTMENT OF THE INTERIOR

Fish and Wildlife Service

50 CFR Part 92

[FWS-R7-MB-2009-0082] [91200-1231-9BPP-L2]

RIN 1018-AW67

Migratory Bird Subsistence Harvest in Alaska; Harvest Regulations for Migratory Birds in Alaska During the 2010 Season

AGENCY: Fish and Wildlife Service, Interior.

ACTION: Proposed rule.

SUMMARY: The U.S. Fish and Wildlife Service (Service or we) proposes migratory bird subsistence harvest regulations in Alaska for the 2010 season. These regulations will enable the continuation of customary and traditional subsistence uses of migratory birds in Alaska and prescribe regional information on when and where the harvesting of birds may occur. These regulations were developed under a co-management process involving the Service, the Alaska Department of Fish and Game, and Alaska Native representatives. The rulemaking is necessary because the regulations governing the subsistence harvest of migratory birds in Alaska are subject to annual review. This rulemaking proposes region-specific regulations that go into effect on April 2, 2010, and expire on August 31, 2010.

DATES: We will accept comments received or postmarked on or before January 19, 2010. We must receive requests for public hearings, in writing, at the address shown in **FOR FURTHER INFORMATION CONTACT** by January 4, 2010.

ADDRESSES: You may submit comments by one of the following methods:

- Federal eRulemaking Portal: <http://www.regulations.gov>. Follow the instructions for submitting comments to Docket No. FWS-R7-MB-2009-0082.

- U.S. mail or hand-delivery: Public Comments Processing, Attn: FWS-R7-MB-2009-0082; Division of Policy and Directives Management; U.S. Fish and Wildlife Service; 4401 N. Fairfax Drive, Suite 222; Arlington, VA 22203.

We will not accept e-mail or faxes. We will post all comments on <http://www.regulations.gov>. This generally means that we will post any personal information you provide us (see the Public Comment Procedures section below for more information).

FOR FURTHER INFORMATION CONTACT: Fred Armstrong, (907) 786-3887, or Donna

Dewhurst, (907) 786-3499, U.S. Fish and Wildlife Service, 1011 E. Tudor Road, Mail Stop 201, Anchorage, AK 99503.

SUPPLEMENTARY INFORMATION:

Public Comment Procedures

To ensure that any final action resulting from this proposed rule will be as accurate and as effective as possible, we request that you send relevant information for our consideration. The comments that will be most useful and likely to influence our decisions are those that you support by quantitative information or studies and those that include citations to, and analyses of, the applicable laws and regulations. Please make your comments as specific as possible and explain the bases for them. In addition, please include sufficient information with your comments to allow us to authenticate any scientific or commercial data you include.

You must submit your comments and materials concerning this proposed rule by one of the methods listed above in the **ADDRESSES** section. We will not accept comments sent by e-mail or fax or to an address not listed in **ADDRESSES**. If you submit a comment via <http://www.regulations.gov>, your entire comment—including any personal identifying information, such as your address, telephone number, or e-mail address—will be posted on the Web site. Please note that comments submitted to this Web site are not immediately viewable. When you submit a comment, the system receives it immediately. However, the comment will not be publicly viewable until we post it, which might not occur until several days after submission.

If you mail or hand-carry a hardcopy comment directly to us that includes personal information, you may request at the top of your document that we withhold this information from public review. However, we cannot guarantee that we will be able to do so. To ensure that the electronic docket for this rulemaking is complete and all comments we receive are publicly available, we will post all hardcopy comments on <http://www.regulations.gov>.

In addition, comments and materials we receive, as well as supporting documentation used in preparing this proposed rule, will be available for public inspection in two ways:

(1) You can view them on <http://www.regulations.gov>. In the Search Documents box, enter FWS-R7-MB-2009-0082, which is the docket number for this rulemaking. Then, in the Search panel on the left side of the screen, select the type of documents you want

to view under the Document Type heading.

(2) You can make an appointment, during normal business hours, to view the comments and materials in person at the Division of Migratory Bird Management, U.S. Fish and Wildlife Service; 4501 N. Fairfax Drive, Room 4107, Arlington, VA 22203-1610.

Public Availability of Comments

As stated above in more detail, before including your address, phone number, e-mail address, or other personal identifying information in your comment, you should be aware that your entire comment—including your personal identifying information—may be made publicly available at any time. Though you can ask us in your comment to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Why Is This Rulemaking Necessary?

This rulemaking is necessary because, by law, the migratory bird harvest season is closed unless opened by the Secretary of the Interior, and the regulations governing subsistence harvest of migratory birds in Alaska are subject to public review and annual approval. This rule proposes regulations for the taking of migratory birds for subsistence uses in Alaska during the spring and summer of 2010. This rule lists proposed migratory bird season openings and closures in Alaska by region.

How Do I Find the History of These Regulations?

Background information, including past events leading to this action, accomplishments since the Migratory Bird Treaties with Canada and Mexico were amended, and a history addressing conservation issues can be found in the following **Federal Register** documents:

Date	Federal Register Citation
August 16, 2002	67 FR 53511
July 21, 2003	68 FR 43010
April 2, 2004	69 FR 17318
April 8, 2005	70 FR 18244
February 28, 2006 ...	71 FR 10404
April 11, 2007	72 FR 18318
March 14, 2008	73 FR 13788
May 19, 2009	74 FR 23336

These final rules setting forth the annual harvest regulations, are available at <http://alaska.fws.gov/ambcc/regulations.htm>.

What Is the Process for Issuing Regulations for the Subsistence Harvest of Migratory Birds in Alaska?

The U.S. Fish and Wildlife Service (Service or we) proposes migratory bird subsistence harvest regulations in Alaska for the 2010 season. These regulations will enable the continuation of customary and traditional subsistence uses of migratory birds in Alaska and prescribe regional information on when and where the harvesting of birds may occur. These regulations were developed under a co-management process involving the Service, the Alaska Department of Fish and Game, and Alaska Native representatives.

We opened the process to establish regulations for the 2010 spring and summer subsistence harvest of migratory birds in Alaska this past spring in a proposed rule published in the **Federal Register** on April 10, 2009 (74 FR 16339). While that proposed rule dealt primarily with the regulatory process for hunting migratory birds for all purposes throughout the United States, we also discussed the background and history of the Alaska subsistence regulations, explained the annual process for their establishment, and requested proposals for the 2010 season. The rulemaking processes for both types of migratory bird harvest are related, and the April 10, 2009, proposed rule explained the connection between the two.

The Alaska Migratory Bird Co-management Council (Co-management Council) held a meeting in April 2009 to develop recommendations for changes effective for the 2010 harvest season. The April 10, 2009, proposed rule set a deadline of June 15, 2009, for the Co-management Council to submit proposals for the 2010 spring and summer migratory bird subsistence harvest season to us and the Flyway Councils. This timeframe allowed the Flyway Councils and the Service to consider the proposals and present recommendations to the Service Regulations Committee at the committee's meeting on July 29 and 30, 2009.

Who Is Eligible To Hunt Under These Regulations?

Eligibility to harvest under the regulations established in 2003 was limited to permanent residents, regardless of race, in villages located within the Alaska Peninsula, Kodiak Archipelago, the Aleutian Islands, and

in areas north and west of the Alaska Range (50 CFR 92.5). These geographical restrictions opened the initial subsistence migratory bird harvest to only about 13 percent of Alaska residents. High-population areas such as Anchorage, the Matanuska-Susitna and Fairbanks North Star boroughs, the Kenai Peninsula roaded area, the Gulf of Alaska roaded area, and Southeast Alaska were excluded from the eligible subsistence harvest areas.

Based on petitions requesting inclusion in the harvest, in 2004, we added 13 additional communities based on criteria set forth in 50 CFR 92.5(c). These communities were Gulkana, Gakona, Tazlina, Copper Center, Mentasta Lake, Chitina, Chistochina, Tatitlek, Chenega, Port Graham, Nanwalek, Tyonek, and Hoonah, with a combined population of 2,766. In 2005, we added three additional communities for glaucous-winged gull egg gathering only, based on petitions requesting inclusion. These southeastern communities were Craig, Hydaburg, and Yakutat, with a combined population of 2,459.

In 2007, we enacted the Alaska Department of Fish and Game's request to expand the Fairbanks North Star Borough excluded area to include the Central Interior area. This action excluded the following communities from participation in this harvest: Big Delta/Fort Greely, Healy, McKinley Park/Village and Ferry, with a combined population of 2,812. These removed communities reduced the percentage of the State population included in the subsistence harvest to 13 percent.

How Will the Service Ensure That the Subsistence Harvest Will Not Raise Overall Migratory Bird Harvest or Threaten the Conservation of Endangered and Threatened Species?

We have monitored subsistence harvest for the past 15 years through the use of annual household surveys in the most heavily used subsistence harvest areas, such as the Yukon-Kuskokwim Delta. Continuation of this monitoring enables tracking of any major changes or trends in levels of harvest and user participation after legalization of the harvest. This rule proposes for the second year to restrict hunting on the North Slope to times of day with sufficient daylight to enable hunters to distinguish and avoid shooting closed species. In addition, three conservation measures, which focus on increased migratory bird hunter outreach prior to hunts, increased regulatory enforcement and in-season harvest verification of Steller's eider mortality, would continue to provide additional protection for

threatened spectacled and Steller's eiders. Finally, we have an emergency closure provision (50 CFR 92.21), which specifies that the harvest may be closed or temporarily suspended upon a finding that a continuation of the regulation allowing the harvest would pose an imminent threat to the conservation of any endangered or threatened species or other migratory bird population.

With regard to Steller's eiders, the proposed regulation at 50 CFR 92.32, carried over from last year, clarifies that we will take action under 50 CFR 92.21 as is necessary to prevent further take of Steller's eiders, which could include temporary or long-term closures of the harvest in all or a portion of the geographic area open to harvest. If mortality of threatened eiders occurs, we will evaluate each mortality event by criteria such as: cause, quantity, sex, age, location, and date. We will consult the Co-management Council when an emergency closure is being considered. Any emergency closure deemed necessary will be designed to minimize its impact on the subsistence harvest.

What Is Different in the Region-Specific Regulations for 2010?

Yellow-billed Loons

Consistent with the request of the North Slope Borough Fish and Game Management Committee and the recommendation of the Co-management Council, this proposed rule continues into 2010 the provisions originally established in 2005 to allow subsistence use of yellow-billed loons (*Gavia adamsii*) inadvertently entangled in subsistence fishing (gill) nets on the North Slope. Yellow-billed loons are culturally important for the Inupiat Eskimo of the North Slope for use in traditional dance regalia. A maximum of 20 yellow-billed loons may be caught in 2010 under this provision. This provision does not authorize intentional harvest of yellow-billed loons, but allows use of those loons inadvertently entangled during normal subsistence fishing activities. Individual reporting to the North Slope Borough Department of Wildlife is required by the end of each season. However, the North Slope Borough has asked fishermen, through announcements on the radio and through personal contact, to report inadvertent entanglements of loons as they occur, to better estimate the level of mortality caused by gill nets. In 2008, one yellow-billed loon was reported to be found dead in a fishing net; one severely injured yellow-billed loon was observed by Borough staff; and two were

released uninjured from fishing nets by Borough staff.

Aleutian and Arctic Terns

We propose to remove the provision that opened a season May 15–June 30 for harvesting Aleutian (*Onychoprion aleutica*) and arctic tern (*Sterna paradisaea*) eggs in the Yakutat Harvest area, from Icy Bay (Icy Cape to Point Riou) and the coastal islands bordering the Gulf of Alaska from Point Manby southeast to and including Dry Bay. The Yakutat Tlingit Tribe requested this regulation be removed at the April 2009 Co-Management Council meeting, stating that they will not be able to adequately monitor the tern subsistence take as requested by the Service, so they would prefer to withdraw the regulation at this time.

Spectacled and Steller's Eiders

Spectacled eiders (*Somateria fischeri*) and the Alaska-breeding population of Steller's eiders (*Polysticta stelleri*) are listed as threatened species, and their migration and breeding distribution overlaps with the spring and summer subsistence harvest on the Yukon-Kuskokwim Delta and the North Slope. Both spectacled and Steller's eiders are closed to hunting in the subsistence harvest, but harvest surveys and Service documentation indicate substantial numbers of both species have been taken during recent subsistence harvests on the North Slope.

The North Slope breeding population of spectacled eiders was estimated to be 12,916 (10,942–14,890, 95% Confidence Limits) individual birds during 2002–06 (Service unpublished data), and they nest relatively widely across the North Slope. It is estimated that 35 (33–40, 95% Confidence Limits) spectacled eiders were taken on the North Slope during the 2005 subsistence season (Service unpublished data, 2006); 99 (44–155, 95% Confidence Limits) were taken during the 2007 subsistence season; and 9 (1–25, 184% confidence limits) were taken during the 2008 subsistence season (Alaska Department of Fish and Game, preliminary data).

The North Slope breeding population of Steller's eider was estimated to be 576 annually (292–859, 90% Confidence Limits) individual birds during 1993–2008 (Service, unpublished data), and most of their nesting appears to be concentrated near Barrow, the northernmost point in Alaska. It is estimated that 19 (9–37, 95% Confidence Limits) Steller's eiders were taken on the North Slope during the 2005 subsistence season; 36 (1–85, 135% Confidence Limits) were taken during the 2007 subsistence season; and

0 were taken during the 2008 subsistence season (Alaska Department of Fish and Game, preliminary data). However, during the 2008 subsistence season, the Service documented 20 Steller's eiders shot at Barrow, with another 7 found dead but too heavily scavenged to determine cause of death.

Therefore, harvest survey estimates and direct observation of shot birds indicated that direct shooting occurs during the subsistence harvest, with impacts probably on the order of tens of each threatened eider species taken per year. Take is not authorized for either species during the subsistence harvest, and, in the case of Steller's eider, this amount of shooting mortality is likely not sustainable for the small Alaska-breeding population. Because of the Steller's eider small breeding population size, their breeding concentration near Barrow, and the relatively high proportion of the estimated population shot during recent subsistence harvests, the Service focused on considering regulations and conservation efforts on the North Slope to benefit the Alaska-breeding population of Steller's eiders.

Several spectacled and Steller's eider management needs are addressed by this proposed rule. It restricts hunting on the North Slope, from Barrow through Point Hope, to time of day with sufficient daylight to ensure hunters can distinguish and avoid shooting species closed for harvest; it clarifies for subsistence users that Service law enforcement personnel have authority to verify species of birds possessed by hunters; it clarifies that it is illegal to possess any bird closed to harvest; and it describes how the Service's existing authority of emergency closure would be implemented, if necessary, to protect Steller's eiders. The regulations, implemented in accordance with conservation measures (described below), are considered the principal way in which shooting mortality of threatened eider will be substantially reduced or eliminated. The emergency closure authority provides an additional level of assurance that, if an unexpected amount of Steller's eider shooting mortality occurs, it will be curtailed to avoid approaching jeopardy to the existence of the species.

The Service developed three conservation measures that are an integral part of the proposed harvest and were approved for implementation by the Alaska Regional Director on April 6, 2009. The conservation measures substantially increased protection for spectacled and, particularly, Steller's eiders on the North Slope in 2009, and described how the Service would detect,

remedy, and quickly curtail any shooting mortality or injury of Steller's eiders that might occur during the harvest. In January 2009, the Service commenced planning for implementation of each measure in anticipation of the subsistence harvest. The three conservation measures were:

1. Increase Migratory Bird Hunter Outreach Prior to the Hunts

The Service with North Slope partners would provide migratory bird hunter outreach in Wainwright, Point Hope, Point Lay, and Barrow prior to each subsistence harvest. The outreach educational objectives included: hunter understanding of the hunting regulations; ability to distinguish among the open and closed species of eiders in flight; the need to reduce crippling loss; and an understanding of the Service's role and obligation for enforcement and monitoring. This was done prior to and during the 2009 subsistence season.

2. Increased Service Enforcement of Migratory Bird Regulations

During the 2009 subsistence season, the Service sustained a law enforcement presence on the North Slope during the migratory bird hunts. The Service believes this action was necessary to increase community understanding and acceptance of the shooting mortality problem, deter violations, and obtain compliance with the regulations. The Service conducted real-time monitoring of the harvest to meet the primary objective of detecting Steller's eider mortality during the hunts so that appropriate and timely corrective action could be taken. Regulatory enforcement objectives will continue to be achieved through a two-part strategy: (i) pre-season community and hunter education and outreach, and (ii) in-season implementation of the law enforcement portion of this plan and enforcement of all Service regulations.

3. In-season Harvest Verification of Steller's Eider Mortality and Injury

Three types of monitoring efforts were used during the 2009 subsistence harvest and fall hunts on the North Slope: (i) Steller's eider breeding surveys to inform the coordination of the conservation measures, (ii) harvest verification by Service law enforcement to meet the objective of detecting Steller's eider mortality during the hunts so appropriate and timely corrective action can be taken to prevent further mortality; and (iii) monitoring for injured and dead birds to begin to quantify crippling rate and loss. We will continue to use all in-season monitoring information to independently evaluate

harvest survey reports, the efficiency of the regulations, conservation measures, and outreach efforts.

To summarize, the Service has dual goals and responsibilities of authorizing a subsistence harvest while protecting migratory birds and threatened species. Although these goals were and continue to be challenging, they are not irreconcilable with sufficient recognition of the need to protect threatened species, measures to remedy documented threats, and commitment from the subsistence community and other conservation partners to work together toward these dual goals. With these dual goals in mind, the Service is proposing to continue the provision that restricts hunting on the North Slope to times of day with sufficient daylight to enable hunters to avoid shooting closed species. Moreover, the Service, working with partners, developed additional measures to eliminate the potential for shooting mortality or injury of the Alaska-breeding population of Steller's eider on the North Slope. These measures include: (1) increased waterfowl hunter outreach and community awareness; (2) increased enforcement of the migratory bird regulations that are protective of listed eiders; and (3) in-season Service verification of the harvest to detect any Steller's eider mortality.

For the 2009 season, the Service and the community planned to immediately address and remedy any detected Steller's eider mortality; and, as a matter of Service policy, any detected Steller's eider shooting mortality was curtailed to an amount estimated to be sustainable by the population. The summer of 2009 was not a breeding year for Steller's eiders in the Barrow area. Even so, the Service conducted an extensive outreach program including eight public meetings in all of the affected eider communities, three radio shows, and five newspaper articles. The Service increased its law enforcement presence in Barrow as well as harvest monitoring. No Steller's eiders were found shot during monitoring of the subsistence harvest.

In 2009, the Service also continued working into July refining and implementing the Memorandum of Understanding (MOU) established between the Service and North Slope government and Native organizations. The reason to initiate an MOU was to increase involvement by Alaska Native organizations in the conservation of Steller's eiders on the North Slope of Alaska. The purposes outlined were to: (1) conserve and manage Steller's eiders; (2) preserve the customary and traditional subsistence hunt, and (3)

reduce or eliminate the take of Steller's eiders.

A review of the conservation strategy implemented by the Service in 2009 based on the conservation measures developed, showed that overall the Service made advancements in several areas. These conservation measures and the subsequent MOU engaged our partners on the North Slope, encouraging local ownership of the conservation goal, and ultimately heightening awareness of what actions were necessary to move in a positive direction to protect Steller's eiders. Based on these successes, the Service proposes to continue these conservation measures into the 2010 season with some modification as to the amount of effort and emphasis each will receive. The Service also proposes to continue the regulatory changes implemented in 2009 for the North Slope through the 2010 subsistence season, including the emergency closure provisions.

Statutory Authority

We derive our authority to issue these regulations from the Migratory Bird Treaty Act of 1918, 16 U.S.C. 712(1), which authorizes the Secretary of the Interior, in accordance with the treaties with Canada, Mexico, Japan, and Russia, to "issue such regulations as may be necessary to assure that the taking of migratory birds and the collection of their eggs, by the indigenous inhabitants of the State of Alaska, shall be permitted for their own nutritional and other essential needs, as determined by the Secretary of the Interior, during seasons established so as to provide for the preservation and maintenance of stocks of migratory birds."

Required Determinations

Regulatory Planning and Review (Executive Order 12866)

The Office of Management and Budget (OMB) has determined that this rule is not significant and has not reviewed this rule under Executive Order 12866 (E.O. 12866). OMB bases its determination upon the following four criteria:

(a) Whether the rule will have an annual effect of \$100 million or more on the economy or adversely affect an economic sector, productivity, jobs, the environment, or other units of the government.

(b) Whether the rule will create inconsistencies with other Federal agencies' actions.

(c) Whether the rule will materially affect entitlements, grants, user fees, loan programs, or the rights and obligations of their recipients.

(d) Whether the rule raises novel legal or policy issues.

Regulatory Flexibility Act

The Department of the Interior certifies that this rule will not have a significant economic impact on a substantial number of small entities as defined under the Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*). An initial regulatory flexibility analysis is not required. Accordingly, a Small Entity Compliance Guide is not required. The rule legalizes a pre-existing subsistence activity, and the resources harvested will be consumed by the harvesters or persons within their local community.

Small Business Regulatory Enforcement Fairness Act

This rule is not a major rule under 5 U.S.C. 804(2), the Small Business Regulatory Enforcement Fairness Act. This rule:

(a) Will not have an annual effect on the economy of \$100 million or more. It will legalize and regulate a traditional subsistence activity. It will not result in a substantial increase in subsistence harvest or a significant change in harvesting patterns. The commodities being regulated under this rule are migratory birds. This rule deals with legalizing the subsistence harvest of migratory birds and, as such, does not involve commodities traded in the marketplace. A small economic benefit from this rule derives from the sale of equipment and ammunition to carry out subsistence hunting. Most, if not all, businesses that sell hunting equipment in rural Alaska would qualify as small businesses. We have no reason to believe that this rule will lead to a disproportionate distribution of benefits.

(b) Will not cause a major increase in costs or prices for consumers; individual industries; Federal, State, or local government agencies; or geographic regions. This rule does not deal with traded commodities and, therefore, does not have an impact on prices for consumers.

(c) Does not have significant adverse effects on competition, employment, investment, productivity, innovation, or the ability of U.S.-based enterprises to compete with foreign-based enterprises. This rule deals with the harvesting of wildlife for personal consumption. It does not regulate the marketplace in any way to generate effects on the economy or the ability of businesses to compete.

Unfunded Mandates Reform Act

We have determined and certified under the Unfunded Mandates Reform Act (2 U.S.C. 1501 *et seq.*) that this rule

will not impose a cost of \$100 million or more in any given year on local, State, or tribal governments or private entities. The rule does not have a significant or unique effect on State, local, or tribal governments or the private sector. A statement containing the information required by the Unfunded Mandates Reform Act is not required. Participation on regional management bodies and the Co-management Council will require travel expenses for some Alaska Native organizations and local governments. In addition, they will assume some expenses related to coordinating involvement of village councils in the regulatory process. Total coordination and travel expenses for all Alaska Native organizations are estimated to be less than \$300,000 per year. In the Notice of Decision (65 FR 16405; March 28, 2000), we identified 12 partner organizations (Alaska Native nonprofits and local governments) to administer the regional programs. The Alaska Department of Fish and Game will also incur expenses for travel to Co-management Council and regional management body meetings. In addition, the State of Alaska will be required to provide technical staff support to each of the regional management bodies and to the Co-management Council. Expenses for the State's involvement may exceed \$100,000 per year, but should not exceed \$150,000 per year. When funding permits, we make annual grant agreements available to the partner organizations and the Alaska Department of Fish and Game to help offset their expenses.

Takings (Executive Order 12630)

Under the criteria in Executive Order 12630, this rule does not have significant takings implications. This rule is not specific to particular land ownership, but applies to the harvesting of migratory bird resources throughout Alaska. A takings implication assessment is not required.

Federalism (Executive Order 13132)

Under the criteria in Executive Order 13132, this rule does not have sufficient federalism implications to warrant the preparation of a Federalism Assessment. We discuss effects of this proposed rule on the State of Alaska in the Unfunded Mandates Reform Act section above. We worked with the State of Alaska to develop these regulations. Therefore, a Federalism Assessment is not required.

Civil Justice Reform (Executive Order 12988)

The Department, in promulgating this rule, has determined that it will not unduly burden the judicial system and that it meets the requirements of sections 3(a) and 3(b)(2) of Executive Order 12988.

Government-to-Government Relations With Native American Tribal Governments

Because eligibility to hunt under these regulations is not limited to tribal members, but rather extends to all indigenous inhabitants of the subsistence harvest areas, we are not required to engage in formal consultation with tribes. However, in keeping with the spirit of the President's memorandum of April 29, 1994, "Government-to-Government Relations With Native American Tribal Governments" (59 FR 22951), and Executive Order 13175 (65 FR 67249; November 6, 2000), concerning consultation and coordination with Indian Tribal Governments, we conducted meetings with the affected tribes and tribal nonprofit organizations to discuss the proposed changes in the regulations for possible effects on tribes or trust resources, and have determined that there are no significant effects. The rule will legally recognize the subsistence harvest of migratory birds and their eggs for indigenous inhabitants including tribal members. In 1998, we began a public involvement process to determine how to structure management bodies in order to provide the most effective and efficient involvement of subsistence users. We began by publishing in the **Federal Register** stating that we intended to establish management bodies to implement the spring and summer subsistence harvest (63 FR 49707, September 17, 1998). We held meetings with the Alaska Department of Fish and Game and the Native Migratory Bird Working Group to provide information regarding the amended treaties and to listen to the needs of subsistence users. The Native Migratory Bird Working Group was a consortium of Alaska Natives formed by the Rural Alaska Community Action Program to represent Alaska Native subsistence hunters of migratory birds during the treaty negotiations. We held forums in Nome, Kotzebue, Fort Yukon, Allakaket, Naknek, Bethel, Dillingham, Barrow, and Copper Center. We led additional briefings and discussions at the annual meeting of the Association of Village Council Presidents in Hooper Bay and

for the Central Council of Tlingit & Haida Indian Tribes in Juneau.

On March 28, 2000, we published in the **Federal Register** (65 FR 16405) the Notice of Decision: "Establishment of Management Bodies in Alaska To Develop Recommendations Related to the Spring/Summer Subsistence Harvest of Migratory Birds." This notice described the way in which management bodies would be established and organized. Based on the wide range of views expressed on the options document, the decision incorporated key aspects of two of the modules. The decision established one statewide management body consisting of 1 Federal member, 1 State member, and 7–12 Alaska Native members, with each component serving as equals.

Paperwork Reduction Act

This rule has been examined under the Paperwork Reduction Act of 1995 and does not contain new collections of information that require Office of Management and Budget approval. OMB has approved our collection of information associated with the voluntary annual household surveys used to determine levels of subsistence take. The OMB control number is 1018–0124, which expires on January 31, 2010. An agency may not conduct or sponsor and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number.

Endangered Species Act Consideration

Prior to issuance of annual spring and summer subsistence regulations, we will consult under section 7 of the Endangered Species Act of 1973, as amended (16 U.S.C. 1531–1543; hereinafter the Act), to ensure that the 2010 subsistence harvest is not likely to jeopardize the continued existence of any species designated as endangered or threatened, or modify or destroy its critical habitats, and that the regulations are consistent with conservation programs for those species. Consultation under section 7 of the Act for the annual subsistence take regulations may cause us to change these regulations. Our biological opinion resulting from the section 7 consultation is a public document available from the person listed under **FOR FURTHER INFORMATION CONTACT**.

National Environmental Policy Act Consideration

The annual regulations and options were considered in the Environmental Assessment, "Managing Migratory Bird Subsistence Hunting in Alaska: Hunting Regulations for the 2010 Spring/

Summer Harvest," issued October 9, 2009. Copies are available from the person listed under **FOR FURTHER INFORMATION CONTACT** or at www.Regulations.gov.

Energy Supply, Distribution, or Use (Executive Order 13211)

Executive Order 13211 requires agencies to prepare Statements of Energy Effects when undertaking certain actions. This is not a significant regulatory action under this Executive Order; it would allow only for traditional subsistence harvest and would improve conservation of migratory birds by allowing effective regulation of this harvest. Further, this rule is not expected to significantly affect energy supplies, distribution, or use. Therefore, this action is not a significant energy action under Executive Order 13211 and no Statement of Energy Effects is required.

List of Subjects in 50 CFR Part 92

Exports, Hunting, Imports, Reporting and recordkeeping requirements, Subsistence, Treaties, Wildlife.

For the reasons set out in the preamble, we propose to amend title 50, chapter I, subchapter G, of the Code of Federal Regulations as follows:

PART 92—MIGRATORY BIRD SUBSISTENCE HARVEST IN ALASKA

1. The authority citation for part 92 continues to read as follows:

Authority: 16 U.S.C. 703–712.

Subpart D—Annual Regulations Governing Subsistence Harvest

2. In subpart D, add § 92.31 to read as follows:

§ 92.31 Region-specific regulations.

The 2010 season dates for the eligible subsistence harvest areas are as follows:

- (a) *Aleutian/Pribilof Islands Region*.
 - (i) Northern Unit (Pribilof Islands):
 - (i) Season: April 2–June 30.
 - (ii) Closure: July 1–August 31.
 - (2) Central Unit (Aleut Region's eastern boundary on the Alaska Peninsula westward to and including Unalaska Island):
 - (i) Season: April 2–June 15 and July 16–August 31.
 - (ii) Closure: June 16–July 15.
 - (iii) Special Black Brant Season Closure: August 16–August 31, only in Izembek and Moffet lagoons.
 - (iv) Special Tundra Swan Closure: All hunting and egg gathering closed in units 9(D) and 10.
 - (3) Western Unit (Umnak Island west to and including Attu Island):
 - (i) Season: April 2–July 15 and August 16–August 31.

(ii) Closure: July 16–August 15.

(b) *Yukon/Kuskokwim Delta Region*.

(1) Season: April 2–August 31.

(2) Closure: 30–day closure dates to be announced by the Service's Alaska Regional Director or his designee, after consultation with local subsistence users, field biologists, and the Association of Village Council President's Waterfowl Conservation Committee. This 30–day period will occur between June 1 and August 15 of each year. A press release announcing the actual closure dates will be forwarded to regional newspapers and radio and television stations and posted in village post offices and stores.

(3) Special Black Brant and Cackling Goose Season Hunting Closure: From the period when egg laying begins until young birds are fledged. Closure dates to be announced by the Service's Alaska Regional Director or his designee, after consultation with field biologists and the Association of Village Council President's Waterfowl Conservation Committee. A press release announcing the actual closure dates will be forwarded to regional newspapers and radio and television stations and posted in village post offices and stores.

(c) *Bristol Bay Region*.

(1) Season: April 2–June 14 and July 16–August 31 (general season); April 2–July 15 for seabird egg gathering only.

(2) Closure: June 15–July 15 (general season); July 16–August 31 (seabird egg gathering).

(d) *Bering Strait/Norton Sound Region*.

(1) Stebbins/St. Michael Area (Point Romanof to Canal Point):

(i) Season: April 15–June 14 and July 16–August 31.

(ii) Closure: June 15–July 15.

(2) Remainder of the region:

(i) Season: April 2–June 14 and July 16–August 31 for waterfowl; April 2–July 19 and August 21–August 31 for all other birds.

(ii) Closure: June 15–July 15 for waterfowl; July 20–August 20 for all other birds.

(e) *Kodiak Archipelago Region*, except for the Kodiak Island roaded area, which is closed to the harvesting of migratory birds and their eggs. The closed area consists of all lands and waters (including exposed tidelands) east of a line extending from Crag Point in the north to the west end of Saltery Cove in the south and all lands and water south of a line extending from Termination Point along the north side of Cascade Lake extending to Anton Larson Bay. Waters adjacent to the closed area are closed to harvest within 500 feet from the water's edge. The offshore islands are open to harvest.

(1) Season: April 2–June 30 and July 31–August 31 for seabirds; April 2–June 20 and July 22–August 31 for all other birds.

(2) Closure: July 1–July 30 for seabirds; June 21–July 21 for all other birds.

(f) *Northwest Arctic Region.*

(1) Season: April 2–June 9 and August 15–August 31 (hunting in general); waterfowl egg gathering May 20–June 9 only; seabird egg gathering May 20–July 12 only; hunting molting/non-nesting waterfowl July 1–July 31 only.

(2) Closure: June 10–August 14, except for the taking of seabird eggs and molting/non-nesting waterfowl as provided in paragraph (f)(1) of this section.

(g) *North Slope Region.*

(1) Southern Unit (Southwestern North Slope regional boundary east to Peard Bay, everything west of the longitude line 158°30'W and south of the latitude line 70°45'N to the west bank of the Ikpiqpuq River, and everything south of the latitude line 69°45'N between the west bank of the Ikpiqpuq River to the east bank of Sagavinirktok River):

(i) Season: April 2–June 29 and July 30–August 31 for seabirds; April 2–June 19 and July 20–August 31 for all other birds.

(ii) Closure: June 30–July 29 for seabirds; June 20–July 19 for all other birds.

(iii) Special Black Brant Hunting Opening: From June 20–July 5. The open area would consist of the coastline, from mean high water line outward to include open water, from Nokotlek Point east to longitude line 158°30'W. This includes Peard Bay, Kugrua Bay, and Wainwright Inlet, but not the Kuk and Kugrua river drainages.

(2) Northern Unit (At Peard Bay, everything east of the longitude line 158°30'W and north of the latitude line 70°45'N to west bank of the Ikpiqpuq River, and everything north of the latitude line 69°45'N between the west bank of the Ikpiqpuq River to the east bank of Sagavinirktok River):

(i) Season: April 6–June 6 and July 7–August 31 for king and common eiders; April 2–June 15 and July 16–August 31 for all other birds.

(ii) Closure: June 7–July 6 for king and common eiders; June 16–July 15 for all other birds.

(3) Eastern Unit (East of eastern bank of the Sagavinirktok River):

(i) Season: April 2–June 19 and July 20–August 31.

(ii) Closure: June 20–July 19.

(4) All Units: yellow-billed loons. Annually, up to 20 yellow-billed loons total for the region may be inadvertently

entangled in subsistence fishing nets in the North Slope Region and kept for subsistence use. Individuals must report each yellow-billed loon inadvertently entangled while subsistence gill net fishing to the North Slope Borough Department of Wildlife Management by the end of the season.

(5) North Coastal Zone (Cape Thompson north to Point Hope and east along the Arctic Ocean coastline around Point Barrow to Ross Point, including Iko Bay, and 5 miles inland).

(i) Migratory bird hunting is permitted from one-half hour before sunrise until sunset, during August.

(ii) No person may at any time, by any means, or in any manner, possess or have in custody any migratory bird or part thereof, taken in violation of subpart C and D of this part.

(iii) Upon request from a Service law enforcement officer, hunters taking, attempting to take, or transporting migratory birds taken during the subsistence harvest season must present them to the officer for species identification.

(h) *Interior Region.*

(1) Season: April 2–June 14 and July 16–August 31; egg gathering May 1–June 14 only.

(2) Closure: June 15–July 15.

(i) *Upper Copper River Region* (Harvest Area: Units 11 and 13) (Eligible communities: Gulkana, Chitina, Tazlina, Copper Center, Gakona, Mentasta Lake, Chistochina and Cantwell).

(1) Season: April 15–May 26 and June 27–August 31.

(2) Closure: May 27–June 26.

(3) The Copper River Basin communities listed above also documented traditional use harvesting birds in Unit 12, making them eligible to hunt in this unit using the seasons specified in paragraph (h) of this section.

(j) *Gulf of Alaska Region.*

(1) Prince William Sound Area (Harvest area: Unit 6 [D]), (Eligible Chugach communities: Chenega Bay, Tatitlek).

(i) Season: April 2–May 31 and July 1–August 31.

(ii) Closure: June 1–30.

(2) Kachemak Bay Area (Harvest area: Unit 15[C] South of a line connecting the tip of Homer Spit to the mouth of Fox River) (Eligible Chugach Communities: Port Graham, Nanwalek).

(i) Season: April 2–May 31 and July 1–August 31.

(ii) Closure: June 1–30.

(k) *Cook Inlet* (Harvest area: portions of Unit 16[B] as specified below) (Eligible communities: Tyonek only).

That portion of (1) Season: April 2–May 31 Unit 16(B) south of the

Skwentna River and west of the Yentna River That, and August 1–31 portion of Unit 16(B) south of the Beluga River, Beluga Lake, and the Triumvirate Glacier.

(2) Closure: June 1–July 31.

(l) *Southeast Alaska.*

(1) Community of Hoonah (Harvest area: National Forest lands in Icy Strait and Cross Sound, including Middle Pass Rock near the Inian Islands, Table Rock in Cross Sound, and other traditional locations on the coast of Yakobi Island. The land and waters of Glacier Bay National Park remain closed to all subsistence harvesting (50 CFR Part 100.3(a)).

(i) Season: glaucous-winged gull egg gathering only: May 15–June 30.

(ii) Closure: July 1–August 31.

(2) Communities of Craig and Hydaburg (Harvest area: small islands and adjacent shoreline of western Prince of Wales Island from Point Baker to Cape Chacon, but also including Coronation and Warren islands).

(i) Season: glaucous-winged gull egg gathering only: May 15–June 30.

(ii) Closure: July 1–August 31.

(3) Community of Yakutat (Harvest area: Icy Bay (Icy Cape to Point Riou), and coastal lands and islands bordering the Gulf of Alaska from Point Manby southeast to Dry Bay).

(i) Season: glaucous-winged gull egg gathering: May 15–June 30.

(ii) Closure: July 1–August 31.

3. In subpart D, add § 92.32 to read as follows:

§ 92.32 Emergency regulations to protect Steller's eiders.

Upon finding that continuation of these subsistence regulations would pose an imminent threat to the conservation of threatened Steller's eiders (*Polysticta stelleri*), the U.S. Fish and Wildlife Service Alaska Regional Director, in consultation with the Co-management Council, will immediately under § 92.21 take action as is necessary to prevent further take. Regulation changes implemented could range from a temporary closure of duck hunting in a small geographic area to large-scale regional or State-wide long-term closures of all subsistence migratory bird hunting. Such closures or temporary suspensions will remain in effect until the Regional Director, in consultation with the Co-management Council, determines that the potential for additional Steller's eiders to be taken no longer exists.

Dated: November 3, 2009.

Thomas L. Strickland,

Assistant Secretary for Fish and Wildlife and Parks.

[FR Doc. E9–27870 Filed 11–19–09; 8:45 am]

BILLING CODE 4310–55–S

Notices

Federal Register

Vol. 74, No. 223

Friday, November 20, 2009

This section of the FEDERAL REGISTER contains documents other than rules or proposed rules that are applicable to the public. Notices of hearings and investigations, committee meetings, agency decisions and rulings, delegations of authority, filing of petitions and applications and agency statements of organization and functions are examples of documents appearing in this section.

DEPARTMENT OF AGRICULTURE

Forest Service

Ochoco National Forest, Lookout Mountain Ranger District; Oregon; Ochoco Summit; OHV Trail EIS

AGENCY: Forest Service, USDA.

ACTION: Notice of Intent to prepare an Environmental Impact Statement.

SUMMARY: The Ochoco National Forest is preparing an Environmental Impact Statement (EIS) to analyze the effects of changing the existing motorized trail system to create and designate a sustainable system of roads, trails and areas open to motor vehicles that will provide legal public access, enhance regulation of unmanaged wheeled motor vehicle travel, protect resources, and decrease conflicts between motorized and non-motorized use on the Ochoco National Forest. Consistent with the Ochoco National Forest Land and Resource Management Plan, as amended, this action is needed to provide to the public a diversity of road and trail opportunities for experiencing a variety of environments and modes of travel.

DATES: Comments concerning the scope of the analysis must be received by December 21, 2009. The draft environmental impact statement is expected to be completed and available for public comment in May 2010. The final environmental impact statement is expected to be completed in August 2010.

ADDRESSES: Send written comments to Ochoco Summit OHV Trail Planning Team, Ochoco National Forest, 3160 NE Third Street, Prineville, Oregon 97754. Alternately, electronic comments may be sent to comments-pacificnorthwest-ochoco@fs.fed.us. Electronic comments must be submitted as part of the actual e-mail message, or as an attachment in plain text (.txt), Microsoft Word (.doc),

rich text format (.rtf), or portable document format (.pdf).

FOR FURTHER INFORMATION CONTACT: Dede Steele, Project Leader, at 3160 NE Third Street, Prineville, Oregon 97754, or at (541) 416-6500, or by e-mail at dsteeler@fs.fed.us.

Responsible Official: The responsible official will be Jeff Walter, Forest Supervisor, Ochoco National Forest, 3160 NE Third Street, Prineville, Oregon 97754.

SUPPLEMENTARY INFORMATION:

Purpose and Need. The Deschutes and Ochoco National Forests are working to complete a Travel Management EIS. If implemented, the two-forest Travel Management EIS would identify specific roads as open for motorized mixed use, and would prohibit off-road travel except where specifically allowed. Opportunities for recreation with off-road vehicles would be reduced. There currently is only one motorized trail in a forested setting on the Ochoco National Forest: The Green Mountain Trail. At just over eight miles, it is not of sufficient length to provide a day of riding to an experienced rider, let alone a weekend of opportunity. As a result, riders are currently venturing off the trail and have created a network of loops. This represents an unauthorized expansion of an undersized trail system. To provide a successful OHV trail system, the system must contain adequate length, diversity, difficulty, loops, alternative routes and other features to provide a quality experience and to keep the use on the designated system.

OHV riders have indicated a desire for additional motorized opportunities other than on mixed use roads. NFS roads are designed primarily for highway-legal vehicles such as passenger cars or log trucks, and are often too wide and too smooth to provide a course with sufficient technical difficulty to keep OHV riders interested and challenged. The intent of providing trails for OHVs is to provide routes with sufficient technical difficulty, diversity of experience and interesting features to keep the riders interested, challenged and engaged with staying on the designated route. The intent of providing mixed use roads is to provide riders with access to a variety of locations on the forest and to provide easy routes for riders who are not

looking for a technically difficult experience.

Proposed Action. The Proposed Action focuses on designating motorized trails and supporting areas, in conjunction with opportunities that would remain on mixed use roads identified in the forest-level Travel Management EIS. The Proposed Action would:

- Designate a system of trails and areas (including staging areas, play areas, riding areas where young riders may be supervised by adults, learner/warm-up loops, picnic and camping areas) by class of vehicle and season of use.
 - Utilize designated open motorized mixed use roads as connectors between trail segments.
 - Designate areas for developed and dispersed camping activities with legal trail access.
 - Implement rehabilitation or restoration activities in previously damaged areas and interconnecting unauthorized or user-created routes to promote recovery, and to prevent confusion about which routes are open and which are not.
 - Establish directional, informational and interpretive signing to: Facilitate proper trail use, safety and enforcement; to provide public information and education; to define trail, riding area, staging area and camp sites locations; to promote recovery of rehabilitation and restoration sites; and to encourage reporting of violations, restoration or maintenance needs.
 - Trails would be designed with width and difficulty appropriate for each intended vehicle type, while roads designated as open in the forest-level Travel Management EIS would not be narrowed to trail standards (*i.e.* designated open roads would remain designated open roads).
- Issues.** Preliminary issues identified include:
- Resource concerns including effects to wildlife, fish, streams, sensitive habitats, forage and weeds.
 - Inadequate quality of experience for off-highway vehicle use on open roads.
 - Retention of non-motorized use experience, potential noise levels.
 - Retention of traditional motorized recreational experience, noise/traffic levels.
 - Concern for increasing off-highway vehicle use on the Forest.

- User conflicts associated with motorized/non-motorized recreation.
- Economic sustainability of road and trail system.
- Monitoring, maintenance and enforcement of appropriate use.
- Potential economic benefits to communities that rely on recreation-tourism.
- Potential impacts to adjacent land owners.
- Potential impacts to livestock and range improvements on permitted allotments.

Comment. Public comments about this proposal are requested in order to assist in identifying issues, determine how to best manage the resources, and to focus the analysis. Comments received to this notice, including names and addresses of those who comment, will be considered part of the public record on this proposed action and will be available for public inspection. Comments submitted anonymously will be accepted and considered; however, those who submit anonymous comments will not have standing to appeal the subsequent decision under 36 CFR parts 215 and 217. Additionally, pursuant to 7 CFR 1.27(d), any person may request the agency to withhold a submission from the public record by showing how the Freedom of Information Act (FOIA) permits such confidentiality. Persons requesting such confidentiality should be aware that, under FOIA, confidentiality may be granted in only very limited circumstances, such as to protect trade secrets. The Forest Service will inform the requester of the agency's decision regarding the request for confidentiality, and where the request is denied, the agency will return the submission and notify the requester that the comments may be resubmitted with or without name and address within a specified number of days.

A draft EIS will be filed with the Environmental Protection Agency (EPA) and available for public review by May, 2010. The EPA will publish a Notice of Availability (NOA) of the draft EIS in the **Federal Register**. The final EIS is scheduled to be available August, 2010.

The comment period on the draft EIS will be 45 days from the date the EPA publishes the notice of availability in the **Federal Register**.

The Forest Service believes, at this early stage, it is important to give reviewers notice of several court rulings related to public participation in the environmental review process. First, reviewers of a draft EIS must structure their participation in the environmental review of the proposal so that it is meaningful and alerts an agency to the

reviewer's position and contentions [*Vermont Yankee Nuclear Power Corp. v. NRDC*, 435 U.S. 519, 553 (1978)]. Also, environmental objections that could be raised at the draft EIS stage but that are not raised until after completion of the final EIS may be waived or dismissed by the courts [*City of Angoon v. Harris*, 490 F. Supp. 1334, 1338 (E.D. Wis. 1980)]. Because of these court rulings, it is very important that those interested in this proposed action participate by the close of the 45-day comment period so that substantive comments and objections are made available to the Forest Service at a time when it can meaningfully consider them and respond to them in the final EIS.

To assist the Forest Service in identifying and considering issues and concerns on the proposed action, comments on the draft EIS should be as specific as possible. It is also helpful if comments refer to specific pages or chapters of the draft statement. Comments may also address the adequacy of the draft EIS of the merits of the alternatives formulated and discussed in the statement. Reviewers may wish to refer to the Council on Environmental Quality Regulations for implementing the procedural provisions of the National Environmental Policy Act at 40 CFR 1503.3 in addressing these points.

In the final EIS, the Forest Service is required to respond to substantive comments received during the comment period for the draft EIS. The Forest Service is the lead agency and the responsible official is the Forest Supervisor, Ochoco National Forest. The responsible official will decide whether and how to change the existing motorized trail system on the Ochoco National Forest. The responsible official will also decide how to mitigate impacts of this action and will determine when and how monitoring of effects will take place.

The Ochoco Summit OHV Trail decision and the reasons for the decision will be documented in the record of decision. That decision will be subject to Forest Service Appeal Regulations (35 CFR Part 215).

Dated: November 12, 2009.

William R. Queen,
District Ranger.

[FR Doc. E9-27801 Filed 11-19-09; 8:45 am]

BILLING CODE 3410-11-M

DEPARTMENT OF COMMERCE

International Trade Administration

[A-570-951]

Certain Woven Electric Blankets From the People's Republic of China: Postponement of Preliminary Determination of Antidumping Duty Investigation

AGENCY: Import Administration, International Trade Administration, Department of Commerce.

DATES: *Effective Date:* November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Drew Jackson or Howard Smith, AD/CVD Operations, Office 4, Import Administration, International Trade Administration, U.S. Department of Commerce, 14th Street and Constitution Avenue, NW., Washington, DC, 20230; telephone: (202) 482-4406 or (202) 482-5193, respectively.

SUPPLEMENTARY INFORMATION:

Postponement of Preliminary Determination

On July 20, 2009, the Department of Commerce (the "Department") initiated the antidumping duty investigation of Certain Woven Electric Blankets from the People's Republic of China. See *Certain Woven Electric Blankets From the People's Republic of China: Initiation of Antidumping Duty Investigation*, 74 FR 37001 (July 27, 2009) ("*Initiation Notice*"). The *Initiation Notice* stated that, "{i}n accordance with section 733(b)(1)(A) of the Act, unless postponed, we will make our preliminary determination no later than 140 days after the date of this initiation." Id. at 37004.

On November 5, 2009, the petitioner made a timely request pursuant to 19 CFR 351.205(e) for a 50-day postponement of the preliminary determination in this investigation. The petitioner requested postponement of the preliminary determination because "the number of factors of production is usually high in this case and will require additional time to research and analyze". There are no compelling reasons to deny the petitioner's request. Therefore, the Department is postponing this preliminary determination under section 733 (c) (1)(A) of the Tariff Act of 1930, as amended (the "Act") by 50 days from December 7, 2009 to January 26, 2010. The deadline for the final determination will continue to be 75 days after the date of the preliminary determination, unless extended.

This notice is issued and published pursuant to sections 733(c) (2) and

777(i)(1) of the Act and 19 CFR 351.205(f)(1).

Dated: November 16, 2009.

John M. Andersen,

Acting Deputy Assistant Secretary for Antidumping and Countervailing Duty Operations.

[FR Doc. E9-27932 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-DS-P

DEPARTMENT OF COMMERCE

International Trade Administration

[A-570-849]

Certain Cut-to-Length Carbon Steel Plate from the People's Republic of China: Notice of Extension of Time Limit for Final Results of Administrative Review

AGENCY: Import Administration, International Trade Administration, Department of Commerce.

EFFECTIVE DATE: November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Demitri Kalogeropoulos, AD/CVD Operations, Office 8, Import Administration, International Trade Administration, U.S. Department of Commerce, 14th Street and Constitution Avenue, NW, Washington, DC 20230; telephone: (202) 482-2623.

Background

On August 10, 2009, the Department of Commerce ("Department") published the preliminary results of the administrative review of the antidumping duty order on certain cut-to-length carbon steel plate from the People's Republic of China, covering the period November 1, 2007, through October 31, 2008. See *Certain Cut-to-Length Carbon Steel Plate From the People's Republic of China: Preliminary Results and Partial Rescission of Antidumping Duty Administrative Review*, 74 FR 39921 (August 10, 2009) ("*Preliminary Results*"). The final results are currently due no later than December 8, 2009.

Extension of Time Limits for Final Results

Pursuant to section 751(a)(3)(A) of the Tariff Act of 1930, as amended ("the Act"), the Department shall issue the final results of an administrative review within 120 days after the date on which the preliminary results are published. However, if it is not practicable to complete the review within this time period, section 751(a)(3)(A) of the Act allows the Department to extend the time period to a maximum of 180 days. Completion of the final results of this

review within the 120-day period is not practicable because the Department needs additional time to analyze and address complicated separate rate and affiliation issues for the final results. Therefore, in accordance with section 751(a)(3)(A) of the Act, given the complexity of issues in this case, we are extending the time limit for completion of the final results by 60 days.

An extension of 60 days from the current deadline of December 8, 2009, would result in a new deadline of February 6, 2010. However, since February 6, 2010, falls on a Saturday, a non-business day, the final results will now be due no later than February 8, 2010, the next business day.

This notice is published in accordance with sections 751(a)(3)(A) and 777(i) of the Act.

Dated: November 16, 2009.

John M. Andersen,

Acting Deputy Assistant Secretary for Antidumping and Countervailing Duty Operations.

[FR Doc. E9-27935 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-DS-S

DEPARTMENT OF COMMERCE

International Trade Administration

[A-533-838]

Carbazole Violet Pigment 23 From India: Extension of Time Limit for Preliminary Results of Antidumping Duty Administrative Review

AGENCY: Import Administration, International Trade Administration, Department of Commerce.

FOR FURTHER INFORMATION CONTACT: Jerrold Freeman or Richard Rimlinger, AD/CVD Operations, Office 5, Import Administration, International Trade Administration, U.S. Department of Commerce, 14th Street and Constitution Avenue, NW, Washington, DC, 20230; telephone: (202) 482-0180 or (202) 482-4477, respectively.

SUPPLEMENTARY INFORMATION:

Background

At the request of interested parties, the Department of Commerce (the Department) initiated an administrative review of the antidumping duty order on carbazole violet pigment 23 from India for the period December 1, 2007, through November 30, 2008. See *Initiation of Antidumping and Countervailing Duty Administrative Reviews and Requests for Revocation in Part*, 74 FR 5821 (February 2, 2009). On September 3, 2009, we extended the due date for the completion of the

preliminary results of review by 75 days. See *Carbazole Violet Pigment 23 From India: Extension of Time Limit for Preliminary Results of Antidumping Duty Administrative Review*, 74 FR 45610 (September 3, 2009). The preliminary results of the review are currently due no later than November 16, 2009.

Extension of Time Limit for Preliminary Results

Section 751(a)(3)(A) of the Tariff Act of 1930, as amended (the Act), requires the Department to make a preliminary determination within 245 days after the last day of the anniversary month of an order for which a review is requested and a final determination within 120 days after the date on which the preliminary determination is published. If it is not practicable to complete the review within these time periods, section 751(a)(3)(A) of the Act allows the Department to extend the time limit for the preliminary determination to a maximum of 365 days after the last day of the anniversary month. See also 19 CFR 351.213(h)(2).

We determine that it is not practicable to complete the preliminary results of this administrative review by the current deadline of November 16, 2009, because we are in the process of analyzing the respondent's recent response to our supplemental questionnaire. Therefore, in accordance with section 751(a)(3)(A) of the Act and 19 CFR 351.213(h)(2), we are extending the time period for issuing the preliminary results of this review by 29 additional days until December 15, 2009. The final results continue to be due 120 days after the publication of the preliminary results.

This notice is published in accordance with sections 751(a)(3)(A) and 777(i)(1) of the Act and 19 CFR 351.213(h)(2).

Dated: November 13, 2009.

John M. Andersen,

Acting Deputy Assistant Secretary for Antidumping and Countervailing Duty Operations.

[FR Doc. E9-27934 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-DS-S

DEPARTMENT OF COMMERCE**Foreign-Trade Zones Board**

[Docket 51–2009]

Foreign-Trade Zone 37—Orange County, NY; Application for Expansion and Reorganization Under Alternative Site Framework

An application has been submitted to the Foreign-Trade Zones (FTZ) Board (the Board) by the County of Orange, New York, grantee of FTZ 37, requesting authority to expand the zone and reorganize under the alternative site framework (ASF) adopted by the Board (74 FR 1170, 01/12/09; correction 74 FR 3987, 01/22/09). The ASF is an option for grantees for the establishment or reorganization of general-purpose zones and can permit significantly greater flexibility in the designation of new “usage-driven” FTZ sites for operators/users located within a grantee’s “service area” in the context of the Board’s standard 2,000-acre activation limit for a general-purpose zone project. The application was submitted pursuant to the provisions of the Foreign-Trade Zones Act, as amended (19 U.S.C. 81a-81u), and the regulations of the Board (15 CFR part 400). It was formally filed on November 12, 2009.

The grantee’s proposed service area under the ASF would be Orange County, New York. If approved, the grantee would be able to serve sites throughout the service area based on companies’ needs for FTZ designation. The proposed service area is adjacent to or within the New York/Newark Customs and Border Protection port of entry.

FTZ 37 was approved by the Board on May 4, 1978 (Board Order 130, 43 FR 20526, 5/12/1978) and expanded on July 9, 1999 (Board Order 1044, 64 FR 38887, 7/20/1999). The applicant is requesting to include its current sites 3 and 7 as “magnet sites”. The applicant proposes that Site 3 be exempt from “sunset” time limits that otherwise apply to sites under the ASF. The applicant is requesting removal of sites 1, 2 and 5. Sites 4 and 6 have lapsed. The applicant is also requesting approval of the following initial “usage-driven” site: *Proposed Site 8* (36 acres)—within the Chester Industrial Park, 29 Elizabeth Drive, Chester, NY.

In accordance with the Board’s regulations, Maureen Hinman of the FTZ staff is designated examiner to evaluate and analyze the facts and information presented in the application and case record and to report findings and recommendations to the Board.

Public comment is invited from interested parties. Submissions (original and 3 copies) shall be addressed to the Board’s Executive Secretary at the address listed below. The closing period for their receipt is January 19, 2010. Rebuttal comments in response to material submitted during the foregoing period may be submitted during the subsequent 15-day period (to February 3, 2010).

A copy of the application will be available for public inspection at the Office of the Executive Secretary, Foreign-Trade Zones Board, Room 2111, U.S. Department of Commerce, 1401 Constitution Avenue, NW., Washington, DC 20230–0002, and in the “Reading Room” section of the Board’s Web site, which is accessible via <http://www.trade.gov/ftz>. For further information, contact Maureen Hinman at maureen.hinman@trade.gov or (202) 482–0627.

Dated: November 12, 2009.

Andrew McGilvray,
Executive Secretary.

[FR Doc. E9–27931 Filed 11–19–09; 8:45 am]

BILLING CODE 3510–DS–P

DEPARTMENT OF COMMERCE**International Trade Administration**

[C–489–502]

Certain Welded Carbon Steel Standard Pipe from Turkey: Extension of Time Limit for Preliminary Results of Countervailing Duty Administrative Review

AGENCY: Import Administration, International Trade Administration, Department of Commerce.

EFFECTIVE DATE: November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Kristen Johnson, AD/CVD Operations, Office 3, Import Administration, International Trade Administration, U.S. Department of Commerce, 14th Street and Constitution Avenue NW, Washington, DC 20230; telephone: (202) 482–4793.

SUPPLEMENTARY INFORMATION:**Background Information**

On April 27, 2009, the U.S. Department of Commerce (the Department) published a notice of initiation of the administrative review of the countervailing duty order on certain welded carbon steel standard pipe from Turkey covering the period of review January 1, 2008, through December 31, 2008. See *Initiation of Antidumping and Countervailing Duty Administrative Reviews*, 74 FR 19042 (April 27, 2009).

The preliminary results are currently due no later than December 1, 2009.

Extension of Time Limit for Preliminary Results

Section 751(a)(3)(A) of the Tariff Act of 1930, as amended (the Act), requires the Department to make a preliminary determination within 245 days after the last day of the anniversary month of an order or finding for which a review is requested. Section 751(a)(3)(A) of the Act further states that if it is not practicable to complete the review within the time period specified, the administering authority may extend the 245-day period to issue its preliminary results by up to 120 days.

The respondents under review are Borusan Mannesmann Boru Sanayi ve Ticaret A.S., Borusan Istikbal Ticaret T.A.S., Tosiya dis Ticaret A.S., Toscelik Profil ve Sac Endustrisi A.S. and the Government of Turkey. In this review, there are 12 programs and new subsidies allegations, which the Department continues to examine. As such, we have determined that it is not practicable to complete the preliminary results of this review within the 245-day period. Therefore, in accordance with section 751(a)(3)(A) of the Act, we are extending the time period for issuing the preliminary results of the review by 120 days. The preliminary results are now due no later than March 31, 2010. The final results continue to be due 120 days after publication of the preliminary results.

This notice is issued and published in accordance with sections 751(a)(3)(A) and 777(i)(1) of the Act.

Dated: November 13, 2009.

John M. Andersen,

Acting Deputy Assistant Secretary for Antidumping and Countervailing Duty Operations.

[FR Doc. E9–27933 Filed 11–19–09; 8:45 am]

BILLING CODE 3510–DS–S

DEPARTMENT OF COMMERCE**International Trade Administration**

[C–580–851]

Dynamic Random Access Memory Semiconductors from the Republic of Korea: Final Results of Countervailing Duty Administrative Review

AGENCY: Import Administration, International Trade Administration, Department of Commerce

SUMMARY: The Department of Commerce has completed an administrative review of the countervailing duty order on dynamic random access memory semiconductors from the Republic of

Korea for the period January 1, 2007, through December 31, 2007. We find that Hynix Semiconductor, Inc. received countervailable subsidies during the period of review, which result in a de minimis subsidy rate.

EFFECTIVE DATE: November 20, 2009.

FOR FURTHER INFORMATION CONTACT: David Neubacher or Shane Subler, Office of AD/CVD Operations, Office 1, Import Administration, International Trade Administration, U.S. Department of Commerce, Room 3069, 14th Street and Constitution Avenue, NW, Washington, DC 20230; telephone: (202) 482-5823 and (202) 482-0189, respectively.

SUPPLEMENTARY INFORMATION:

Background

On August 4, 2009, we published the *Preliminary Results* of the review. See *Dynamic Random Access Memory Semiconductors from the Republic of Korea: Preliminary Results of Countervailing Duty Administrative Review*, 74 FR 38579 (August 4, 2009) (“Preliminary Results”). No interested parties requested a hearing or submitted briefs.

Scope of the Order

The products covered by the order are dynamic random access memory semiconductors (“DRAMs”) from the Republic of Korea (“ROK”), whether assembled or unassembled. Assembled DRAMs include all package types. Unassembled DRAMs include processed wafers, uncut die, and cut die. Processed wafers fabricated in the ROK, but assembled into finished semiconductors outside the ROK are also included in the scope. Processed wafers fabricated outside the ROK and assembled into finished semiconductors in the ROK are not included in the scope.

The scope of the order additionally includes memory modules containing DRAMs from the ROK. A memory module is a collection of DRAMs, the sole function of which is memory. Memory modules include single in-line processing modules, single in-line memory modules, dual in-line memory modules, small outline dual in-line memory modules, Rambus in-line memory modules, and memory cards or other collections of DRAMs, whether unmounted or mounted on a circuit board. Modules that contain other parts that are needed to support the function of memory are covered. Only those modules that contain additional items which alter the function of the module to something other than memory, such as video graphics adapter boards and

cards, are not included in the scope. The order also covers future DRAMs module types.

The scope of the order additionally includes, but is not limited to, video random access memory and synchronous graphics random access memory, as well as various types of DRAMs, including fast page-mode, extended data-out, burst extended data-out, synchronous dynamic RAM, Rambus DRAM, and Double Data Rate DRAM. The scope also includes any future density, packaging, or assembling of DRAMs. Also included in the scope of the order are removable memory modules placed on motherboards, with or without a central processing unit, unless the importer of the motherboards certifies with U.S. Customs and Border Protection (“CBP”) that neither it, nor a party related to it or under contract to it, will remove the modules from the motherboards after importation. The scope of the order does not include DRAMs or memory modules that are re-imported for repair or replacement.

The DRAMs subject to the order are currently classifiable under subheadings 8542.21.8005, 8542.21.8020 through 8542.21.8030, and 8542.32.0001 through 8542.32.0023 of the Harmonized Tariff Schedule of the United States (“HTSUS”). The memory modules containing DRAMs from the ROK, described above, are currently classifiable under subheadings 8473.30.1040, 8473.30.1080, 8473.30.1140, and 8473.30.1180 of the HTSUS. Removable memory modules placed on motherboards are classifiable under subheadings 8443.99.2500, 8443.99.2550, 8471.50.0085, 8471.50.0150, 8517.30.5000, 8517.50.1000, 8517.50.5000, 8517.50.9000, 8517.61.0000, 8517.62.0010, 8517.62.0050, 8517.69.0000, 8517.70.0000, 8517.90.3400, 8517.90.3600, 8517.90.3800, 8517.90.4400, 8542.21.8005, 8542.21.8020, 8542.21.8021, 8542.21.8022, 8542.21.8023, 8542.21.8024, 8542.21.8025, 8542.21.8026, 8542.21.8027, 8542.21.8028, 8542.21.8029, 8542.21.8030, 8542.31.0000, 8542.33.0000, 8542.39.0000, 8543.89.9300, and 8543.89.9600 of the HTSUS. Although the HTSUS subheadings are provided for convenience and customs purposes, the Department’s written description of the scope of this order remains dispositive.

Scope Rulings

On December 29, 2004, the Department received a request from Cisco Systems, Inc., to determine

whether removable memory modules placed on motherboards that are imported for repair or refurbishment are within the scope of the order. See *Notice of Countervailing Duty Order: Dynamic Random Access Memory Semiconductors from the Republic of Korea*, 68 FR 47546 (August 11, 2003) (“CVD Order”). The Department initiated a scope inquiry pursuant to 19 CFR 351.225(e) on February 4, 2005. On January 12, 2006, the Department issued a final scope ruling, finding that removable memory modules placed on motherboards that are imported for repair or refurbishment are not within the scope of the CVD Order provided that the importer certifies that it will destroy any memory modules that are removed for repair or refurbishment. See Memorandum from Stephen J. Claeys to David M. Spooner, regarding Final Scope Ruling, Countervailing Duty Order on DRAMs from the Republic of Korea (January 12, 2006).

Period of Review

The period for which we are measuring subsidies, *i.e.*, the period of review (“POR”), is January 1, 2007, through December 31, 2007.

Final Results of Review

In accordance with 19 CFR 351.221(b)(4)(i), in the *Preliminary Results* we calculated an individual subsidy rate for Hynix Semiconductor, Inc. (“Hynix”), the producer/exporter covered by this administrative review. Neither the petitioner, Micron Technology, Inc., nor the respondent commented on the *Preliminary Results*, and we find that no changes were warranted.

Listed below are the programs we examined in the review and our findings with respect to each of these programs. For a complete analysis of the programs found to be countervailable, not countervailable, and terminated, see *Preliminary Results*.

- I. Programs Determined to Confer Subsidies During the POR
 - A. GOK Entrustment or Direction Prior to 2004
 - B. Operation G-7/HAN Program
 - C. 21st Century Frontier R&D Program
 - D. Import Duty Reduction Program for Certain Factory Automation Items
 - E. Import-Export Bank of Korea Import Financing
- II. Programs Found Not to Have Been Used or Provided No Benefits During the POR
 - A. Short-Term Export Financing
 - B. Reserve for Research and Human Resources Development (formerly Technological Development Reserve) (Article 9 of RSTA /

- formerly, Article 8 of TERCL)
 - C. Tax Credit for Investment in Facilities for Productivity Enhancement (Article 24 of RSTA / Article 25 of TERCL)
 - D. Tax Credit for Investment in Facilities for Special Purposes (Article 25 of RSTA)
 - E. Reserve for Overseas Market Development (formerly, Article 17 of TERCL)
 - F. Reserve for Export Loss (formerly, Article 16 of TERCL)
 - G. Tax Exemption for Foreign Technicians (Article 18 of RSTA)
 - H. Reduction of Tax Regarding the Movement of a Factory That Has Been Operated for More Than Five Years (Article 71 of RSTA)
 - I. Tax Reductions or Exemption on Foreign Investments under Article 9 of the Foreign Investment Promotion Act ("FIPA")/ FIPA (formerly, Foreign Capital Inducement Law)
 - J. Duty Drawback on Non-Physically Incorporated Items and Excessive Loss Rates
 - K. Export Insurance
 - L. Electricity Discounts Under the RLA Program
 - M. Import Duty Reduction for Cutting Edge Products
 - N. System IC 2010 Project
- The calculations will be disclosed to the interested parties in accordance with 19 CFR 351.224(b).

We determine that the total estimated net countervailable subsidy rate for Hynix for calendar year 2007 is 0.06 percent *ad valorem*, which is *de minimis* in accordance with 19 CFR 351.106(c)(1). The Department will instruct CBP to liquidate shipments of DRAMS by Hynix entered or withdrawn from warehouse, for consumption from January 1, 2007, through December 31, 2007, without regard to countervailing duties. See 19 CFR 351.106(c)(1). We intend to issue these instructions 15 days after publication of these final results of review.

On October 3, 2008, the Department published a **Federal Register** notice that, *inter alia*, revoked this order, effective August 11, 2008. See *Dynamic Random Access Memory Semiconductors From the Republic of Korea: Final Results of Sunset Review and Revocation of Order*, 73 FR 57594 (October 3, 2008). As a result, CBP is no longer suspending liquidation for entries of subject merchandise occurring after the revocation. Therefore, there is no need to issue new cash deposit instructions for these final results of review.

This notice serves as a reminder to parties subject to administrative

protective order ("APO") of their responsibility concerning the disposition of proprietary information disclosed under APO in accordance with 19 CFR 351.305(a)(3). Timely written notification of return or destruction of APO materials or conversion to judicial protective order is hereby requested. Failure to comply with the regulations and the terms of an APO is a sanctionable violation.

We are issuing and publishing these results in accordance with sections 751(a)(1) and 777(i)(1) of the Act.

Dated: November 13, 2009.

Ronald K. Lorentzen,

Acting Assistant Secretary for Import Administration.

[FR Doc. E9-27937 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-DS-S

DEPARTMENT OF COMMERCE

Patent and Trademark Office

[Docket No. PTO-P-2009-0052]

Grant of Interim Extension of the Term of U.S. Patent No. 5,407,914; SURFAXIN® (lucinactant)

AGENCY: United States Patent and Trademark Office.

ACTION: Notice of interim patent term extension.

SUMMARY: The United States Patent and Trademark Office has issued an order granting interim extension under 35 U.S.C. 156(d)(5) for a one-year interim extension of the term of U.S. Patent No. 5,407,914.

FOR FURTHER INFORMATION CONTACT:

Mary C. Till by telephone at (571) 272-7755; by mail marked to her attention and addressed to the Commissioner for Patents, Mail Stop Hatch-Waxman PTE, P.O. Box 1450, Alexandria, VA 22313-1450; by fax marked to her attention at (571) 273-7755, or by e-mail to Mary.Till@uspto.gov.

SUPPLEMENTARY INFORMATION: Section 156 of Title 35, United States Code, generally provides that the term of a patent may be extended for a period of up to five years if the patent claims a product, or a method of making or using a product, that has been subject to certain defined regulatory review, and that the patent may be extended for interim periods of up to one year if the regulatory review is anticipated to extend beyond the expiration date of the patent.

On October 6, 2009, Discovery Laboratories Inc., on behalf of patent owner Scripps Research Institute, timely filed an application under 35 U.S.C.

156(d)(5) for an interim extension of the term of U.S. Patent No. 5,407,914. The patent claims the human drug product, SURFAXIN® (lucinactant) and a method of using SURFAXIN® (lucinactant). The application indicates that a New Drug Application, NDA No. 21-746, for the human drug product SURFAXIN® (lucinactant) has been filed, and is currently undergoing regulatory review before the Food and Drug Administration for permission to market or use the product commercially.

Review of the application indicates that except for permission to market or use the product commercially, the subject patent would be eligible for an extension of the patent term under 35 U.S.C. 156, and that the patent should be extended for one year as required by 35 U.S.C. 156(d)(5)(B). Because it is apparent that the regulatory review period will continue beyond the original expiration date of the patent November 17, 2009, interim extension of the patent term under 35 U.S.C. 156(d)(5) is appropriate.

An interim extension under 35 U.S.C. 156(d)(5) of the term of U.S. Patent No. 5,407,914 is granted for a period of one year from the original expiration date of the patent, *i.e.*, until November 17, 2010.

Dated: November 16, 2009.

David J. Kappos,

Under Secretary of Commerce for Intellectual Property and Director of the United States Patent and Trademark Office.

[FR Doc. E9-27903 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-16-P

DEPARTMENT OF COMMERCE

National Oceanic and Atmospheric Administration

RIN 0648-XT03

South Atlantic Fishery Management Council; Public Meetings

AGENCY: National Marine Fisheries Service (NMFS), National Oceanic and Atmospheric Administration (NOAA), Commerce.

ACTION: Notice of public meetings.

SUMMARY: The South Atlantic Fishery Management Council (Council) will hold meetings of its Scientific and Statistical Committee (SSC), Spiny Lobster Committee, Law Enforcement Committee, a joint meeting of its Executive and Finance Committees, Protected Resources Committee, Ecosystem-Based Management Committee, Personnel Committee (Closed Session), Dolphin/Wahoo Committee, Mackerel Committee,

Snapper Grouper Committee, and a meeting of the full Council. The Council will hold an informal public question and answer session with the NMFS Regional Administrator and Council Chairman as well as open public comment periods relative to agenda items. See **SUPPLEMENTARY INFORMATION** for additional details.

DATES: The meeting will be held December 6 - 11, 2009. See

SUPPLEMENTARY INFORMATION for specific dates and times.

ADDRESSES: The meeting will be held at the Sheraton Atlantic Beach Oceanfront Hotel, 2717 W. Fort Macon Road, Atlantic Beach, NC 28512; telephone: (800) 624-8875 or (252) 240-1155; fax: (252) 240-1452. Copies of documents are available from Kim Iverson, Public Information Officer, South Atlantic Fishery Management Council, 4055 Faber Place Drive, Suite 201, North Charleston, SC 29405.

FOR FURTHER INFORMATION CONTACT: Kim Iverson, Public Information Officer; telephone: (843) 571-4366 or toll free at (866) SAFMC-10; fax: (843) 769-4520; email: kim.iverson@safmc.net.

SUPPLEMENTARY INFORMATION:

Meeting Dates:

1. Scientific and Statistical Committee (SSC) Meeting: December 6, 2009, 3 p.m. until 6 p.m., December 7, 2009, 8 a.m. until 5:30 p.m., and December 8, 2009, 8 a.m. until 3 p.m.

The SSC will review and provide recommendations as appropriate on administrative and planning documents relative to the Southeast Data, Assessment, and Review (SEDAR) stock assessment program, and review and finalize terms of reference for SEDAR assessments as appropriate. The Committee will receive updates on the status of Amendments 15B and 16 to the Snapper Grouper Fishery Management Plan (FMP), the Fishery Ecosystem Plan (FEP), and Comprehensive Ecosystem-Based Amendment 1. The SSC will review and provide recommendations as appropriate relative to draft Amendment 18 to the Coastal Migratory Pelagics (mackerel) FMP to address requirements of the reauthorized Magnuson-Stevens Fisheries Conservation and Management Act, Amendments 17A, 17B, 18, and 20 to the Snapper Grouper FMP, Amendment 5 to the Golden Crab FMP, and draft Amendment 10 to the Joint Gulf of Mexico and South Atlantic Spiny Lobster FMP. The SSC will also review and provide recommendations as appropriate on the Comprehensive Annual Catch Limit (ACL) Amendment and Comprehensive Ecosystem-Based Amendment (CE-BA) 2.

Amendment 17A to the Snapper Grouper FMP addresses requirements of the reauthorized Magnuson-Stevens Act to end overfishing and rebuild the red snapper stock in the South Atlantic and establishes a monitoring program, and Amendment 17B addresses requirements of the Act regarding nine other species in the snapper grouper fishery management complex currently listed as undergoing overfishing. Amendment 18 to the Snapper Grouper FMP addresses several management measures relative to the management complex, including expansion of the management unit northward of the Council's current jurisdiction, limiting participation in the commercial fishery for golden tilefish, modifications of management for the black sea bass pot fishery, allocations, changes to the golden tilefish fishing year, improvements to fisheries statistics, and designation of Essential Fish Habitat in northern areas. Amendment 20 to the Snapper Grouper FMP addresses changes to the Wreckfish commercial fishery Individual Transferable Quota (ITQ) program. Amendment 5 to the Golden Crab FMP addresses requirements of the reauthorized Magnuson-Stevens Act to establish ACLs and Accountability Measures for the golden crab fishery. The draft joint Spiny Lobster Amendment 10 currently includes measures addressing tailing permits, the use of "shorts" or undersized lobster in the fishery, and management authority issues. The Comprehensive ACL Amendment addresses requirements to establish ACLs and AMs for all species under the Council's jurisdiction not currently listed as undergoing overfishing as required by the reauthorized Magnuson-Stevens Act. Comprehensive Ecosystem-Based Amendment 2 addresses management measures relative to the harvest of octocorals.

2. Spiny Lobster Committee Meeting: December 7, 2009, 1 p.m. until 2 p.m.

The Spiny Lobster Committee will provide recommendations for the appointment of individuals to serve on the SEDAR stock assessment update for spiny lobster and terms of reference. The Committee will also review results of the Gulf of Mexico Fishery Management Council's scoping meetings and actions, review NOAA Fisheries Service's Biological Opinion, and provide guidance to staff regarding draft Amendment 10 to the Spiny Lobster FMP.

3. Law Enforcement Committee: December 7, 2009, 2 p.m. until 3 p.m.

The Law Enforcement Committee will discuss criteria for a Law Enforcement Officer of the Year award and provide recommendations, receive briefings on the Surveillance and Enforcement of Remote Marine Protected Areas (SERMA) Workshop and a Coral Program FY 2010 proposal, and discuss alternatives related to mackerel nets and provide recommendations as appropriate.

4. Joint Executive and Finance Committees Meeting: December 7, 2009, 3 p.m. until 4 p.m.

The Committees will review the status of the Calendar Year (CY) 2009 budget, the FMP/Amendment timelines, and the Congressional 2010 budget. The Committees will also review the Council's proposed CY 2010 budget.

5. Protected Resources Committee Meeting: December 7, 2009, 4 p.m. until 5:30 p.m.

The Protected Resources Committee will receive presentations regarding the bycatch of shortnose sturgeon, and an update from NOAA Fisheries Southeast Regional Office regarding the following: Petition to list shortnose sturgeon under the Endangered Species Act (ESA); sea turtle take in the large mesh flounder fishery in NC; Biological Opinion for the shrimp fishery due to the incidental take of smalltooth sawfish; and the Center for Biological Diversity's petition to list 80 species of coral under the ESA.

6. Ecosystem-Based Management Committee Meeting: December 8, 2009, 8:30 a.m. until 11 a.m.

The Ecosystem-Based Management Committee will receive an overview of changes to Comprehensive Ecosystem-Based Amendment 2 options paper, the SSC's recommendations on CE-BA 2, a briefing on Florida's Marine Life regulations, and provide guidance to staff regarding CE-BA 2. The Committee will also receive updates from the SERMA Workshop, Coral Reef Conservation Program FY2010 proposal, and Ecosystem Coordination Activities.

7. Dolphin Wahoo Committee Meeting: December 8, 2009, 11 a.m. until 12 noon

The Dolphin Wahoo Committee will review SSC input on the draft ACL Amendment and provide recommendations to staff.

8. Mackerel Committee Meeting: December 8, 2009, 1:30 p.m. until 2:30 p.m.

The Mackerel Committee will review the results from the Gulf of Mexico

Fishery Management Council's public scoping meetings and the Amendment 18 decision document addressing requirements of the Magnuson-Stevens Act, discuss alternatives to address cutting mackerel nets/ trip limits, and discuss legal issues relative to the State of South Carolina regulating dolphin prior to federal regulations.

9. Snapper Grouper Committee Meeting: December 8, 2009, 2:30 p.m. until 5 p.m., December 9, 2009, 8:30 a.m. until 5:30 p.m., and December 10, 2009, 9:30 a.m. until 12 noon

The Snapper Grouper Committee will receive updates on Oculina Bank outreach, status of the Red Snapper Interim Rule request, and a presentation on red snapper rebuilding projections. The Committee will also receive a report relative to snapper grouper management from the SSC. The Committee will review management alternatives in Amendment 17A and modify the document as necessary, including the selection of preferred alternatives. The Committee will review Amendment 17B and the Proposed Rule, modify the document as necessary and develop recommendations on submission of the amendment to the Secretary of Commerce. The Committee will review Amendments 18 and 20, modify the documents as necessary and provide guidance as appropriate. The Committee will review the Comprehensive ACL Amendment alternatives, modify the document as necessary and provide guidance to staff. The Committee will also recommend participants and terms of reference for the black sea bass 2010 SEDAR stock assessment update.

NOTE: There will be an informal public question and answer session with NOAA Fisheries Services' Regional Administrator and the Council Chairman, on December 9, 2009 beginning at 5:30 p.m.

10. Personnel Committee Meeting: December 10, 2009, 8:30 a.m. until 9:30 a.m. (CLOSED SESSION)

The Personnel Committee will meet in a closed session to discuss personnel issues.

11. Council Session: December 10, 2009, 1:30 p.m. until 6 p.m. and December 11, 2009, 8:30 a.m. until 12 noon

Council Session: December 10, 1:30 p.m. until 6 p.m.

From 1:30 p.m. - 1:45 p.m., the Council will call the meeting to order, adopt the agenda, and approve the September 2009 meeting minutes.

From 1:45 p.m. - 2 p.m., the Council will receive a report from the SSC.

NOTE: Interested persons will be provided the opportunity to present oral or written statements regarding matters on the Council agenda beginning at 2 p.m. on Thursday, December 10, 2009. The amount of time provided to individuals will be determined by the Chairman based on the number of individuals wishing to comment.

The Council will also take public comment regarding Amendment 17B to the Snapper Grouper FMP during this time period.

From 4:30 p.m. - 5:15 p.m., the Council will receive a report from the Snapper Grouper Committee, consider recommendations, and take action as appropriate.

From 5:15 p.m. - 5:30 p.m., the Council will receive a report from the SSC Selection Committee.

From 5:30 p.m. - 5:45 p.m., the Council will receive a report from the Spiny Lobster Committee, consider recommendations and take action as appropriate.

From 5:45 p.m. - 6 p.m., the Council will receive a report from the Law Enforcement Committee, consider recommendations and take action as appropriate.

Council Session: December 11, 2009, 8:30 a.m. until 12 noon

From 8:30 a.m. - 9 a.m., the Council will receive legal briefing on litigation (CLOSED SESSION)

From 9 a.m. - 9:15 a.m., the Council will receive a report from the Joint Executive Finance Committee, consider recommendations, and take action as appropriate.

From 9:15 a.m. - 9:30 a.m., the Council will receive a report from the Protected Resources Committee, consider recommendations and take action as appropriate.

From 9:30 a.m. - 9:45 a.m., the Council will receive a report from the Ecosystem-Based Management Committee, consider recommendations and take action as appropriate.

From 9:45 a.m. - 10 a.m., the Council will receive a report from the Dolphin Wahoo Committee, consider recommendations and take action as appropriate.

From 10 a.m. - 10:15 a.m., the Council will receive a report from the Mackerel Committee, consider recommendations and take action as appropriate.

From 10:15 a.m. - 10:30 a.m., the Council will review and develop recommendations on Experimental Fishing Permits as necessary.

From 10:30 a.m. - 12 noon, the Council will receive a status report from NOAA Fisheries Service on commercial quotas by fishing year for: Atlantic king

mackerel, Gulf king mackerel (eastern zone), Atlantic Spanish mackerel, snowy grouper, golden tilefish, wreckfish, greater amberjack, South Atlantic Octocorals and dolphin (soft quota ratios). The Council will also receive a status report on Snapper Grouper Amendment 13C quotas, status of data collection programs to address the black sea bass stock assessment update, and the status of recreational catches versus allocations for Atlantic king mackerel, Atlantic Spanish mackerel, black sea bass, golden tilefish, snowy grouper, red pogy, greater amberjack, and dolphin. The Council will also receive agency and liaison reports, discuss other business, and upcoming meetings.

Documents regarding these issues are available from the Council office (see **ADDRESSES**).

Although non-emergency issues not contained in this agenda may come before this Council for discussion, those issues may not be the subjects of formal final Council action during this meeting. Council action will be restricted to those issues specifically listed in this notice and any issues arising after publication of this notice that require emergency action under section 305 (c) of the Magnuson-Stevens Act, provided the public has been notified of the Council's intent to take final action to address the emergency.

Except for advertised (scheduled) public hearings and public comment, the times and sequence specified on this agenda are subject to change.

Special Accommodations

These meetings are physically accessible to people with disabilities. Requests for sign language interpretation or other auxiliary aids should be directed to the Council office (see **ADDRESSES**) by December 4, 2009.

Dated: November 17, 2009.

Tracey L. Thompson,

Acting Director, Office of Sustainable Fisheries, National Marine Fisheries Service.

[FR Doc. E9-27944 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-22-S

DEPARTMENT OF COMMERCE

International Trade Administration

[A-427-801]

Ball Bearings and Parts Thereof From France: Preliminary Results of Changed-Circumstances Review

AGENCY: Import Administration, International Trade Administration, Department of Commerce.

SUMMARY: The Department of Commerce is conducting a changed-circumstances review of the antidumping duty order on ball bearings and parts thereof from France pursuant to section 751(b) of the Tariff Act of 1930, as amended. We preliminarily determine that, after acquisition by NTN Corporation, SNR Roulements S.A. is the successor-in-interest to pre-acquisition SNR Roulements S.A. Interested parties are invited to comment on these preliminary results.

DATES: *Effective Date:* November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Thomas Schauer or Richard Rimlinger, AD/CVD Operations, Office 5, Import Administration, International Trade Administration, U.S. Department of Commerce, 14th Street and Constitution Avenue, NW., Washington, DC 20230; (202) 482-0410 or (202) 482-4477, respectively.

SUPPLEMENTARY INFORMATION:

Background

On August 21, 2009, SNR Roulements S.A. (SNR) requested that, because NTN Corporation (NTN) acquired a 51-percent interest in SNR, the Department of Commerce (the Department) initiate a changed-circumstances review to determine whether post-acquisition SNR is the successor-in-interest to pre-acquisition SNR.

On September 18, 2009, we initiated a changed-circumstances review. See *Ball Bearings and Parts Thereof From France: Initiation of Antidumping Duty Changed-Circumstances Review*, 74 FR 47920 (September 18, 2009) (*CCR Initiation*).

On September 22, 2009, we sent a questionnaire to SNR. After granting SNR an extension of the deadline, SNR submitted a response on October 23, 2009.

Since the initiation of the review, no other interested party has submitted comments.

Scope of the Order

The products covered by the order are ball bearings and parts thereof. These products include all antifriction bearings that employ balls as the rolling element. Imports of these products are classified under the following categories: Antifriction balls, ball bearings with integral shafts, ball bearings (including radial ball bearings) and parts thereof, and housed or mounted ball bearing units and parts thereof.

Imports of these products are classified under the following Harmonized Tariff Schedule of the

United States (HTSUS) subheadings: 3926.90.45, 4016.93.10, 4016.93.50, 6909.19.50.10, 8431.20.00, 8431.39.00.10, 8482.10.10, 8482.10.50, 8482.80.00, 8482.91.00, 8482.99.05, 8482.99.35, 8482.99.25.80, 8482.99.65.95, 8483.20.40, 8483.20.80, 8483.30.40, 8483.30.80, 8483.50.90, 8483.90.20, 8483.90.30, 8483.90.70, 8708.50.50, 8708.60.50, 8708.60.80, 8708.93.30, 8708.93.60.00, 8708.99.06, 8708.99.31.00, 8708.99.40.00, 8708.99.49.60, 8708.99.58, 8708.99.80.15, 8708.99.80.80, 8803.10.00, 8803.20.00, 8803.30.00, 8803.90.30, 8803.90.90.

As a result of changes to the HTSUS, effective February 2, 2007, the subject merchandise is also classifiable under the following additional HTSUS item numbers: 8708.30.50.90, 8708.40.75, 8708.50.79.00, 8708.50.89.00, 8708.50.91.50, 8708.50.99.00, 8708.70.60.60, 8708.80.65.90, 8708.93.75.00, 8708.94.75, 8708.95.20.00, 8708.99.55.00, 8708.99.68, 8708.99.81.80.

Although the HTSUS item numbers above are provided for convenience and customs purposes, the written description of the scope of the order remains dispositive.

Preliminary Results

In conducting this changed-circumstances review pursuant to section 751(b) of the Tariff Act of 1930, as amended (the Act), the Department has conducted a successor-in-interest analysis. In making a successor-in-interest determination, the Department examines several factors including, but not limited to, changes in the following: (1) Management; (2) production facilities; (3) supplier relationships; (4) customer base. See, e.g., *Brake Rotors From the People's Republic of China: Final Results of Changed Circumstances Antidumping Duty Administrative Review*, 70 FR 69941 (November 18, 2005), and *Notice of Final Results of Changed Circumstances Antidumping Duty Administrative Review: Polychloroprene Rubber From Japan*, 67 FR 58 (January 2, 2002). While no single factor or combination of factors will necessarily provide a dispositive indication of a successor-in-interest relationship, the Department will generally consider the new company to be the successor to the previous company if the new company's resulting operation is not materially dissimilar to that of its predecessor. See *Fresh and Chilled Atlantic Salmon From Norway: Final Results of Changed Circumstances Antidumping Duty Administrative Review*, 64 FR 9979 (March 1, 1999), and *Industrial Phosphoric Acid From*

Israel; Final Results of Antidumping Duty Changed Circumstances Review, 59 FR 6944 (February 14, 1994).

Thus, if the evidence demonstrates that, with respect to the production and sale of subject merchandise, the new company operates as the same business entity as the former company, the Department will accord the new company the same antidumping treatment as its predecessor.

We preliminarily determine that post-acquisition SNR is the successor-in-interest to pre-acquisition SNR. In its August 21, 2009, and October 23, 2009, submissions, SNR provided evidence supporting its claim to be the successor-in-interest to pre-acquisition SNR. Specifically, SNR demonstrated that there were no changes in corporate structure or product mix and only minor changes in management, production facilities,¹ supplier base, or customer base. Moreover, NTN stated that it does not plan to make any significant changes to the pre-acquisition SNR production facilities, management personnel, sources of supply, and customer bases. NTN stated further that it intends to maintain, market, and promote the NTN and SNR brands separately in all markets and for all applications.

In summary, post-acquisition SNR has presented evidence to establish a *prima facie* case of its successorship status. The record indicates that the acquisition of SNR by NTN has not changed the operations of the company in a meaningful way. SNR's management, production facilities, supplier relationships, and customer base are substantially unchanged from their status or circumstances prior to the acquisition. The record evidence demonstrates that the new entity operates essentially in the same manner as the predecessor company. Consequently, we preliminarily determine that post-acquisition SNR should be assigned the same antidumping-duty treatment as pre-acquisition SNR.

Public Comment

Case briefs from interested parties may be submitted not later than 30 days after the date of publication of this notice of preliminary results of changed-circumstances review. Rebuttal briefs from interested parties, limited to the issues raised in the case briefs, may be submitted not later than five days after the time limit for filing the case briefs or comments. Parties who submit case briefs or rebuttal briefs in this

¹ The only changes in production facilities were minor changes in production capacity.

proceeding are requested to submit with each argument a statement of the issue, a summary of the arguments not exceeding five pages, and a table of statutes, regulations, and cases cited.

Interested parties who wish to request a hearing or to participate in a hearing if a hearing is requested must submit a written request to the Assistant Secretary for Import Administration within 30 days of the date of publication of this notice. See 19 CFR 351.310(c). Such requests should contain the following information: (1) The party's name, address, and telephone number; (2) the number of participants; (3) a list of issues to be discussed. Issues raised in the hearing will be limited to those discussed in the case briefs. If requested, any hearing will be held two days after the scheduled date for submission of rebuttal briefs.

The Department will publish in the **Federal Register** a notice of the final results of this changed-circumstances review, including the results of its analysis of issues raised in any written briefs or at the hearing if requested.

As indicated in the *CCR Initiation*, during the course of this changed-circumstances review we will not change any cash-deposit requirements on entries of merchandise subject to the antidumping duty order unless a change is determined to be warranted pursuant to the final results of this changed-circumstances review.

We are issuing and publishing these preliminary results and notice in accordance with sections 751(b) and 777(i)(1) of the Act and 19 CFR 351.216.

Dated: November 16, 2009.

John M. Andersen,

Acting Deputy Assistant Secretary for Antidumping and Countervailing Duty Operations.

[FR Doc. E9-27929 Filed 11-19-09; 8:45 am]

BILLING CODE 3510-DS-P

CONSUMER PRODUCT SAFETY COMMISSION

Sunshine Act Meetings

TIME AND DATE: Wednesday, November 18, 2009, 10 a.m.–12 noon.

PLACE: Hearing Room 420, Bethesda Towers, 4330 East West Highway, Bethesda, Maryland.

STATUS: Closed to the Public.

Matter To Be Considered

Compliance Weekly and Monthly Reports—Commission Briefing

The staff will brief the Commission on various compliance matters.

For a recorded message containing the latest agenda information, call (301) 504-7948.

CONTACT PERSON FOR MORE INFORMATION:

Todd A. Stevenson, Office of the Secretary, U.S. Consumer Product Safety Commission, 4330 East West Highway, Bethesda, MD 20814 (301) 504-7923.

Dated: November 13, 2009.

Todd A. Stevenson,

Secretary.

[FR Doc. E9-27818 Filed 11-19-09; 8:45 am]

BILLING CODE 6355-01-M

DEPARTMENT OF DEFENSE

Department of the Navy

Notice of Availability of Record of Decision for the Swimmer Interdiction Security System

AGENCY: Department of the Navy, DoD.

ACTION: Notice.

SUMMARY: The Department of the Navy (Navy), after carefully weighing the operational and environmental consequences of the proposed action, announces its decision to construct and operate a Swimmer Interdiction Security System at Naval Base Kitsap-Bangor, Washington to find, identify, and interdict surface and underwater intruders for engagement by harbor security forces, in furtherance of the Navy's statutory obligations under Title 10 of the United States Code governing the roles and responsibilities of the Navy. In its decision, the Navy considered applicable laws and executive orders, including an analysis of the effects of its actions in compliance with the Endangered Species Act, the Coastal Zone Management Act, and the National Historic Preservation Act, and the requirements of Executive Order (EO) 12898, *Federal Actions to Address Environmental Justice in Minority Populations and Low Income Populations and EO 13045, Protection of Children from Environmental Health Risks and Safety Risks.*

The proposed action will be accomplished as set out in Alternative 1, described in the Final Environmental Impact Statement (FEIS) as the preferred alternative. Implementation of the preferred alternative could begin immediately.

SUPPLEMENTARY INFORMATION: The Record of Decision (ROD) has been distributed to all those individuals who requested a copy of the FEIS and agencies and organizations that received

a copy of the FEIS. The complete text of the Navy's ROD is available for public viewing on the project Web site at <http://www.nbkeis.gcsaic.com> along with copies of the FEIS and supporting documents. Single copies of the ROD will be made available upon request by contacting Ms. Shannon Kasa, 619-553-3889.

Dated: November 16, 2009.

A.M. Vallandingham,

Lieutenant Commander, Judge Advocate General's Corps, U.S. Navy, Federal Register Liaison Officer.

[FR Doc. E9-27959 Filed 11-19-09; 8:45 am]

BILLING CODE 3810-FF-P

DEPARTMENT OF DEFENSE

Department of the Navy

Notice of Availability and Notice of Public Hearings for the Draft Environmental Impact Statement/ Overseas Environmental Impact Statement for the Guam and Commonwealth of the Northern Mariana Islands Military Relocation

AGENCY: Department of the Navy, DoD.

ACTION: Notice.

SUMMARY: Pursuant to section 102(2)(c) of the National Environmental Policy Act (NEPA); the Council of Environmental Quality (CEQ) Regulations for implementing the procedural provisions of NEPA (Title 40 Code of Federal Regulations [CFR] Parts 1500-1508) and the Department of Navy (DON) regulations for implementing NEPA (32 CFR 775), DON announces the availability of the Draft Environmental Impact Statement/ Overseas Environmental Impact Statement (hereafter called the Draft EIS) to evaluate the potential environmental impacts associated with relocating Marines from Okinawa, Japan to Guam, constructing transient nuclear aircraft carrier berthing facilities, and establishing a U.S. Army Air and Missile Defense Task Force (AMDTF) on Guam.

The DON is the lead Federal agency for development of the Draft EIS. The agencies that have accepted the invitation to participate as cooperating agencies are U.S. Fish and Wildlife Service, Federal Highways Administration, Federal Aviation Administration, U.S. Environmental Protection Agency Region 9, U.S. Office of Insular Affairs, U.S. Department of Agriculture, U.S. Army Corps of Engineers, and U.S. Air Force.

The Draft EIS examines potential environmental impacts from the three

proposed actions included in the Guam and Commonwealth of the Northern Mariana Islands (CNMI) Military Relocation. The Draft EIS also examines off base mission critical, mission support, and community support infrastructure improvements needed to ensure that Joint Region Marianas can provide expanded direct support of the DoD strategic mission and operational readiness in the Western Pacific Region.

The Draft EIS considers reasonable alternatives for siting operational, training, and support facilities on Guam and CNMI in addition to the no-action alternative.

The DON will conduct six (6) public hearings to receive oral and written comments on the Draft EIS. Federal agencies, territorial/local governmental agencies, and interested individuals are invited to be present or represented at the public hearings. This notice announces the dates and locations of the public hearings for this Draft EIS.

A Notice of Intent for this Draft EIS/OEIS was published in the **Federal Register** on March 7, 2007 (72 FR 10186).

DATES: The public comment period for the Draft EIS will start at 8:45 a.m. (Eastern Standard Time) on November 20, 2009, with the publication of a Notice of Availability in the **Federal Register** by the U.S. Environmental Protection Agency and will end at midnight (Eastern Standard Time) on February 17, 2010. All comments on the Draft EIS must be postmarked or submitted by midnight (Eastern Standard Time) on February 17, 2010.

The DON will hold six (6) public hearings to receive oral and written comments from the public on the Draft EIS. These six (6) public hearings will include a two-hour open house session at the beginning of the public hearings where the public can learn more about the proposed actions and potential environmental impacts from project team members and subject matter experts.

Public hearings and open house sessions will be held as follows: Thursday, January 7, 2010, open house from 5 p.m. to 7 p.m. and public hearing from 7 p.m. to 9 p.m., Southern High School, Santa Rita, Guam; Saturday, January 9, 2010, open house from 1 p.m. to 3 p.m. and public hearing from 3 p.m. to 5 p.m., University of Guam Field House, Mangilao, Guam; Monday, January 11, 2010, open house from 5 p.m. to 7 p.m. and public hearing from 7 p.m. to 9 p.m., Yigo Gymnasium, Yigo, Guam; Tuesday, January 12, 2010, open house from 5 p.m. to 7 p.m. and public hearing from 7 p.m. to 9 p.m., Okkodo

High School, Dededo, Guam; Thursday, January 14, 2010, open house from 5 p.m. to 7 p.m. and public hearing from 7 p.m. to 9 p.m., Tinian Elementary School, San Jose, Tinian; and Friday, January 15, 2010, open house from 5 p.m. to 7 p.m. and public hearing from 7 p.m. to 9 p.m., Multi-Purpose Center, Susupe, Saipan.

More information about the public hearings and open house sessions can be found on the official project Web site at <http://www.guambuildupeis.us/>.

ADDRESSES: The public can provide comments during the open houses/public hearings through the Web site at <http://www.guambuildupeis.us/>, or by mail at: Joint Guam Program Office, c/o Naval Facilities Engineering Command Pacific, 258 Makalapa Drive, Suite 100, Pearl Harbor, Hawaii 96860-3134, *Attention:* GPMO.

Electronic copies of the Draft EIS can be downloaded from the official project Web site at <http://www.guambuildupeis.us/>. Copies of the Draft EIS are available for public review at the following libraries: University of Guam Robert F. Kennedy Memorial Library, Government Documents, Tan Siu Lin Building, UOG Station, Mangilao, GU 96923; Nieves M. Flores Memorial Library, 254 Martyr Street, Hagåtña, GU 96910; Joeten-Kiyu Public Library, P.O. Box 501092, Saipan, MP 96950; Olympio T. Borja Memorial Library, P.O. Box 501250, Saipan, MP 96950 and Tinian Public Library, P.O. Box 520704, Tinian, MP 96952.

Electronic copies of the Draft EIS and copies of the executive summary can also be obtained on the project Web site at <http://www.guambuildupeis.us/>, or by contacting the Joint Guam Program Office, c/o Naval Facilities Engineering Command Pacific, 258 Makalapa Drive, Suite 100, Pearl Harbor, Hawaii 96860-3134, *Attention:* GPMO.

SUPPLEMENTARY INFORMATION: The overarching purpose for the proposed actions is to locate U.S. military forces to meet international agreement and treaty obligations and fulfill U.S. government national security to provide mutual defense, deter aggression, and dissuade coercion in the Western Pacific Region. The need of the proposed actions is to meet various criteria based upon U.S. policy, international agreements, and treaties, including but not limited to positioning forces to defend the homeland and U.S. Pacific territories; respond within a timely response range; maintain regional stability; provide flexibility to respond to regional threats; defend U.S., Japan, and allied interests, and other defense related criteria. Guam's location as the

westernmost part of the United States is critical to U.S. national security interests. In implementing U.S. national security policy, the Department of Defense would increase the role of Guam and the CNMI through the relocation of Marines to Guam, increased presence of a transient aircraft carrier, and enhanced capability to protect the U.S. homeland, Pacific territories, forces, and its allies from ballistic missile attacks.

As a result of reviews of the U.S. defense posture in the Pacific region and a parallel review of U.S. force posture supporting the U.S. alliance with Japan, a portion of U.S. Marine Corps forces currently located in Okinawa, Japan would be relocated to Guam. This relocation is proposed to occur concurrent with proposed wharf construction in Guam's Apra Harbor to support the Navy transiting nuclear aircraft carriers. An Army AMDTF is also proposed for Guam to protect the U.S. homeland, Pacific territories, forces and its allies against the threat of harm from ballistic missile attacks.

The Draft EIS was prepared to support inform decision making based on an understanding of the environmental impacts of the proposed Guam and CNMI military relocation and take measures to protect, restore, and enhance the environment. The decisions to be made are whether and how to implement the proposed actions.

The three proposed actions are briefly stated as follows: (1) Develop and construct facilities and infrastructure to support approximately 8,600 Marines and their 9,000 dependents relocated from Okinawa to Guam and to support training and operations on Guam and CNMI for those relocated Marines; (2) Develop and construct a new deep-draft wharf with shore side infrastructure improvements creating the capability in Apra Harbor, Guam to support a transient nuclear-powered aircraft carrier; and (3) Develop and construct facilities and infrastructure on Guam to support approximately 600 military personnel and their 900 dependents and to establish and operate an Army AMDTF.

The project locations addressed in Draft EIS are on Guam and CNMI. Guam and CNMI are part of the Mariana Islands archipelago.

Facilities construction and improvements would be necessary to accommodate the proposed actions. The proposed actions would entail increased operational activities associated with Marine Corps and Army basing, more frequent ship berthing, and the establishment of aviation maintenance operations and facilities. There would

also be increased opportunities for additional military personnel to meet critical training requirements. Training could take the form of communications, command and control, combat skills, aviation, logistics, amphibious vehicle maneuvers, and weapons firing activities. Thus, required construction would include the facilities and infrastructure for maintaining a permanent Marine Corps and Army presence on Guam, and the creation of new training ranges to accommodate training a larger population of military personnel. In summary, implementation of the proposed actions would result in the following: a temporary increase in population related to the construction-related work force; a permanent increase in number of military and civilian personnel and dependents on Guam; an increase in transient presence on Guam and Tinian; an increase in number and type of major equipment assets to support military personnel and operations (e.g., aircraft, ships, amphibious watercraft); an increase in number and type of training activities; construction of new facilities; improvements to existing facilities; improvements to existing infrastructure (including roads and utilities); and the potential acquisition or long-term leasing of additional land to support the Marine Corps main cantonment area and live fire training ranges on Guam.

To accomplish the Guam and CNMI proposed actions, the DoD has considered developmental and operational alternatives as required by NEPA. The Draft EIS analyzes a range of alternatives for the proposed actions including the no action alternative, which represents the baseline.

The Draft EIS provides information on the affected environment and impacts of the proposed actions for eighteen distinct resource areas. Volume 1 of the Draft EIS provides an overview of the proposed actions and alternatives. Volumes 2 through 5 of the Draft EIS provide details on the impacts of individual proposed Marine Corps, Navy and Army actions while Volume 6 addresses island-wide impacts of utilities and proposed roadway improvement projects. Volume 7 provides a summary of the impacts of all of the proposed actions should the preferred alternatives be implemented as well as a discussion of cumulative impacts. The Draft EIS evaluates the following resource areas: groundwater, marine water quality, coral, terrestrial biology, socioeconomic, cultural resources, recreation, roadways, air quality, noise, and utilities (including water, power and wastewater), among others.

FOR FURTHER INFORMATION CONTACT: Joint Guam Program Office in Guam at (671) 333-2302 or in Washington, DC at (703) 602-4716.

Dated: November 16, 2009.

A.M. Vallandingham

Lieutenant Commander, Judge Advocate General's Corps, U.S. Navy, Federal Register Liaison Officer.

[FR Doc. E9-27960 Filed 11-19-09; 8:45 am]

BILLING CODE 3810-FF-P

DEPARTMENT OF EDUCATION

**Submission for OMB Review;
Comment Request**

AGENCY: Department of Education.

SUMMARY: The Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management invites comments on the submission for OMB review as required by the Paperwork Reduction Act of 1995.

DATES: Interested persons are invited to submit comments on or before December 21, 2009.

ADDRESSES: Written comments should be addressed to the Office of Information and Regulatory Affairs, *Attention:* Education Desk Officer, Office of Management and Budget, 725 17th Street, NW., Room 10222, New Executive Office Building, Washington, DC 20503, be faxed to (202) 395-5806 or send e-mail to oir_submission@omb.eop.gov.

SUPPLEMENTARY INFORMATION: Section 3506 of the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35) requires that the Office of Management and Budget (OMB) provide interested Federal agencies and the public an early opportunity to comment on information collection requests. OMB may amend or waive the requirement for public consultation to the extent that public participation in the approval process would defeat the purpose of the information collection, violate State or Federal law, or substantially interfere with any agency's ability to perform its statutory obligations. The Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management, publishes that notice containing proposed information collection requests prior to submission of these requests to OMB. Each proposed information collection, grouped by office, contains the following: (1) Type of review requested, e.g., new, revision, extension, existing

or reinstatement; (2) Title; (3) Summary of the collection; (4) Description of the need for, and proposed use of, the information; (5) Respondents and frequency of collection; and (6) Reporting and/or Recordkeeping burden. OMB invites public comment.

Dated: November 16, 2009.

Sheila Carey,

Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management.

**Office of Special Education and
Rehabilitative Services**

Type of Review: Extension.

Title: Annual Performance Report for the State Grant for Assistive Technology Program.

Frequency: Annually.

Affected Public: Federal Government; Not-for-profit institutions; State, Local, or Tribal Gov't, SEAs or LEAs.

Reporting and Recordkeeping Hour Burden:

Responses: 56.

Burden Hours: 26,768.

Abstract: Section 4 of the Assistive Technology (AT) Act of 1998, as amended, requires states to submit annual data reports. This instrument helps the grantees report annual data related to the required activities implemented by the State under the AT Act. This data is used by Rehabilitation Services Administration (RSA) in order to prepare required annual reports to Congress. RSA calls this data collection an Annual Progress Report.

Requests for copies of the information collection submission for OMB review may be accessed from <http://edicsweb.ed.gov>, by selecting the "Browse Pending Collections" link and by clicking on link number 4131. When you access the information collection, click on "Download Attachments" to view. Written requests for information should be addressed to U.S. Department of Education, 400 Maryland Avenue, SW., LBJ, Washington, DC 20202-4537. Requests may also be electronically mailed to the Internet address ICDocketMgr@ed.gov or faxed to 202-401-0920. Please specify the complete title of the information collection when making your request.

Comments regarding burden and/or the collection activity requirements should be electronically mailed to ICDocketMgr@ed.gov. Individuals who use a telecommunications device for the deaf (TDD) may call the Federal Information Relay Service (FIRS) at 1-800-877-8339.

[FR Doc. E9-27901 Filed 11-19-09; 8:45 am]

BILLING CODE 4000-01-P

DEPARTMENT OF EDUCATION**Notice of Proposed Information Collection Requests**

AGENCY: Department of Education.

SUMMARY: The Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management, invites comments on the proposed information collection requests as required by the Paperwork Reduction Act of 1995.

DATES: Interested persons are invited to submit comments on or before January 19, 2010.

SUPPLEMENTARY INFORMATION: Section 3506 of the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35) requires that the Office of Management and Budget (OMB) provide interested Federal agencies and the public an early opportunity to comment on information collection requests. OMB may amend or waive the requirement for public consultation to the extent that public participation in the approval process would defeat the purpose of the information collection, violate State or Federal law, or substantially interfere with any agency's ability to perform its statutory obligations. The Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management, publishes that notice containing proposed information collection requests prior to submission of these requests to OMB. Each proposed information collection, grouped by office, contains the following: (1) Type of review requested, e.g., new, revision, extension, existing or reinstatement; (2) Title; (3) Summary of the collection; (4) Description of the need for, and proposed use of, the information; (5) Respondents and frequency of collection; and (6) Reporting and/or Recordkeeping burden. OMB invites public comment.

The Department of Education is especially interested in public comment addressing the following issues: (1) Is this collection necessary to the proper functions of the Department; (2) will this information be processed and used in a timely manner; (3) is the estimate of burden accurate; (4) how might the Department enhance the quality, utility, and clarity of the information to be collected; and (5) how might the Department minimize the burden of this collection on the respondents, including through the use of information technology.

Dated: November 17, 2009.

James Hylar,

Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management.

Federal Student Aid

Type of Review: Extension.

Title: Federal Family Education Loan (FFEL) Program: Federal Consolidation Loan Application and Promissory Note and Related Documents.

Frequency: On occasion.

Affected Public: Individuals or households; Private Sector.

Reporting and Recordkeeping Hour Burden:

Responses: 84,705.

Burden Hours: 117,527.

Abstract: The Federal Consolidation Loan Application and Promissory Note serves as the means by which a borrower applies for a Federal Consolidation Loan and promises to repay the loan. Related documents included as part of this collection are (1) Additional Loan Listing Sheet (provides additional space for a borrower to list loans that he or she wishes to consolidate, if there is insufficient space on the Federal Consolidation Loan Application and Promissory Note); (2) Request to Add Loans (serves as the means by which a borrower may add other loans to an existing Federal Consolidation Loan within a specified time period); and (3) Loan Verification Certificate (serves as the means by which a consolidating lender obtains the information needed to pay off the holders of the loans that the borrower wants to consolidate).

Requests for copies of the proposed information collection request may be accessed from <http://edicsweb.ed.gov>, by selecting the "Browse Pending Collections" link and by clicking on link number 4175. When you access the information collection, click on "Download Attachments" to view. Written requests for information should be addressed to U.S. Department of Education, 400 Maryland Avenue, SW., LBJ, Washington, DC 20202-4537. Requests may also be electronically mailed to ICDocketMgr@ed.gov or faxed to 202-401-0920. Please specify the complete title of the information collection when making your request.

Comments regarding burden and/or the collection activity requirements should be electronically mailed to ICDocketMgr@ed.gov. Individuals who use a telecommunications device for the deaf (TDD) may call the Federal

Information Relay Service (FIRS) at 1-800-877-8339.

[FR Doc. E9-27958 Filed 11-19-09; 8:45 am]

BILLING CODE 4000-01-P

DEPARTMENT OF EDUCATION**Submission for OMB Review; Comment Request**

AGENCY: Department of Education.

SUMMARY: The Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management invites comments on the submission for OMB review as required by the Paperwork Reduction Act of 1995.

DATES: Interested persons are invited to submit comments on or before December 21, 2009.

ADDRESSES: Written comments should be addressed to the Office of Information and Regulatory Affairs, *Attention:* Education Desk Officer, Office of Management and Budget, 725 17th Street, NW., Room 10222, New Executive Office Building, Washington, DC 20503, be faxed to (202) 395-5806 or send e-mail to oir_submission@omb.eop.gov.

SUPPLEMENTARY INFORMATION: Section 3506 of the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35) requires that the Office of Management and Budget (OMB) provide interested Federal agencies and the public an early opportunity to comment on information collection requests. OMB may amend or waive the requirement for public consultation to the extent that public participation in the approval process would defeat the purpose of the information collection, violate State or Federal law, or substantially interfere with any agency's ability to perform its statutory obligations. The Acting Director, Information Collection Clearance Division, Regulatory Information Management Services, Office of Management, publishes that notice containing proposed information collection requests prior to submission of these requests to OMB. Each proposed information collection, grouped by office, contains the following: (1) Type of review requested, e.g. new, revision, extension, existing or reinstatement; (2) Title; (3) Summary of the collection; (4) Description of the need for, and proposed use of, the information; (5) Respondents and frequency of collection; and (6) Reporting and/or Recordkeeping burden. OMB invites public comment.

Dated: November 16, 2009.

Sheila Carey,

*Acting Director, Information Collection
Clearance Division, Regulatory Information
Management Services, Office of Management.*

**Office of Elementary and Secondary
Education**

Type of Review: New.

Title: Guidance on Title I, Part A
Waivers.

Frequency: One time.

Affected Public: Business or other for-
profit; State, Local, or Tribal Gov't,
SEAs or LEAs.

*Reporting and Recordkeeping Hour
Burden:*

Responses: 947.

Burden Hours: 29,640.

Abstract: The U.S. Department of Education (ED) plans to issue guidance inviting waivers related to the use of FY 2009 Title I, Part A funds available through the America Recovery and Reinvestment Act of 2009 (ARRA), waivers related to certain Title I, Part A statutory and regulatory provisions related to public school choice and supplemental educational services, and waivers related to maintenance of effort requirements. The guidance provides instructions for State educational agencies (SEA) on how they may apply to ED for waivers and instructions for local educational agencies (LEAs) on how they may implement the waivers obtained by their SEA.

Requests for copies of the information collection submission for OMB review may be accessed from <http://edicsweb.ed.gov>, by selecting the "Browse Pending Collections" link and by clicking on link number 4002. When you access the information collection, click on "Download Attachments" to view. Written requests for information should be addressed to U.S. Department of Education, 400 Maryland Avenue, SW., LBJ, Washington, DC 20202-4537. Requests may also be electronically mailed to the Internet address ICDocketMgr@ed.gov or faxed to 202-401-0920. Please specify the complete title of the information collection when making your request.

Comments regarding burden and/or the collection activity requirements should be electronically mailed to ICDocketMgr@ed.gov. Individuals who use a telecommunications device for the deaf (TDD) may call the Federal Information Relay Service (FIRS) at 1-800-877-8339.

[FR Doc. E9-27902 Filed 11-19-09; 8:45 am]

BILLING CODE 4000-01-P

DEPARTMENT OF ENERGY

**Environmental Management Site-
Specific Advisory Board, NV**

AGENCY: Department of Energy.

ACTION: Notice of open meeting.

SUMMARY: This notice announces a meeting of the Environmental Management Site-Specific Advisory Board (EM SSAB), Nevada Test Site. The Federal Advisory Committee Act (Pub. L. 92-463, 86 Stat. 770) requires that public notice of this meeting be announced in the **Federal Register**.

DATES: Wednesday, December 16, 2009, 5 p.m.

ADDRESSES: Centennial Hills Library, 6711 North Buffalo Drive, Las Vegas, Nevada 89131.

FOR FURTHER INFORMATION CONTACT:

Denise Rupp, Board Administrator, 232 Energy Way, M/S 505, North Las Vegas, Nevada 89030. *Phone:* (702) 657-9088; *Fax* (702) 295-5300 or *E-mail:* ntscab@nv.doe.gov.

SUPPLEMENTARY INFORMATION:

Purpose of the Board: The purpose of the Board is to make recommendations to DOE in the areas of environmental restoration, waste management, and related activities.

Tentative Agenda:

Sub-Committee Updates

- Industrial Sites Committee
- Membership Committee
- Outreach Committee
- Soils Committee
- Transportation/Waste Committee
- Underground Test Area Committee

Public Participation: The EM SSAB, Nevada Test Site, welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Denise Rupp at least seven days in advance of the meeting at the phone number listed above. Written statements may be filed with the Board either before or after the meeting. Individuals who wish to make oral presentations pertaining to agenda items should contact Denise Rupp at the telephone number listed above. The request must be received five days prior to the meeting and reasonable provision will be made to include the presentation in the agenda. The Deputy Designated Federal Officer is empowered to conduct the meeting in a fashion that will facilitate the orderly conduct of business. Individuals wishing to make public comment will be provided a

maximum of five minutes to present their comments.

Minutes: Minutes will be available by writing to Denise Rupp at the address listed above or at the following Web site: <http://www.ntscab.com/MeetingMinutes.htm>.

Issued at Washington, DC on November 17, 2009.

Rachel Samuel,

Deputy Committee Management Officer.

[FR Doc. E9-27906 Filed 11-19-09; 8:45 am]

BILLING CODE 6450-01-P

DEPARTMENT OF ENERGY

State Energy Advisory Board (STEAB)

AGENCY: Department of Energy.

ACTION: Notice of open teleconference.

SUMMARY: This notice announces a meeting of the State Energy Advisory Board (STEAB). The Federal Advisory Committee Act (Pub. L. 2-463; 86 Stat. 770) requires that public notice of these meetings be announced in the **Federal Register**.

DATES: December 17, 2009, 1 to 2 p.m. EDT.

FOR FURTHER INFORMATION CONTACT: Gary Burch, STEAB Designated Federal Officer, Office of Commercialization and Project Management, Energy Efficiency Division, Golden Field Office, U.S. Department of Energy, 1617 Cole Boulevard, Golden, CO 80401, Telephone 303-275-4801.

SUPPLEMENTARY INFORMATION:

Purpose of the Board: To make recommendations to the Assistant Secretary for the Office of Energy Efficiency and Renewable Energy regarding goals and objectives, programmatic and administrative policies, and to otherwise carry out the Board's responsibilities as designated in the State Energy Efficiency Programs Improvement Act of 1990 (Pub. L. 101-440).

Tentative Agenda: Discuss ways STEAB can support DOE's implementation of the Economic Recovery Act and update members of the Board on routine business matters.

Public Participation: The meeting is open to the public. Written statements may be filed with the Board either before or after the meeting. Members of the public who wish to make oral statements pertaining to agenda items should contact Gary Burch at the address or telephone number listed above. Requests to make oral comments must be received five days prior to the meeting; reasonable provision will be made to include requested topic(s) on

the agenda. The Chair of the Board is empowered to conduct the meeting in a fashion that will facilitate the orderly conduct of business.

Minutes: The minutes of the meeting will be available for public review and copying within 60 days on the STEAB Web site, <http://www.steab.org>.

Issued at Washington, DC, on November 17, 2009.

Rachel Samuel,

Deputy Committee Management Officer.

[FR Doc. E9-27907 Filed 11-19-09; 8:45 am]

BILLING CODE 6450-01-P

DEPARTMENT OF ENERGY

Senior Executive Service; Performance Review Board

AGENCY: U.S. Department of Energy.

ACTION: SES Performance Review Board Standing Register.

SUMMARY: This notice provides the Performance Review Board Standing Register for the Department of Energy. This listing supersedes all previously published lists of PRB members.

DATES: These appointments are effective as of September 30, 2009.

ADAMS, VINCENT NMN

ALLISON, JEFFREY M

AMARAL, DAVID M

ANDERSON, CYNTHIA V

ANDERSON, MARGOT H

AOKI, STEVEN NMN

ARMSTRONG, DAVID J

ASCANIO, XAVIER NMN

BAKER, KENNETH E

BARKER JR, WILLIAM L

BARWELL, OWEN F

BASHISTA, JOHN R

BATTERSHELL, CAROL J

BAUER, CARL O

BEAMON, JOSEPH A

BEARD, JEANNE M

BEARD, SUSAN F

BEAUDRY-LOSQUE, JACQUES A

BEAUSOLEIL, GEOFFREY L

BEKKEDAHL, LARRY N

BELL, MELODY C

BERKOWITZ, BARRY E

BIENIAWSKI, ANDREW J

BISCONTI, GIULIA R

BLACK, RICHARD L

BLACK, STEVEN K

BOARDMAN, KAREN L

BONILLA, SARAH J

BORGSTROM, CAROL M

BORGSTROM, HOWARD G

BOSCO, PAUL NMN

BOULDEN III, JOHN S

BOYD, DAVID O

BOYD, GERALD G

BOYKO, THOMAS R

BOYLE, WILLIAM J

BRESE, ROBERT F

BREWER, STEPHANIE J

BROCKMAN, DAVID A

BROMBERG, KENNETH M

BROTT, MATTHEW J

BROWN III, ROBERT J

BROWN, FRED L

BROWN, STEPHANIE H

BROWN, THOMAS E

BRYAN, WILLIAM N

BURCH, LINDA C

BURNS, ALLEN L

BURROWS, CHARLES W

BUTTRESS, LARRY D

BUZZARD, CHRISTINE M

CADIEUX, GENA E

CALLAHAN, SAMUEL N

CAMPBELL II, HUGH T

CANNON, SCOTT C

CARABETTA, RALPH A

CAROSINO, ROBERT M

CARY, STEVEN V

CAVANAGH, JAMES J

CERVENY, THELMA J

CHALK, STEVEN G

CHARBONEAU, STACY L

CHECK, PETER L

CHOI, JOANNE Y

CHUNG, DAE Y

CLAPPER, DANIEL R

CLARK, DIANA D

CLARK, LARRY W

COHEN, DANIEL NMN

COLLARD, GEORGE W

COLLAZO, YVETTE T

COMO, ANTHONY J

CONNOR, MICHAEL A

CONTI, JOHN J

COOK, JOHN S

COOKE JR, KEVIN R

CORBIN, ROBERT F

COREY, RAY J

COSTLOW, BRIAN D

CRAIG JR, JACKIE R

CRANDALL, DAVID H

CRAWFORD, DAVID W

CRAWFORD, GLEN D

CROUTHER, DESI A

CUGINI, ANTHONY V

DAUB, VERNON NMN

DAVENPORT, SHARI T

DAVIS, KIMBERLY A

DAVIS, PATRICK B

DEAROLPH, DOUGLAS J

DECKER, ANITA J

DEDIK, PATRICIA NMN

DEENEY, CHRISTOPHER NMN

DEHAVEN, DARREL S

DEHMER, PATRICIA M

DEHORATIIS JR, GUIDO NMN

DELWICHE, GREGORY K

DER, VICTOR K

DIAMOND, BRUCE M

DICAPUA, MARCO S

DIFIGLIO, CARMEN NMN

DOWELL, JONATHAN A

DUKE JR, RICHARD D

DYER, J RUSSELL

ECKROADE, WILLIAM A

EDWARDS JR, ROBERT H

EGGER, MARY H

EHLI, CATHY L

EKIMOFF, LANA NMN

ELKIND, JONATHAN H

ELY, LOWELL V

ERHART, STEVEN C

ERICKSON, LEIF NMN

ESCHENBERG, JOHN R

FAUL, JERRY W

FERRARO, PATRICK M

FIGLIO, JAMES J

FOLEY, KATHLEEN Y

FOLEY, THOMAS C

FRANCO JR., JOSE R

FRANKLIN, RITA R

FRANTZ, DAVID G

FREMONT, DOUGLAS E

FRESCO, MARY ANN E

FURRER, ROBIN R

FURSTENAU, RAYMOND V

FYGLI, ERIC J

GARCIA, DONALD J

GASPEROW, LESLEY A

GEISER, DAVID W

GELLES, CHRISTINE M

GENDRON, MARK O

GERRARD, JOHN E

GIBBS, ROBERT C

GIBSON JR, WILLIAM C

GILBERTSON, MARK A

GILLO, JEHANNE E

GIST, WALTER J

GOLAN, PAUL M

GOLDSMITH, ROBERT NMN

GOLUB, SAL JOSEPH

GOODRUM, WILLIAM S

GOODWIN, KARL E

GORDON, THEANNE E

GOTTLIEB, PAUL A

GREENAUGH, KEVIN C

GREENWOOD, JOHNNIE D

GRUENSPECHT, HOWARD K

GUEVARA, ARNOLD E

GUEVARA, KAREN C

HANDSCHY, MARK A

HANDWERKER, ALAN I

HANNIGAN, JAMES J

HARDWICK JR, RAYMOND J

HARMS, TIMOTHY C

HARRELL, JEFFREY P

HARRINGTON, PAUL G

HARRIS, ROBERT J

HARTMAN, JOHN R

HARVEY, STEPHEN J

HASS, RICKEY R

HEGBURG, ALAN S

HENNEBERGER, KAREN O

HENNEBERGER, MARK W

HENRY, EUGENE A

HERRERA, C ROBERT D

HILL, JOANNE NMN

HINE, SCOTT E

HINTZE, DOUGLAS E

HOFFMAN, DENNIS J

HOFFMAN, PATRICIA A

HOLLAND, MICHAEL D

HOLLAND, MICHAEL J

HOLLAND, WENDOLYN S

HOLLRITH, JAMES W

HORTON, LINDA L
HOWARD, MICHAEL F
HUFFER, WARREN L
HUIZENGA, DAVID G
HUNTEMAN, WILLIAM J
HYNDMAN, JOHN E
JENKINS, ROBERT G
JOHNSON, DAVID F
JOHNSON, ROBERT SHANE
JOHNSON, SANDRA L
JOHNSTON, MARC NMN
JONAS, DAVID S
JONES, GREGORY A
JONES, MARCUS E
JONES, WAYNE NMN
JUAREZ, LIOVA D
KAEMPF, DOUGLAS E
KANE, MICHAEL C
KEARNEY, JAMES H
KELLY, HENRY C
KELLY, LARRY C
KENCHINGTON, HENRY S
KENDELL, JAMES M
KETCHAM, TIMOTHY E
KIDD IV, RICHARD G
KIGHT, GENE H
KILROY, EDWARD F
KLARA, SCOTT M
KLAUSING, KATHLEEN A
KLING, JON NMN
KNOELL, THOMAS C
KNOLL, WILLIAM S
KOLB, INGRID A C
KONOPNICKI, THAD T
KOURY, JOHN F
KOUTS, CHRISTOPHER A
KOVAR, DENNIS G
KRAHN, STEVEN L
KROL, JOSEPH J
KUNG, HUIJOU HARRIET
KUSNEZOV, DIMITRI F
LAGDON JR, RICHARD H
LAMBERT, JAMES B
LANGE, ROBERT G
LANTHRUM, J GARY
LAWRENCE, ANDREW C
LAWRENCE, STEVEN J
LAY, WILLIAM G
LEATHLEY, KIMBERLY A
LECKEY, THOMAS J
LEE, STEVEN NMN
LEE, TERRI TRAN
LEGG, KENNETH E
LEHMAN, DANIEL R
LEIFHEIT, KEVIN R
LEISTIKOW, DANIEL A
LEMPKE, MICHAEL K
LERSTEN, CYNTHIA A
LEV, SEAN A
LEVITAN, WILLIAM M
LEWIS III, CHARLES B
LEWIS JR, WILLIAM A
LEWIS, ROGER A
LINGAN, ROBERT M
LISOWSKI, PAUL W
LIVENGOOD, JOANNA M
LOWE, OWEN W
LOYD, RICHARD NMN
LUCZAK, JOANN H
LUSHETSKY, JOHN M
LUTHA, RONALD J
LUTZE, NEILE MILLER
LYONS, PETER B
MACINTYRE, DOUGLAS M
MAINZER, ELLIOT E
MALOSH, GEORGE J
MARCINOWSKI III, FRANK NMN
MARLAY, ROBERT C
MARMOLEJOS, POLI A
MARTINEZ, ELOY DENNIS
MCARTHUR, BILLY R
MCCLOUD, FLOYD R
MCCLUER, MEGAN S
MCCONNELL, JAMES J
MCCORMICK, MATTHEW S
MCCRACKEN, STEPHEN H
MCGINNIS, EDWARD G
MCGUIRE, PATRICK W
MCKEE, BARBARA N
MCKENZIE, JOHN M
MCRAE, JAMES BENNETT
MEACHAM, A AVON
MEEKS, TIMOTHY J
MELLINGTON, STEPHEN A
MELLINGTON, SZAPANNE P
MILLER, CLARENCE L
MILLER, DEBORAH C
MILLER, WENDY L
MILLIKEN, JOANN NMN
MIOTLA, DENNIS M
MOE, DARRICK C
MONETTE, DEBORAH D
MONTANO, PEDRO A
MONTTOYA, ANTHONY H
MOODY III, DAVID C
MOORE, JOHN W O
MOORER, RICHARD F
MOREDOCK, J EUN
MORTENSON, VICTOR A
MUELLER, TROY J
MURPHIE, WILLIAM E
MUSTIN, TRACY P
NAPLES, ELMER M
NASSIF, ROBERT J
NEUHOFF, JON W
NEWMAN, LARRY NMN
NICOLL, ERIC G
NIEDZIELSKI-EICHNER, PHILL
NOLAN, ELIZABETH A
NORMAN, PAUL E
NOUSEN, DOUGLAS L
O'CONNOR, J RODERICK
O'CONNOR, STEPHEN C
O'CONNOR, THOMAS J
O'KONSKI, PETER J
OLENCZ, JOSEPH NMN
OLINGER, SHIRLEY J
OLIVER, STEPHEN R
OOSTERMAN, CARL H
OSHEIM, ELIZABETH L
OTT, MERRIE CHRISTINE
OWENDOFF, JAMES M
PALMISANO, ANNA C
PARNES, SANFORD J
PAVETTO, CARL S
PEASE, HARRISON G
PENRY, JUDITH M
PERSON JR, GEORGE L
PETERSON, BRADLEY A
PHAN, THOMAS H
PHOEBE, CHRISTINE A
PODONSKY, GLENN S
PORTER, STEVEN A
POWERS, KENNETH W
PROCARIO, MICHAEL P
PROVENCER, RICHARD B
PURUCKER, ROXANNE E
PYKE JR, THOMAS N
RAINES, ROBERT B
RAMSEY, CLAY HARRISON
RHODERICK, JAY E
RICHARDS, AUNDRA M
RICHARDSON, HERBERT NMN
RICHARDSON, SUSAN S
ROACH, RANDY A
RODGERS, DAVID E
RODGERS, STEPHEN J
ROEGE, WILLIAM H
ROGERS, MATTHEW C
ROHLFING, ERIC A
RUSSIAL, THOMAS J
RUSSO, FRANK B
SALMON, JEFFREY T
SAVAGE, CARTER D
SCHEINMAN, ADAM M
SCHOENBAUER, MARTIN J
SCHWIER, JEAN F
SCOTT, RANDAL S
SEDILLO, DAVID NMN
SEWARD, LACHLAN W
SHAFIK, CHRISTINE M
SHEELY, KENNETH B
SHEPPARD, CATHERINE M
SHERRY, THEODORE D
SHOOP, DOUG S
SHORT, STEPHANIE A
SILVERSTEIN, BRIAN L
SIMONSON, STEVEN C
SIMPSON, EDWARD R
SITZER, SCOTT B
SKUBEL, STEPHEN C
SMITH-KEVERN, REBECCA F
SMITH, KEVIN W
SMITH, THOMAS Z
SNIDER, ERIC S
SNIDER, LINDA J
SNYDER, ROGER E
SPEARS, TERREL J
SPERLING, GILBERT P
STAKER, THOMAS R
STALLMAN, ROBERT M
STARK, RICHARD M
STARNES, ALBERT J
STATON, CARL P
STENSETH, WILLIAM LYNN
STONE, BARBARA R
STRAYER, MICHAEL R
STREIT, LISA D
SURASH, JOHN E
SWEETNAM, GLEN E
SYKES, MERLE L
SYNAKOWSKI, EDMUND J
TALBOT JR, GERALD L
TAYLOR, HUGH N
TAYLOR, WILLIAM J
THOMPSON, MICHAEL A
THRESS JR, DONALD F

TOCZKO, JAMES E
 TOMER, BRADLEY J
 TORKOS, THOMAS M
 TOWNE, LAWRENCE H
 TRAUTMAN, STEPHEN J
 TUCKER, CRAIG A
 TURL, JAMES A
 TURNBULL, WILLIAM THOMAS
 TURNER, SHELLEY P
 TYNER, TERESA M
 UNDERWOOD, WILLIAM R
 UTECH, DAN G
 VALDEZ, WILLIAM J
 VAVOSO, THOMAS G
 VENUTO, KENNETH T
 WADDELL, JOSEPH F
 WAGNER, M PATRICE
 WAISLEY, SANDRA L
 WALL, EDWARD JAMES
 WARD, GARY K
 WARNICK, WALTER L
 WEATHERWAX, SHARLENE C
 WEEBER, DANIEL M
 WEEDALL, MICHAEL J
 WEIS, MICHAEL J
 WELLING, DAVID CRAIG
 WELLS, RITA L
 WESTON-DAWKES, ANDREW P
 WHITAKER JR, MARK B
 WHITNEY, JAMES M
 WHITTED, LINDA F
 WILBANKS, LINDA R
 WILBER, DEBORAH A
 WILCHER, LARRY D
 WILKEN, DANIEL H
 WILLIAMS, ALICE C
 WILLIAMS, MARK H
 WILLIAMS, RHYS M
 WILSON JR, THOMAS NMN
 WINCHELL JR, DONALD L
 WORLEY, MICHAEL N
 WORTHINGTON, JON C
 WORTHINGTON, PATRICIA R
 WRIGHT, STEPHEN J
 WU, CHUAN-FU NMN
 WYKA JR, THEODORE A
 YOSHIDA, PHYLLIS G
 YUAN-SOO HOO, CAMILLE C
 ZABRANSKY, DAVID K
 ZAMORSKI, MICHAEL J
 ZEH, CHARLES M

Issued in Washington, DC on November 16, 2009.

Sarah J. Bonilla,

Director, Office of Human Capital Management.

[FR Doc. E9-27908 Filed 11-19-09; 8:45 am]

BILLING CODE 6450-01-P

DEPARTMENT OF ENERGY

Senior Executive Service; Performance Review Board

AGENCY: U.S. Department of Energy.

ACTION: Designation of Performance Review Board Chair.

SUMMARY: This notice provides the Performance Review Board Chair designee for the Department of Energy.

DATES: This appointment is effective as of September 30, 2009.

Susan F. Beard.

Issued in Washington, DC, November 16, 2009.

Sarah J. Bonilla,

Director, Office of Human Capital Management.

[FR Doc. E9-27909 Filed 11-19-09; 8:45 am]

BILLING CODE 6450-01-P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

Combined Notice of Filings #1

November 10, 2009.

Take notice that the Commission received the following electric corporate filings:

Docket Numbers: EC10-15-000.

Applicants: West Georgia Generating Company, LLC; DeSoto County Generating Company, LLC; Broadway Gen Funding, LLC; Southern Power Company.

Description: West Georgia Generating Co, LLC *et al.* (Joint Applicants) submits the joint application for authorization under Section 203 of the Federal Power Act and Request for Expedited and Privileged Treatment.

Filed Date: 11/02/2009.

Accession Number: 20091105-0154.

Comment Date: 5 p.m. Eastern Time on Thursday, November 30, 2009.

Take notice that the Commission received the following electric rate filings:

Docket Numbers: ER01-3121-022; ER02-2085-017; ER02-417-021; ER02-418-021; ER03-1326-020; ER03-296-024; ER03-416-024; ER03-951-024; ER04-94-021; ER05-1146-021; ER05-1262-024; ER05-332-021; ER05-365-021; ER05-481-022; ER06-1093-020; ER06-200-020; ER07-1378-013; ER07-195-016; ER07-242-015; ER07-254-014; ER07-287-014; ER07-460-011; ER08-387-011; ER08-912-008; ER08-933-008; ER09-1284-003; ER09-1285-002; ER09-1723-002; ER09-279-004; ER09-281-003; ER09-282-004; ER09-30-005; ER09-31-005; ER09-32-006; ER09-382-004; ER08-934-009; ER07-240-015.

Applicants: Klamath Energy LLC; Northern Iowa Windpower LLC; Phoenix Wind Power LLC; Klamath Generation LLC; Colorado Green Holdings, LLC; Flying Cloud Power Partners, LLC; Klondike Wind Power

LLC; Moraine Wind LLC; Mountain View Power Partners III, LLC; Shiloh I Wind Project LLC; Flat Rock Windpower LLC; Klondike Wind Power II LLC; Elk River Windfarm LLC; Trimont Wind I LLC; Flat Rock Windpower II LLC; Big Horn Wind Project LLC; Providence Heights Wind, LLC; Locust Ridge Wind Farm, LLC; MinnDakota Wind LLC; Casselman Windpower, LLC; Klondike Wind Power III LLC; Dillon Wind LLC; Atlantic Renewables Projects II LLC; Iberdrola Renewables MBR Sellers; Lempster Wind, LLC; Rugby Wind, LLC; Stretator-Cayuga Ridge Wind Power, LLC; Dry Lake Wind Power, LLC; Buffalo Ridge I LLC; Pebble Springs Wind, LLC; Moraine Wind II LLC; Elm Creek Wind, LLC; Farmers City Wind, LLC; Barton Windpower LLC; Hay Canyon Wind LLC; Locust Ridge Wind Farm II, LLC; Twin Buttes Wind LLC.

Description: Quarterly Report Pursuant to 18 CFR 35.42(d) of Iberdrola Renewables MBR Sellers.

Filed Date: 10/30/2009.

Accession Number: 20091030-5183.

Comment Date: 5 p.m. Eastern Time on Friday, November 20, 2009.

Docket Numbers: ER05-764-005.

Applicants: MATL LLP, Montana Alberta Tie Ltd.

Description: Notice of Non-Material Change in Facts of Montana Alberta Tie Ltd. and MATL LLP.

Filed Date: 11/09/2009.

Accession Number: 20091109-5086.

Comment Date: 5 p.m. Eastern Time on Monday, November 30, 2009.

Docket Numbers: ER10-149-000.

Applicants: Elk City Wind, LLC.

Description: Request for authorization to sell energy and capacity at market-based rates, and waiver of the 60-day notice requirement re Elk City Wind, LLC.

Filed Date: 11/02/2009.

Accession Number: 20091104-0032.

Comment Date: 5 p.m. Eastern Time on Monday, November 23, 2009.

Docket Numbers: ER10-224-000.

Applicants: E. ON. U.S. LLC.

Description: E. ON. U.S. LLC submits a pro forma Emergency Energy Transaction Protocol Agreement.

Filed Date: 11/05/2009.

Accession Number: 20091106-0090.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Docket Numbers: ER10-227-000.

Applicants: Nevada Power Company.

Description: Nevada Power Company submits amendments to the Related Power Purchase Agreement between Sierra Pacific Power Company.

Filed Date: 11/05/2009.

Accession Number: 20091106-0093.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Docket Numbers: ER10–230–000.

Applicants: Kansas City Power & Light Company, KCP&L Greater Missouri Operations Company.

Description: Kansas City Power & Light Company *et al.* submits revised tariff sheets for the GMO open access transmission tariff, revised tariff sheets for schedule 1 *etc.*

Filed Date: 11/06/2009.

Accession Number: 20091110–0047.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Docket Numbers: ER10–234–000.

Applicants: Arizona Public Service Company.

Description: Arizona Public Service Company submits Amendment 1 and 2 Contract 87–BCA–10031, as Restated and Superseded *et al.* designated as APS' FERC Electric Rate 217 *etc.*

Filed Date: 11/10/2009.

Accession Number: 20091110–0046.

Comment Date: 5 p.m. Eastern Time on Tuesday, December 01, 2009.

Docket Numbers: ER10–236–000.

Applicants: Ohms Energy Company, LLC.

Description: Ohms Energy Company, LLC submits Petition for Acceptance of Initial Tariff, Waivers and Blanket Authority.

Filed Date: 11/05/2009.

Accession Number: 20091106–0089.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Take notice that the Commission received the following electric securities filings:

Docket Numbers: ES10–2–001.

Applicants: Trans-Allegheny Interstate Line Company.

Description: Trans-Allegheny Interstate Line Company, Additional Information to Section 204 Application (2009).

Filed Date: 11/09/2009.

Accession Number: 20091109–5150.

Comment Date: 5 p.m. Eastern Time on Thursday, November 19, 2009.

Take notice that the Commission received the following public utility holding company filings:

Docket Numbers: PH10–2–000.

Applicants: ITC Holdings Corp., *et al.*
Description: FERC–65B Joint Waiver Notification of ITC Holdings Corp.

Filed Date: 11/06/2009.

Accession Number: 20091106–5150.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Any person desiring to intervene or to protest in any of the above proceedings must file in accordance with Rules 211 and 214 of the Commission's Rules of

Practice and Procedure (18 CFR 385.211 and 385.214) on or before 5 p.m. Eastern time on the specified comment date. It is not necessary to separately intervene again in a subdocket related to a compliance filing if you have previously intervened in the same docket. Protests will be considered by the Commission in determining the appropriate action to be taken, but will not serve to make protestants parties to the proceeding. Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant. In reference to filings initiating a new proceeding, interventions or protests submitted on or before the comment deadline need not be served on persons other than the Applicant.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission, 888 First St., NE., Washington, DC 20426.

The filings in the above proceedings are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov or call (866) 208–3676 (toll free). For TTY, call (202) 502–8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9–27923 Filed 11–19–09; 8:45 am]

BILLING CODE 6717–01–P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

Combined Notice of Filings #1

November 13, 2009.

Take notice that the Commission received the following electric corporate filings:

Docket Numbers: EC10–17–000.

Applicants: Astoria Energy LLC, SUEZ Energy Astoria LLC, CDPQ Investments (U.S.) Inc., WE Energy Investment LLC, USPF Holdings, LLC, Robert O. Gurman, JEMB Family LP, EIF Astoria III, LLC, AE Bowery Bay LLC.

Description: Astoria Energy LLC *et al.* submits application for authorization of indirect disposition of jurisdictional facilities and acquisition of securities, and request for expedited consideration.

Filed Date: 11/04/2009.

Accession Number: 20091106–0038.

Comment Date: 5 p.m. Eastern Time on Wednesday, November 25, 2009.

Docket Numbers: EC10–18–000.

Applicants: First Solar, Inc., FSE Blythe 1, LLC, NRG Energy, Inc.

Description: NRG Energy, Inc. *et al.* submits Joint Application for Authorization under Section 203 of the Federal Power Act, requesting expedited treatment and request for confidential treatment *etc.*

Filed Date: 11/04/2009.

Accession Number: 20091106–0091.

Comment Date: 5 p.m. Eastern Time on Wednesday, November 25, 2009.

Docket Numbers: EC10–19–000.

Applicants: Illinois Power Company.

Description: Illinois Power Company *et al.* submits request for approvals pursuant to section 203 of the Federal Power Act and expedited consideration and shortened notice period.

Filed Date: 11/10/2009.

Accession Number: 20091112–0104.

Comment Date: 5 p.m. Eastern Time on Tuesday, December 01, 2009.

Docket Numbers: EC10–20–000.

Applicants: Locust Ridge Wind Farm, LLC, Fortis Energy Marketing & Trading GP.

Description: Application for Authorization under Section 203 of the Federal Power Act of Fortis Energy Marketing & Trading GP and Locust Ridge Wind Farm, LLC.

Filed Date: 11/10/2009.

Accession Number: 20091110–5119.

Comment Date: 5 p.m. Eastern Time on Tuesday, December 01, 2009.

Take notice that the Commission received the following electric rate filings:

Docket Numbers: ER08–374–004;

EL08–38–003.

Applicants: Atlantic Path 15, LLC.
Description: Atlantic Path 15, LLC's Refund Report.

Filed Date: 11/12/2009.

Accession Number: 20091112-5164.

Comment Date: 5 p.m. Eastern Time on Thursday, December 03, 2009.

Docket Numbers: ER09-153-001; ER09-154-001; ER09-155-001; ER09-156-001.

Applicants: Southern Company Services, Inc.

Description: Refund Report of Southern Company Services, Inc., in Compliance with Settlement.

Filed Date: 11/12/2009.

Accession Number: 20091112-5236.

Comment Date: 5 p.m. Eastern Time on Thursday, December 03, 2009.

Docket Numbers: ER10-191-001; EC06-4-003.

Applicants: LG&E Energy Corporation, E.ON U.S. LLC.

Description: E.ON U.S. LLC seeks approval to change Applicants method of (i) complying with Order 888, 889 and 890.

Filed Date: 11/05/2009.

Accession Number: 20091106-0086.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Docket Numbers: ER10-210-000.

Applicants: VIRIDIAN ENERGY PA LLC.

Description: Viridian Energy PA LLC submits petition for acceptance of initial tariff, waivers and blanket authority.

Filed Date: 11/09/2009.

Accession Number: 20091110-0036.

Comment Date: 5 p.m. Eastern Time on Monday, November 30, 2009.

Docket Numbers: ER10-225-000.

Applicants: Major Energy Electric Services.

Description: Major Energy Electric Services, LLC submits Petition for Acceptance of Initial Rate Schedule, Waivers and Blanket Authority.

Filed Date: 11/10/2009.

Accession Number: 20091113-0135.

Comment Date: 5 p.m. Eastern Time on Tuesday, December 01, 2009.

Docket Numbers: ER10-226-000.

Applicants: Clean Currents, LLC.

Description: Clean Currents, LLC submits petition for acceptance of initial rate schedule, waivers and blanket authority.

Filed Date: 11/09/2009.

Accession Number: 20091110-0037.

Comment Date: 5 p.m. Eastern Time on Monday, November 30, 2009.

Docket Numbers: ER10-228-000.

Applicants: Star Point Wind Project LLC.

Description: Star Point Wind Project LLC submits FERC Electric Tariff, Original Volume No 1.

Filed Date: 11/06/2009.

Accession Number: 20091110-0038.

Comment Date: 5 p.m. Eastern Time on Friday, November 27, 2009.

Take notice that the Commission received the following PURPA 210(m)(3) filings:

Docket Numbers: QM09-7-001.

Applicants: Old Dominion Electric Cooperative, Inc.

Description: Old Dominion Electric Cooperative Supplemental Information in support of its Application to Terminate Purchase Obligation.

Filed Date: 11/13/2009.

Accession Number: 20091113-5042.

Comment Date: 5 p.m. Eastern Time on Friday, December 11, 2009.

Take notice that the Commission received the following electric reliability filings:

Docket Numbers: RR10-2-000.

Applicants: North American Electric Reliability Corp.

Description: Petition of the North American Electric Reliability Corporation for Approval of Amendment to its Delegation Agreements with Regional Entities to Extend Initial Term of Agreements to May 2011.

Filed Date: 11/12/2009.

Accession Number: 20091112-5187.

Comment Date: 5 p.m. Eastern Time on Thursday, December 03, 2009.

Docket Numbers: RR10-3-000.

Applicants: North American Electric Reliability Corp.

Description: Petition of the North American Electric Reliability Corporation for Approval of Amendments to its Rules of Procedure to Reflect Elimination of the Reliability Readiness Evaluation and Improvement Program.

Filed Date: 11/12/2009.

Accession Number: 20091112-5188.

Comment Date: 5 p.m. Eastern Time on Thursday, December 03, 2009.

Any person desiring to intervene or to protest in any of the above proceedings must file in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and 385.214) on or before 5 p.m. Eastern time on the specified comment date. It is not necessary to separately intervene again in a subdocket related to a compliance filing if you have previously intervened in the same docket. Protests will be considered by the Commission in determining the appropriate action to be taken, but will not serve to make protestants parties to the proceeding. Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant. In reference

to filings initiating a new proceeding, interventions or protests submitted on or before the comment deadline need not be served on persons other than the Applicant.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission, 888 First St., NE., Washington, DC 20426.

The filings in the above proceedings are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov or call (866) 208-3676 (toll free). For TTY, call (202) 502-8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27916 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Project No. 2306-143-VT]

Great Bay Hydro Corporation; Notice of Availability of Environmental Assessment

November 10, 2009.

In accordance with the National Environmental Policy Act of 1969 and the Federal Energy Regulatory Commission's (Commission) regulations, 18 CFR Part 380 (Order No. 486, 52 FR 47879), the Office of Energy Projects has reviewed the application filed by Great Bay Hydro Corporation on February 19, 2009, requesting Commission approval to amend the license for the Clyde River Hydroelectric Project. The project is

located on the Clyde River in the City of Newport, in Orleans County, Vermont.

The Commission has prepared an environmental assessment (EA) that evaluates the environmental impacts of the licensee's proposal. The EA finds that approval of the application would not constitute a major federal action significantly affecting the quality of the human environment.

The EA is attached to a Commission order titled "Order Amending License", issued November 9, 2009, and is available for review and reproduction at the Commission's Public Reference Room, located at 888 First Street, NE., Room 2-A, Washington, DC 20426. The EA may also be viewed on the Commission's Web site at <http://www.ferc.gov> using the "eLibrary" link. Enter the docket number (P-2306) excluding the last three digits in the docket number field to access the document. For assistance, contact FERC Online Support at FERCOnlineSupport@ferc.gov or toll-free at 1-866-208-3676, or for TTY, (202) 502-8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27922 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Docket No. ER10-226-000]

Clean Currents, L.L.C.; Supplemental Notice That Initial Market-Based Rate Filing Includes Request for Blanket Section 204 Authorization

November 13, 2009.

This is a supplemental notice in the above-referenced proceeding of Clean Currents, L.L.C.'s application for market-based rate authority, with an accompanying rate tariff, noting that such application includes a request for blanket authorization, under 18 CFR Part 34, of future issuances of securities and assumptions of liability.

Any person desiring to intervene or to protest should file with the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426, in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and § 385.214). Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant.

Notice is hereby given that the deadline for filing protests with regard

to the applicant's request for blanket authorization, under 18 CFR Part 34, of future issuances of securities and assumptions of liability, is December 3, 2009.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426.

The filings in the above-referenced proceeding are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov or call (866) 208-3676 (toll free). For TTY, call (202) 502-8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27918 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Docket No. ER10-210-000]

Viridian Energy P; LLC Supplemental Notice That Initial Market-Based Rate Filing Includes Request for Blanket Section 204 Authorization

November 13, 2009.

This is a supplemental notice in the above-referenced proceeding of Viridian Energy PA LLC's application for market-based rate authority, with an accompanying rate tariff, noting that such application includes a request for blanket authorization, under 18 CFR Part 34, of future issuances of securities and assumptions of liability.

Any person desiring to intervene or to protest should file with the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426, in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and 385.214). Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant.

Notice is hereby given that the deadline for filing protests with regard to the applicant's request for blanket authorization, under 18 CFR Part 34, of future issuances of securities and assumptions of liability, is December 3, 2009.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426.

The filings in the above-referenced proceeding are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov or call (866) 208-3676 (toll free). For TTY, call (202) 502-8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27920 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY**Federal Energy Regulatory Commission**

[Docket No. ER10-228-000]

Star Point Wind Project LLC; Supplemental Notice That Initial Market-Based Rate Filing Includes Request for Blanket Section 204 Authorization

November 13, 2009.

This is a supplemental notice in the above-referenced proceeding of Star Point Wind Project LLC's application for market-based rate authority, with an accompanying rate tariff, noting that such application includes a request for blanket authorization, under 18 CFR part 34, of future issuances of securities and assumptions of liability.

Any person desiring to intervene or to protest should file with the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426, in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and 385.214). Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant.

Notice is hereby given that the deadline for filing protests with regard to the applicant's request for blanket authorization, under 18 CFR part 34, of future issuances of securities and assumptions of liability, is December 3, 2009.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426.

The filings in the above-referenced proceeding are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed

docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov, or call (866) 208-3676 (toll free). For TTY, call (202) 502-8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27919 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY**Federal Energy Regulatory Commission**

[Docket No. ER10-236-000]

Ohms Energy Company, LLC; Supplemental Notice That Initial Market-Based Rate Filing Includes Request for Blanket Section 204 Authorization

November 13, 2009.

This is a supplemental notice in the above-referenced proceeding of Ohms Energy Company, LLC's application for market-based rate authority, with an accompanying rate tariff, noting that such application includes a request for blanket authorization, under 18 CFR part 34, of future issuances of securities and assumptions of liability.

Any person desiring to intervene or to protest should file with the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426, in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and 385.214). Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant.

Notice is hereby given that the deadline for filing protests with regard to the applicant's request for blanket authorization, under 18 CFR part 34, of future issuances of securities and assumptions of liability, is December 3, 2009.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission,

888 First Street, NE., Washington, DC 20426.

The filings in the above-referenced proceeding are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov or call (866) 208-3676 (toll free). For TTY, call (202) 502-8659.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27917 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY**Federal Energy Regulatory Commission**

[Docket No. ER08-83-003]

Gilberton Power Company; Supplemental Notice That Initial Market-Based Rate Filing Includes Request for Blanket Section 204 Authorization

November 13, 2009.

This is a supplemental notice in the above-referenced proceeding of Gilberton Power Company's application for market-based rate authority, with an accompanying rate tariff, noting that such application includes a request for blanket authorization, under 18 CFR part 34, of future issuances of securities and assumptions of liability.

Any person desiring to intervene or to protest should file with the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426, in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and 385.214). Anyone filing a motion to intervene or protest must serve a copy of that document on the Applicant.

Notice is hereby given that the deadline for filing protests with regard to the applicant's request for blanket authorization, under 18 CFR part 34, of future issuances of securities and assumptions of liability, is December 3, 2009.

The Commission encourages electronic submission of protests and interventions in lieu of paper, using the FERC Online links at <http://www.ferc.gov>. To facilitate electronic

service, persons with Internet access who will eFile a document and/or be listed as a contact for an intervenor must create and validate an eRegistration account using the eRegistration link. Select the eFiling link to log on and submit the intervention or protests.

Persons unable to file electronically should submit an original and 14 copies of the intervention or protest to the Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426.

The filings in the above-referenced proceeding are accessible in the Commission's eLibrary system by clicking on the appropriate link in the above list. They are also available for review in the Commission's Public Reference Room in Washington, DC. There is an eSubscription link on the Web site that enables subscribers to receive e-mail notification when a document is added to a subscribed docket(s). For assistance with any FERC Online service, please e-mail FERCOnlineSupport@ferc.gov, or call (866) 208-3676 (toll free). For TTY, call (202) 502-8659.

Nathaniel J. Davis, Sr.,
Deputy Secretary.

[FR Doc. E9-27915 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY

Western Area Power Administration

Boulder Canyon Project—Post-2017 Application of the Energy Planning and Management Program Power Marketing Initiative

AGENCY: Western Area Power Administration, DOE.

ACTION: Notice of proposal.

SUMMARY: The Western Area Power Administration (Western), Desert Southwest Region, a Federal power marketing agency of the Department of Energy (DOE) announces its Post-2017 remarketing effort for the Boulder Canyon Project (BCP). Current BCP long-term contracts will expire on September 30, 2017. In 1995, Western adopted the Power Marketing Initiative (PMI) in Subpart C of the Energy Planning and Management Program (Program) (10 CFR part 905). The Record of Decision for the Program states that application of the PMI will be done on a project-specific basis. If, by means of a public process, Western applies the PMI to the BCP, the current long-term contractors of the project would receive an extension of a major portion of the

resources available to them at the time their contracts expire. Western now proposes to apply the PMI to the long-term power contracts of the BCP.

DATES: Western will hold three public information forums regarding the BCP remarketing on the following dates: December 1, 2009, 1 p.m., PST, Las Vegas, Nevada; December 2, 2009, 1 p.m., PST, Ontario, California; December 3, 2009, 1 p.m., MST, Phoenix, Arizona.

Western will also hold three public comment forums. The dates for these forums are: January 19, 2010, 1 p.m., PST, Las Vegas, Nevada; January 20, 2010, 1 p.m., MST, Phoenix, Arizona; January 21, 2010, 1 p.m., PST, Ontario, California. Western will accept written comments on or before January 29, 2010. Western reserves the right to not consider any comments received after the prescribed date and time.

ADDRESSES: Comments may be submitted to: Mr. Darrick Moe, Western Area Power Administration, Desert Southwest Regional Manager, P.O. Box 6457, Phoenix, AZ 85005-6457. Comments may also be faxed to (602) 605-2490 or e-mailed to Post2017BCP@wapa.gov.

The public information and public comment forum locations are: Las Vegas Tropicana, 3801 Las Vegas Boulevard, South Las Vegas, Nevada; Western Area Power Administration, Desert Southwest Regional Office, 615 S. 43rd Ave, Phoenix, Arizona; DoubleTree Ontario Airport, 222 N. Vineyard, Ontario, California.

As access to Western facilities is controlled, any U.S. citizen wishing to attend any meeting held at Western must present an official form of picture identification, such as a U.S. driver's license, U.S. passport, U.S. Government ID, or U.S. Military ID, at the time of the meeting. Foreign nationals shall contact Western at least 45 days in advance of the meeting to obtain the necessary admittance to Western.

FOR FURTHER INFORMATION CONTACT: Mr. Mike Simonton, Remarketing Specialist, Desert Southwest Region, Western Area Power Administration, P.O. Box 6457, Phoenix, AZ 85005, telephone (602) 605-2675, e-mail Post2017BCP@wapa.gov. Program information and the Conformed General Consolidated Power Marketing Criteria or Regulations for Boulder City Area Projects (Conformed Criteria) published in the **Federal Register** (49 FR 50582) on December 28, 1984, are available at <http://www.wapa.gov/dsw/pwrnkt>.

SUPPLEMENTARY INFORMATION: In 1987, Western marketed the power resources of the BCP and entered into 30-year

term contracts with the current BCP contractors in accordance with the Hoover Power Plant Act of 1984, and the Conformed Criteria. These events resulted in the allocation of 1,951 megawatts (MW) of contingent capacity with an associated 4,527,001 megawatt-hours (MWh) of annual firm energy. Pursuant to 43 U.S.C. 619a(4), these long-term contracts will expire on September 30, 2017. Western must determine if the PMI, as outlined in the Program, will be applied to the BCP for commitments beyond that date, the quantity of resources to be extended to existing contractors, the size of the proposed resource pool, excess energy provisions, and the term of the contract.

Western first proposed the Program on April 19, 1991 (56 FR 16093). The goals of the Program were to encourage efficient energy use by Western's power customers by requiring Integrated Resource Planning and to extend Western's firm power resource commitments. In the final rule of the Program, Western stated that application of the PMI, including the amount of resources extended, would initially apply only to the Pick-Sloan Missouri Basin Program-Eastern Division and the Loveland Area Projects. Applicability to other projects would be determined through future, project-specific public processes. Specific to the BCP, Western stated that it would evaluate application of the PMI to the BCP no more than 10 years before existing contracts expire. 60 FR 54151, 54157 (Oct. 20, 1995).

Consistent with the application of the PMI to other recent Western marketing efforts, Western proposes to apply the PMI (10 CFR parts 905.30 through 905.37) to the BCP. In consultation with the Bureau of Reclamation and referencing the most recent hydrologic studies, Western proposes to market 2,044 MW of contingent capacity with an associated 4,116,000 MWh of annual firm energy from the BCP. Western proposes to extend 100 percent of the existing contractors' contingent capacity allocations, totaling 1,951 MW, and 95 percent of the proposed marketable firm energy, totaling 3,910,200 MWh annually to the existing contractors based proportionally upon their existing allocations of marketed annual firm energy. This proposal would result in the creation of a single, one-time resource pool consisting of 93 MW of contingent capacity with an associated 205,800 MWh of annual firm energy. Western proposes that all contract terms be a length of 30 years commencing on October 1, 2017.

The marketing area is generally defined as consisting of southern

California, southern Nevada, most of Arizona, and a small part of New Mexico; and is more specifically defined in the Conformed Criteria. New customers meeting the requirements established in the BCP marketing criteria and qualifying Native American Tribes within the BCP marketing area will be eligible to request an allocation of capacity and energy from the BCP resource pool. Native American Tribes need not have utility status to qualify for an allocation. As provided in the current BCP Implementation Agreement, new contractors, or contractors who receive an increased allocation will be required to reimburse existing BCP contractors for replacement capital advances to the extent existing contractors' allocations are reduced as a result of creating the resource pool.

Western is seeking comments regarding the applicability of the PMI to the BCP, the quantity of resources to be extended to existing contractors, the size of the proposed resource pool, excess energy provisions, and the term of contracts.

Under their respective State laws the Colorado River Commission (CRC) and the Arizona Power Authority (APA) have been the designated agents for acquiring and remarketing BCP power within Nevada and Arizona respectively. Western seeks comments regarding CRC and APA's role in Western's allocation process.

Following the public comment period, Western will analyze the comments received and publish its policy regarding the application of PMI to BCP in the **Federal Register**.

Regulatory Procedure Requirements

Determination Under Executive Order 12866

Western has an exemption from centralized regulatory review under Executive Order 12866; accordingly, no clearance of this notice by the Office of Management and Budget is required.

Environmental Compliance

In accordance with the DOE National Environmental Policy Act Implementing Procedures (10 CFR 1021), Western has determined that this action fits within class of action B4.1 Contracts/marketing plans/policies for excess electric power, in Appendix B to Subpart D to part 1021—Categorical Exclusions Applicable to Specific Agency Actions.

Dated: November 10, 2009.

Timothy J. Meeks,

Administrator.

[FR Doc. E9-27910 Filed 11-19-09; 8:45 am]

BILLING CODE 6450-01-P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Project No. 13571-000]

Goshen Powerhouse, LLC: Notice of Preliminary Permit Application Accepted for Filing and Soliciting Comments, Motions To Intervene, and Competing Applications

November 10, 2009.

On August 24, 2009, Goshen Powerhouse, LLC filed an application for a preliminary permit, pursuant to section 4(f) of the Federal Power Act, proposing to study the feasibility of the Goshen Dam Hydroelectric Project No. 13571, to be located at the existing Goshen Dam, on the Elkhart River, in Elkhart County, Indiana. The sole purpose of a preliminary permit, if issued, is to grant the permit holder priority to file a license application during the permit term. A preliminary permit does not authorize the permit holder to perform any land disturbing activities or otherwise enter upon lands or waters owned by others without the owners' express permission.

The existing dam is owned and operated by Elkhart County, Indiana and the existing powerhouse and headrace canal is owned and operated by the City of Goshen, Indiana. The proposed project would consist of: (1) The existing 12.5-foot-high concrete dam equipped with a 200-foot-long ogee spillway; (2) an existing 765-acre impoundment with a normal water surface elevation of 790.9 feet mean sea level; (3) an existing 25-foot-long by 49-foot-wide powerhouse to contain two new hydrokinetic turbine-generator systems for a total installed capacity of 80 kilowatts; (4) a new 60-foot-long, 12.5-kilovolt transmission line; and (5) appurtenant facilities. The proposed project would operate in a run-of-river mode and generate an estimated average annual generation of 578,160 kilowatt-hours.

Applicant Contact: Edward L. Kurth, Goshen Powerhouse, LLC, San Antonio Zanesville, Texas 78216, (210) 496-5902.

FERC Contact: Michael Watts, (202) 502-6123.

Deadline for filing comments, motions to intervene, competing applications (without notices of intent), or notices of intent to file competing applications: 60 days from the issuance of this notice.

Comments, motions to intervene, notices of intent, and competing applications may be filed electronically via the Internet. See 18 CFR 385.2001(a)(1)(iii) and the instructions

on the Commission's Web site under the "eFiling" link. If unable to be filed electronically, documents may be paper-filed. To paper-file, an original and eight copies should be mailed to: Kimberly D. Bose, Secretary, Federal Energy Regulatory Commission, 888 First Street, NE., Washington, DC 20426. For more information on how to submit these types of filings please go to the Commission's Web site located at <http://www.ferc.gov/filing-comments.asp>. More information about this project can be viewed or printed on the "eLibrary" link of Commission's Web site at <http://www.ferc.gov/docs-filing/elibrary.asp>. Enter the docket number (P-13571) in the docket number field to access the document. For assistance, call toll-free 1-866-208-3372.

Nathaniel J. Davis, Sr.,

Deputy Secretary.

[FR Doc. E9-27921 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Docket No. RM98-1-000]

Records Governing Off-the-Record Communications: Public Notice

November 10, 2009.

This constitutes notice, in accordance with 18 CFR 385.2201(b), of the receipt of prohibited and exempt off-the-record communications.

Order No. 607 (64 FR 51222, September 22, 1999) requires Commission decisional employees, who make or receive a prohibited or exempt off-the-record communication relevant to the merits of a contested proceeding, to deliver to the Secretary of the Commission, a copy of the communication, if written, or a summary of the substance of any oral communication.

Prohibited communications are included in a public, non-decisional file associated with, but not a part of, the decisional record of the proceeding. Unless the Commission determines that the prohibited communication and any responses thereto should become a part of the decisional record, the prohibited off-the-record communication will not be considered by the Commission in reaching its decision. Parties to a proceeding may seek the opportunity to respond to any facts or contentions made in a prohibited off-the-record communication, and may request that the Commission place the prohibited

communication and responses thereto in the decisional record. The Commission will grant such a request only when it determines that fairness so requires. Any person identified below as having made a prohibited off-the-record communication shall serve the document on all parties listed on the official service list for the applicable proceeding in accordance with Rule 2010, 18 CFR 385.2010.

Exempt off-the-record communications are included in the

decisional record of the proceeding, unless the communication was with a cooperating agency as described by 40 CFR 1501.6, made under 18 CFR 385.2201(e)(1)(v).

The following is a list of off-the-record communications recently received by the Secretary of the Commission. The communications listed are grouped by docket numbers in ascending order. These filings are available for review at the Commission in the Public Reference Room or may be

viewed on the Commission's Web site at <http://www.ferc.gov> using the eLibrary link. Enter the docket number, excluding the last three digits, in the docket number field to access the document. For assistance, please contact FERC, Online Support at FERCOnlineSupport@ferc.gov or toll free at (866) 208-3676, or for TTY, contact (202) 502-8659.

EXEMPT

Docket No.	File date	Presenter or requester
1. CP04-36-005	11-2-09	David Swearingen ¹ .
2. CP09-444-000	11-2-09	David Hanobic ² .

Nathaniel J. Davis, Sr.,
Deputy Secretary.

[FR Doc. E9-27924 Filed 11-19-09; 8:45 am]

BILLING CODE 6717-01-P

ENVIRONMENTAL PROTECTION AGENCY

[EPA-HQ-OPPT-2008-0717; FRL-8983-7; EPA ICR No. 2328.01; OMB Control No. 2070-TBD]

Agency Information Collection Activities; Submission to OMB for Review and Approval; Comment Request; Pressed Wood Manufacturing Industry Survey

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice.

SUMMARY: In compliance with the Paperwork Reduction Act (44 U.S.C. 3501 *et seq.*), this document announces that an Information Collection Request (ICR) has been forwarded to the Office of Management and Budget (OMB) for review and approval: Pressed Wood Manufacturing Industry Survey; ICR No. 2328.01, OMB No. 2070-TBD. The ICR, which is abstracted below, describes the nature of the information collection activity and its expected burden and costs.

DATES: Additional comments may be submitted on or before December 21, 2009.

ADDRESSES: Submit your comments, referencing docket ID Number EPA-HQ-OPPT-2008-0717 to (1) EPA online using <http://www.regulations.gov> (our preferred method), by e-mail to oppt.ncic@epa.gov or by mail to: Document Control Office (DCO), Office of Pollution Prevention and Toxics (OPPT), Environmental Protection Agency, Mail Code: 7407T, 1200 Pennsylvania Ave., NW., Washington, DC 20460, and (2) OMB at: Office of Information and Regulatory Affairs, Office of Management and Budget (OMB), Attention: Desk Officer for EPA, 725 17th Street, NW., Washington, DC 20503.

FOR FURTHER INFORMATION CONTACT: Barbara Cunningham, Director, Environmental Assistance Division, Office of Pollution Prevention and Toxics, Environmental Protection Agency, Mailcode: 7408-M, 1200 Pennsylvania Ave., NW., Washington, DC 20460; telephone number: 202-554-1404; e-mail address: TSCA-Hotline@epa.gov.

SUPPLEMENTARY INFORMATION: EPA has submitted the following ICR to OMB for review and approval according to the procedures prescribed in 5 CFR 1320.12. On December 24, 2008 (73 FR 79083), EPA sought comments on this ICR pursuant to 5 CFR 1320.8(d). EPA received four comments during the comment period, which are addressed in the ICR. Any comments related to this ICR should be submitted to EPA and OMB within 30 days of this notice.

EPA has established a public docket for this ICR under Docket ID No. EPA-HQ-OPPT-2008-0717, which is available for online viewing at <http://www.regulations.gov>, or in person inspection at the OPPT Docket in the EPA Docket Center (EPA/DC), EPA

West, Room 3334, 1301 Constitution Ave., NW., Washington, DC. The EPA/DC Public Reading Room is open from 8 a.m. to 4:30 p.m., Monday through Friday, excluding legal holidays. The telephone number for the Reading Room is 202-566-1744, and the telephone number for the Pollution Prevention and Toxics Docket is 202-566-0280.

Use EPA's electronic docket and comment system at <http://www.regulations.gov> to submit or view public comments, access the index listing of the contents of the public docket, and to access those documents in the public docket that are available electronically. Once in the system, select "docket search," then key in the docket ID number identified above. Please note that EPA's policy is that public comments, whether submitted electronically or in paper, will be made available for public viewing in <http://www.regulations.gov> as EPA receives them and without change, unless the comment contains copyrighted material, confidential business information (CBI), or other information whose public disclosure is restricted by statute. When EPA identifies a comment containing copyrighted material, EPA will provide a reference to that material in the version of the comment that is placed in <http://www.regulations.gov>. The entire printed comment, including the copyrighted material, will be available in the public docket. Although identified as an item in the official docket, information claimed as CBI, or whose disclosure is otherwise restricted by statute, is not included in the official public docket, and will not be available for public viewing in <http://www.regulations.gov>. For further information about the electronic docket, go to <http://www.regulations.gov>.

¹ Memo to file from staff attaching correspondence—seven responsive letters from Captain Raymond J. Perry, U.S. Coast Guard, with respect to Weaver's Cove Bay Berth Amendment.

² Telephone communication record.

Title: Pressed Wood Manufacturing Industry Survey.

ICR Numbers: EPA ICR No. 2328.01, OMB Control No. 2070-TBD.

ICR Status: This ICR is for a new information collection activity. An Agency may not conduct or sponsor, and a person is not required to respond to, a collection of information, unless it displays a currently valid OMB control number. The OMB control numbers for EPA's regulations in title 40 of the CFR, after appearing in the **Federal Register** when approved, are listed in 40 CFR part 9, are displayed either by publication in the **Federal Register** or by other appropriate means, such as on the related collection instrument or form, if applicable. The display of OMB control numbers in certain EPA regulations is consolidated in 40 CFR part 9.

Abstract: EPA has initiated a proceeding to investigate whether and what type of regulatory or other action might be appropriate to control the levels of formaldehyde emitted from pressed wood products, as described in EPA's Advanced Notice of Proposed Rulemaking (ANPR) for Formaldehyde Emissions from Pressed Wood Products, which published in the **Federal Register** on December 3, 2008 (73 FR 73620). As part of this investigation, EPA seeks to survey U.S. pressed wood manufacturers to collect information on the categories and volume of pressed wood manufactured; the types of resins used in the manufacturing process; the formaldehyde emissions levels from the pressed wood; recent or planned changes to reduce formaldehyde emissions and the resulting costs; and any issues that may affect the ability to reduce formaldehyde emissions. The survey will be sent to all U.S. pressed wood manufacturers identified by EPA (*i.e.*, it will be a census). This survey asks for readily obtainable information, so the plant does not have to generate new information (for example, by testing product emissions) to complete the survey.

When EPA sought comments on the ICR on December 24, 2008 (73 FR 79083), the Agency intended to use the same questionnaire for all pressed wood manufacturers. EPA has now modified the survey so that there are two different questionnaires. One questionnaire is for manufacturers of hardwood plywood, particleboard, and medium density fiberboard (which are the products subject to the CARB rule). The second questionnaire is for manufacturers of hardboard or structural composites (glued laminated timber, I-joists, oriented strandboard, softwood plywood, and structural composite

lumber) that are not subject to the CARB rule. These manufacturers are unlikely to have as much readily available information to use responding to the survey, this questionnaire asks for a more limited set of information.

EPA will request that all U.S. pressed wood manufacturers voluntarily complete the survey. If EPA does not achieve a sufficient survey response rate to accurately characterize the industry, EPA will consider whether to exercise the authority available to it under TSCA section 11(c), 15 U.S.C. 2610(c). TSCA section 11(c) provides EPA with the authority to issue subpoenas requiring the production of reports, papers, documents, answers to questions, and other information that the Administrator deems necessary. EPA could potentially use its TSCA section 11(c) authority to issue subpoenas requiring recipients (*i.e.*, non-respondents) to complete and return the survey.

Respondents may elect to claim certain submitted information as confidential business information (CBI) if there is a legitimate need to do so as described in EPA's regulations at 40 CFR part 2. These claims will be handled according to EPA procedures described in the regulation at 40 CFR part 2. EPA will disclose information that is covered by a claim of confidentiality only to the extent permitted by, and in accordance with, the procedures in TSCA section 14, 15 U.S.C. 2613, and the regulation at 40 CFR part 2.

The information collected through the survey will allow EPA to predict a future baseline for the types of resins that will be used in pressed wood, and the levels of formaldehyde that will be emitted from them. EPA will also use this information to assess the incremental benefits and costs of potential actions at the national level on formaldehyde emissions from pressed wood products. This information is necessary to inform Agency decisionmaking about the need for and scope of regulatory or other actions to control the levels of formaldehyde emitted from pressed wood products. If this survey is not conducted, EPA will not have this data to establish the baseline for its actions nor insight into the adjustments that plants in the industry are planning to make. Therefore, this survey is necessary for the proper performance of Agency functions.

Burden Statement: The annual public reporting and recordkeeping burden for this collection of information is estimated to range between 14 and 20 hours per response, depending upon the nature of the respondent. Burden means

the total time, effort or financial resources expended by persons to generate, maintain, retain or disclose or provide information to or for a Federal agency. This includes the time needed to review instructions; develop, acquire, install and utilize technology and systems for the purposes of collecting, validating and verifying information, processing and maintaining information, and disclosing and providing information; adjust the existing ways to comply with any previously applicable instructions and requirements; train personnel to be able to respond to a collection of information; search data sources; complete and review the collection of information; and transmit or otherwise disclose the information.

Respondents/Affected Entities: Entities potentially affected by this action are facilities engaged in the manufacturing of pressed wood products, including glued laminated timber, hardboard, medium density fiberboard, oriented strandboard, particleboard, hardwood and softwood plywood, prefabricated I-joists, and structural composite lumber (which includes laminated veneer lumber, laminated strand lumber, parallel strand lumber, and oriented strand lumber).

Frequency of Collection: Once.

Estimated average number of responses for each respondent: 1.

Estimated No. of Respondents: 343.

Estimated Total Annual Burden on Respondents: 5,804 hours.

Estimated Total Annual Costs: \$324,220.

Changes in Burden Estimates: This is a new ICR. This estimated burden for this new ICR is estimated to be 5,804 hours and is a program change.

Dated: November 16, 2009.

John Moses,

Acting Director, Collection Strategies Division.

[FR Doc. E9-27941 Filed 11-19-09; 8:45 am]

BILLING CODE 6560-50-P

ENVIRONMENTAL PROTECTION AGENCY

[ER-FRL-8599-5]

Environmental Impact Statements and Regulations; Availability of EPA Comments

Availability of EPA comments prepared pursuant to the Environmental Review Process (ERP), under Section 309 of the Clean Air Act and Section 102(2)(c) of the National Environmental Policy Act as amended. Requests for copies of EPA comments can be directed

to the Office of Federal Activities at 202-564-7146 or <http://www.epa.gov/compliance/nepa/>.

An explanation of the ratings assigned to draft environmental impact statements (EISs) was published in FR dated July 17, 2009 (74 FR 34754).

Draft EISs

EIS No. 20090312, ERP No. D-COE-F39043-OH, Cleveland Harbor Dredged Material Management Plan, Operations and Maintenance, Cuyahoga County, OH.

Summary: EPA expressed environmental concerns about air quality impacts, and recommended implementation of mitigation measures. Rating EC2.

EIS No. 20090315, ERP No. D-FTA-K40273-CA, Crenshaw Transit Corridor Project, Proposes to Improve Transit Services, Funding, Los Angeles County Metropolitan Transportation Authority (LACMTA), Los Angeles County, CA.

Summary: EPA expressed environmental concerns about the air quality analysis, and offered additional suggestions for water quality impact analysis and mitigation. Rating EC2.

EIS No. 20090382, ERP No. D-IBR-K39123-CA, Central Valley Project Water Supply Contracts Under Public Law 101-514 (Section 206), Proposed Water Service Contracts with the El Dorado County Water Agency, El Dorado County, CA.

Summary: EPA does not object to the proposed action. Rating LO.

Final EISs

EIS No. 20090340, ERP No. F-AFS-J65499-UT, Pockets Resource Management Project, Additional Information on Analysis and Disclosure on the Effect of the PA and Alternatives on Three Unroaded and Undeveloped Areas Identified on a 2005 Draft Map, Proposes to Salvage Dead and Dying Spruce/Fir, Regenerate Aspen, and Manage Travel, Escalante Ranger District, Dixie National Forest, Garfield County, UT.

Summary: No formal comment letter was sent to the preparing agency.

EIS No. 20090375, ERP No. F-AFS-L65575-OR, Deadlog Vegetation Management Project, To Implement Treatments that would Reduce the Risk of High Intensity, Stand Replacement Wildlife and the Risk of Heavy Tree Mortality from Insects and Disease, Deschutes National Forest Lands, Deschutes County, OR.

Summary: EPA does not object to the proposed action.

Dated: November 17, 2009.

Robert W. Hargrove,

Director, NEPA Compliance Division, Office of Federal Activities.

[FR Doc. E9-27966 Filed 11-20-09; 8:45 am]

BILLING CODE 6560-50-P

ENVIRONMENTAL PROTECTION AGENCY

[ER-FRL-8799-4]

Environmental Impacts Statements; Notice of Availability

Responsible Agency: Office of Federal Activities, General Information (202) 564-1399 or <http://www.epa.gov/compliance/nepa/>.

Weekly receipt of Environmental Impact Statements.

Filed 11/09/2009 through 11/13/2009. Pursuant to 40 CFR 1506.9.

EIS No. 20090391, Final EIS, AFS, CA, Salt Timber Harvest and Fuel Hazard Reduction Project, Proposing Vegetation Management in the Salt Creek Watershed, South Fork Management Unit, Hayfork Ranger District, Shasta-Trinity National Forest, Trinity County, CA, Wait Period Ends: 12/21/2009, Contact: Sand Mack 406-375-2638.

EIS No. 20090392, Draft Supplement, FHW, HI, Saddle Road (HI-200) Improvements Project. Proposed Improvement from Mamalahoa Highway (HI-190) to Milepost 41, Hawaii County, HI, Comment Period Ends: 01/07/2010, Contact: Melissa Dickard 720-963-3691.

EIS No. 20090393, Final EIS, DOE, MN, Mesaba Energy Project, Proposes to Design, Construct and Operate a Coal-Based Integrated Gasification Cycle (IGCC) Electric Power Generating Facility, Located in the Taconite Tax Relief Area (TTRA), Itasca and St. Louis Counties, MN, Wait Period Ends: 12/21/2009, Contact: Richard A. Hargis, Jr 412-386-6065.

EIS No. 20090394, Draft EIS, USN, GU, Guam and Commonwealth of the Northern Mariana Islands (CNMI) Military Relocation, Proposed Relocating Marines from Okinawa, Visiting Aircraft Carrier Berthing, and Army Air and Missile Defense Task Force, Implementation, GU, Comment Period Ends: 01/04/2010, Contact: Kyle Fujimoto 808-472-1442.

EIS No. 20090395, Draft EIS, FTA, CO, North Metro Corridor Project, Proposed a Commuter Rail Transit from downtown Denver, Colorado, north to State Highway (SH) 7, in the Cities of Denver, Commerce City, Thornton, Northglenn, and Adams

County, CO, Comment Period Ends: 01/15/2010, Contact: David Beckhouse 720-963-3306.

EIS No. 20090396, Draft Supplement, CGD, AL, Bienville Offshore Energy Terminal (BOET) Deepwater Port License Application Amendment (Docket # USCG-2006-24644), Proposes to Construct and Operate a Liquefied Natural Gas Receiving and Regasification Facility, Outer Continental Shelf of the Gulf of Mexico, South of Fort Morgan, AL, Comment Period Ends: 01/04/2010, Contact: Lt. Hannah Kawamoto 202-372-1438.

EIS No. 20090397, Draft EIS, USA, 00, Programmatic—Louisiana Coastal Area (LCA) Beneficial Use of Dredged Material (BUDMAT) Program Study, To Establish the Structure and Management Architecture of the BUDMAT Program, Implementation, MS, TX and LA, Comment Period Ends: 01/04/2010, Contact: Elizabeth McCasland 504-862-2021.

Amended Notices.

EIS No. 20090324, Draft EIS, AFS, 00, Nebraska National Forests and Grassland Travel Management Project, Proposes to Designate Routes and Areas Open to Motorized Travel, Buffalo Gap National Grassland, Oglala National Grassland, Samuel R. McKelvie National Forest, and the Pine Ridge and Bessey Units of the Nebraska National Forest, Fall River, Custer, Pennington, Jackson Counties; SD and Sioux, Dawes, Cherry, Thomas and Blaine Counties, NE., Comment Period Ends: 12/24/2009, Contact: Mark Reichert 530-841-4422.

Revision to FR Published 09/25/2009: Extending Comment Period from 11/09/2009 to 12/24/2009.

EIS No. 20090366, Final EIS, FHW, CO, US-36 Corridor, Multi-Modal Transportation Improvements between I-25 in Adams County and Foothills Parkway/Table Mesa Drive in Boulder, Adams, Denver, Broomfield, Boulder and Jefferson Counties, CO, Wait Period Ends: 12/14/2009, Contact: Monica Pavlik 720-963-3012.

Revision to FR Notice Published 10/30/2009: Extending Comment Period from 11/30/2009 to 12/14/2009.

Dated: November 17, 2009.

Robert W. Hargrove,

Director, NEPA Compliance Division, Office of Federal Activities.

[FR Doc. E9-27968 Filed 11-19-09; 8:45 am]

BILLING CODE 6560-50-P

ENVIRONMENTAL PROTECTION AGENCY

[FRL-8983-8]

National Drinking Water Advisory Council's Climate Ready Water Utilities Working Group Meeting Announcement**AGENCY:** Environmental Protection Agency.**ACTION:** Notice.

SUMMARY: The Environmental Protection Agency (EPA or Agency) is announcing the first in-person meeting of the Climate Ready Water Utilities (CRWU) Working Group of the National Drinking Water Advisory Council (NDWAC). The purpose of this meeting is for the Working Group to begin addressing its charge, including defining the attributes of climate ready water utilities.

DATES: The first in-person CRWU Working Group meeting will take place on December 3, 2009, from 8:30 a.m. to 5 p.m., Eastern Standard Time (EST) and on December 4, 2009, from 8:30 am to 1:30 p.m., EST.

ADDRESSES: The meeting will take place at the Marriott Washington, which is located at 1221 22nd Street, NW., Washington, DC 20037.

FOR FURTHER INFORMATION CONTACT: Interested participants from the public should contact Lauren Wisniewski, Designated Federal Officer, U.S. Environmental Protection Agency, Office of Ground Water and Drinking Water, Water Security Division (Mail Code 4608T), 1200 Pennsylvania Avenue, NW., Washington, DC 20460. Please contact Lauren Wisniewski at wisniewski.lauren@epa.gov or call 202-564-2918 to receive additional details.

SUPPLEMENTARY INFORMATION:

Public Participation: There will be an opportunity for public comment during the CRWU Working Group meeting. Oral statements will be limited to five minutes, and it is preferred that only one person present the statement on behalf of a group or organization. Any person who wishes to file a written statement can do so before or after the CRWU Working Group meeting. Written statements received prior to the meeting will be distributed to all members of the Working Group before any final discussion or vote is completed. Any statements received after the meeting will become part of the permanent meeting file and will be forwarded to the CRWU Working Group members for their information. Any person needing special accommodations for this meeting, including wheelchair access, should contact the Designated Federal

Officer at the number or e-mail listed under the **FOR FURTHER INFORMATION CONTACT** section, at least five business days before the meeting so that appropriate arrangements can be made.

Background: The Agency's *National Water Program Strategy: Response to Climate Change* (2008) identified the need to provide drinking water and wastewater utilities with easy-to-use resources to assess the risk associated with climate change and to identify potential adaptation strategies. The NDWAC, established under the Safe Drinking Water Act, as amended (42 U.S.C. 300f *et seq.*), provides practical and independent advice, consultation and recommendations to the Agency on the activities, functions and policies related to the implementation of the Safe Drinking Water Act. On May 28, 2009, the NDWAC voted on and approved the formation of the CRWU Working Group. EPA anticipates that the Working Group will have five face-to-face meetings over the course of the next year in addition to conference calls and/or video conferencing on an as needed basis. After the Working Group completes its charge, it will make recommendations to the full NDWAC. The full NDWAC will, in turn, make appropriate recommendations to the EPA.

Working Group Charge: The charge for the CRWU Working Group is to evaluate the concept of "Climate Ready Water Utilities" and provide recommendations to the full NDWAC on the development of an effective program for drinking water and wastewater utilities, including recommendations to:

- (1) Define and develop a baseline understanding of how to use available information to develop climate change adaptation and mitigation strategies, including ways to integrate this information into existing complementary programs such as the Effective Utility Management and Climate Ready Estuaries Program;
- (2) Identify climate change-related tools, training, and products that address short-term and long-term needs of water and wastewater utility managers, decision makers, and engineers, including ways to integrate these tools and training into existing programs; and
- (3) Incorporate mechanisms to provide recognition or incentives that facilitate broad adoption of climate change adaptation and mitigation strategies by the water sector into existing EPA Office of Water recognition and awards programs or new recognition programs.

Dated: November 16, 2009.

Cynthia C. Dougherty,*Director, Office of Ground Water and Drinking Water.*

[FR Doc. E9-27940 Filed 11-19-09; 8:45 am]

BILLING CODE 6560-50-P

EQUAL EMPLOYMENT OPPORTUNITY COMMISSION**Sunshine Act Notice Meeting****AGENCY HOLDING THE MEETING:** Equal Employment Opportunity Commission.**FEDERAL REGISTER CITATION OF PREVIOUS ANNOUNCEMENT:** 74 FR 58626, Friday, November 13, 2009.**PREVIOUSLY ANNOUNCED TIME AND DATE OF MEETING:** Wednesday, November 18, 2009, 2 p.m. (Eastern Time).**CHANGE IN THE MEETING:** The meeting has been cancelled.**FOR FURTHER INFORMATION CONTACT:** Stephen Llewellyn, Executive Officer on (202) 663-4070.

This Notice Issued November 17, 2009.

Stephen Llewellyn,*Executive Officer, Executive Secretariat.*

[FR Doc. E9-28030 Filed 11-18-09; 4:15 pm]

BILLING CODE 6570-01-P

FEDERAL ELECTION COMMISSION

[Notice 2009-25]

Filing Dates for the Florida Special Election in the 19th Congressional District**AGENCY:** Federal Election Commission.**ACTION:** Notice of filing dates for special election.

SUMMARY: Florida has scheduled elections on February 2, 2010, and April 6, 2010, to fill the U.S. House seat in the 19th Congressional District being vacated by Representative Robert Wexler.

Committees required to file reports in connection with the Special Primary Election on February 2, 2010, shall file a 12-day Pre-Primary Report. Committees required to file reports in connection with both the Special Primary and Special General Election on April 6, 2010, shall file a 12-day Pre-Primary Report, a 12-day Pre-General Report, and a 30-day Post-General Report.

FOR FURTHER INFORMATION CONTACT: Mr. Kevin R. Salley, Information Division, 999 E Street, NW., Washington, DC 20463; *Telephone:* (202) 694-1100; Toll Free (800) 424-9530.

SUPPLEMENTARY INFORMATION:

Principal Campaign Committees

All principal campaign committees of candidates who participate in the Florida Special Primary and Special General Elections shall file a 12-day Pre-Primary Report on January 21, 2010; a 12-day Pre-General Report on March 25, 2010; and a 30-day Post-General Report on May 6, 2010. (See chart below for the closing date for each report).

All principal campaign committees of candidates participating *only* in the Special Primary Election shall file a 12-day Pre-Primary Report on January 21, 2010. (See chart below for the closing date for each report).

Note that these reports are in addition to the campaign committee's quarterly filings in April and July. (See chart below for the closing date for each report).

Unauthorized Committees (PACs and Party Committees)

Political committees filing on a quarterly basis in 2010 are subject to special election reporting if they make previously undisclosed contributions or expenditures in connection with the

Florida Special Primary or Special General Elections by the close of books for the applicable report(s). (See chart below for the closing date for each report).

Since disclosing financial activity from two different calendar years on one report would conflict with the calendar year aggregation requirements stated in the Commission's disclosure rules, unauthorized committees that trigger the filing of the Pre-Primary Report will be required to file this report on two separate forms: One form to cover 2009 activity, labeled as the Year-End Report; and the other form to cover only 2010 activity, labeled as the Pre-Primary Report. Both forms must be filed by January 21, 2010.

Committees filing monthly that make contributions or expenditures in connection with the Florida Special Primary or Special General Elections will continue to file according to the monthly reporting schedule.

Additional disclosure information in connection with the Florida Special Election may be found on the FEC Web site at http://www.fec.gov/info/report_dates.shtml.

Disclosure of Lobbyist Bundling Activity

Campaign committees, party committees and Leadership PACs that are otherwise required to file reports in connection with the special elections must simultaneously file FEC Form 3L if they receive two or more bundled contributions from lobbyists/registrants or lobbyist/registrant PACs that aggregate in excess of the lobbyist bundling disclosure threshold during the special election reporting periods (see charts below for closing date of each period). 11 CFR 104.22(a)(5)(v).

The lobbyist bundling disclosure threshold for calendar year 2009 is \$16,000. This threshold amount may increase in 2010 based upon the annual cost of living adjustment (COLA). As soon as the adjusted threshold amount is available, the Commission will publish it in the **Federal Register** and post it on its Web site. 11 CFR 104.22(g) and 110.17(e)(2). For more information on these requirements, see **Federal Register** Notice 2009-03, 74 FR 7285 (February 17, 2009).

CALENDAR OF REPORTING DATES FOR FLORIDA SPECIAL ELECTION

Report	Close of books ¹	Reg./cert. & overnight mailing deadline	Filing deadline
Committees Involved in <i>Only</i> the Special Primary (02/02/10) Must File:			
Year-End		—WAIVED—	
Pre-Primary	01/13/10	² 01/18/10	01/21/10
April Quarterly	03/31/10	04/15/10	04/15/10
Committees Involved in Both the Special Primary (02/02/10) and Special General (04/06/10) Must File:			
Year-End		—WAIVED—	
Pre-Primary	01/13/10	² 01/18/10	01/21/10
Pre-General	03/17/10	03/22/10	03/25/10
April Quarterly	03/31/10	04/15/10	04/15/10
Post-General	04/26/10	05/06/10	05/06/10
July Quarterly	06/30/10	07/15/10	07/15/10
Committees Involved in <i>Only</i> the Special General (04/06/10) Must File:			
Pre-General	03/17/10	03/22/10	03/25/10
April Quarterly	03/31/10	04/15/10	04/15/10
Post-General	04/26/10	05/06/10	05/06/10
July Quarterly	06/30/10	07/15/10	07/15/10

¹ The reporting period always begins the day after the closing date of the last report filed. If the committee is new and has not previously filed a report, the first report must cover all activity that occurred before the committee registered as a political committee with the Commission up through the close of books for the first report due.

² Notice that the registered/certified & overnight mailing deadline falls on a federal holiday. The report should be postmarked on or before that date.

Dated: November 16, 2009.
On behalf of the Commission.

Steven T. Walther,
Chairman, Federal Election Commission.
[FR Doc. E9-27869 Filed 11-19-09; 8:45 am]

BILLING CODE 6715-01-P

FEDERAL RESERVE SYSTEM

Change in Bank Control Notices; Acquisition of Shares of Bank or Bank Holding Companies

The notificants listed below have applied under the Change in Bank

Control Act (12 U.S.C. 1817(j)) and § 225.41 of the Board's Regulation Y (12 CFR 225.41) to acquire a bank or bank holding company. The factors that are considered in acting on the notices are set forth in paragraph 7 of the Act (12 U.S.C. 1817(j)(7)).

The notices are available for immediate inspection at the Federal Reserve Bank indicated. The notices also will be available for inspection at the office of the Board of Governors. Interested persons may express their views in writing to the Reserve Bank indicated for that notice or to the offices of the Board of Governors. Comments must be received not later than December 4, 2009.

A. Federal Reserve Bank of Philadelphia (Michael E. Collins, Senior Vice President) 100 North 6th Street, Philadelphia, Pennsylvania 19105-1521:

1. *Patriot Financial Partners, GP, L.P., Patriot Financial Partners, L.P., Patriot Financial Partners Parallel, L.P., Patriot Financial Partners, GP, LLC, Patriot Financial Managers, L.P., and Ira M. Lubert, W. Kirk Wycoff and James J. Lynch*, all of Philadelphia, Pennsylvania; to acquire voting shares of Square 1 Financial, Inc., and thereby indirectly acquire voting shares of Square 1 Bank, both of Durham, North Carolina.

B. Federal Reserve Bank of Atlanta (Steve Foley, Vice President) 1000 Peachtree Street, N.E., Atlanta, Georgia 30309:

1. *Serguei Kouzmine*, Atlanta, Georgia; to acquire voting shares of UCB Financial Group, Inc., and thereby indirectly acquire voting shares of Atlanta Business Bank, both of Atlanta, Georgia.

C. Federal Reserve Bank of Kansas City (Todd Offenbacher, Assistant Vice President) 1 Memorial Drive, Kansas City, Missouri 64198-0001:

1. *The Estate of Catherine G. Eisemann*, Trinidad, Colorado, and Roger Dean Eisemann, Houston, Texas, individually, as a member of the Eisemann Family and as Co-Executor of the Estate of Catherine G. Eisemann, to retain control of Republic Trinidad Corporation, Houston, Texas, and thereby indirectly retain control of First National Bank in Trinidad, Trinidad, Colorado.

Board of Governors of the Federal Reserve System, November 16, 2009.

Robert deV. Frierson,

Deputy Secretary of the Board.

[FR Doc. E9-27873 Filed 11-19-09; 8:45 am]

BILLING CODE 6210-01-S

FEDERAL RESERVE SYSTEM

Formations of, Acquisitions by, and Mergers of Bank Holding Companies

The companies listed in this notice have applied to the Board for approval, pursuant to the Bank Holding Company

Act of 1956 (12 U.S.C. 1841 *et seq.*) (BHC Act), Regulation Y (12 CFR Part 225), and all other applicable statutes and regulations to become a bank holding company and/or to acquire the assets or the ownership of, control of, or the power to vote shares of a bank or bank holding company and all of the banks and nonbanking companies owned by the bank holding company, including the companies listed below.

The applications listed below, as well as other related filings required by the Board, are available for immediate inspection at the Federal Reserve Bank indicated. The applications also will be available for inspection at the offices of the Board of Governors. Interested persons may express their views in writing on the standards enumerated in the BHC Act (12 U.S.C. 1842(c)). If the proposal also involves the acquisition of a nonbanking company, the review also includes whether the acquisition of the nonbanking company complies with the standards in section 4 of the BHC Act (12 U.S.C. 1843). Unless otherwise noted, nonbanking activities will be conducted throughout the United States. Additional information on all bank holding companies may be obtained from the National Information Center website at www.ffiec.gov/nic/.

Unless otherwise noted, comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors not later than December 14, 2009.

A. Federal Reserve Bank of Dallas (E. Ann Worthy, Vice President) 2200 North Pearl Street, Dallas, Texas 75201-2272:

1. *Community Trust Financial Corporation*, Ruston, Louisiana; to merge with Madison Financial Corporation, and thereby indirectly acquire Madison County Bank, both of Madison, Mississippi.

Board of Governors of the Federal Reserve System, November 16, 2009.

Robert deV. Frierson,

Deputy Secretary of the Board.

[FR Doc. E9-27872 Filed 11-19-09; 8:45 am]

BILLING CODE 6210-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

[Document Identifier OS-0990-New]

Agency Information Collection Request; 60-Day Public Comment Request

AGENCY: Office of the Secretary, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the

Paperwork Reduction Act of 1995, the Office of the Secretary (OS), Department of Health and Human Services, is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden. To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, e-mail your request, including your address, phone number, OMB number, and OS document identifier, to Sherette.funncoleman@hhs.gov, or call the Reports Clearance Office on (202) 690-6162. Written comments and recommendations for the proposed information collections must be directed to the OS Paperwork Clearance Officer at the above e-mail address within 60 days.

Proposed Project: Evaluation of the Parents Speak Up National Campaign (PSUNC): Focus Groups with Adolescents. OMB No. 0990-NEW-Office of the Secretary/Office of Public Health and Science/Office Adolescent Pregnancy Programs.

Abstract: The data collection will take place once, over a three day period, in early 2010. An estimated 2000 adults will be screened to identify parents who are willing for their child to participate in the study and whose child is eligible. Screening will take an estimated 3 minutes, on average. Study participants will total 160 adolescents ages 13-15. Participation in the study will take an estimated 2 hours on average; including time spent responding to a mini-questionnaire and participating in a bulletin board focus group. Participants will self-administer the mini-questionnaire at home on personal computers and will also participate in the focus group online. The specific aim of this study is to assess qualitatively what kinds of information about sex adolescents want to hear from their parents and their perspectives on the factors that either hinder or facilitate effective communication.

ESTIMATED ONE-YEAR ANNUALIZED BURDEN TABLE

Forms (if necessary)	Type of respondent	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Screener	Adults	2,000	1	3/60	100
Focus group discussion guide and mini-questionnaire.	Adolescents ages 13–15.	160	1	2	320
Total	2,160	420

Seleda M. Perryman,
Office of the Secretary, Paperwork Reduction Act Reports Clearance Officer.
 [FR Doc. E9–27881 Filed 11–19–09; 8:45 am]
BILLING CODE 4150–30–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Delegation of Authority

Notice is hereby given that I have delegated to the Assistant Secretary for Aging the authorities vested in the Secretary of Health and Human Services under Section 1701(a)(3)(A–B), Section 1701(a)(4), and Section 1703(a)(4) of the Public Health Service Act (42 U.S.C. 300u(a)(3)(A–B), 300u(a)(4), and 300u–2(a), as amended), as they pertain to the exercise of the funds transferred by the Secretary to the Administration on Aging under the “Prevention and Wellness Fund” of the American Recovery and Reinvestment Act of 2009, Public Law 111–5 (Feb. 17, 2009) to carry out evidence-based clinical and community-based prevention and wellness strategies through chronic disease self-management programs targeted to improving the health of seniors under the “Communities Putting Prevention to Work” initiative.

These authorities may be redelegated.

Exercise of these authorities is concurrent to and does not supplant existing delegations of authority from the Secretary. Exercise of these authorities shall be in accordance with established policies, procedures, guidelines, and regulations as prescribed by the Secretary.

I hereby affirm and ratify any actions taken by the Assistant Secretary for Aging or his or her subordinates, which involved the exercise of the authorities delegated herein prior to the effective date of this delegation. This delegation is effective immediately.

Dated: November 12, 2009.

Kathleen Sebelius,
Secretary.
 [FR Doc. E9–27863 Filed 11–19–09; 8:45 am]
BILLING CODE 4150–03–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Indian Health Service

Request for Public Comment: 30-Day Notice; Proposed Information Collection: Indian Health Service Forms

AGENCY: Indian Health Service, HHS.
ACTION: Request for Public Comment: 30-day Proposed Information Collection: Indian Health Service Forms to Implement the Privacy Rule (45 CFR parts 160 & 164).

SUMMARY: The Indian Health Service (IHS), as part of its continuing effort to reduce paperwork and respondent burden, conducts a pre-clearance consultation program to provide the general public and Federal agencies with an opportunity to comment on proposed and/or continuing collections of information in accordance with the Paperwork Reduction Act of 1995 (PRA95) (44 U.S.C. 3506(c)(2)(A)). This program helps to ensure that requested data can be provided in the desired format, reporting burden (time and financial resources) is minimized, collection instruments are clearly understood, and the impact of collection requirements on respondents can be properly assessed. As required by section 3507(a)(1)(D) of the PRA95, the proposed information collection has been submitted to the Office of Management and Budget (OMB) for review and approval.

The IHS received no comments in response to the 60-day **Federal Register** notice (74 FR 30095) published on June 24, 2009. The purpose of this notice is to allow an additional 30 days for public comment to be submitted directly to OMB.

Proposed Collection: Title: 0917–0030, “Indian Health Service Forms to Implement the Privacy Rule (45 CFR parts 160 & 164)”. *Type of Information Collection Request:* Extension, with revisions, of currently approved information collection, 0917–0030, “Indian Health Service Forms to Implement the Privacy Rule (45 CFR

parts 160 & 164)”. *Form Number:* IHS–810, IHS–912–1, IHS–912–2, IHS–913, and IHS–917. *Need and Use of Information Collection:* The IHS will use the following data collection instructions to continue the implementation of the information collection requirements contained in the Privacy Rule.

Request for Comments: Your written comments and/or suggestions are invited on one or more of the following points: (a) Whether the information collection activity is necessary to carry out an agency function; (b) whether the agency processes the information collected in a useful and timely fashion; (c) the accuracy of public burden estimate (the estimated amount of time needed for individual respondents to provide the requested information); (d) whether the methodology and assumptions used to determine the estimate are logical; (e) ways to enhance the quality, utility, and clarity of the information being collected; and (f) ways to minimize the public burden through the use of automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

Direct Comments to OMB: Send your written comments and suggestions regarding the proposed information collection contained in this notice, especially regarding the estimated public burden and associated response time, directly to: Office of Management and Budget, Office of Regulatory Affairs, New Executive Office Building, Room 10235, Washington, DC 20503, Attention: Allison Eydt, Desk Officer for IHS.

For Further Information: Send requests for more information on the proposed collection or to obtain a copy of the data collection instrument(s) and instructions to: Ms. Betty Gould, IHS Reports Clearance Officer, 801 Thompson Avenue, TMP, Suite 450, Rockville, MD 20852–1601, call non-toll free (301) 443–7899, send via facsimile to (301) 443–9879, or send your e-mail requests, comments, and return address to: betty.gould@ihs.gov.

Comment Due Date: Your comments regarding this information collection are best assured of having full effect if received within 30 days of the date of this publication.

Dated: November 6, 2009.

Yvette Roubideaux,

Director, Indian Health Service.

[FR Doc. E9-27541 Filed 11-19-09; 8:45 am]

BILLING CODE 4165-16-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2009-N-0556]

Agency Information Collection Activities; Proposed Collection; Comment Request; Records and Reports Concerning Experience With Approved New Animal Drugs; Proposed New Data Elements for Adverse Event Reports on Revised Forms FDA 1932 and 1932a

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information and to allow for public comment in response to the notice. This notice solicits comments on requirements for recordkeeping and reports concerning experience with approved new animal drugs, specifically on new data elements to be used in revised versions of Forms FDA 1932 and 1932a. The information contained in the reports required by this regulation enables FDA to monitor the use of new animal drugs after approval and to ensure their continued safety and efficacy.

DATES: Submit written or electronic comments on the collection of information by December 21, 2009.

ADDRESSES: Submit electronic comments on the collection of information to <http://www.regulations.gov>. Submit written comments on the collection of information to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the

docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Denver Presley, Office of Management Programs (HFA-710), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-796-3793.

SUPPLEMENTARY INFORMATION:

I. Background

Under the PRA (44 U.S.C. 3501-3520), Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. "Collection of information" is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal agencies to provide a notice in the **Federal Register** concerning each proposed collection of information before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on these topics: (1) Whether the proposed collection of information is necessary for the proper performance of FDA's functions, including whether the information will have practical utility; (2) the accuracy of FDA's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.

II. Records and Reports Concerning Experience With Approved New Animal Drugs; Proposed New Data Elements for Adverse Event Reports on Revised Forms FDA 1932 and 1932a; 21 CFR 514.80 (OMB Control No. 0910-0645)—Revision

Section 512(l) of the Federal Food, Drug and Cosmetic Act (the act) (21 U.S.C. 360b(l)) and § 514.80(b) of FDA regulations (21 CFR 514.80) require applicants of approved new animal drug applications (NADAs) and approved abbreviated new animal drug applications (ANADAs) to report adverse drug experiences and product/manufacturing defects.

This continuous monitoring of approved NADAs and ANADAs affords the primary means by which FDA obtains information regarding potential problems with the safety and efficacy of marketed approved new animal drugs as well as potential product/manufacturing problems. Postapproval marketing surveillance is important because data previously submitted to FDA may no longer be adequate, as animal drug effects can change over time and less apparent effects may take years to manifest.

An applicant must report adverse drug experiences and product/manufacturing defects on Form FDA 1932, "Veterinary Adverse Drug Reaction, Lack of Effectiveness, Product Defect Report." Periodic drug experience reports and special drug experience reports must be accompanied by a completed Form FDA 2301, "Transmittal of Periodic Reports and Promotional Material for New Animal Drugs" (see § 514.80(d)). Form FDA 1932a, "Veterinary Adverse Drug Reaction, Lack of Effectiveness or Product Defect Report," allows for voluntary reporting of adverse drug experiences or product/manufacturing defects.

Collection of information using existing paper forms FDA 2301, 1932, and 1932a is currently approved under OMB control number 0910-0284, set to expire on January 31, 2010. FDA currently is seeking renewal of that information collection.

FDA recently proposed to collect information using electronic versions of Forms FDA 1932 and 1932a as part of the agency-wide information collection (MedWatch^{Plus} Portal and Rational Questionnaire) that was announced for public comment in the **Federal Register** on October 23, 2008 (73 FR 63153). The MedWatch^{Plus} Portal and Rational Questionnaire are components of a new electronic system for collecting, submitting, and processing adverse event reports and other safety information for all FDA-regulated products.

In this 30-day notice, FDA is requesting public comment on data elements associated with revisions to forms FDA 1932 and 1932a (both paper and electronic) under revised OMB control number 0910-0645, described below. We will publish separately in the **Federal Register** a 30-day notice to complete the renewal of OMB control number 0910-0284, the collection of information using existing paper forms FDA 2301, 1932, and 1932a, to provide time for development of the revised FDA Forms 1932 and 1932a and their incorporation into the MedWatch OMB

control number 0910–0645. After these forms have been incorporated under MedWatch OMB control number 0910–0645, they will cease to exist under OMB control number 0910–0284. FDA Form 2301 will continue without revision under OMB control number 0910–0284.

This 30-day notice lists the data elements associated with revised versions of both paper and electronic forms 1932 and 1932a under a revision

to OMB control number 0910–0645. It is estimated that, during the first 3 years that the MedWatch^{Plus} Portal is in use, half of the reports will be submitted in paper format and half will be submitted electronically.

The reporting and recordkeeping burden estimates, including the total number of annual responses, are based on the submission of reports to the Division of Surveillance, Center for Veterinary Medicine. The hours per

response for both paper and electronic versions of revised Forms FDA 1932 and 1932a are assumed to be the time it will take to gather the required information and complete each form. The annual frequency of responses was calculated as the total annual responses divided by the number of respondents.

FDA estimates the burden of this collection of information as follows:

TABLE 1.—ESTIMATED ANNUAL REPORTING BURDEN¹

21 CFR Section or Section of the Act	FDA Form No.	No. of Respondents	Annual Frequency per Response	Total Annual Responses	Hours per Response	Total Hours
514.80(b)(1), (b)(2)(i), (b)(2)(ii), and (b)(3); Paper Version	1932 ²	404	44.264	17,882.5	1.5	26,824
514.80(b)(1), (b)(2)(i), (b)(2)(ii), and (b)(3); Electronic Version	1932 ²	404	44.264	17,882.5	1	17,882.5
Voluntary reporting FDA Form 1932a for public; Paper Version	1932a ³	81.5	1	81.5	1	81.5
Voluntary reporting FDA Form 1932a for public; Electronic Version	1932a ³	81.5	1	81.5	0.6	48.9
Total Hours						44,836.9

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

² FDA received 35,765 mandatory reports (Form FDA 1932) during 2007 from 808 respondents. Based on this experience, and taking into account the data element revisions, we estimate that CVM will receive 35,765 mandatory reports from 808 respondents annually. We estimate that one half of the respondents (404) will use the paper form, while the other half (404) will submit electronically; that is, we will receive 17,882.5 reports in paper form, and 17,882.5 reports electronically. We estimate the reporting burden for mandatory reporting to be: Paper form: 26,824 hours (404 respondents x 44.264 annual frequency of response x 1.5 hours ≈ 26,824 hours). Electronic form: 17,882.5 hours (404 respondents x 44.264 annual frequency of response x 1 hour ≈ 17,882.5 hours).

³ FDA received 163 voluntary reports (Form FDA 1932a) during 2007. Based on this experience, and taking into account the data element revisions, we estimate that CVM will receive 163 voluntary reports from 163 respondents annually. We estimate that one half of the respondents (81.5) will use the paper form, while the other half (81.5) will submit electronically; that is, we will receive 81.5 reports in paper form, and 81.5 reports electronically. We estimate the reporting burden for voluntary reporting to be: Paper form: 81.5 hours (81.5 respondents x 1 annual frequency of response x 1 hour per report = 81.5 hours). Electronic form: 48.9 hours (81.5 respondents x 1 annual frequency of response x 0.6 hours per report = 48.9 hours).

TABLE 2.—ESTIMATED ANNUAL RECORDKEEPING BURDEN¹

21 CFR Section	No. of Recordkeepers	Annual Frequency per Recordkeeping	Total Annual Records	Hours per Record	Total Hours
514.80(e) ²	90	55	4,949	0.5	2,475 ³

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

² Section 514.80(e) covers recordkeeping hours for adverse event reporting on revised forms 1932 and 1932a.

³ The annual frequency of responses was calculated as the total annual responses divided by the number of respondents.

III. Revisions to Forms FDA 1932 and 1932a and Request for Comments

A. Background on Revisions

Section 514.80(d) of FDA’s regulations requires applicants of approved NADAs and ANADAs to report adverse drug experiences and product and manufacturing defects associated with their new animal drug products using Form FDA 1932. For voluntary reporting, Form FDA 1932a should be used instead.

As part of FDA’s ongoing effort to harmonize the agency’s adverse event (AE) regulatory reporting requirements

with those of other nations and streamline reporting for product and manufacturing defects, FDA is contemplating changes to the data elements reported on Forms FDA 1932 and 1932a. Furthermore, the contemplated changes to Forms FDA 1932 and 1932a are based on FDA’s experience in determining the safety and effectiveness of product(s) and need for efficient data capture and entry.

The contemplated changes to the AE reporting requirements for Form FDA 1932 are the product of discussions undertaken between the United States, Japan, and the European Union as part

of the International Cooperation on Harmonisation of Technical Requirements for Registration of Veterinary Medicinal Products (VICH). FDA is considering revisions to Form FDA 1932 that would bring the AE reporting data elements on the form more in line with the data elements developed as a result of the VICH discussions.¹ The agency also is

¹ FDA will implement all of the VICH data elements verbatim from the draft guidance document entitled “Pharmacovigilance of Veterinary Medicinal Products Data Elements for Submission of Adverse Event Reports” (VICH GL–42), in Form FDA 1932. VICH GL–42 is currently

contemplating the inclusion of additional new data elements that would gather information specific only to FDA. Collecting this FDA-specific information is essential for the processing, review, and regulatory disposition of the electronic and paper reports. Inclusion of some of the new data elements is necessitated by the Rational Questionnaire.

In addition, the agency is considering adding new data elements for product and manufacturing defect reports on Form FDA 1932 and 1932a. These changes are the product of internal FDA discussions and are intended to capture additional pertinent product and manufacturing defect information.

B. Proposed Revisions

1. Form FDA 1932

This section describes data elements on the current Form FDA 1932,

proposed new data elements, and data elements we propose to delete from the current form. These AE and product and manufacturing defect data elements will be collected electronically, through the MedWatch^{Plus} Portal and Rational Questionnaire (currently under development), and in the paper form.

Table 3 of this document, entitled "Data Elements for Form FDA 1932," presents the data elements for the collection of animal drug adverse event reports and manufacturing and product defect reports. The data elements are listed in the column entitled "Data Elements." The column entitled "Current, New, or Deleted Data Element" indicates whether the data element is currently being collected (Current)², is a proposed new data element (New), or is a data element FDA proposes to delete (Deleted).

As previously mentioned in this document, the agency has had

discussions with VICH regarding the data elements to be collected for animal drug adverse events. As a result, the agency is proposing new data elements that have been negotiated with VICH. The column entitled "VICH-Negotiated or FDA-Proposed Data Element" differentiates between VICH-negotiated and FDA-proposed data elements.

The agency intends to allow the regulated industry to submit this information collection in three different submission/transmission formats. Industry will be able to submit these reports using a paper form, the Web-based Rational Questionnaire, or an electronic file through the FDA electronic Gateway-to-Gateway transmission. The column entitled "Submission/Transmission Format" presents the submission/transmission format(s) that will be used with each particular data element.

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
1	United States-Only Specific Information, including:			
2	Report Identifier (The Report Identifier is the FDA application or file number of the AER being sent.)	Current	FDA Proposed	All Formats
3	Domestic vs. Foreign Category (This is a list of values describing whether the product is an FDA-approved product, a foreign-approved product, or other type of product, e.g., an unapproved drug.)	New	FDA Proposed	All Formats
4	United States Pharmacovigilance Contact Person for the Applicant or Nonapplicant (This is the person within the United States acting on behalf of the applicant or nonapplicant and is the contact person for the FDA for any pharmacovigilance issues about the report.), including:			
5	Title, First and Last Name	Current	FDA Proposed	All Formats
6	Telephone Number, Fax Number, and E-Mail Address	New	FDA Proposed	All Formats
7	Message Sender Identifier (Name and contact information of person responsible for any corresponding communications regarding the whole batch electronic transmission.), including:			
8	Street Address, City, State/County, and Mail/Zip Code	New	FDA Proposed	EGG Only
9	Three-character Country Code (This is the list of country codes from the International Organization for Standardization (ISO) 3166 standard.)	New	FDA Proposed	EGG Only
10	First and Last Name	New	FDA Proposed	EGG Only
11	Telephone Number, Fax Number, and E-Mail Address	New	FDA Proposed	EGG Only
12	Profile Identifier Code (This information indicates the type of report contained in the electronic message.)	New	FDA Proposed	EWBRQ and EGG Only

under discussion at Step 6. This guidance is available on the Internet at <http://www.fda.gov/RegulatoryInformation/Guidances/default.htm>.

²In general, the information being collected is the same, but the data element has been renamed or restructured to facilitate data collection.

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932—Continued

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
13	Batch ID (This information identifies the reports in this batch as a whole electronic message.)	New	FDA Proposed	EGG Only
14	Message Date (This information indicates the date this batch report is created.)	New	FDA Proposed	EGG Only
15	Message Version Number & Release Number (This information indicates the Health Level Seven, Inc. (HL7) "Message Version" and "Release Number" on which this batch report electronic submission is based.)	New	FDA Proposed	EGG Only
16	Adverse Event Report (AER) Information, including:			
17	Unique AER Identification Number (This globally unique AER identification number is created by and assigned by the applicant or nonapplicant.)	New	VICH Negotiated	All Formats
18	Original Receive Date (The original receive date is the date on which the first full communication of the AER was received by the applicant or nonapplicant responsible for reporting the AER to the FDA.)	Current	VICH Negotiated	All Formats
19	Date of Current Submission (This is the date the current AER was submitted to the Regulatory Authority (RA).)	Current	VICH Negotiated	All Formats
20	Type of Report, including:			
21	Type of Submission (This is a list of values describing the regulatory type of report being submitted to the RA, e.g., 15-day NADA/ANADA alert report, 3-day NADA/ANADA field alert report, followup report, nullification report, periodic drug experience report, and other report.)	Current	VICH Negotiated	All Formats
22	Reason for Nullification Report (This is a text description of why this AER is being nullified.)	Current	VICH Negotiated	All Formats
23	Type of Information in Report (This is a list of values for the categorization of the type of information in the AER, e.g., spontaneous safety and lack of expected effectiveness information, clinical study safety information, product and manufacturing defect information, product and manufacturing defect with safety and lack of expected effectiveness information, and other type of information.)	New	VICH Negotiated	All Formats
24	Regulatory Authority (RA) Information (This is the RA to which this AE report (AER) is to be initially submitted based on the RA that has authority to regulate the product.), including:			
25	RA Name	Current	VICH Negotiated	All Formats
26	Street Address, City, State/County, and Mail/Zip Code	Current	VICH Negotiated	All Formats
27	Three-character Country Code	Current	VICH Negotiated	All Formats
28	Marketing Authorization Holder (MAH) information. (The MAH is the applicant or the nonapplicant who is responsible for reporting the AER to the RA.), including:			
29	Business Name	Current	VICH Negotiated	All Formats
30	Street Address, City, State/County, and Mail/Zip Code	Current	VICH Negotiated	All Formats
31	Three-character Country Code	Current	VICH Negotiated	All Formats
32	Person Acting on Behalf of the MAH information, including:			
33	Title, First and Last Name	Current	VICH Negotiated	All Formats

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932—Continued

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
34	Telephone Number, Fax Number, and E-Mail Address	New	VICH Negotiated	All Formats
35	Primary Reporter's information (The primary reporter is the person or organization, as determined by the MAH, which holds or provides the most pertinent information related to this AER.), including:			
36	First and Last Name	Current	VICH Negotiated	All Formats
37	Telephone and Fax Number	Current	VICH Negotiated	All Formats
38	E-Mail Address	New	VICH Negotiated	All Formats
39	Business Name	Current	VICH Negotiated	All Formats
40	Street Address, City, State/County, and Mail/Zip Code	Current	VICH Negotiated	All Formats
41	Three-character Country Code	Current	VICH Negotiated	All Formats
42	Primary Reporter Category (This is a list of values describing the role/involvement of the primary reporter, e.g., animal owner, physician, et cetera.)	New	VICH Negotiated	All Formats
43	Other Reporter's information (The other reporter is the person or organization, determined by the MAH, who also possesses pertinent information related to this AER.), including:			
44	First and Last Name	Current	VICH Negotiated	All Formats
45	Telephone and Fax Number	Current	VICH Negotiated	All Formats
46	E-Mail Address	New	VICH Negotiated	All Formats
47	Business Name	Current	VICH Negotiated	All Formats
48	Street Address, City, State/County, and Mail/Zip Code	Current	VICH Negotiated	All Formats
49	Three-character Country Code	Current	VICH Negotiated	All Formats
50	Other Reporter Category (This is a list of values describing the role/involvement of the other reporter, e.g., animal owner, physician, et cetera.)	New	VICH Negotiated	All Formats
51	Veterinary Medical Product (VMP) and Data Usage (for all VMPs), including:			
52	Registered or Brand Name (This is the name by which the product is presented by the MAH, also known as the Proprietary Name or Trade Name of the product.)	Current	VICH Negotiated	All Formats
53	Product Code (The product code is the National Drug Code (NDC) number for U.S. FDA-regulated products.)	New	VICH Negotiated	All Formats
54	Registration Identifier (The Registration Identifier is the code for where the VMP is approved, what RA is responsible for regulating VMP, and the registration number of the VMP.)		VICH Negotiated	All Formats
55	ATCvet Code (ATCvet stands for Anatomic Therapeutic Chemical System for Veterinary Medicine. It is used for the classification of substances intended for therapeutic use and can serve as a tool for the classification of veterinary medicinal products. More information about the ATCvet code is available at http://www.whocc.no/atcvet/)	New	VICH Negotiated	All Formats
56	Who Administered the VMP (This is a list of values describing the person who administered the VMP(s) to the animal involved in the AE, e.g., veterinarian, animal owner, et cetera.)	Current	VICH Negotiated	All Formats

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932—Continued

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
57	Company or MAH (This is the name(s) of the company or MAH that owns the VMP(s) involved in the AE.)	Current	VICH Negotiated	All Formats
58	MAH Assessment (This is the assessment by the MAH of the association between the use of the VMP and the AE.)	Current	VICH Negotiated	All Formats
59	FDA, Office of Regulatory Affairs (ORA) District Field Office (This is a list of values identifying the ORA District Field Office or local FDA residence post to which the product and manufacturing defect information was submitted. This field is used for product and manufacturing defect reports and if the report is both an AE and a product and manufacturing defect report.)	New	FDA Proposed	All Formats
60	Use According to Label (This element requests information regarding whether the VMP(s) was used according to its label.)	Current	VICH Negotiated	All Formats
61	Explanation for Off-Label Use Code (This is the list of values describing how the VMP was used in an off-label (extralabel) manner.)	New	VICH Negotiated	All Formats
62	Active Ingredient information, including:			
63	Active Ingredient(s) (These are the names of the pharmaceutical substances that comprise the active component of the VMP.)	Current	VICH Negotiated	All Formats
64	Strength and Strength Unit (Numerator and Denominator) (Strength is the concentration of the active ingredient.)	Current	VICH Negotiated	All Formats
65	Active Ingredient Code (The active ingredient code is the Unique Ingredient Identifier (UNII) code. The UNII code is generated by the joint FDA/United States Pharmacopeia (USP) Substance Registration System (SRS).)	New	VICH Negotiated	All Formats
66	Dosage Form (This is a selection for a list of values for the labeled dosage form of the VMP(s).)	Current	VICH Negotiated	All Formats
67	Dosing Information, including:			
68	Date of First Exposure (Day, Month, Year) (This is the date on which the animal was first treated with the VMP.)	Current	VICH Negotiated	All Formats
69	Date of Last Exposure (Day, Month, Year) (This is the date on which the animal was last treated with the VMP.)	Current	VICH Negotiated	All Formats
70	Numeric Value and Unit for Interval of Administration (This is the frequency of administration of the VMP(s).)	Current	VICH Negotiated	All Formats
71	Numeric Value and Unit for Dose (This is the actual quantity of the dose administered.)	Current	VICH Negotiated	All Formats
72	Route of Exposure (This is a selection from a list of values for the route by which the VMP was administered.)	Current	VICH Negotiated	All Formats
73	Lot Number Information, including:			
74	Lot Number (This is the lot number associated with the VMP in this AER.)	Current	VICH Negotiated	All Formats
75	Expiration Date (Day, Month, Year) (This is the expiration date associated with the lot number.)	New	VICH Negotiated	All Formats
76	Manufacturing Site Identifier Number (This is the FDA Establishment Number (FEI Number) or the Data Universal Number System (D-U-N-S® Number).)	New	FDA Proposed	All Formats

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932—Continued

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
77	Manufacturer's Identifier Type (This is a list of values describing the type of manufacturing site identifier number, i.e., FEI Number or D-U-N-S® Number.)	New	FDA Proposed	All Formats
78	Manufacturing Date (Day, Month, Year) (This is the date the VMP was manufactured.)	New	FDA Proposed	All Formats
79	Number of Defective Units (This is the number of defective units associated with this VMP.)	New	FDA Proposed	All Formats
80	Number of Units Returned (This is the number of defective units associated with this VMP returned to the applicant or non-applicant.)	New	FDA Proposed	All Formats
81	Adverse Event Information, including:			
82	Attending Veterinarian's Assessment (This is a list of values describing the assessment of the attending veterinarian regarding the association between the VMP(s) and the AE (other than human).)	Current	VICH Negotiated	All Formats
83	Previous Exposure to the VMP (Was the animal previously exposed to the VMP(s)?)	Current	VICH Negotiated	All Formats
84	Previous AE to the VMP (Did the animal have a previous AE to the VMP(s)?)	Current	VICH Negotiated	All Formats
85	Duration and Time Units (This is the length of time the AE lasted.)	Current	VICH Negotiated	All Formats
86	Serious AE (Was the AE serious?)	Current	VICH Negotiated	All Formats
87	Treatment of AE (Was the AE treated?)	Current	VICH Negotiated	All Formats
88	Outcome to Date, including: (number of)			
89	Recovered/Normal, Ongoing, Recovered with Sequela, and Unknown	Current	VICH Negotiated	All Formats
90	Euthanized	New	VICH Negotiated	All Formats
91	Died	Current	VICH Negotiated	All Formats
92	Length of Time Between Exposure to VMP(s) and Onset of AE (This is a list of values describing the length of time between the first exposure to the VMP and the onset of the AE.)	Current	VICH Negotiated	All Formats
93	Date of Onset of AE (Day, Month, Year) (This is the date of the first clinical manifestation of the AE.)	Current	VICH Negotiated	All Formats
94	Adverse Clinical Manifestations (This is a list of values describing the clinical signs that occurred during the AE.)	Current	VICH Negotiated	All Formats
95	Narrative of AE (open text field) (This is a detailed description of the case, regardless of the type of information contained in the report.)	Current	VICH Negotiated	All Formats
96	Did the AE Abate After Stopping the VMP?	Current	VICH Negotiated	All Formats
97	Did the AE Reappear After Re-Introduction of the VMP?	Current	VICH Negotiated	All Formats
98	Animal Data, including:			
99	Species (This is a list of values describing the species of the animal(s) involved in the AER.)	Current	VICH Negotiated	All Formats

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932—Continued

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
100	Breeds and Crossbreed Information (This is a list of values describing the breed(s) of animal(s) involved in the AER.)	Current	VICH Negotiated	All Formats
101	Gender (This is a list of values for the selection of the gender(s) of animal(s) involved in the AER.)	Current	VICH Negotiated	All Formats
102	Reproductive Status (This is a list of values describing if the animal is intact, neutered, etc.)	Current	VICH Negotiated	All Formats
103	Female Physiological Status. (This is a list of values describing the animal's pregnancy and lactation status.)	Current	VICH Negotiated	All Formats
104	Age (Measured, Estimated, Unknown), including:			
105	Precision Value for Age (Measured, Estimated, Unknown Age. This is a list of values describing whether the age(s) provided are measured or estimated, or if age is not known.)	New	VICH Negotiated	All Formats
106	Minimum Age Value and Units.	Current	VICH Negotiated	All Formats
107	Maximum Age Value and Units.	Current	VICH Negotiated	All Formats
108	Weight, including:			
109	Precision Value for Weight (Measured, Estimated, Unknown Weights) (This is a list of values describing whether the weight(s) provided are measured or estimated, or if weight is not known.)	New	VICH Negotiated	All Formats
110	Minimum Weight	Current	VICH Negotiated	All Formats
111	Maximum Weight	Current	VICH Negotiated	All Formats
112	Attending Veterinarian's Assessment of Animal Health Status Prior to VMP. (This is a list of values describing the attending veterinarian's assessment of the health status of the animal(s) involved in the AE prior to their exposure to the VMP.)	Current	VICH Negotiated	All Formats
113	Number of Animals Treated (This is the number of animal(s) being directly treated by the VMP(s).)	Current	VICH Negotiated	All Formats
114	Number of Animals Affected (This is the total number of animals affected in the AER, whether by direct or indirect exposure.)	Current	VICH Negotiated	All Formats
115	Supplemental Documents, including:			
116	Attached Document (These are additional documents containing information relevant to the AE, such as medical record, radiology, clinical chemistry reports, newspaper articles, and letters.)	Current	VICH Negotiated	All Formats
117	Attached Document Filename (This is the name of the document for paper documents or the electronic file for electronic transmissions.)	Current	VICH Negotiated	All Formats
118	Attached Document Type (This is a list of values describing the type of document that is attached, e.g., necropsy report)	Current	VICH Negotiated	All Formats
119	The following data elements are being deleted from the information collection:			
120	2c. Number of Days Between 2a and b:	Deleted		
121	11. Illness/reason for use of this drug	Deleted		

TABLE 3.—DATA ELEMENTS FOR FORM FDA 1932—Continued

Line No.	Data Elements	Current, New, or Deleted Data Element	VICH-Negotiated or FDA-Proposed Data Element	Submission/Transmission Format (Paper Form, Electronic Web-based Rational Questionnaire (EWBRQ), and/or Electronic Gateway-to-Gateway (EGG))
122	17. Did any new illness develop or did initial diagnosis change after suspect drug started?	Deleted		
123	25. Outcome of Reaction to Date - Died	Deleted		
124	26. When reaction appeared, treatment with suspect drug: has already been completed, discontinued, replaced with another drug; continued at altered dose, other (explain)—and the reaction: continued, stopped, recurred, or other (explain)	Deleted		
125	29. Had animal(s) previously reacted to other drugs?	Deleted		
126	30. Has the attending veterinarian seen similar reactions to this drug in any other animals?	Deleted		
127	32. Signature of individual responsible for accuracy of reported information	Deleted		

2. Form FDA 1932a

This section describes data elements on the current Form FDA 1932a and the proposed new data elements. These AE and product and manufacturing defect data elements will be collected electronically, through the MedWatch^{Plus} Portal Rational Questionnaire, and in the paper form. All the data elements will be captured

using the MedWatch^{Plus} Portal Rational Questionnaire or the paper form.

Table 4 of this document, entitled “Data Element Information Collection for Form FDA 1932a,” presents the data elements the agency is proposing for the collection of animal drug adverse events reports and manufacturing and product problem reports for individuals who choose to report information voluntarily to FDA. The current and proposed new

data elements are listed in the column entitled “Data Elements.” In general, the information being collected is the same, but the data element has been renamed or restructured to facilitate data collection. As stated previously in this document, the proposed changes are based on FDA’s experience in determining the safety and effectiveness of product(s) and need for efficient data capture and entry.

TABLE 4.—DATA ELEMENT INFORMATION COLLECTION FOR FORM FDA 1932A

Line No.	Data Elements	Current or New Data Element
1	Individual Case Safety Report Number (FDA-Assigned Number)	New
2	Date of Initial Report. (This is the date the sender sent the first report of the information.)	New
3	Date Reported (This is the date of this current report.)	Current
4	Submission Type (This is a list of values describing the type of submission, e.g., Initial or Followup Report)	New
5	Report Type (This is a list of values describing the type of information in the report, e.g., adverse event, product problem, or both)	New
6	Manufacturer’s Case Number. (The manufacturer’s case number is given to the sender by the manufacturer of the product if the sender contacted the manufacturer.)	Current
7	Sender Information (The sender is the person or organization which fills out the report and submits or transmits the report to FDA.), including:	
8	Sender First and Last Name	New
9	Sender Street Address, City, State/Province, Postal/Zip Code, and Country	New
10	Sender Primary and Other Telephone Number, E-Mail Address, and Fax Number	New
11	Sender Category. (This is a list of values describing the role/involvement of the sender, e.g., animal owner, physician, etc.)	New

TABLE 4.—DATA ELEMENT INFORMATION COLLECTION FOR FORM FDA 1932A—Continued

Line No.	Data Elements	Current or New Data Element
12	Did the sender report to other sources?	New
13	Sender also reported to other sources. (This is a list of values describing the sources to which the sender reported the AE or product problem, e.g., manufacturer, distributor, etc.)	New
14	No identity disclosure (This data element indicates whether the sender wants their identity disclosed to the manufacturer.)	Current
15	Preferred Method of Contact. (This is a list of values describing the preferred method of contacting the sender, e.g., telephone, e-mail.)	New
16	Healthcare Professional Information, including:	
17	Healthcare Professional First and Last Name.	Current
18	Healthcare Professional Street Address, City, State/Province, and Postal/Zip Code	Current
19	Healthcare Professional Primary and Other Phone Number	Current
20	Healthcare Professional e-mail address	New
21	Healthcare Professional Country	New
22	Owner's Information (This is the owner of the animal involved in the case.), including:	
23	Owner First and Last Name.	Current
24	Owner Primary and Other Phone Number, and E-Mail Address	New
25	Owner Street Address, City, State/Province, Postal/Zip Code, and Country	New
26	Product Information:	
27	Name of Suspected Product. (This is the name of the product suspected of causing the AE or the product with the product problem.)	Current
28	Name of Manufacturer	Current
29	Lot Number	Current
30	Expiration Date	Current
31	Diagnosis and/or Reason for Use of the Product	Current
32	Product Use Information: Dose Administered (amount of product administered), Interval of Administration (frequency of administration—every 12 hours or for 5 days), and Route of Administration (oral, injection, topical, etc.).	Current
33	Dosage Form. (This is how the product was supplied to the animal, e.g., chewable tablet, topical, injection)	Current
34	Date of First and Last Exposure. (This is the date the product(s) was first administered and last administered to the animal.)	Current
35	Duration of Product Use (Number) and Units of Measurement. (This is the duration the product was given, e.g., 2 weeks.)	New
36	Product Administered By (This is a list of values describing who administered the product(s), e.g., veterinarian/veterinary staff, Owner)	Current
37	Concurrent Drugs Administered (Were concurrent product(s) given to the animal(s)?)	Current
38	Concurrent Products Names. (This is the name of all concurrent products involved in the case.)	Current
39	Animal Information:	
40	Species. (This is a list of values for selecting the species of the animal(s) involved in the case.)	Current
41	Breed and Crossbreed (This is the breed(s) of animal(s) involved in the report.)	Current
42	Gender. (This a list of values for the selection of the gender(s) of animal(s) involved in the AER.)	Current

TABLE 4.—DATA ELEMENT INFORMATION COLLECTION FOR FORM FDA 1932A—Continued

Line No.	Data Elements	Current or New Data Element
43	Reproductive Status. (This is a list of values describing whether the animal is intact, neutered, et cetera.)	Current
44	Age and Age Units	Current
45	Weight and Weight Units	Current
46	Overall Health Status When Suspected Product Given. (This is a list of values describing the health status of the animal(s) involved in the AE prior to their exposure to the product(s).)	Current
47	Number of Animals Treated (This is the number of animal(s) being directly treated by the product(s).)	New
48	Number of Animals Affected. (This is the total number of animals affected in the AER, whether by direct or indirect exposure.)	New
49	Adverse Event Information:	
50	Veterinarian's Level of Suspicion that Product Caused the AE. (This is a list of values describing the veterinarian's level of suspicion, e.g., high, medium, low, or unknown.)	Current
51	Treatment of AE. (This is a description of how the AE was treated.)	Current
52	Did the AE Abate After Stopping the Product?	Current
53	Did the AE Reappear After Reintroduction of the product?	Current
54	Outcome. (This is a list of values describing the overall animal health status after exposure to the product.)	Current
55	Length of Time Between Initial Exposure to Suspected Product and Onset of AE, numeric value and units of measurement	Current
56	Length of Time Between Last Administration of Suspected Product and Onset of AE, numeric value and units of measurement	Current
57	Date of Onset of AE. (This is the date that the first adverse clinical sign(s) occurred.)	New
58	Date of Product Problem Discovery. (This is the date that the product problem was discovered.)	New
59	When the AE Occurred, Treatment with Suspected Product. (This is a list of values describing the use of the suspected product after the AE occurred)	Current
60	Other Relevant Clinical Information:	
61	Concurrent Clinical Problem (Does the animal(s) have concurrent clinical problems?)	Current
62	List Concurrent Clinical Problem(s)	Current
63	AE/Product Problem (Long Narrative) (This is a detailed description of the case.)	Current
64	Supplemental Documents:	
65	Attached Document Name/File name (if electronic) (This is the name of the document for paper documents or the name of the electronic file for electronic transmissions.)	Current
66	Attached Document Type (This is a list of values describing the type of document that is attached, e.g., necropsy report)	Current
67	Attached Document(s) (These are additional documents containing information relevant to the AE, e.g., medical record, radiology, clinical chemistry reports, newspaper articles, and letters.)	Current
68	Attached Document Description. (This is a description of the document.)	New

C. Request for Comments

FDA invites comments on all aspects of the revised collection of the data elements for Forms FDA 1932 and 1932a as set forth in section III.B of this notice, including whether such lists

incorporate all data elements necessary to report an adverse event and a product or manufacturing defect, and whether certain data elements should be deleted or modified. Interested persons may submit to the Division of Dockets Management (see **ADDRESSES**) electronic

or written comments regarding the proposed changes. Submit a single copy of electronic comments or two paper copies of any mailed comments, except that individuals may submit one paper copy. Comments are to be identified with the docket number found in

brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

Dated: November 16, 2009.

David Horowitz,

Assistant Commissioner for Policy.

[FR Doc. E9-27956 Filed 11-19-09; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Government-Owned Inventions; Availability for Licensing

AGENCY: National Institutes of Health, Public Health Service, HHS.

ACTION: Notice.

SUMMARY: The inventions listed below are owned by an agency of the U.S. Government and are available for licensing in the U.S. in accordance with 35 U.S.C. 207 to achieve expeditious commercialization of results of Federally-funded research and development. Foreign patent applications are filed on selected inventions to extend market coverage for companies and may also be available for licensing.

ADDRESSES: Licensing information and copies of the U.S. patent applications listed below may be obtained by writing to the indicated licensing contact at the Office of Technology Transfer, National Institutes of Health, 6011 Executive Boulevard, Suite 325, Rockville, Maryland 20852-3804; *telephone:* 301/496-7057; *fax:* 301/402-0220. A signed Confidential Disclosure Agreement will be required to receive copies of the patent applications.

Phage Display Plasmids With Improved Expression Properties for Human and Chimeric Nonhuman/Human Fab Libraries

Description of Invention: The Fab molecule was the first generated antibody fragment and still dominates basic research and clinical applications. New phage display vectors were designed to generate and select Fab libraries with human constant domains. These vectors facilitate bacterial expression of human, humanized, and chimeric nonhuman/human Fab antibody fragments. They differ from currently available pComb3H and pComb3X phage display vectors by assembling human and chimeric nonhuman/human Fab libraries in two rather than three PCR steps. As a result,

these novel constructs retain the initial variable light and heavy chain sequences and improve the resulting Fab library's complexity in terms of number, diversity, and affinity. These constructs were developed with and without a His tag and yield approximately 100 µg to 2 mg of protein, which can be used for evaluation and characterization of Fab binding properties such as affinity and specificity. Notably, the His tag provides a handle to easily purify Fab.

Applications

- Generation of human, humanized, and chimeric nonhuman/human Fab antibody fragments.
- Research tool to characterize Fab antibody fragments.

Advantages

- Improved Fab library with complexity and number, diversity, and affinity.
 - His tag construct allows for simplified purification assays.
- Inventor:* Christoph Rader (NCI).

Relevant Publications

1. KY Kwong and C Rader. E. coli expression and purification of Fab antibody fragments. *Curr Protoc Protein Sci.* 2009 Feb;Chapter 6:Unit 6.10.
2. T Hofer *et al.* Chimeric rabbit/human Fab and IgG specific for members of the Nogo-66 receptor family selected for species cross-reactivity with an improved phage display vector. *J Immunol Methods.* 2007 Jan 10;318(1-2):75-87.

Patent Status: HHS Reference No. E-008-2010/0—Research Tool. Patent protection is not being pursued for this technology.

Licensing Status: Available for licensing.

Licensing Contact: Jennifer Wong; 301-435-4633; wongje@mail.nih.gov.

Potent and Selective Inhibitors of Human Lipoxygenase for Prostate Cancer Therapy

Description of Invention: With more than \$2 billion in revenues in the US in 2007, the market for diagnostic and therapeutic products for prostate cancer is substantial. More than 2,000,000 American men currently live with prostate cancer and more than 200,000 new cases are diagnosed each year.

Researchers led by Dr. David Maloney at the National Human Genome Research Institute (NHGRI) have discovered several novel compounds that selectively and potently inhibit lipoxygenase (LOX), an enzyme that metabolizes polyunsaturated fatty acids which has been implicated in the

pathogenesis of prostate cancers. These novel compounds are small molecules, and as such have an advantage over antibody-based technologies in this market. As prostate cancer is the most commonly diagnosed malignancy among men in the USA and Europe, the significant need for new therapies suggests that these novel LOX inhibitor compounds have a strong potential of reaching the marketplace.

Applications

- Therapeutics for prostate cancer.
- Therapeutics for several other LOX-associated pathologies including atherosclerosis, asthma, other cancers, glomerulonephritis, osteoporosis, and Alzheimer's disease.

Advantages

- Potent and selective inhibitory activity to reduce negative side effects.
- Compounds are small molecules (less immunogenic than antibodies).

Development Status: Pre-clinical.

Inventors: David Maloney *et al.* (NHGRI).

Relevant Publications

1. V Kenyon *et al.* Novel human lipoxygenase inhibitors discovered using virtual screening with homology models. *J Med Chem.* 2006 Feb 23;49(4):1356-1363.

2. JD Deschamps *et al.* Baicalein is a potent in vitro inhibitor against both reticulocyte 15-human and platelet 12-human lipoxygenases. *Bioorg Med Chem.* 2006 Jun 15;14(12):4295-4301.

3. Y Vasquez-Martinez *et al.* Structure-activity relationship studies of flavonoids as potent inhibitors of human platelet 12-hLO, reticulocyte 15-hLO-1, and prostate epithelial 15-hLO-2. *Bioorg Med Chem.* 2007 Dec 1;15(23):7408-7425.

4. J Inglese *et al.* Quantitative high-throughput screening: a titration-based approach that efficiently identifies biological activities in large chemical libraries. *Proc Natl Acad Sci USA.* 2006 Aug 1;103(31): 11473-11478.

Patent Status: U.S. Provisional Application No. 61/238,972 filed 01 Sep 2009 (HHS Reference No. E-252-2009/0-US-01).

Licensing Status: Available for licensing.

Licensing Contact: Patrick P. McCue, Ph.D.; 301-435-5560; mccuepat@mail.nih.gov.

Collaborative Research Opportunity: The NIH Chemical Genomics Center, NHGRI, is seeking statements of capability or interest from parties interested in collaborative research to further develop, evaluate, or commercialize this technology. Please

contact Claire Driscoll at cdriscoll@mail.nih.gov or 301-594-2235 for more information.

Dated: November 13, 2009.

Richard U. Rodriguez,

Director, Division of Technology Development and Transfer, Office of Technology Transfer, National Institutes of Health.

[FR Doc. E9-27925 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2008-D-0614]

Guidance for Industry on Changes to Approved New Animal Drug Applications—New Animal Drug Applications Versus Category II Supplemental New Animal Drug Applications; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a guidance for industry #191 entitled “Changes to Approved NADAs—New NADAs vs. Category II Supplemental NADAs.” This guidance is intended to assist sponsors who wish to apply for approval of changes to approved new animal drugs that require FDA to reevaluate safety and/or effectiveness data. The goal of this guidance is to create greater consistency in how such applications are handled by sponsors and by FDA’s Center for Veterinary Medicine (CVM).

DATES: Submit written or electronic comments on agency guidances at any time.

ADDRESSES: Submit written requests for single copies of the guidance to the Communications Staff (HFV-12), Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855. Send one self-addressed adhesive label to assist that office in processing your requests.

Submit written comments on the guidance to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to <http://www.regulations.gov>. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the guidance document.

FOR FURTHER INFORMATION CONTACT: Suzanne J. Sechen, Center for Veterinary Medicine (HFV-126), Food and Drug

Administration, 7500 Standish Pl., Rockville, MD 20855, 240-276-8105, e-mail: suzanne.sechen@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is announcing the availability of a guidance for industry #191 entitled “Changes to Approved NADAs—New NADAs vs. Category II Supplemental NADAs.” This guidance is intended to assist sponsors who wish to apply for approval of changes to approved new animal drugs that require FDA to reevaluate safety and/or effectiveness data. The guidance explains how the Office of New Animal Drug Evaluation (ONADE) categorizes possible changes to approved new animal drugs that require reevaluation of safety and/or effectiveness data and explains which administrative vehicle—a new original new animal drug application (NADA) (new NADA) or a Category II supplemental application to the original new animal drug application (Category II supplemental NADA)—a sponsor should use when applying for approval of these changes. The goal of this guidance is to create greater consistency in how such applications are handled by sponsors and by ONADE.

In the **Federal Register** of December 16, 2008 (73 FR 76363), FDA published the notice of availability for a draft guidance entitled “Changes to Approved NADAs—New NADAs vs. Category II Supplemental NADAs,” which gave interested persons until February 17, 2009, to comment on the draft guidance. FDA received a few comments on the draft guidance and those comments were considered as the guidance was finalized. In addition to some of the changes based on the comments received, CVM made a few minor changes to the guidance to add clarity and accuracy. The guidance announced in this notice finalizes the draft guidance dated December 16, 2008.

II. Significance of Guidance

This level 1 guidance is being issued consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The guidance represents the agency’s current thinking on the topic. It does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirements of the applicable statutes and regulations.

III. Paperwork Reduction Act of 1995

This guidance refers to previously approved collections of information found in FDA regulations. These

collections of information are subject to review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501–3520). The collections of information have been approved under OMB control no. 0910-0032 (expiration date April 30, 2010).

IV. Comments

Submit written requests for single copies of the guidance to the Communications Staff (HFV-12), Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855. Send one self-addressed adhesive label to assist that office in processing your requests.

Interested persons may submit to the Division of Dockets Management (see **ADDRESSES**) written or electronic comments regarding this document. Submit a single copy of electronic comments or two paper copies of any mailed comments, except that individuals may submit one paper copy. Comments are to be identified with the docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

V. Electronic Access

Persons with access to the Internet may obtain the guidance at either <http://www.fda.gov/AnimalVeterinary/default.htm> or <http://www.regulations.gov>.

Dated: November 13, 2009.

David Horowitz,

Assistant Commissioner for Policy.

[FR Doc. E9-27926 Filed 11-19-09; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute on Aging; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The contract proposals and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the contract proposals, the disclosure of which

would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institutes on Aging, Special Emphasis Panel Exceptional Aging.

Date: December 10, 2009.

Time: 9 a.m. to 12 p.m.

Agenda: To review and evaluate contract proposals.

Place: National Institutes of Health, Gateway Building, 7201 Wisconsin Avenue 2C212, Bethesda, MD 20982 (Telephone Conference Call).

Contact Person: Alicja L. Markowska, PhD, DSC, Scientific Review Officer, Scientific Review Branch, National Institute on Aging, 7201 Wisconsin Avenue, Suite 2C212, Bethesda, MD 20892, 301-496-9666, markowska@nia.nih.gov.

Name of Committee: National Institute on Aging, Special Emphasis Panel SYNTHESIS.

Date: December 14, 2009.

Time: 4:30 p.m. to 6 p.m.

Agenda: To review and evaluate contract proposals.

Place: National Institutes of Health, Gateway Building, 7201 Wisconsin Avenue 2C212, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Ramesh Vemuri, PhD, Chief, Scientific Review Office, National Institute on Aging, National Institutes of Health, 7201 Wisconsin Avenue, Bethesda, MD 20892, 301-402-7700, rv23r@nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.866, Aging Research, National Institutes of Health, HHS)

Dated: November 13, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-27800 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute on Drug Abuse; Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute on Drug Abuse Special Emphasis Panel; Conference Grant Application Review.

Date: December 3, 2009.

Time: 10 a.m. to 1 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6101 Executive Boulevard, Rockville, MD 20852, (Virtual Meeting).

Contact Person: Gerald L. McLaughlin, PhD, Scientific Review Administrator, Office of Extramural Affairs, National Institute on Drug Abuse, NIH, DHHS, Room 220, MSC 8401, 6101 Executive Blvd., Bethesda, MD 20892-8401, 301-402-6626, gm145a@nih.gov.

(Catalogue of Federal Domestic Assistance Program No. 93279, Drug Abuse and Addiction Research Programs, National Institutes of Health, HHS)

Dated: November 12, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-27727 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of General Medical Sciences; Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute of General Medical Sciences Special Emphasis Panel; Minority Biomedical Research Support.

Date: December 3, 2009.

Time: 9:30 a.m. to 10:30 a.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, Natcher Building, Room 3AN12, 45 Center Drive, Bethesda, MD 20892, (Telephone Conference Call).

Contact Person: Helen R. Sunshine, PhD, Chief, Office of Scientific Review, National Institute of General Medical Sciences, National Institutes of Health, Natcher Building, Room 3AN12F, Bethesda, MD 20892, 301-594-2881, sunshinh@nigms.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.375, Minority Biomedical Research Support; 93.821, Cell Biology and Biophysics Research; 93.859, Pharmacology, Physiology, and Biological Chemistry Research; 93.862, Genetics and Developmental Biology Research; 93.88, Minority Access to Research Careers; 93.96, Special Minority Initiatives, National Institutes of Health, HHS)

Dated: November 12, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-27732 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Center for Scientific Review Special Emphasis Panel; ZRG1 IFCN-C (02) M Member Conflicts: Cognition and Neurotoxicology.

Date: December 10, 2009.

Time: 1:30 p.m. to 4 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892, (Telephone Conference Call).

Contact Person: Michael Selmanoff, PhD, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 3134, MSC 7844, Bethesda, MD 20892, 301-435-1119, mselemanoff@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; RFA-OD-09-008 BRDG-SPAN and RFA-OD-09-009 Catalyst ARRA Review Panel 17.

Date: December 10, 2009.

Time: 2 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892, (Telephone Conference Call).

Contact Person: Lawrence E. Boerboom, PhD, Scientific Review Officer, Center for

Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 5156, MSC 7814, Bethesda, MD 20892, (301) 435-8367, boerboom@nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Fellowships: Cell Biology Specials.

Date: December 10, 2009.

Time: 2:30 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892, (Telephone Conference Call).

Contact Person: Alessandra M. Bini, PhD, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 5142, MSC 7840, Bethesda, MD 20892, 301-435-1024, binia@csr.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.306, Comparative Medicine; 93.333, Clinical Research, 93.306, 93.333, 93.337, 93.393-93.396, 93.837-93.844, 93.846-93.878, 93.892, 93.893, National Institutes of Health, HHS)

Dated: November 16, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-27952 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Center for Scientific Review Special Emphasis Panel, Member Conflict Panel: Population Sciences and Epidemiology IRG.

Date: December 9, 2009.

Time: 12 p.m. to 2 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Fungai Chanetsa, MPH, PhD, Scientific Review Officer, Center for

Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 3135, MSC 7770, Bethesda, MD 20892. 301-435-1262. fungai.chanetsa@nih.hhs.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel, BDCN Member Conflict Special Emphasis Panel.

Date: December 11, 2009.

Time: 2 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Samuel C. Edwards, PhD, Chief, BDCN IRG, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 4200, MSC 7812, Bethesda, MD 20892. (301) 435-1152. edwardss@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel, Musculoskeletal Engineering.

Date: December 11, 2009.

Time: 5 p.m. to 7 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Jo Pelham, BA, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 4102, MSC 7814, Bethesda, MD 20892. (301) 435-1786. pelhamj@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel, Topics in Microbiology.

Date: December 15, 2009.

Time: 1 p.m. to 2:30 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Liangbiao Zheng, PhD, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 3214, MSC 7808, Bethesda, MD 20892. 301-402-5671. zhengli@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel, Shared Instrumentation.

Date: December 15-16, 2009.

Time: 3 p.m. to 6 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Virtual Meeting).

Contact Person: Michael A. Marino, PhD, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 2216, MSC 7890, Bethesda, MD 20892. (301) 435-0601. marinomi@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel, LCMI Member Conflict Applications.

Date: December 16, 2009.

Time: 3 p.m. to 4:30 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Telephone Conference Call).

Contact Person: Everett E. Sinnett, PhD, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 2178, MSC 7818, Bethesda, MD 20892. 301-435-1016. sinnett@nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.306, Comparative Medicine; 93.333, Clinical Research, 93.306, 93.333, 93.337, 93.393-93.396, 93.837-93.844, 93.846-93.878, 93.892, 93.893, National Institutes of Health, HHS)

Dated: November 16, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-27950 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Cancer Institute; Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Cancer Institute Initial Review Group; Subcommittee J—Population and Patient-Oriented Training.

Date: February 10, 2010.

Time: 7:45 a.m. to 6 p.m.

Agenda: To review and evaluate grant applications.

Place: Hilton Alexandria Old Town, 1767 King Street, Alexandria, VA 22314.

Contact Person: Ilda M. McKenna, PhD, Scientific Review Officer, Research Training Review Branch, Division of Extramural Activities, National Cancer Institute, 6116 Executive Boulevard, Room 8111, Bethesda, MD 20892, 301-496-7481, mckennai@mail.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.392, Cancer Construction; 93.393, Cancer Cause and Prevention Research; 93.394, Cancer Detection and Diagnosis Research; 93.395, Cancer Treatment Research; 93.396, Cancer Biology Research; 93.397, Cancer Centers Support; 93.398, Cancer Research Manpower; 93.399,

Cancer Control, National Institutes of Health, HHS)

Dated: November 16, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-27949 Filed 11-19-09; 8:45 am]

BILLING CODE 4140-01-P

ADVISORY COUNCIL ON HISTORIC PRESERVATION

Notice of Program Comment for the Rural Utilities Service, the National Telecommunications and Information Administration, and the Federal Emergency Management Agency To Avoid Duplicative Section 106 Reviews for Wireless Communication Facilities Construction and Modification

AGENCY: Advisory Council on Historic Preservation.

ACTION: The Advisory Council on Historic Preservation has issued a Program Comment for the Rural Utilities Service, the National Telecommunications and Information Administration, and the Federal Emergency Management Agency to avoid duplicative Section 106 reviews for wireless communication facilities construction and modification.

SUMMARY: The Advisory Council on Historic Preservation has issued a Program Comment for the Rural Utilities Service, the National Telecommunications and Information Administration, and the Federal Emergency Management Agency to relieve them of the need to conduct duplicate reviews under Section 106 of the National Historic Preservation Act when these agencies assist a telecommunications project that is exempt from, or subject to, Section 106 review by the Federal Communications Commission under existing nationwide programmatic Agreements.

DATES: The Program Comment went into effect on October 23, 2009.

ADDRESSES: Address all questions concerning the Program Comment to Blythe Semmer, Office of Federal Agency Programs, Advisory Council on Historic Preservation, 1100 Pennsylvania Avenue, NW., Suite 803, Washington, DC 20004. Fax (202) 606-8647. You may submit electronic questions to: bsemmer@achp.gov.

FOR FURTHER INFORMATION CONTACT: Blythe Semmer, (202) 606 8552, bsemmer@achp.gov.

SUPPLEMENTARY INFORMATION: Section 106 of the National Historic Preservation Act requires Federal

agencies to consider the effects of their undertakings on historic properties and to provide the Advisory Council on Historic Preservation (ACHP) a reasonable opportunity to comment with regard to such undertakings. The ACHP has issued the regulations that set forth the process through which Federal agencies comply with these duties. Those regulations are codified under 36 CFR part 800 (Section 106 regulations)

Under Section 800.14(e) of those regulations, agencies can request the ACHP to provide a "Program Comment" on a particular category of undertakings in lieu of conducting individual reviews of each individual undertaking under such category, as set forth in 36 CFR 800.3 through 800.7. An agency can meet its Section 106 responsibilities with regard to the effects of particular aspects of those undertakings by taking into account ACHP's Program Comment and following the steps set forth in that comment.

I. Background

The ACHP has issued a Program Comment to the U.S. Department of Agriculture Rural Utilities Service (RUS), the U.S. Department of Commerce National Telecommunications and Information Administration (NTIA), and the Federal Emergency Management Agency (FEMA) to relieve them from conducting duplicate reviews under Section 106 of the National Historic Preservation Act when these agencies assist a telecommunications project subject to Section 106 review by the Federal Communications Commission (FCC). The ACHP membership voted in favor of issuing the Program Comment via an unassembled vote on October 23, 2009.

The American Recovery and Reinvestment Act (ARRA) provides NTIA and RUS with \$7.2 billion to expand access to broadband services in the United States. NTIA will implement the Broadband Technology Opportunities Program (BTOP), which will award grants to expand public computer capacity, encourage sustainable adoption of broadband of broadband service, and deploy broadband infrastructure to unserved and underserved areas. RUS, through its Broadband Initiatives Program (BIP), will use loan and grant combinations to support broadband deployment in rural areas.

Broadband deployment can include the construction and placement of communication towers and antennas. Some of those towers and antennas are also regulated by the FCC, and therefore undergo, or are exempted from, Section 106 review under the Nationwide

Programmatic Agreement for Review of Effect on Historic Properties for Certain Undertakings Approved by the FCC and the Nationwide Programmatic Agreement for the Collocation of Wireless Antennas (FCC NPAs). RUS, NTIA, or FEMA will be relieved by the Program Comment of the need to conduct a separate Section 106 review for undertakings subject to review under the FCC NPAs.

The ACHP took steps to inform the public and stakeholders about the proposed Program Comment, including an e-mail distribution, posting on the agency Web site, and a notice published in the **Federal Register**. ACHP also sent a letter to the Indian tribal leaders requesting their comments on the Program Comment. Public comments resulting from the September 17, 2009 public notice in the **Federal Register** (74 FR 47807-47809) were received by the ACHP by October 8, 2009.

Various substantive comments from the public were received and considered by the ACHP, as noted below.

FEMA requested inclusion in the provisions of the Program Comment given that its grant programs provide funding for emergency communications facilities that are also subject to review by FCC under the FCC NPAs. FEMA's request would not expand the types of undertakings covered by the Program Comment, so FEMA has been added to the Program Comment.

Two comments objected to how tribal consultation appeared to have been coordinated for the Program Comment, but the characterization of early coordination with intertribal organizations by RUS and NTIA prior to the agencies' formal request to the ACHP did not constitute ACHP's tribal consultation on this program alternative.

Two comments expressed concern about how State and Tribal Historic Preservation Officers (SHPOs and THPOs) and Indian tribes will be notified when the Program Comment is applied. SHPOs and THPOs and Indian tribes will be notified according to the regular FCC NPAs review processes. There is no change to the FCC NPAs procedures.

Two comments expressed objections or concerns about the FCC NPAs and two comments expressed positive views on the functioning of the FCC NPAs. Nothing in the Program Comment will alter the FCC NPAs, but these comments will be referred to FCC for their consideration on the operation of their NPAs. One comment expressed concerns about towers that may have been constructed before undergoing a Section 106 review. The Program

Comment deals with the construction of towers and collocation on existing towers. It does not address or affect pre-existing Section 106 issues. Those issues should be referred to the FCC.

Four comments expressed support for the efficiencies the Program Comment will offer in Section 106 reviews.

Two comments offered views on a concept plan for a nationwide programmatic agreement circulated separately by RUS and NTIA. Those comments will be considered in the context of that program initiative.

The Colorado Historical Society requested clarification about the 6-year term of the Program Comment. This time period recognizes that ARRA-assisted communications facilities construction may be ongoing for several years. The ACHP and others will be able to reevaluate the Program Comment, and whether to extend its duration prior to the conclusion of those 6 years.

The Texas Historical Commission questioned what would happen should an FCC NPA Section 106 review yield a finding of adverse effect within a larger RUS or NTIA undertaking of multiple components. As explicitly stated in the Program Comment, RUS, NTIA, or FEMA will be conducting its own Section 106 review for the larger undertaking, but will not have to consider the effects of the FCC-regulated component of that larger undertaking. RUS, NTIA, or FEMA will make effect determinations based on the non-tower components of the undertaking. Since it is possible that the larger undertaking may not be able to proceed until the FCC review of the tower component has concluded, it is expected that RUS, NTIA, FEMA, and the FCC will coordinate their review efforts accordingly and keep consulting parties appraised.

II. Final Text of the Program Comment

The text of the issued Program Comment is included below:

Program Comment for Streamlining Section 106 Review for Wireless Communication Facilities Construction and Modification Subject to Review Under the FCC Nationwide Programmatic Agreement and/or the Nationwide Programmatic Agreement for the Collocation of Wireless Antennas.

I. Background

The Rural Utilities Service (RUS), the National Telecommunications and Information Administration (NTIA), and the Federal Emergency Management Agency (FEMA) provide financial assistance to applicants for various undertakings, including broadband deployment, which can involve the construction and placement of communications towers and antennas. RUS,

NTIA, and FEMA must therefore comply with Section 106 of the National Historic Preservation Act, 16 U.S.C. 470f, and its implementing regulations at 36 CFR part 800 (Section 106) for these undertakings. Some of those communications towers and antennas are also regulated by the Federal Communications Commission (FCC), and therefore undergo, or are exempted from, Section 106 review under the Nationwide Programmatic Agreement for Review of Effects on Historic Properties for Certain Undertakings Approved by the FCC (FCC Nationwide PA) and the Nationwide Programmatic Agreement for the Collocation of Wireless Antennas (FCC Collocation PA). The FCC Nationwide PA was executed by the FCC, the Advisory Council on Historic Preservation (ACHP), and the National Conference of State Historic Preservation Officers (NCSHPO) on October 4, 2004. The FCC Collocation PA was executed by the FCC, ACHP, and NCSHPO on March 16, 2001. The undertakings addressed by the FCC Nationwide PA primarily include the construction and modification of communication towers. The undertakings addressed by the FCC Collocation PA include the collocation of communications equipment on existing structures and towers.

This Program Comment is intended to streamline Section 106 review of the construction and modification of communication towers and antennas for which FCC and RUS, NTIA, or FEMA share Section 106 responsibility.

Nothing in this Program Comment alters or modifies the FCC Nationwide PA or the FCC Collocation PA, or imposes Section 106 responsibilities on the FCC for elements of a RUS, NTIA, or FEMA undertaking that are unrelated to a communications facility within the FCC's jurisdiction or are beyond the scope of the FCC Nationwide PA.

II. Establishment and Authority

This Program Comment was issued by the ACHP on October 23, 2009 pursuant to 36 CFR 800.14(e).

III. Date of Effect

This Program Comment went into effect on October 23, 2009.

IV. Use of This Program Comment To Comply With Section 106 for the Effects of Facilities Construction or Modification Reviewed Under the FCC Nationwide PA and/or the FCC Collocation PA

RUS, NTIA and FEMA will not need to comply with Section 106 with regard to the effects of communication facilities construction or modification that has either undergone or will undergo Section 106 review, or is exempt from Section 106 review, by the FCC under the FCC Nationwide PA and/or the FCC Collocation PA. For purposes of this program comment, review under the FCC Nationwide PA means the historic preservation review that is necessary to complete the FCC's Section 106 responsibility for an undertaking that is subject to the FCC Nationwide PA.

When an RUS, NTIA, or FEMA undertaking includes both communications facilities construction or modification covered by the FCC Nationwide PA or

Collocation PA and components in addition to such communication facilities construction or modification, RUS, NTIA, or FEMA, as applicable, will comply with Section 106 in accordance with the process set forth at 36 CFR 800.3 through 800.7, or 36 CFR 800.8(c), or another applicable alternate procedure under 36 CFR 800.14, but will not have to consider the effects of the communication facilities construction or modification component of the undertaking on historic properties. Whenever RUS, NTIA, or FEMA uses this Program Comment for such undertakings, RUS, NTIA or FEMA will apprise the relevant State Historic Preservation Officer (SHPO) or Tribal Historic Preservation Officer (THPO) of the use of this Program Comment for the relevant communication facilities construction or modification component.

V. Amendment

The ACHP may amend this Program Comment after consulting with FCC, RUS, NTIA, FEMA, and other parties as appropriate, and publishing notice in the **Federal Register** to that effect.

VI. Sunset Clause

This Program Comment will terminate on September 30, 2015, unless it is amended to extend the period in which it is in effect.

VII. Termination

The ACHP may terminate this Program Comment by publication of a notice in the **Federal Register** thirty (30) days before the termination takes effect.

Authority: 36 CFR 800.14(e).

Dated: November 10, 2009.

Reid Nelson,

Acting Executive Director.

[FR Doc. E9-27798 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-K6-M

DEPARTMENT OF HOMELAND SECURITY

Customs and Border Protection

Agency Information Collection Activities: Visa Waiver Program Carrier Agreement (Form I-775)

AGENCY: U.S. Customs and Border Protection (CBP), Department of Homeland Security.

ACTION: 60-Day notice and request for comments; Revision of an existing information collection: 1651-0110.

SUMMARY: As part of its continuing effort to reduce paperwork and respondent burden, CBP invites the general public and other Federal agencies to comment on the Visa Waiver Program Carrier Agreement (Form I-775). This request for comment is being made pursuant to the Paperwork Reduction Act of 1995 (Pub. L. 104-13; 44 U.S.C. 3505(c)(2)).

DATES: Written comments should be received on or before January 19, 2010, to be assured of consideration.

ADDRESSES: Direct all written comments to U.S. Customs and Border Protection, Attn: Tracey Denning, Office of Regulations and Rulings, 799 9th Street, NW., 7th Floor, Washington, DC 20229-1177.

FOR FURTHER INFORMATION CONTACT: Requests for additional information should be directed to Tracey Denning, U.S. Customs and Border Protection, Office of Regulations and Rulings, 799 9th Street, NW., 7th Floor, Washington, DC 20229-1177, at 202-325-0265.

SUPPLEMENTARY INFORMATION: CBP invites the general public and other Federal agencies to comment on proposed and/or continuing information collections pursuant to the Paperwork Reduction Act of 1995 (Pub. L. 104-13; 44 U.S.C. 3505(c)(2)). The comments should address: (a) Whether the collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimates of the burden of the collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; (d) ways to minimize the burden including the use of automated collection techniques or the use of other forms of information technology; and (e) estimates of capital or start-up costs and costs of operations, maintenance, and purchase of services to provide information. The comments that are submitted will be summarized and included in the request for Office of Management and Budget (OMB) approval. All comments will become a matter of public record. In this document the CBP is soliciting comments concerning the following information collection:

Title: Visa Waiver Program Carrier Agreement.

OMB Number: 1651-0110.

Form Number: I-775.

Abstract: Pursuant to section 217 of the Immigration and Nationality Act (INA), paragraphs (a) and (e) and 8 CFR 217.6, all carriers must enter into an agreement with CBP in order to transport passengers to the United States under the Visa Waiver Program (VWP). Form I-775 functions as the agreement between CBP and carriers, serving to hold the carriers liable for transportation costs and to ensure the completion of required forms. CBP is proposing to adjust the burden hours for this collection of information because

the estimated response time has decreased from 2 hours to 30 minutes.

CBP is also proposing to add new provisions to this Agreement including: (1) A prohibition on transporting any alien who is not authorized by the Electronic System for Travel Authorization (ESTA) to travel to the United States under the VWP; (2) a requirement that carriers applying to become signatory to a visa waiver contract with CBP have must have paid all their User Fee obligations and any previous penalties under the INA or U.S. Customs laws; (3) a requirement that carriers applying to become signatory to the VWP with CBP must post a bond sufficient to cover the total penalty amounts for violations that were imposed against the carrier during the previous fiscal year; (4) a provision that if the carrier ceases operations in the United States, then the agreement becomes null and void; and, (5) a provision that the Agreement must be renewed every seven years. In addition, CBP proposes to add a statement to Form I-775 regarding the submission of electronic arrival and departure manifests by carriers, which is an existing requirement provided under 8 CFR 217.7(a) and (b).

Current Actions: This submission is being made to extend the expiration date with a revision to the burden hours.

Type of Review: Extension (with change).

Affected Public: Businesses.

Estimated Number of Respondents: 400.

Estimated Time per Respondent: 30 minutes.

Estimated Total Annual Burden Hours: 200.

Dated: November 17, 2009.

Tracey Denning,

Agency Clearance Officer, U.S. Customs and Border Protection.

[FR Doc. E9-27904 Filed 11-19-09; 8:45 am]

BILLING CODE 9111-14-P

DEPARTMENT OF HOMELAND SECURITY

U.S. Citizenship and Immigration Services

Agency Information Collection Activities: Form N-400, Extension of an Existing Information Collection; Comment Request

ACTION: 30-Day Notice of Information Collection Under Review; Form N-400, Application for Naturalization; OMB Control No. 1615-0052.

The Department of Homeland Security, U.S. Citizenship and Immigration Services (USCIS) has submitted the following information collection request to the Office of Management and Budget (OMB) for review and clearance in accordance with the Paperwork Reduction Act of 1995. The information collection was previously published in the **Federal Register** on September 3, 2009, at 74 FR 45648, announcing a revision to the form and instructions. However, USCIS has decided not to revise the form or instructions at this time. Should USCIS decide to revise the form and instructions in the near future it will once again publish a 60-day notice in the **Federal Register** and allow the public 60-days to submit comments. USCIS did receive two comments on the September 3, 2009, notice. USCIS responded to those two comments in item 8 of the supporting statement that will be posted on <http://www.regulations.gov>.

The purpose of this notice is to allow an additional 30 days for public comments on the extension. Comments are encouraged and will be accepted until December 21, 2009. This process is conducted in accordance with 5 CFR 1320.10.

Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the Department of Homeland Security (DHS), and to the Office of Information and Regulatory Affairs, Office of Management and Budget (OMB), USCIS Desk Officer. Comments may be submitted to: USCIS, Chief, Regulatory Products Division, Clearance Office, 111 Massachusetts Avenue, Washington, DC 20529-2210. Comments may also be submitted to DHS via facsimile to 202-272-8352 or via e-mail at rfs.regs@dhs.gov, and to the OMB USCIS Desk Officer via facsimile to 202-395-5806 or via e-mail at oir_submission@omb.eop.gov.

When submitting comments by e-mail, please make sure to add OMB Control No. 1615-0052 in the subject box. Written comments and suggestions from the public and affected agencies should address one or more of the following four points:

(1) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;

(2) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information,

including the validity of the methodology and assumptions used;

(3) Enhance the quality, utility, and clarity of the information to be collected; and

(4) Minimize the burden of the collection of information on those who are to respond including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses.

Overview of This Information Collection

(1) *Type of Information Collection:* Extension of an existing information collection.

(2) *Title of the Form/Collection:* Application for Naturalization.

(3) *Agency form number, if any, and the applicable component of the Department of Homeland Security sponsoring the collection:* Form N-400; U.S. Citizenship and Immigration Services.

(4) *Affected public who will be asked or required to respond, as well as a brief abstract:* Primary: Individuals or households. USCIS uses the information on this form to determine an applicant's eligibility for naturalization.

(5) *An estimate of the total number of respondents and the amount of time estimated for an average respondent to respond:* 700,000 responses at 6 hours and 8 minutes (6.13 hours) per response.

(6) *An estimate of the total public burden (in hours) associated with the collection:* 4,291,000 annual burden hours.

If you need a copy of the information collection instrument, please visit the Web site at: <http://www.regulations.gov/>.

We may also be contacted at: USCIS, Regulatory Products Division, 111 Massachusetts Avenue, NW., Washington, DC 20529-2210, Telephone number 202-272-8377.

Dated: November 17, 2009.

Stephen Tarragon,

Deputy Chief, Regulatory Products Division, U.S. Citizenship and Immigration Services.
[FR Doc. E9-27905 Filed 11-19-09; 8:45 am]

BILLING CODE 9111-97-P

DEPARTMENT OF HOMELAND SECURITY

Coast Guard

[Docket No. USCG-2009-0800]

Notification of the Imposition of Conditions of Entry for Certain Vessels Arriving to the United States, Madagascar

AGENCY: Coast Guard, DHS.

ACTION: Notice.

SUMMARY: The Coast Guard announces that it will impose conditions of entry on vessels arriving to the United States from Madagascar, with the exception of vessels arriving from the port of Toamasina (also known as Tamatave).

DATES: The requirements announced in this notice will become effective December 4, 2009.

ADDRESSES: This notice will be available for inspection and copying at the Docket Management Facility at the U.S. Department of Transportation, Room W12-140 on the Ground Floor of the West Building, 1200 New Jersey Avenue, SE., Washington, DC 20590, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays. The telephone number is 202-366-9329.

FOR FURTHER INFORMATION CONTACT: If you have questions on this notice, call Mr. Michael Brown, International Port Security Evaluation Division, Coast Guard, telephone 202-372-1081. If you have questions on viewing or submitting material to the docket, call Renee V. Wright, Program Manager, Docket Operations, telephone 202-366-9826.

SUPPLEMENTARY INFORMATION:

Background and Purpose

Section 70110 of the Maritime Transportation Security Act of 2002 (Pub. L. 107-295, Nov. 25, 2002) (46 U.S.C. 70110) provides that the Secretary of Homeland Security may impose conditions of entry on vessels arriving from ports that are not maintaining effective anti-terrorism measures, may deny entry into the United States to any vessel that does not meet such conditions set forth herein, and shall provide public notice for passengers of the ineffective antiterrorism measures. The Coast Guard has been delegated the authority by the Secretary to carry out the provisions of this section. Previous notices have imposed or removed conditions of entry on vessels arriving from certain countries and those conditions of entry and the countries

they pertain to remain in effect unless modified by this notice.

Based on an assessment conducted pursuant to the provisions of 46 U.S.C. 70108 and the International Ship and Port Facility Security (ISPS) Code, the Coast Guard has determined that ports in Madagascar are not maintaining effective anti-terrorism measures. Inclusive to this determination is an assessment that Madagascar presents significant risk of introducing instruments of terror into international maritime commerce.

Consistent with 46 U.S.C. 70109, the United States notified Madagascar of this determination on May 17, 2007, and identified steps necessary to improve the antiterrorism measures in Madagascar. To date, the United States cannot confirm that the identified deficiencies have been corrected.

Accordingly, effective December 4, 2009 the Coast Guard will impose the following conditions of entry on vessels that visited ports in Madagascar, with the exception of vessels arriving from the port of Toamasina (also known as Tamatave) during their last five port calls. Vessels must:

- Implement measures per the ship's security plan equivalent to "Security Level 2" while in a port in Madagascar. As defined in the ISPS Code and incorporated herein, "Security Level 2" refers to the "level for which appropriate additional protective security measures shall be maintained for a period of time as a result of heightened risk of a security incident."
- Ensure that each access point to the ship is guarded and that the guards have total visibility of the exterior (both landside and waterside) of the vessel while the vessel is in ports in Madagascar. Guards may be provided by the ship's crew, however additional crewmembers should be placed on the ship if necessary to ensure that limits on maximum hours of work are not exceeded and/or minimum hours of rest are met. Guards may also be provided by outside security forces approved by the ship's master and "Company Security Officer." As defined in the ISPS Code and incorporated herein, "Company Security Officer" refers to the "person designated by the Company for ensuring that a ship security assessment is carried out; that a ship security plan is developed, submitted for approval, and thereafter implemented and maintained and for liaison with port facility security officers and the ship security officer."

- Attempt to execute a Declaration of Security while in port in Madagascar;
- Log all security actions in the ship's log; and

• Report actions taken to the cognizant Coast Guard Captain of the Port prior to arrival into U.S. waters. In addition, based on the findings of a Coast Guard boarding or examination, vessels may be required to ensure that each access point to the ship is guarded by armed security guards and that they have total visibility of the exterior (both landside and waterside) of the vessel while in U.S. ports. The number and position of the guards has to be acceptable to the cognizant Coast Guard Captain of the Port prior to the vessel's arrival. Consistent with 46 U.S.C. 70110, the United States may deny entry into the United States to any vessel that does not meet the conditions set forth herein. This notice also informs passengers of the ineffective antiterrorism measures at ports in Madagascar with the exception of the Port of Toamasina, also known as Tamatave.

This notice is issued under authority of 46 U.S.C. 70110(a).

Dated: October 15, 2009.

Sally Brice-O'Hara,

USCG, Deputy Commandant for Operations.

[FR Doc. E9-27876 Filed 11-19-09; 8:45 am]

BILLING CODE 9110-04-P

DEPARTMENT OF HOUSING AND URBAN DEVELOPMENT

[Docket No. FR-5280-N-45]

Federal Property Suitable as Facilities To Assist the Homeless

AGENCY: Office of the Assistant Secretary for Community Planning and Development, HUD.

ACTION: Notice.

SUMMARY: This Notice identifies unutilized, underutilized, excess, and surplus Federal property reviewed by HUD for suitability for possible use to assist the homeless.

FOR FURTHER INFORMATION CONTACT: Kathy Ezzell, Department of Housing and Urban Development, 451 Seventh Street, SW., Room 7266, Washington, DC 20410; telephone (202) 708-1234; TTY number for the hearing- and speech-impaired (202) 708-2565 (these telephone numbers are not toll-free), or call the toll-free Title V information line at 800-927-7588.

SUPPLEMENTARY INFORMATION: In accordance with 24 CFR part 581 and section 501 of the Stewart B. McKinney Homeless Assistance Act (42 U.S.C. 11411), as amended, HUD is publishing this Notice to identify Federal buildings and other real property that HUD has reviewed for suitability for use to assist

the homeless. The properties were reviewed using information provided to HUD by Federal landholding agencies regarding unutilized and underutilized buildings and real property controlled by such agencies or by GSA regarding its inventory of excess or surplus Federal property. This Notice is also published in order to comply with the December 12, 1988 Court Order in *National Coalition for the Homeless v. Veterans Administration*, No. 88-2503-OG (D.D.C.).

Properties reviewed are listed in this Notice according to the following categories: Suitable/available, suitable/unavailable, suitable/to be excess, and unsuitable. The properties listed in the three suitable categories have been reviewed by the landholding agencies, and each agency has transmitted to HUD: (1) Its intention to make the property available for use to assist the homeless, (2) its intention to declare the property excess to the agency's needs, or (3) a statement of the reasons that the property cannot be declared excess or made available for use as facilities to assist the homeless.

Properties listed as suitable/available will be available exclusively for homeless use for a period of 60 days from the date of this Notice. Where property is described as for "off-site use only" recipients of the property will be required to relocate the building to their own site at their own expense. Homeless assistance providers interested in any such property should send a written expression of interest to HHS, addressed to Theresa Rita, Division of Property Management, Program Support Center, HHS, room 5B-17, 5600 Fishers Lane, Rockville, MD 20857; (301) 443-2265. (This is not a toll-free number.) HHS will mail to the interested provider an application packet, which will include instructions for completing the application. In order to maximize the opportunity to utilize a suitable property, providers should submit their written expressions of interest as soon as possible. For complete details concerning the processing of applications, the reader is encouraged to refer to the interim rule governing this program, 24 CFR part 581.

For properties listed as suitable/to be excess, that property may, if subsequently accepted as excess by GSA, be made available for use by the homeless in accordance with applicable law, subject to screening for other Federal use. At the appropriate time, HUD will publish the property in a Notice showing it as either suitable/available or suitable/unavailable.

For properties listed as suitable/unavailable, the landholding agency has decided that the property cannot be declared excess or made available for use to assist the homeless, and the property will not be available.

Properties listed as unsuitable will not be made available for any other purpose for 20 days from the date of this Notice. Homeless assistance providers interested in a review by HUD of the determination of unsuitability should call the toll free information line at 1-800-927-7588 for detailed instructions or write a letter to Mark Johnston at the address listed at the beginning of this Notice. Included in the request for review should be the property address (including zip code), the date of publication in the **Federal Register**, the landholding agency, and the property number.

For more information regarding particular properties identified in this Notice (*i.e.*, acreage, floor plan, existing sanitary facilities, exact street address), providers should contact the appropriate landholding agencies at the following addresses: *Army:* Ms. Veronica Rines, Department of the Army, Office of the Assistant Chief of Staff for Installation Management, DAIM-ZS, Room 8536, 2511 Jefferson Davis Hwy., Arlington, VA 22202; (703) 601-2545; (These are not toll-free numbers).

Dated: November 12, 2009.

Mark R. Johnston,

Deputy Assistant Secretary for Special Needs.

TITLE V, FEDERAL SURPLUS PROPERTY PROGRAM FEDERAL REGISTER REPORT FOR 11/20/2009

Suitable/Available Properties

Building

New York
Bldg. 1230
U.S. Army Garrison
Orange NY 10996
Landholding Agency: Army
Property Number: 21200940014
Status: Unutilized
Comments: 4538 sq. ft., possible asbestos/
lead paint, most recent use—clubhouse,
off-site use only

Land

Tennessee
Parcel No. 6
Fort Campbell
Hwy 79
Montgomery TN 42223
Landholding Agency: Army
Property Number: 21200940013
Status: Excess
Comments: 4.55 acres, wooded w/dirt road/
fire break

Unsuitable Properties*Building*

Alabama

11 Bldgs.

Redstone Arsenal

Madison AL 35898

Landholding Agency: Army

Property Number: 21200940015

Status: Unutilized

Directions: 99, 119, 1421, 1422, 3496, 3614,
3648, 3768, 3772, 3773, 3774Reasons: Extensive deterioration; Secured
Area

8 Bldgs.

Redstone Arsenal

Madison AL 35898

Landholding Agency: Army

Property Number: 21200940016

Status: Unutilized

Directions: 4501, 5132, 7108, 7288, 7380,
7523, 7524, 7525Reasons: Extensive deterioration; Secured
Area

4 Bldgs.

Redstone Arsenal

Madison AL 35898

Landholding Agency: Army

Property Number: 21200940017

Status: Unutilized

Directions: 7554, 7567, 7592, 7907

Reasons: Secured Area; Extensive
deterioration

Alaska

7 Bldgs.

Fort Richardson

Anchorage AK 99505

Landholding Agency: Army

Property Number: 21200940018

Status: Unutilized

Directions: 992, 992C1, 992C2, 39002, 39199,
39228, 39600

Reasons: Extensive deterioration

5 Bldgs.

Fort Richardson

Anchorage AK 99505

Landholding Agency: Army

Property Number: 21200940019

Status: Unutilized

Directions: 57112, 57451, 57452, 57453,
57458

Reasons: Extensive deterioration

6 Bldgs.

Fort Richardson

Anchorage AK 99505

Landholding Agency: Army

Property Number: 21200940020

Status: Unutilized

Directions: RANMS, RANNL, RANOS,
RANPS, RANZS, WOLT2

Reasons: Extensive deterioration

Bldg. XTENA

Fort Greely

Fort Greely AK 99731

Landholding Agency: Army

Property Number: 21200940021

Status: Unutilized

Reasons: Within 2000 ft. of flammable or
explosive material; Secured Area;
Extensive deterioration

California

Bldg. 00053

Moffett Community Housing

Santa Clara CA 94035

Landholding Agency: Army

Property Number: 21200940022

Status: Unutilized

Reasons: Extensive deterioration

Bldg. 00005

Los Alamitos Joint Force

Training Base

Orange CA 90720

Landholding Agency: Army

Property Number: 21200940023

Status: Excess

Reasons: Extensive deterioration

8 Bldgs.

Fort Hunter Liggett

Monterey CA 93928

Landholding Agency: Army

Property Number: 21200940024

Status: Unutilized

Directions: 410, 413, 414, 415, 416, 417, 419,
420

Reasons: Extensive deterioration

Georgia

7 Bldgs.

Fort Stewart

Liberty GA 31314

Landholding Agency: Army

Property Number: 21200940025

Status: Excess

Directions: 918, 1076, 1103, 1268, 7803,
7804, 7805

Reasons: Extensive deterioration

Bldgs. 240, 701, 719

Hunter Army Airfield

Savannah GA 31409

Landholding Agency: Army

Property Number: 21200940026

Status: Excess

Reasons: Extensive deterioration

6 Bldgs.

Fort Benning

Ft. Benning GA 31905

Landholding Agency: Army

Property Number: 21200940027

Status: Unutilized

Directions: 1189, 1199, 2603, 2605, 8621,
8622

Reasons: Secured Area

Hawaii

Bldgs. 00039, 01650

Fort Shafter

Honolulu HI 96858

Landholding Agency: Army

Property Number: 21200940039

Status: Unutilized

Reasons: Extensive deterioration

Bldg. 01007

Wheeler Army Airfield

Honolulu HI 96786

Landholding Agency: Army

Property Number: 21200940040

Status: Unutilized

Reasons: Extensive deterioration

Kansas

Bldgs. 2351, 6420, 8999

Fort Riley

Geary KS 66441

Landholding Agency: Army

Property Number: 21200940041

Status: Unutilized

Reasons: Extensive deterioration

Kentucky

4 Bldgs.

Fort Knox

Ft. Knox KY 40121

Landholding Agency: Army

Property Number: 21200940042

Status: Unutilized

Directions: 1172, 6898, 7737, 7739

Reasons: Extensive deterioration

Maryland

7 Bldgs.

Aberdeen Proving Ground

Harford MD 21005

Landholding Agency: Army

Property Number: 21200940028

Status: Unutilized

Directions: 4210, 4211, 4212, 4213, 4302,
4303, 5650

Reasons: Secured Area

9 Bldgs.

Aberdeen Proving Grounds

Harford MD 21005

Landholding Agency: Army

Property Number: 21200940029

Status: Unutilized

Directions: E3220, E4405, E4410, E4430,
E4435, E4445, E4455, E4460, E4475

Reasons: Secured Area

9 Bldgs.

Aberdeen Proving Ground

Harford MD 21005

Landholding Agency: Army

Property Number: 21200940030

Status: Unutilized

Directions: E5641, E5642, E5684, E5685,
E5686, E5687, E5910, E5911, E5912

Reasons: Secured Area

Bldgs. 718, 917

Fort Detrick

Frederick MD 21702

Landholding Agency: Army

Property Number: 21200940043

Status: Unutilized

Reasons: Secured Area

Missouri

13 Bldgs.

Fort Leonard Wood

Pulaski MO 65473

Landholding Agency: Army

Property Number: 21200940044

Status: Unutilized

Directions: 401, 761, 762, 766, 790, 791, 792,
793, 794, 795, 796, 797, 798

Reasons: Secured Area

7 Bldgs.

Fort Leonard Wood

Pulaski MO 65473

Landholding Agency: Army

Property Number: 21200940045

Status: Unutilized

Directions: 851, 852, 853, 854, 857, 859, 2305

Reasons: Secured Area

9 Bldgs.

Fort Leonard Wood

Pulaski MO 65473

Landholding Agency: Army

Property Number: 21200940046

Status: Unutilized

Directions: 9004, 9005, 9007, 9009, 9011,
9013, 9015, 9017, 9029

Reasons: Secured Area

9 Bldgs.

Fort Leonard Wood

Pulaski MO 65473

Landholding Agency: Army

Property Number: 21200940047

Status: Unutilized
 Directions: 9031, 9033, 9035, 9037, 9039, 9041, 9043, 9045, 9047
 Reasons: Secured Area
 6 Bldgs.
 Fort Leonard Wood
 Pulaski MO 65473
 Landholding Agency: Army
 Property Number: 21200940048
 Status: Unutilized
 Directions: 9057, 9059, 9061, 9063, 9071, 12315
 Reasons: Secured Area
 New Jersey
 7 Bldgs.
 Picatinny Arsenal
 Dover NJ 07806
 Landholding Agency: Army
 Property Number: 21200940031
 Status: Unutilized
 Directions: 80, 80C, 81, 82, 83, 948, 949
 Reasons: Within 2000 ft. of flammable or explosive material; Secured Area
 5 Bldgs.
 Picatinny Arsenal
 Dover NJ 07806
 Landholding Agency: Army
 Property Number: 21200940032
 Status: Unutilized
 Directions: 3710, 3711, 3712, 3713, 3714
 Reasons: Secured Area; Within 2000 ft. of flammable or explosive material
 New York
 14 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940001
 Status: Unutilized
 Directions: 192, 401, 403, 404, 405, 406, 407, 408, 410, 411, 412, 416, 417, 418
 Reasons: Extensive deterioration
 5 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940002
 Status: Unutilized
 Directions: 420, 421, 422, 423, 424
 Reasons: Extensive deterioration
 9 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940003
 Status: Unutilized
 Directions: 440, 441, 442, 443, 444, 445, 446, 447, 448
 Reasons: Extensive deterioration
 7 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940004
 Status: Unutilized
 Directions: 450, 451, 452, 453, 454, 456, 458
 Reasons: Extensive deterioration
 7 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940005
 Status: Unutilized
 Directions: 470, 471, 472, 473, 474, 477, 478

Reasons: Extensive deterioration
 11 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940006
 Status: Unutilized
 Directions: 501, 503, 513, 514, 515, 521, 522, 523, 527, 528, 529
 Reasons: Extensive deterioration
 11 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940007
 Status: Unutilized
 Directions: 532, 540, 541, 542, 543, 544, 545, 546, 547, 548, 549
 Reasons: Extensive deterioration
 6 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940008
 Status: Unutilized
 Directions: 550, 551, 552, 553, 557, 558, 559
 Reasons: Extensive deterioration
 7 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940009
 Status: Unutilized
 Directions: 560, 561, 562, 571, 572, 573, 574
 Reasons: Extensive deterioration
 6 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940010
 Status: Unutilized
 Directions: 1190, 1714, 10181, 10183, 10287, 11457
 Reasons: Extensive deterioration
 18 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940011
 Status: Unutilized
 Directions: M425A, M425B, M426A, M426B, M427A, M427B, M430A, M430B, M431A, M431B, M432A, M432B, M433A, M433B, M434A, M434B, M435A, M435B
 Reasons: Extensive deterioration
 17 Bldgs.
 Fort Drum
 Jefferson NY 13602
 Landholding Agency: Army
 Property Number: 21200940012
 Status: Unutilized
 Directions: M449A, M455A, M457B, M459B, M460A, M460B, M461A, M461B, M462A, M462B, M463A, M463B, M464B, M465A, M475A, M476B, M479B
 Reasons: Extensive deterioration
 North Carolina
 Bldgs. T3354, T3361
 Fort Bragg
 Camp Mackall NC 28373
 Landholding Agency: Army
 Property Number: 21200940033
 Status: Unutilized
 Reasons: Extensive deterioration; Secured Area

Oklahoma
 7 Bldgs.
 McAlester Army Ammo Plant
 Pittsburg OK 74501
 Landholding Agency: Army
 Property Number: 21200940049
 Status: Excess
 Directions: 81, 82, 98, 213, 449, 628, 643
 Reasons: Within 2000 ft. of flammable or explosive material; Secured Area
 Pennsylvania
 4 Bldgs.
 Letterkenny Army Depot
 Franklin PA 17201
 Landholding Agency: Army
 Property Number: 21200940034
 Status: Unutilized
 Directions: S3627, 03811, S4344, S5298
 Reasons: Secured Area
 Tennessee
 Bldgs. ZZ001, ZZ002
 Milan AAP
 Gibson TN 38358
 Landholding Agency: Army
 Property Number: 21200940035
 Status: Excess
 Reasons: Within 2000 ft. of flammable or explosive material; Secured Area
 Texas
 Bldgs. 11284, 11304
 Fort Bliss
 El Paso TX 79916
 Landholding Agency: Army
 Property Number: 21200940036
 Status: Unutilized
 Reasons: Secured Area
 Virginia
 Bldgs. 1218, 1296
 Fort A.P. Hill
 Bowling Green VA 22427
 Landholding Agency: Army
 Property Number: 21200940037
 Status: Unutilized
 Reasons: Extensive deterioration
 Bldgs. 1000, 2000, 2010
 Radford AAP
 Montgomery VA 24143
 Landholding Agency: Army
 Property Number: 21200940038
 Status: Unutilized
 Reasons: Secured Area; Within 2000 ft. of flammable or explosive material
 [FR Doc. E9-27566 Filed 11-19-09; 8:45 am]
BILLING CODE 4210-67-P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[F-14846-B, F-14846-C, F-14846-B2, F-14846-F2, F-14846-H2; LLAK964000-L1410000-KC0000-P]

Alaska Native Claims Selection

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of decision approving lands for conveyance.

SUMMARY: As required by 43 CFR 2650.7(d), notice is hereby given that an appealable decision approving the surface estate in certain lands for conveyance pursuant to the Alaska Native Claims Settlement Act will be issued to Chalkyitsik Native Corporation. The lands are in the vicinity of Chalkyitsik, Alaska, and are located in:

Lot 2, U.S. Survey No. 4133, Alaska.
Containing 2.34 acres.

Fairbanks Meridian, Alaska

T. 23 N., R. 16 E.,

Sec. 29;

Secs. 32 to 36, inclusive.

Containing approximately 2,580 acres.

T. 22 N., R. 17 E.,

Secs. 25 to 36, inclusive.

Containing approximately 6,192 acres.

T. 23 N., R. 18 E.,

Secs. 4 to 8, inclusive;

Secs. 18, 19, and 30.

Containing approximately 3,891 acres.

T. 22 N., R. 19 E.,

Secs. 11 to 14, inclusive;

Secs. 23 to 36, inclusive.

Containing approximately 9,086 acres.

T. 21 N., R. 20 E.,

Secs. 1 to 6, inclusive;

Secs. 9 to 14, inclusive.

Containing approximately 6,335 acres.

Aggregating approximately 28,087 acres.

The subsurface estate in these lands will be conveyed to Doyon, Limited, when the surface estate is conveyed to Chalkyitsik Native Corporation. Notice of the decision will also be published four times in the Fairbanks Daily News-Miner.

DATES: The time limits for filing an appeal are:

1. Any party claiming a property interest which is adversely affected by the decision shall have until December 21, 2009 to file an appeal.

2. Parties receiving service of the decision by certified mail shall have 30 days from the date of receipt to file an appeal.

Parties who do not file an appeal in accordance with the requirements of 43 CFR part 4, subpart E, shall be deemed to have waived their rights.

ADDRESSES: A copy of the decision may be obtained from: Bureau of Land Management, Alaska State Office, 222 West Seventh Avenue, #13, Anchorage, Alaska 99513-7504.

FOR FURTHER INFORMATION CONTACT: The Bureau of Land Management by phone at 907-271-5960, or by e-mail at ak.blm.conveyance@ak.blm.gov. Persons who use a telecommunication device (TTD) may call the Federal Information Relay Service (FIRS) at 1-800-877-8339, 24 hours a day, seven days a

week, to contact the Bureau of Land Management.

Barbara Opp Waldal,

Land Law Examiner, Land Transfer Adjudication I Branch.

[FR Doc. E9-27864 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-JA-P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[F-19328-B; LLA964 000-L1410000-KC0000-P]

Alaska Native Claims Selection

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of Decision approving lands for conveyance.

SUMMARY: As required by 43 CFR 2650.7(d), notice is hereby given that an appealable decision approving the surface estate in certain lands for conveyance pursuant to the Alaska Native Claims Settlement Act will be issued to Evansville, Incorporated. The lands are in the vicinity of Evansville, Alaska, and are located in:

Fairbanks Meridian, Alaska

T. 26 N., R. 17 W.,

Secs. 35 and 36.

Containing approximately 1,094 acres.

T. 25 N., R. 18 W.,

Secs. 16, 21, 28 and 29.

Containing approximately 2,437 acres.

T. 24 N., R. 19 W.,

Secs. 7 and 8.

Containing approximately 1,230 acres.

Aggregating approximately 4,761 acres.

The subsurface estate in these lands will be conveyed to Doyon, Limited when the surface estate is conveyed to Evansville, Incorporated. Notice of the decision will also be published four times in the Fairbanks Daily News-Miner.

DATES: The time limits for filing an appeal are:

1. Any party claiming a property interest which is adversely affected by the decision shall have until December 21, 2009 to file an appeal.

2. Parties receiving service of the decision by certified mail shall have 30 days from the date of receipt to file an appeal.

Parties who do not file an appeal in accordance with the requirements of 43 CFR Part 4, Subpart E, shall be deemed to have waived their rights.

ADDRESSES: A copy of the decision may be obtained from: Bureau of Land Management, Alaska State Office, 222

West Seventh Avenue, #13, Anchorage, Alaska 99513-7504.

FOR FURTHER INFORMATION, CONTACT: The Bureau of Land Management by phone at 907-271-5960, or by e-mail at: ak.blm.conveyance@ak.blm.gov. Persons who use a telecommunication device (TTD) may call the Federal Information Relay Service (FIRS) at 1-800-877-8339, 24 hours a day, seven days a week, to contact the Bureau of Land Management.

Jenny M. Anderson,

Land Law Examiner, Land Transfer Adjudication I Branch.

[FR Doc. E9-27868 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-JA-P

DEPARTMENT OF THE INTERIOR

Fish and Wildlife Service

[FWS-R2-ES-2009-N217; 20124-1113-0000-F5]

Endangered and Threatened Species Permit Applications

AGENCY: Fish and Wildlife Service, Interior.

ACTION: Notice of receipt of applications; request for public comment.

SUMMARY: The following applicants have applied for scientific research permits to conduct certain activities with endangered species under the Endangered Species Act of 1973, as amended (Act). The Act requires that we invite public comment on these permit applications.

DATES: To ensure consideration, written comments must be received on or before December 21, 2009.

ADDRESSES: Written comments should be submitted to the Chief, Endangered Species Division, Ecological Services, P.O. Box 1306, Room 6034, Albuquerque, NM 87103. Documents and other information submitted with these applications are available for review, subject to the requirements of the Privacy Act and Freedom of Information Act. Documents will be available for public inspection, by appointment only, during normal business hours at the U.S. Fish and Wildlife Service, 500 Gold Ave., SW., Room 6034, Albuquerque, NM. Please refer to the respective permit number for each application when submitting comments.

FOR FURTHER INFORMATION CONTACT: Susan Jacobsen, Chief, Endangered Species Division, P.O. Box 1306, Albuquerque, NM 87103; (505) 248-6920.

SUPPLEMENTARY INFORMATION:

Public Availability of Comments

Before including your address, phone number, e-mail address, or other personal identifying information in your comment, you should be aware that your entire comment—including your personal identifying information—may be made publicly available at any time. While you can ask us in your comment to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Permit TE-226979

Applicant: Christopher Murray, San Antonio, Texas.

Applicant requests a new permit for research and recovery purposes to conduct presence/absence surveys for golden-cheeked warbler (*Dendroica chrysoparia*) and black-capped vireo (*Vireo atricapilla*) within Texas.

Permit TE-227505

Applicant: Kathleen O'Connor, Georgetown, Texas.

Applicant requests a new permit for research and recovery purposes to conduct presence/absence surveys for the following species: Texas blind salamander (*Typhlomolge rathbuni*), San Marcos salamander (*Eurycea sosorum*), Peck's Cave amphipod (*Stygobromus pecki*), Comal Springs dryopid beetle (*Stygoparnus comalensis*), Comal Springs riffle beetle (*Heterelmis comalensis*), Coffin Cave mold beetle (*Batrisodes texanus*), Helotes mold beetle (*Batrisodes venyivi*), Kretschmarr Cave mold beetle (*Texamaurops redelli*), ground beetle (*Rhadine exilis*), ground beetle (*Rhadine infernalis*), Tooth Cave ground beetle (*Rhadine persephone*), Robber Baron Cave meshweaver (*Cicurina baronia*), Madla Cave meshweaver (*Cicurina madla*), Braken Bat Cave meshweaver (*Cicurina venii*), Government Canyon Bat Cave meshweaver (*Cicurina vespera*), Government Canyon Bat Cave spider (*Neoleptoneta microps*), Tooth Cave spider (*Neoleptoneta myopica*), Tooth Cave psuedoscorpion (*Tartarocreagriss texanus*), Bee Creek Cave harvestman (*Texella reddelli*), Bone Cave harvestman (*Texella reyesi*), golden-cheeked warbler (*Dendroica chrysoparia*), and black-capped vireo (*Vireo atricapilla*) within Texas.

Permit TE-230274

Applicant: David Keller, Los Alamos, New Mexico.

Applicant requests a new permit for research and recovery purposes to conduct presence/absence surveys for southwestern willow flycatcher

(*Empidonax traillii extimus*) within New Mexico.

Permit TE-055419

Applicant: Turner Biological Consulting, LLC., Buffalo Gap, Texas.

Applicant requests an amendment to a current permit for research and recovery purposes to conduct presence/absence surveys for Eskimo curlew (*Numenius borealis*) within Texas.

Permit TE-230679

Applicant: David Black, Lakehills, Texas.

Applicant requests a new permit for research and recovery purposes to conduct presence/absence surveys for golden-cheeked warbler (*Dendroica chrysoparia*) within Texas.

Permit TE-819475

Applicant: Bureau of Reclamation, Denver, Colorado.

Applicant requests an amendment to a current permit for research and recovery purposes to conduct presence/absence surveys for interior least tern (*Sternula antillarum athalassos*) within New Mexico.

Permit TE-231653

Applicant: Daniel Allen, Austin, Texas.

Applicant request a new permit for research and recovery purposes to conduct presence/absence surveys for the following species: brown pelican (*Pelecanus occidentalis*), Gulf Coast jaguarundi (*Herpailurus yagouaroundi cacomitli*), ocelot (*Leopardus pardalis*), northern aplomado falcon (*Falco femoralis septentrionalis*), piping plover (*Charadrius melodus*), South Texas ambrosia (*Ambrosia cheiranthifolia*), Texas ayenia (*Ayenia limitaris*), and West Indian manatee (*Trichechus manatus*) within Texas.

Authority: 16 U.S.C. 1531 *et seq.*

Dated: November 10, 2009.

Brian Millsap,

Regional Director, Southwest Region, Fish and Wildlife Service.

[FR Doc. E9-27887 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-55-P

DEPARTMENT OF THE INTERIOR**Fish and Wildlife Service**

[FWS-R4-R-2009-N173; 40136-1265-0000-S3]

Bayou Teche National Wildlife Refuge, St. Mary Parish, LA

AGENCY: Fish and Wildlife Service, Interior.

ACTION: Notice of availability: final comprehensive conservation plan and finding of no significant impact.

SUMMARY: We, the Fish and Wildlife Service (Service), announce the availability of our final comprehensive conservation plan (CCP) and finding of no significant impact (FONSI) for the environmental assessment for Bayou Teche National Wildlife Refuge (NWR). In the final CCP, we describe how we will manage this refuge for the next 15 years.

ADDRESSES: You may obtain a copy of the CCP by writing to: Mr. Paul Yakupzack, Refuge Manager, Mandalay NWR, 3599 Bayou Black Drive, Houma, LA 70360. You may also access and download the document from the Service's Web site: <http://southeast.fws.gov/planning>.

FOR FURTHER INFORMATION CONTACT: Mr. Paul Yakupzack; *telephone:* 985/853-1078; *fax:* 985/853-1079; *e-mail:* paul_yakupzack@fws.gov.

SUPPLEMENTARY INFORMATION:**Introduction**

With this notice, we finalize the CCP process for Bayou Teche NWR. We started this process through a notice in the **Federal Register** on March 19, 2007 (72 FR 12811). For more about the process, see that notice.

Bayou Teche NWR is located near the town of Franklin in St. Mary Parish, Louisiana. The refuge contains 9,028 acres and is composed of wet bottomland hardwood forests laced with bayous and canals. The refuge was established on October 31, 2001, on lands important to the coastal subpopulation of the Louisiana black bear. The refuge consists of six separate units, ranging in size from 3,724 acres to 80 acres.

We announce our decision and the availability of the final CCP and FONSI for Bayou Teche NWR in accordance with the National Environmental Policy Act (NEPA) [40 CFR 1506.6(b)] requirements. We completed a thorough analysis of impacts on the human environment, which we included in the draft comprehensive conservation plan and environmental assessment (Draft CCP/EA). The CCP will guide us in managing and administering Bayou Teche NWR for the next 15 years.

The compatibility determinations for boating, recreational fishing, recreational hunting, wildlife observation/photography, and environmental education/interpretation are available in the CCP.

Background

The National Wildlife Refuge System Administration Act of 1966 (16 U.S.C. 668dd–668ee) (Administration Act), as amended by the National Wildlife Refuge System Improvement Act of 1997, requires us to develop a CCP for each national wildlife refuge. The purpose for developing a CCP is to provide refuge managers with a 15-year plan for achieving refuge purposes and contributing toward the mission of the National Wildlife Refuge System, consistent with sound principles of fish and wildlife management, conservation, legal mandates, and our policies. In addition to outlining broad management direction on conserving wildlife and their habitats, CCPs identify wildlife-dependent recreational opportunities available to the public, including opportunities for hunting, fishing, wildlife observation, wildlife photography, and environmental education and interpretation. We will review and update the CCP at least every 15 years in accordance with the Administration Act.

Comments

Approximately 100 copies of the Draft CCP/EA were made available for a 30-day public review period as announced in the **Federal Register** on June 8, 2009 (74 FR 27174). Several comments were received from local citizens and the Louisiana Department of Wildlife and Fisheries.

Selected Alternative

After considering the comments we received, and based on the sound professional judgment of the planning team, we selected Alternative B to implement the CCP. The primary focuses of the CCP are to optimize Louisiana black bear and wetland habitats, monitor targeted flora and fauna representative of the lower Atchafalaya Basin, and provide quality public use programs and wildlife-dependent recreational activities. Based on the mission of the National Wildlife Refuge System, the purposes for which Bayou Teche NWR was established, and the focus of the Lower Mississippi River Ecosystem priorities, we believe Alternative B best fits the goals of the refuge.

Authority

This notice is published under the authority of the National Wildlife Refuge System Improvement Act of 1997, Public Law 105–57.

Dated: August 24, 2009.

Patrick Leonard,

Acting Regional Director.

[FR Doc. E9–27888 Filed 11–19–09; 8:45 am]

BILLING CODE 4310–55–P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

**[LLAZC010.L51010000.ER0000.
LVRWA09A2310.241A; AZA 32315]**

Notice of Intent to Prepare an Environmental Impact Statement and Initiate Public Scoping for the Proposed Mohave County Wind Farm Project, Mohave County, AZ

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of intent.

SUMMARY: In compliance with the National Environmental Policy Act (NEPA) of 1976, as amended, the Bureau of Land Management (BLM) Kingman Field Office, Kingman, Arizona, intends to prepare an Environmental Impact Statement (EIS) for the Proposed Mohave County Wind Farm Project and by this notice is announcing the beginning of the scoping process to solicit public comments and identify issues.

DATES: This notice initiates the public scoping process for the EIS. Comments may be submitted in writing until January 4, 2010. The date(s) and location(s) of any scoping meetings will be announced at least 15 days in advance through local news media, newspapers, and the BLM–Arizona Web site at: <http://www.blm.gov/az/st/en.html>. In order to be included in the Draft EIS, all comments must be received prior to the close of the scoping period or 15 days after the last public meeting, whichever is later. We will provide additional opportunities for public participation upon publication of the Draft EIS.

ADDRESSES: You may submit comments related to the Proposed Mohave County Wind Farm Project, Mohave County, Arizona by any of the following methods:

- *Web site:* <http://www.blm.gov/az/st/en.html>.
- *E-mail:* KFO_WindEnergy@blm.gov.
- *Fax:* (928) 718–3761.
- *Mail:* Ruben Sanchez, Field Manager, BLM, Kingman Field Office, 2755 Mission Boulevard, Kingman, Arizona 86401.

Documents pertinent to this proposal may be examined at the Bureau of Land Management Kingman Field Office, Kingman, Arizona.

FOR FURTHER INFORMATION CONTACT: For further information and/or to have your name added to our mailing list, contact Jerry Crockford, BLM-contracted Project Manager at (505) 360–0473 or e-mail KFO_WindEnergy@blm.gov.

SUPPLEMENTARY INFORMATION: On November 1, 2002, the BLM received a right-of-way (ROW) application from BP Wind Energy North America (BPWE) for meteorological testing and monitoring for a wind energy project area. The BLM issued BPWE a ROW grant (AZA 32315) for a wind energy testing and monitoring project area of approximately 41,900 acres. Having gathered wind data for more than four years, BPWE is now moving forward to develop the project area, including an additional 3,520 acres, with a proposal to construct, operate, maintain, and decommission facilities and improvements associated with wind energy generation on the site, including wind turbine generators (WTG), access roads, operations and maintenance buildings, power lines, substations and other ancillary facilities and improvements, and an interconnection with one of two transmission lines which transect the project area. BPWE also proposes installing WTGs on approximately 4,360 acres of private lands adjacent to the ROW over which BPWE holds or anticipates holding wind development leases or easements. Zoning approval for development on private lands will be sought from Mohave County. The project area ROW includes approximately 45,420 acres of public land under jurisdiction of the BLM Kingman Field Office and potentially 4,360 acres of private land in the White Hills area approximately 40 miles northwest of Kingman, Arizona, approximately nine miles south of the Colorado River, and approximately 20 miles southeast of Hoover Dam. The project area is generally located within Townships 27 through 29 North, Ranges 18 and 19 West, and Townships 28 and 29 North, Range 20 West.

Total electric generation capacity of the project is anticipated to be up to 500 megawatts (MW). The project will consist of up to 335 WTGs and consist of construction in multiple phases.

Phase I is proposed to be located on the northwest portion of the BLM project area ROW, and may consist of up to 235 WTGs, access roads, and ancillary facilities. The WTGs are anticipated to range in size from 1.5 to 3.0 MW each. To the extent possible, existing roads would be used for access to the project, supplemented with internal access/service roads to each WTG. Ancillary facilities may include

pad-mounted transformers, an underground 34.5 kilovolt (kV) electrical collection system between the turbines, either a 345 or 500 kV electrical substation, and either a 345 or 500 kV overhead transmission line from the substation to a new switchyard where the project would interconnect to one of the two major existing transmission lines in the area. Up to 10 WTGs could be installed on adjacent private lands during Phase I.

Subsequent phases are proposed to include comparable facilities with additional wind generation capacity of up to 150 MW on the balance of the area within the ROW and the private lands adjacent to the ROW area. A total of 50 to 100 WTGs may be installed on public or adjacent private lands in the subsequent phases of the project. These turbines also are anticipated to range in size from 1.5 to 3.0 MW.

A map of the proposed project area with the news release announcing the public meetings, is available on the BLM-Arizona Web site at: <http://www.blm.gov/az/st/en.html>.

The EIS will consider the impacts of the proposed action, alternatives, and a no action alternative.

The public is invited to submit comments and resource information and identify issues or concerns to be considered in the Draft EIS. Public comments will aid the BLM in identifying alternatives and mitigating measures and will help assure all relevant issues are considered in the EIS.

Preliminary issues that have been identified by the BLM for analysis include: access requirements; air quality during construction; cultural and historical resources; areas with high mineral potential; noise; sensitive soils and geology; recreation resources; socioeconomic; threatened and endangered species; visual resources; water resources; and wildlife habitats.

The BLM will use and coordinate the NEPA commenting process to satisfy the public involvement process for Section 106 of the National Historic Preservation Act (16 U.S.C. 470f) as provided for in 36 CFR 800.2(d)(3). Native American Tribal consultations will be conducted in accordance with policy and Tribal concerns and will be given due consideration, including impacts on Indian trust assets. Federal, State, and local agencies, as well as individuals, organizations, or tribes that may be interested or affected by the BLM's decision on this project are invited to participate in the scoping process and, if eligible, may request or be requested by the BLM to participate as a cooperating agency.

Before including your phone number, e-mail address, or other personal identifying information in your comment, you should be aware that your entire comment—including your personal identifying information—may be made publicly available at any time. While you can ask us in your comment to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Authority: 43 CFR 2800.

James G. Kenna,
State Director.

[FR Doc. E9-27867 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-32-P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[LLNVL0000 L51010000.ER0000
LVRWF09F1640; N-82076; 09-08807;
MO4500009275; TAS:14X5017]

Notice of Availability of the Draft Supplemental Environmental Impact Statement for the One Nevada Transmission Line, NV

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of Availability.

SUMMARY: In accordance with the National Environmental Policy Act of 1969 (NEPA), the Bureau of Land Management (BLM) has prepared a Draft Supplemental Environmental Impact Statement (SEIS) for the One Nevada Transmission Line and by this Notice is announcing the opening of the comment period.

DATES: To ensure comments will be considered, the BLM must receive written comments on the Draft SEIS for the One Nevada Transmission Line within 60 days following the date the Environmental Protection Agency publishes its Notice of Availability in the **Federal Register**. The BLM will announce future meetings or hearings and any other public involvement activities at least 15 days in advance through public notices, media news releases, and/or mailings.

ADDRESSES: You may submit comments by any of the following methods:

- *Web site:* www.blm.gov/nv.
- *E-mail:* michael_dwyer@blm.gov.
- *Fax:* (775) 289-1910.
- *Mail:* Michael Dwyer, BLM, HC 33 Box 33500, Ely, NV 89301.

Copies of the Draft SEIS for the One Nevada Transmission Line are available at the following locations in Nevada:

—BLM Ely District Office, 702 North Industrial Way, Ely.

—White Pine County Library, 950 Campton Street, Ely.

—BLM Nevada State Office, 1340 Financial Blvd., Reno.

—BLM Caliente Field Station, U.S. Highway 93, Caliente.

—Caliente Branch Library, 100 Depot Avenue, Caliente.

—BLM Southern Nevada District Office, 4701 North Torrey Pines, Las Vegas.

—North Las Vegas Library, 2300 Civic Center Drive, North Las Vegas.

FOR FURTHER INFORMATION CONTACT: Michael Dwyer, (702) 821-7102; e-mail: michael_dwyer@blm.gov.

SUPPLEMENTARY INFORMATION: On March 30, 2009, the BLM received an amended right-of-way application and Plan of Development from NV Energy for the One Nevada Transmission Line Project (ON Line Project). The Draft SEIS analyzes the construction, operation, and abandonment of a 236-mile, 500 kilovolt transmission line and telecommunication facilities running generally from Ely to Las Vegas, one new substation near Ely, and an expansion of one existing substation on private land near Battle Mountain, Nevada. The Notice of Intent to Prepare a Supplemental Environmental Impact Statement for the Proposed One Nevada Transmission Line, Nevada was published in the **Federal Register** on July 29, 2009 (74 FR 37728).

The components of the ON Line Project had been part of the original Ely Energy Center (EEC) proposal. In February 2009, during the public comment period for the EEC Draft Environmental Impact Statement (EIS), NV Energy made public their intention to postpone including the coal-fired power generation facilities associated with the EEC in their proposal until carbon capture technology becomes commercially feasible.

Two north-south utility corridors exist in Nevada that could accommodate a transmission line linking the northern and southern grids: one on the eastern side of the state and the other on the Western side of the State. The westerly corridor was considered as a potential location for the ON Line Project, but was eliminated because it would not provide access to transmission infrastructure for renewable energy resource areas in Eastern Nevada. Two alternative alignments within the eastern corridor (except in a few locations) are assessed in the SEIS. The "action" alternative generally follows the western boundary of the corridor and is the proponent's preferred

alternative for engineering and operational reasons. The other alternative generally follows the eastern boundary of the corridor, but would be more costly to construct. A no-action alternative is also assessed.

Applicable comments collected during the public comment period on the EEC Draft EIS were carried forward into the SEIS process.

Please note that public comments and information submitted, including names, street addresses, and e-mail addresses of persons who submit comments will be available for public review and disclosure at the above address during regular business hours (8 a.m. to 4 p.m.), Monday through Friday, except holidays.

Before including your address, phone number, e-mail address, or other personal identifying information in your comment, you should be aware that your entire comment—including your personal identifying information—may be made publicly available at any time. While you can ask us in your comment to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Authority: 40 CFR 1506.6 and 1506.10.

Rosemary Thomas,

District Manager, Ely District Office.

[FR Doc. E9-27891 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-HC-P

DEPARTMENT OF THE INTERIOR

National Park Service

Intent To Prepare an Environmental Impact Statement for the Comprehensive Management Plan, Lewis and Clark National Historic Trail, in Illinois, Missouri, Kansas, Nebraska, Iowa, South Dakota, North Dakota, Montana, Idaho, Washington, and Oregon

AGENCY: National Park Service, DOI.

ACTION: Notice.

SUMMARY: Pursuant to Section 102(2)(C) of the National Environmental Policy Act of 1969 (42 U.S.C. 4332(2)(C)), the National Park Service (NPS) is announcing its intent to prepare an environmental impact statement (EIS) for a comprehensive management plan (CMP) for the Lewis and Clark National Historic Trail (Trail) in the states of Illinois, Missouri, Kansas, Nebraska, Iowa, South Dakota, North Dakota, Montana, Idaho, Washington, and Oregon. The EIS will be approved by the Regional Director, Midwest Region.

The CMP will prescribe the resource conditions and visitor experiences that are to be achieved and maintained for the Trail over the next 15 to 20 years. The clarification of what must be achieved according to law and policy will be based on review of the Trail's purpose, significance, special mandates, and the body of laws and policies that direct Trail administration. Based on determinations of desired conditions, the CMP will outline the kinds of resource management activities, visitor activities, and development that would be appropriate in the future. A range of reasonable management alternatives will be developed through this planning process and will include, at a minimum, no-action and the preferred alternative.

Major issues to be addressed in the CMP include: issues surrounding preserving Trail resources (such as developing management strategies to preserve and maintain historic structures and cultural landscapes, and protect archeological sites in the face of a predicted increase in visitation); issues surrounding visitor understanding; education and appreciation of park resources; and issues surrounding ensuring organizational effectiveness (such as identifying existing and potential partnerships with the state, federal, local, tribal, nonprofit organizations).

DATES: Any comments on the scope of issues to be addressed in the EIS can be received at any time after the publication of this notice in the FR. Public meetings regarding the CMP will be held during the scoping period. Specific dates, times, and locations will be made available in the local media, on the Trail Web site (<http://www.nps.gov/lecl>), on the NPS Planning, Environment and Public Comment (PEPC) Web site (<http://parkplanning.nps.gov/lecl>), or by contacting the Superintendent at the address and telephone number below.

ADDRESSES: Information on the planning process will be available from Mark Weekley, Superintendent, Lewis and Clark National Historic Trail, 601 Riverfront Drive, Omaha, Nebraska 68182, telephone 402-661-1806.

SUPPLEMENTARY INFORMATION: If you wish to comment on any issues associated with the CMP, you may submit your comments by any one of several methods. You may mail comments to Lewis and Clark National Historic Trail, 601 Riverfront Drive, Omaha, Nebraska 68182. You may provide comments electronically by entering them into the PEPC Web site at the address above. Finally, you may deliver comments to the Trail located at

601 Riverfront Drive, Omaha, Nebraska 68182.

Before including your address, telephone number, electronic mail address, or other personal identifying information in your comments, you should be aware that your entire comment (including your personal identifying information) may be made publicly available at any time. While you can ask us in your comments to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Dated: September 30, 2009.

Martin A. Sterkel,

Acting Regional Director, Midwest Region.

[FR Doc. E9-27519 Filed 11-19-09; 8:45 am]

BILLING CODE P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[LLCAD0000L14300000.DS0000]

Notice of Intent To Prepare an Environmental Impact Statement for the Proposed Desert Renewable Energy Conservation Plan, Kern, Los Angeles, San Bernardino, Inyo, Riverside, Imperial, San Diego, and Tulare Counties, CA and Possible Land Use Plan Amendment

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of Intent.

SUMMARY: In compliance with the National Environmental Policy Act (NEPA) of 1969, as amended, and the Federal Land Policy and Management Act of 1976, as amended, the Bureau of Land Management (BLM) California Desert District, Moreno Valley, California, intends to prepare an Environmental Impact Statement (EIS), which may include an amendment to the California Desert Conservation Area (CDCA) Plan of 1980, as amended, and by this notice is announcing the beginning of the scoping process to solicit public comments and identify issues.

DATES: This notice initiates the public scoping process for the EIS and possible plan amendment. Comments on issues may be submitted in writing until December 21, 2009. The dates and locations of any scoping meetings will be announced at least 15 days in advance through local media, newspapers, and the BLM Web site at: <http://www.blm.gov/ca/st/en/fo/cdd.html>. In order to be considered in the Draft EIS, all comments must be

received before the close of the scoping period or 15 days after the last public meeting, whichever is later. We will provide additional opportunities for public participation upon publication of the Draft EIS.

ADDRESSES: You may submit comments on issues and planning criteria related to the Desert Renewable Energy Conservation Plan (DRECP) by any of the following methods:

- *Web site:* <http://www.blm.gov/ca/st/en/fo/cdd.html>.

- *E-mail:* DRECP@blm.gov.

- *Fax:* (916) 978-4657.

- *Mail or hand delivery:* ATTN:

DRECP, BLM California State Office, 2800 Cottage Way, Suite W-1623, Sacramento, California 95825.

Documents pertinent to this proposal may be examined at the BLM California State office or the BLM California Desert District office, 22835 Calle San Juan de Los Lagos, Moreno Valley, California 92553-9046.

FOR FURTHER INFORMATION CONTACT: For further information or to have your name added to our mailing list, contact Amy Fesnock, Project Manager, telephone (916) 978-4646; address BLM California State Office, 2800 Cottage Way, Suite W-1623, Sacramento, California 95825; e-mail DRECP@blm.gov.

SUPPLEMENTARY INFORMATION: The BLM, along with the California Department of Fish and Game, the California Energy Commission, and the U.S. Fish and Wildlife Service, propose to develop the DRECP to advance State and Federal conservation goals in the Mojave and Colorado desert regions in California, while also facilitating the timely permitting of renewable energy projects under applicable State and Federal laws. Specifically, the planning goals for the DRECP include, but are not limited to, the following:

- Provide for the long-term conservation and management of identified species in the planning area;
 - Preserve, restore, and enhance natural communities and ecosystems that support identified species in the planning area;

- Build on the Competitive Renewable Energy Zones identified by the State's Renewable Energy Transmission Initiative that depict areas where renewable energy generation project permitting may be expedited;

- Identify the most appropriate locations in the planning area for the development of utility-scale renewable energy projects, taking into account potential impacts to threatened and endangered species, sensitive natural communities, and cultural resources;

- Coordinate and standardize mitigation and compensation requirements for renewable energy activities in the planning area; and
- Develop an efficient process for authorizing renewable energy projects in the planning area that results in greater conservation values than the process provided by project-by-project or species-by-species reviews.

The purpose of the public scoping process is to determine relevant issues that will influence the scope of the environmental analysis, including alternatives, and guide the process for developing the EIS. The BLM has identified the following preliminary issues: Special status species, mitigation measures for special status species, vegetation communities, cultural resources, special area designations, and areas of high potential for renewable energy development.

Authorization of this proposal may require amendment of the CDCA Plan. By this notice, the BLM is complying with requirements in 43 CFR 1610.2(c) to notify the public of potential amendments to land use plans, predicated on the findings of the EIS. If a land use plan amendment is necessary, the BLM will integrate the land use planning process with the NEPA process for this proposal.

The BLM will use the NEPA commenting process to satisfy the public involvement process for Section 106 of the National Historic Preservation Act (16 U.S.C. 470f), as provided for in 36 CFR 800.2(d)(3). Native American Tribal consultations will be conducted in accordance with policy and Tribal concerns will be given due consideration, including impacts on Indian trust assets. Federal, State, Tribes, and local agencies, along with other stakeholders that may be interested or affected by the BLM's decision on this project, are invited to participate in the scoping process and, if eligible, may request or be requested by the BLM to participate as a cooperating agency.

Before including your address, phone number, e-mail address, or other personal identifying information in your comment, you should be aware that your entire comment—including your personal identifying information—may be made publicly available at any time. While you can ask us in your comment to withhold your personal identifying information from public review, we cannot guarantee that we will be able to do so.

Authority: 40 CFR 1501.7 and 43 CFR 1610.2.

Tom Pogacnik,

Deputy State Director for Natural Resources.

[FR Doc. E9-27862 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-40-P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[LLCON01000 L0777 XX]

Notice of Public Meeting, BLM Colorado Northwest Resource Advisory Council, Correction, Cancellation of Meeting

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of public meeting; cancellation.

SUMMARY: The Bureau of Land Management (BLM) published a document in the **Federal Register** of April 28, 2009, notifying the public regarding meeting dates and locations for the BLM Colorado Northwest Resource Advisory Council (RAC). The meeting on December 3, 2009 has been cancelled.

SUPPLEMENTARY INFORMATION: The RAC meets in accordance with the Federal Land Policy and Management Act and the Federal Advisory Committee Act of 1972 (FACA), 5 U.S.C.

FOR FURTHER INFORMATION CONTACT: Jamie Connell, BLM Northwest Colorado District Manager, 2815 H Road, Grand Junction, CO 81506, 970-244-3000; or David Boyd, Public Affairs Specialist, 2300 River Frontage Road, Silt, CO 81652, 970-876-9008.

Dated: November 16, 2009.

Jamie Connell,

Designated Federal Official.

[FR Doc. E9-27938 Filed 11-19-09; 8:45 am]

BILLING CODE P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[L12100000.PH0000 CO 912]

Notice of Intent To Solicit Nominations for the Dominguez-Escalante National Conservation Area Advisory Council

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice.

SUMMARY: The Secretary of the Interior (Secretary) is directed by the Omnibus Public Lands Act of 2009 to establish the Dominguez-Escalante National

Conservation Area (D-E NCA) Advisory Council (Council). The Secretary is requesting nominations for 10 members to sit on the Council. The Council will advise the Secretary, through the Bureau of Land Management (BLM), on matters regarding the preparation and implementation of the D-E NCA Resource Management Plan (RMP).

DATES: Submit nomination packages on or before December 21, 2009.

ADDRESSES: Send completed Council nominations to D-E NCA Interim Manager, Grand Junction Field Office, 2815 H Road, Grand Junction, Colorado 81506. Nomination forms may be obtained at the Grand Junction Field Office at the above address or at the Uncompahgre Field Office, Bureau of Land Management, 2465 S. Townsend Avenue, Montrose, Colorado 81401.

FOR FURTHER INFORMATION CONTACT: Katie A. Stevens, D-E NCA Interim Manager, (970) 244-3049, Katie_A_Stevens@blm.gov.

SUPPLEMENTARY INFORMATION: The D-E NCA and Dominguez Canyon Wilderness Area, located within the D-E NCA, was established by the Omnibus Public Land Management Act of 2009, Public Law 111-11 (Act). The D-E NCA is comprised of approximately 209,610 acres of public land, including approximately 66,280 acres of wilderness, located in Delta, Montrose, and Mesa counties. The purposes of the D-E NCA are to conserve and protect, for the benefit and enjoyment of present and future generations, the unique and important resources and values of the land. These resources and values include the geological, cultural, archaeological, paleontological, natural, scientific, recreational, wilderness, wildlife, riparian, historical, educational, and scenic resources of the public lands, and the water resources of area streams that are necessary to support aquatic, riparian, and terrestrial species and communities. The Act also calls for the establishment of the D-E NCA Council, comprised of 10 members, to advise the Secretary, through the BLM, on matters regarding the preparation and implementation of an RMP for the area. These 10 members shall include, to the extent practicable:

- (1) One member appointed after considering the recommendations of the Mesa County Commission;
- (2) One member appointed after considering the recommendations of the Montrose County Commission;
- (3) One member appointed after considering the recommendations of the Delta County Commission;
- (4) One member appointed after considering the recommendations of the

permittees holding grazing allotments within the D-E NCA; and

(5) Six members who reside in, or within reasonable proximity to Mesa, Delta, or Montrose Counties, Colorado, with backgrounds that reflect:

(A) The purposes for which the D-E NCA was established; and

(B) The interests of the stakeholders that are affected by the planning and management of the D-E NCA.

Any individual or organization may nominate one or more persons to serve on the Council. Individuals may nominate themselves for Council membership. Nomination forms may be obtained from the BLM Grand Junction or Uncompahgre Field Offices, or may be downloaded from the following Web site: <http://www.blm.gov/co/st/en/fo/denca.html>.

Nomination packages must include a completed nomination form, letters of reference from the represented interests or organizations, as well as any other information relevant to the nominee's qualifications.

The Grand Junction and Uncompahgre Field Offices will review the nomination packages in coordination with the affected counties and the Governor of Colorado before forwarding its recommendations to the Secretary, who will make the appointments.

The Council shall be subject to the Federal Advisory Committee Act, 5 U.S.C. App. 2; and the Federal Land Policy and Management Act of 1976, 43 U.S.C. 1701 *et seq.*

Authority: Sec. 2407 of Public Law 111-11.

Dated: November 2, 2009.

David B. Hunsaker,

Acting State Director.

[FR Doc. E9-27865 Filed 11-19-09; 8:45 am]

BILLING CODE 4310-JB-P

DEPARTMENT OF JUSTICE

Notice of Lodging of Consent Decree Under the Comprehensive Environmental Response Compensation, and Liability Act

Notice is hereby given that on November 13, 2009, a proposed Consent Decree (the "Decree") in *United States v. Cabot Corporation, et al.*, Civil Action No. 1:09-cv-5783, was lodged with the United States District Court for the District of New Jersey.

In a complaint, filed simultaneously with the Decree, the United States alleges that Cabot Corporation, KB Alloys, Inc., Shieldalloy Metallurgical Corporation, E.I. du pont de Nemours and Company, International Wire

Group, Inc. and its subsidiary Omega Wire, Inc., are liable pursuant to Section 107(a)(3) of the Comprehensive Environmental Response, Compensation and Liability Act ("CERCLA"), 42 U.S.C. 9607(a)(3), for response costs incurred by the Environmental Protection Agency ("EPA") in cleaning up the Pioneer Smelting Superfund Site located at Factory Road, Route 532, in Chatsworth, New Jersey.

Pursuant to the Decree, the parties will jointly and severally be responsible for paying the United States \$750,000 to resolve any claim the United States has associated with costs incurred by EPA at the Pioneer Smelting Superfund Site.

The Department of Justice will receive, for a period of thirty (30) days from the date of this publication, comments relating to the Decree. Comments should be addressed to the Assistant Attorney General, Environment and Natural Resources Division, and either e-mailed to pubcomment-ees.enrd@usdoj.gov or mailed to P.O. Box 7611, U.S. Department of Justice, Washington, DC 20044-7611, and should refer to *United States v. Cabot Corporation et al.*, D.J. Ref. 90-11-2-09344/1.

During the public comment period, the Decree may be examined on the following Department of Justice Web site, http://www.usdoj.gov/enrd/Consent_Decrees.html. A copy of the Decree may also be obtained by mail from the Consent Decree Library, P.O. Box 7611, U.S. Department of Justice, Washington, DC 20044-7611 or by faxing or e-mailing a request to Tonia Fleetwood (tonia.fleetwood@usdoj.gov), fax no. (202) 514-0097, phone confirmation number (202) 514-1547. In requesting a copy from the Consent Decree Library, please enclose a check in the amount of \$7.75 (25 cents per page reproduction cost) payable to the U.S. Treasury or, if by e-mail or fax, forward a check in that amount to the Consent Decree Library at the stated address.

Maureen Katz,

Assistant Section Chief, Environmental Enforcement Section, Environment and Natural Resources Division.

[FR Doc. E9-27866 Filed 11-19-09; 8:45 am]

BILLING CODE P

DEPARTMENT OF JUSTICE**Drug Enforcement Administration****[Docket No. DEA-326F]****Established Assessment of Annual Needs for the List I Chemicals Ephedrine, Pseudoephedrine, and Phenylpropanolamine for 2010****AGENCY:** Drug Enforcement Administration (DEA), Justice.**ACTION:** Notice of Assessment of Annual Needs for 2010.

SUMMARY: This notice establishes the initial 2010 Assessment of Annual Needs for certain List I chemicals in accordance with the Combat Methamphetamine Epidemic Act of 2005 (CMEA), enacted on March 9, 2006.

DATES: *Effective Date:* November 20, 2009.

FOR FURTHER INFORMATION CONTACT: Christine A. Sannerud, Ph.D., Chief, Drug and Chemical Evaluation Section, Drug Enforcement Administration, 8701 Morrisette Drive, Springfield, Virginia 22152, *Telephone:* (202) 307-7183.

SUPPLEMENTARY INFORMATION: Section 713 of the Combat Methamphetamine Epidemic Act of 2005 (Title VII of Pub. L. 109-177) (CMEA) amended Section 306 of the Controlled Substances Act (CSA) (21 U.S.C. 826) by adding ephedrine, pseudoephedrine, and phenylpropanolamine to existing language to read as follows: "The Attorney General shall determine the total quantity and establish production quotas for each basic class of controlled substance in schedules I and II and for ephedrine, pseudoephedrine, and phenylpropanolamine to be manufactured each calendar year to provide for the estimated medical, scientific, research, and industrial needs of the United States, for lawful export requirements, and for the establishment and maintenance of reserve stocks." Further, section 715 of the CMEA amended 21 U.S.C. 952 "Importation of Controlled Substances" by adding the same List I chemicals to the existing language in paragraph (a), and by adding a new paragraph (d) to read as follows:

(a) Controlled substances in schedule I or II and narcotic drugs in schedule III, IV, or V; exceptions

It shall be unlawful to import into the customs territory of the United States from any place outside thereof (but within the United States), or to import into the United

States from any place outside thereof, any controlled substance in schedule I or II of subchapter I of this chapter, or any narcotic drug in schedule III, IV, or V of subchapter I of this chapter, or ephedrine, pseudoephedrine, and phenylpropanolamine, except that—

(1) such amounts of crude opium, poppy straw, concentrate of poppy straw, and coca leaves, and of ephedrine, pseudoephedrine, and phenylpropanolamine, as the Attorney General finds to be necessary to provide for medical, scientific, or other legitimate purposes * * * may be so imported under such regulations as the Attorney General shall prescribe.

(d)(1) With respect to a registrant under section 958 who is authorized under subsection (a)(1) to import ephedrine, pseudoephedrine, or phenylpropanolamine, at any time during the year the registrant may apply for an increase in the amount of such chemical that the registrant is authorized to import, and the Attorney General may approve the application if the Attorney General determines that the approval is necessary to provide for medical, scientific, or other legitimate purposes regarding the chemical.

Editor's Note: This excerpt of the amendment is published for the convenience of the reader. The official text is published at 21 U.S.C. 952(a) and (d)(1).

Background and Legal Authority

Section 713 of the CMEA of 2005 (Title VII of Pub. L. 109-177) amended section 306 of the CSA (21 U.S.C. 826) to require that the Attorney General establish quotas to provide for the annual needs for ephedrine, pseudoephedrine, and phenylpropanolamine. Section 715 of the CMEA amended 21 U.S.C. 952 by adding ephedrine, pseudoephedrine, and phenylpropanolamine to the existing language concerning importation of controlled substances.

The 2010 Assessment of Annual Needs represents those quantities of ephedrine, pseudoephedrine, and phenylpropanolamine which may be manufactured domestically and/or imported into the United States in 2010 to provide adequate supplies of each chemical for: The estimated medical, scientific, research, and industrial needs of the United States; lawful export requirements; and the establishment and maintenance of reserve stocks.

The responsibility for establishing the assessment has been delegated to the Administrator of the DEA by 28 CFR 0.100. The Administrator, in turn, has re-delegated this function to the Deputy Administrator, pursuant to 28 CFR 0.104.

On September 14, 2009, a notice entitled, "Assessment of Annual Needs

for the List I Chemicals Ephedrine, Pseudoephedrine, and Phenylpropanolamine for 2010" was published in the **Federal Register** (74 FR 47021). That notice proposed the 2010 Assessment of Annual Needs for ephedrine (for sale), ephedrine (for conversion), pseudoephedrine (for sale), phenylpropanolamine (for sale) and phenylpropanolamine (for conversion). All interested persons were invited to comment on or object to the assessments on or before October 14, 2009.

Comments Received

DEA did not receive any comments to the Assessment of Annual Needs for ephedrine (for sale), ephedrine (for conversion), pseudoephedrine (for sale), phenylpropanolamine (for sale) and phenylpropanolamine (for conversion). DEA is finalizing the assessments for these List I chemicals based on information contained in additional applications for 2010 import, manufacturing and procurement quotas provided by DEA registered importers and manufacturers whose quota applications were received as of October 21, 2009. DEA is providing the data used in developing the established assessments for each of the listed chemicals. DEA also notes that the Assessment of Annual Needs may be adjusted at a later date pursuant to 21 CFR 1315.13.

Underlying Data and DEA's Analysis

In determining the final 2010 assessments, DEA has considered the total net disposals (i.e., sales) of the List I chemicals for the current and preceding two years, actual and estimated inventories, projected demand (2010), industrial use, and export requirements from data provided by DEA registered manufacturers and importers in procurement quota applications (DEA 250), from manufacturing quota applications (DEA 189), and from import quota applications (DEA 488).¹

DEA further considered trends as derived from information provided in applications for import, manufacturing, and procurement quotas and in import and export declarations. DEA notes that the inventory, acquisitions (purchases) and disposition (sales) data provided by DEA registered manufacturers and importers reflects the most current information available.

Ephedrine Data

¹ Applications and instructions for procurement, import and manufacturing quotas can be found at

http://www.deadiversion.usdoj.gov/quotas/quota_apps.htm.

EPHEDRINE (FOR SALE) DATA FOR 2010 ASSESSMENT OF ANNUAL NEEDS
[Kilograms]

Ephedrine	2007	2008	2009	2010 Request
Sales* (DEA 250)	2,743	2,508	2,431	2,861
Imports** (DEA 488)	9,595	1,686	2,160	1,552
Export Declarations (DEA 486)	168	91	10	n/a
Inventory* (DEA 250)	1,332	592	181	n/a
IMS*** (NSP)	1,235	1,460	n/a	n/a

* Reported sales and inventory from applications for 2010 procurement quotas (DEA 250).

** Reported imports from applications for 2010 import quotas (DEA 488).

*** IMS Health, IMS National Sales Perspectives™, January 2007 to December 2008, Retail and Non-Retail Channels, Data Extracted October 21, 2009.

Ephedrine Analysis

DEA calculated the established 2010 Assessment of Annual Needs for ephedrine using the calculation developed to determine the 2009 Assessment of Annual Needs. This calculation considers the criteria defined in 21 U.S.C. 826: estimated medical, scientific, research, and industrial needs of the United States; lawful export requirements; and the establishment and maintenance of reserve stocks.

As of October 21, 2009, DEA registered manufacturers of dosage form products containing ephedrine requested the authority to purchase a total of 2,861 kg ephedrine (for sale) in 2010. DEA registered manufacturers of ephedrine reported sales totaling approximately 2,508 kg in 2008 and 2,431 kg in 2009; this represents a 3% decrease in sales reported by these firms from 2008 to 2009. Additionally, exports of ephedrine products from the United States as reported on export

declarations (DEA 486) totaled 91 kg in 2008 and 10 kg in 2009; this represents a 90% decrease from levels observed in 2008. The average of the 2008 and 2009 exports of ephedrine products is approximately 51 kg. DEA also considered information on trends in the national rate of net disposals from sales data provided by IMS Health's National Sales Perspective™ (NSP) database. IMS NSP data reported the average sales volume of ephedrine for the calendar years 2007 and 2008 to be approximately 1,348 kg. DEA notes that the 2009 sales figure reported by manufacturers (2,431 kg) is higher than the average sales reported by IMS for the previous two years (1,348 kg). This is expected because a manufacturer's reported sales include quantities which are necessary to provide reserve stocks for distributors and retailers. DEA, in considering the manufacturer's reported sales, thus believes that 2,431 kg fairly represents the U.S. sales of ephedrine for 2010 and that 51 kg fairly represents the export requirements of ephedrine.

For the establishment and maintenance of reserve stocks, DEA notes that 21 CFR 1315.24 allows for an inventory allowance (reserve stock) of 50% of a manufacturer's estimated sales. DEA also considered the estimated 2009 year end inventory as reported by DEA registrants in determining the inventory allowance.

DEA calculated the ephedrine (for sale) assessment by the following methodology:

2009 sales + reserve stock + export requirement – existing inventory = AAN

$2,431 + (50\% * 2,431) + 51 - 181 = 3,517$
kg ephedrine (for sale) for 2010

This calculation suggests that DEA's Assessment of Annual Needs for ephedrine should be established as 3,600 kg. Accordingly, DEA is establishing the 2010 Assessment of Annual Needs for ephedrine (for sale) at 3,600 kg.

Phenylpropanolamine (for Sale) Data

PHENYLPROPANOLAMINE (FOR SALE) DATA FOR 2010 ASSESSMENT OF ANNUAL NEEDS
[Kilograms]

Phenylpropanolamine (for sale)	2007	2008	2009	2010 Request
Sales* (DEA 250)	3,770	4,274	4,638	6,288
Imports** (DEA 488)	73	79	134	263
Export Declarations (DEA 486)	1,002	0	3	n/a
Inventory* (DEA 250)	3,597	2,093	596	n/a

* Reported sales and inventory from applications for 2010 procurement quotas (DEA 250) received as of October 21, 2009.

** Reported imports from applications for 2010 import quotas (DEA 488) received as of October 21, 2009.

Phenylpropanolamine (for Sale) Analysis

DEA utilized the same general methodology and calculation to establish the assessment for phenylpropanolamine (for sale) as was described for the assessment of ephedrine (for sale), above.

As of October 21, 2009, DEA registered manufacturers of dosage form products containing

phenylpropanolamine requested the authority to purchase 6,288 kg phenylpropanolamine (for sale) in 2010. DEA registered manufacturers of phenylpropanolamine reported sales totaling approximately 4,274 kg in 2008 and 4,638 kg in 2009; this represents an 8% increase in sales reported by these firms from 2008 to 2009. Additionally, exports of phenylpropanolamine products from the U.S. as reported on

export declarations (DEA 486) totaled 0 kg in 2008 and 3 kg in 2009; this represents a 3 kg increase from levels observed in 2008. The average of the 2008 and 2009 exports of phenylpropanolamine products is approximately 2 kg. DEA thus believes that 4,638 kg fairly represents the U.S. sales of phenylpropanolamine for 2010 and that 2 kg fairly represents the export requirements of phenylpropanolamine.

DEA notes that phenylpropanolamine is sold primarily as a veterinary product for the treatment for canine incontinence and is not approved for human consumption. IMS Health's NSP Data does not capture sales of phenylpropanolamine to these channels and is therefore not included. DEA calculated the phenylpropanolamine (for sale)

assessment by the following methodology:
 2009 sales + reserve stock + export requirement - existing inventory = AAN
 $4,638 + (50\% * 4,638) + 2 - 596 = 6,363$ kg phenylpropanolamine (for sale) for 2010

This calculation suggests that DEA's 2010 Assessment of Annual Needs for phenylpropanolamine (for sale) should be established as 6,400 kg. Accordingly, DEA is establishing the 2010 Assessment of Annual Needs for phenylpropanolamine (for sale) at 6,400 kg.

Pseudoephedrine (for Sale) Data

PSEUDOEPHEDRINE (FOR SALE) DATA FOR 2010 ASSESSMENT OF ANNUAL NEEDS

[Kilograms]

Pseudoephedrine (for sale)	2007	2008	2009	2010 Request
Sales* (DEA 250)	238,608	223,196	286,516	225,116
Sales* (DEA 189)	100,300	64,781	33,600	32,760
Imports** (DEA 488)	232,822	170,995	267,808	233,569
Export Declarations (DEA 486)	42,132	47,194	25,526	n/a
Inventory* (DEA 250)	135,097	119,515	62,748	n/a
IMS*** (NSP)	180,204	149,159	n/a	n/a

* Reported sales and inventory from applications for 2010 procurement quotas (DEA 250) and manufacturing quotas (DEA 189) received as of October 21, 2009.

** Reported imports from applications for 2010 import quotas (DEA 488) received as of October 21, 2009.

*** IMS Health, IMS National Sales Perspectives™, January 2007 to December 2008, Retail and Non-Retail Channels, Data Extracted October 21, 2009.

Pseudoephedrine (for Sale) Analysis

DEA utilized the same general methodology and calculations to establish the assessment for pseudoephedrine (for sale) as were described for the assessment of ephedrine (for sale), above.

As of October 21, 2009, DEA registered manufacturers of dosage form products containing pseudoephedrine requested the authority to purchase 225,116 kg pseudoephedrine. DEA registered manufacturers of pseudoephedrine reported sales totaling approximately 223,196 kg in 2008 and 286,516 kg in 2009; this represents a 22% increase in sales reported by these firms from 2008 to 2009. During the same period exports of pseudoephedrine products from the U.S. as reported on export declarations (DEA 486) totaled 47,194 kg in 2008 and

25,526 kg in 2009; this represents a 54% decrease from levels observed in 2008. The average of the 2008 and 2009 exports is 36,360 kg. Additionally, DEA considered information on trends in the national rate of net disposals from sales data provided by IMS Health. IMS NSP data reported the average retail sales volume of pseudoephedrine for the calendar years 2007 and 2008 to be approximately 164,682 kg. DEA thus believes that 286,516 kg of sales reported by manufacturers fairly represents the U.S. sales of pseudoephedrine for 2010 and that 36,360 kg fairly represents the export requirements of pseudoephedrine. DEA notes that manufacturer reported sales for 2009 (286,516 kg) are higher than the average retail sales reported by IMS for the previous two years (164,682 kg). This is expected because a manufacturer's reported sales include

quantities which are necessary to provide reserve stocks for distributors and retailers.

DEA calculated the pseudoephedrine (for sale) assessment by the following methodology:

2009 sales + reserve stock + export requirement - existing inventory = AAN
 $286,516 + (50\% * 286,516) + 36,360 - 62,748 = 403,386$ kg pseudoephedrine (for sale) for 2010.

This calculation suggests that DEA's 2010 Assessment of Annual Needs for pseudoephedrine (for sale) should be established at 404,000 kg. Accordingly, DEA is establishing the 2010 Assessment of Annual Needs for pseudoephedrine (for sale) at 404,000 kg.

Phenylpropanolamine (for Conversion) Data

PHENYLPROPANOLAMINE (FOR CONVERSION) DATA FOR 2010 ASSESSMENT OF ANNUAL NEEDS

[Kilograms]

Phenylpropanolamine (for conversion)	2007	2008	2009	2010 Request
Sales* (DEA 250)	3,621	10,834	13,582	14,900
Imports** (DEA 488)	1,000	3,225	6,514	7,108
Export Declarations (DEA 486)	0	0	0	n/a
Inventory* (DEA 250)	3,581	5,533	4,103	n/a
APQ Amphetamine***	17,000	22,000	22,000	n/a

* Reported sales and inventory from applications for 2010 procurement quotas (DEA 250) received as of October 21, 2009.

** Reported imports from applications for 2010 import quotas (DEA 488) received as of October 21, 2009.

*** Amphetamine Aggregate Production Quota History http://www.deadiversion.usdoj.gov/quotas/quota_history.htm.

Phenylpropanolamine (for Conversion) Analysis

As of October 21, 2009, DEA registered manufacturers of phenylpropanolamine (for conversion) requested the authority to purchase a total of 14,900 kg phenylpropanolamine for the manufacture of amphetamine. DEA registered manufacturers of phenylpropanolamine reported sales of phenylpropanolamine totaling approximately 10,834 kg in 2008 and 13,582 kg in 2009; this represent a 20% increase in sales reported by these firms from 2008 to 2009. There were no reported exports of phenylpropanolamine (for conversion). DEA has not received any requests to synthesize phenylpropanolamine in

2010. DEA has concluded that the 2009 sales of phenylpropanolamine (for conversion), 13,582 kg, fairly represents U.S. requirements for 2010 and zero kg fairly represents the export requirements of phenylpropanolamine (for conversion).

Phenylpropanolamine is used in the production of legitimate amphetamine products. DEA has established an Aggregate Production Quota (APQ) for amphetamine of 22,000 kg for 2009. DEA notes amphetamine is primarily manufactured by the conversion of the schedule II controlled substance phenylacetone to amphetamine. DEA did not consider this alternative synthesis route in the 2009 Assessment of Annual Needs for phenylpropanolamine (for conversion).

DEA calculated the phenylpropanolamine (for conversion) for the manufacture of amphetamine as follows:

$$(2009 \text{ sales}) + \text{reserve stock} + \text{export requirement} - \text{inventory} = \text{AAN}$$

$$(13,582) + 50\% * (13,582) + 0 - 4,103 = 16,270 \text{ kg PPA (for conversion) for 2010}$$

This calculation suggests that DEA's 2010 Assessment of Annual Needs for phenylpropanolamine (for conversion) should be established at 16,500 kg. Accordingly, DEA is establishing the 2010 Assessment of Annual Needs for phenylpropanolamine (for conversion) at 16,500 kg.

Ephedrine (for Conversion) Data

EPHEDRINE (FOR CONVERSION) DATA FOR 2010 ASSESSMENT OF ANNUAL NEEDS
[Kilograms]

Ephedrine (for conversion)	2007	2008	2009	2010 Request
Sales* (DEA 250)	99,622	64,522	40,403	40,646
Imports** (DEA 488)	99,594	64,128	39,897	40,000
Inventory* (DEA 250)	13	160	254	n/a
APQ Methamphetamine***	3,130	3,130	3,130	n/a

* Reported sales and inventory from applications for 2010 procurement quotas (DEA 250) and manufacturing quotas (DEA 189) received as of October 21, 2009.

** Reported imports from applications for 2010 import quotas (DEA 488) received as of October 21, 2009.

*** Methamphetamine Aggregate Production Quota History http://www.deadiversion.usdoj.gov/quotas/quota_history.htm.

Ephedrine (for Conversion) Analysis

As of October 21, 2009, DEA registered manufacturers of ephedrine (for conversion) requested the authority to purchase a total of 40,646 kg ephedrine (for conversion) for the manufacture of two substances: methamphetamine and pseudoephedrine.

DEA considered the ephedrine (for conversion) requirements for the manufacture of methamphetamine and pseudoephedrine. DEA has determined that the established assessments for the manufacture of these two substances are the best indicators of the need for ephedrine (for conversion). The assessment of need for methamphetamine was determined by DEA as the Aggregate Production Quota (APQ) for methamphetamine. DEA determined that the estimated sales of pseudoephedrine, as referenced in the Assessment of Annual Needs (AAN) for pseudoephedrine, represents the need for pseudoephedrine. Reported sales of ephedrine (for conversion) are included as reference to DEA's methodology.

DEA further considered the reported conversion yields of these substances. DEA registered manufacturers reported a conversion yield of 39% for the

synthesis of methamphetamine from ephedrine. DEA cannot disclose the conversion yield for the synthesis of pseudoephedrine because this information is proprietary to the one manufacturer involved in this type of manufacturing.

DEA calculated the ephedrine (for conversion) assessment by the following methodology:

$$\text{methamphetamine requirement} + \text{pseudoephedrine requirement} = \text{AAN}$$

DEA calculated the ephedrine (for conversion) requirement for the manufacture of methamphetamine as follows:

$$(2009 \text{ APQ methamphetamine}/39\% \text{ yield}) + \text{reserve stock} - \text{inventory} = \text{ephedrine (for manufacture of methamphetamine)}$$

$$(3,130/39\% \text{ yield}) + 50\% * (3,130/39\% \text{ yield}) - 46 = 11,993 \text{ kg}$$

The calculation for the ephedrine (for conversion) requirement for the manufacture of pseudoephedrine leads to a result of 63,157 kg. DEA cannot provide the details of the calculation because this would reveal the conversion yield for the synthesis of pseudoephedrine, which is proprietary to the one manufacturer involved in this

type of manufacturing. Therefore, the assessment for ephedrine was determined by the sum total of the ephedrine (for conversion) requirements as described by the following methodology:

$$\text{methamphetamine requirement} + \text{pseudoephedrine requirement} = \text{AAN}$$

$$11,993 + 63,157 = 75,150 \text{ kg ephedrine (for conversion) for 2010}$$

This calculation suggests that DEA's 2010 Assessment of Annual Needs for ephedrine (for conversion) should be established at 75,000 kg. Accordingly, DEA is establishing the 2010 Assessment of Annual Needs for ephedrine (for conversion) at 75,000 kg.

Conclusion

DEA did not receive any comments on its Assessment of Annual Needs for ephedrine (for sale), ephedrine (for conversion), pseudoephedrine (for sale), phenylpropanolamine (for sale) and phenylpropanolamine (for conversion). DEA is finalizing the assessments for these List I chemicals based on information contained in additional applications for 2010 import, manufacturing and procurement quotas provided by DEA registered importers

and manufacturers whose quota applications were received as of October 21, 2009.

Therefore, under the authority vested in the Attorney General by Section 306 of the CSA (21 U.S.C. 826), and delegated to the Administrator of the DEA by 28 CFR 0.100, and redelegated to the Deputy Administrator pursuant to 28 CFR 0.104, the Deputy Administrator hereby orders that the 2010 Assessment of Annual Needs for ephedrine, pseudoephedrine, and phenylpropanolamine, expressed in kilograms of anhydrous acid or base, be established as follows:

List I chemical	Established 2010 assessment of annual needs
Ephedrine (for sale)	3,600
Phenylpropanolamine (for sale)	6,400
Pseudoephedrine (for sale)	404,000
Phenylpropanolamine (for conversion)	16,500
Ephedrine (for conversion)	75,000

The Office of Management and Budget has determined that notices of quotas are not subject to centralized review under Executive Order 12866.

This action does not preempt or modify any provision of state law; nor does it impose enforcement responsibilities on any state; nor does it diminish the power of any state to enforce its own laws. Accordingly, this action does not have any federalism implications warranting the application of Executive Order 13132.

The Deputy Administrator hereby certifies that this action will not have a significant economic impact upon a substantial number of small entities whose interests must be considered under the Regulatory Flexibility Act, 5 U.S.C. 601–612. The establishment of Assessment of Annual Needs for ephedrine, pseudoephedrine, and phenylpropanolamine is mandated by law. The assessments are necessary to provide for the estimated medical, scientific, research and industrial needs of the United States; for lawful export requirements; and the establishment and maintenance of reserve stocks. Accordingly, the Deputy Administrator has determined that this action does not require a regulatory flexibility analysis.

This action meets the applicable standards set forth in Sections 3(a) and 3(b)(2) of Executive Order 12988 Civil Justice Reform.

This action will not result in the expenditure by State, local, and tribal governments, in the aggregate, or by the

private sector, of \$120,000,000 or more (adjusted for inflation) in any one year, and will not significantly or uniquely affect small governments. Therefore, no actions were deemed necessary under the provisions of the Unfunded Mandates Reform Act of 1995.

This action is not a major rule as defined by Section 804 of the Small Business Regulatory Enforcement Fairness Act of 1996 (Congressional Review Act). This action will not result in an annual effect on the economy of \$100,000,000 or more; a major increase in costs or prices; or significant adverse effects on competition, employment, investment, productivity, innovation, or on the ability of United States-based companies to compete with foreign-based companies in domestic and export markets.

Dated: November 11, 2009.

Michele M. Leonhart,

Deputy Administrator.

[FR Doc. E9–27890 Filed 11–19–09; 8:45 am]

BILLING CODE 4410–09–P

DEPARTMENT OF LABOR

Bureau of Labor Statistics

Solicitation of Comments on a Proposal To Revise Method for Estimation of Monthly Labor Force Statistics for Certain Subnational Areas

AGENCY: Bureau of Labor Statistics, Labor.

ACTION: Notice of solicitation of comments.

SUMMARY: The Department of Labor, through the Bureau of Labor Statistics (BLS), is responsible for the development and publication of local area labor force statistics. This program includes the issuance of monthly estimates of the labor force, employment, unemployment, and the unemployment rate for each State and labor market area in the nation. A hierarchy of estimation methods is used to produce the 7,300 estimates covered by the Local Area Unemployment Statistics (LAUS) program (<http://thomas.loc.gov/bss/d111/d111laws.html>), based on the availability and quality of data from the Current Population Survey (CPS). The strongest estimating method—signal-plus-noise models with real-time benchmarking for current estimation and historical benchmarking—is employed for all States and the District of Columbia, the Los Angeles-Long Beach-Glendale, CA metropolitan division, New York City, and the

respective balances of New York and California. Models are also employed for five additional substate areas and their State balances. The areas are: the Chicago-Naperville-Joliet, IL metropolitan division; the Cleveland-Elyria-Mentor, OH metropolitan area; the Detroit-Warren-Livonia, MI metropolitan area; the Miami-Miami Beach-Kendall, FL metropolitan division; and the Seattle-Bellevue-Everett, WA metropolitan division.

As part of a program of continuing improvements in LAUS methodology, BLS is proposing the implementation of smoothed-seasonally-adjusted series for current and historical estimates. This approach is an innovative alternative to an annual historical benchmark for seasonally-adjusted State estimates that will address longstanding issues related to end-of-year revision, and also will enhance the analytical capability of the estimates.

BLS proposes to implement the revised methodology beginning with January 2010 current estimates, with historical estimates revised to 1976 for States, the District of Columbia, Los Angeles, New York City, and the respective balances of California and New York. The five other substate model estimates will be revised back to 1983.

DATES: Submit written comments on or before December 21, 2009.

ADDRESSES: Send comments to Sharon Brown, Division Chief, Division of Local Area Unemployment Statistics, Bureau of Labor Statistics, Room 4675, 2 Massachusetts Avenue, NE., Washington, DC 20212, by FAX at 202–691–6459, or by e-mail at Brown.Sharon@bls.gov.

FOR FURTHER INFORMATION CONTACT: Sharon Brown, Division Chief, Division of Local Area Unemployment Statistics, Bureau of Labor Statistics, Room 4675, 2 Massachusetts Avenue, NE., Washington DC 20212, by telephone at 202–691–6390, or by e-mail at LAUSRM@bls.gov.

SUPPLEMENTARY INFORMATION:

I. Background

The Department of Labor, through the BLS, has been responsible for the development and publication of local area labor force statistics since 1972. In 1978, the BLS broadened the use data from the CPS in the LAUS program by extending the annual reliability criterion to monthly data. Monthly CPS levels were used directly for the 10 largest States, two substate areas (New York City, Los Angeles), and the respective balances of New York and California. In 1985, the sample redesign

and other efficiencies improved the reliability of CPS data at the State level, resulting in the current criterion on monthly and annual average data of an 8 percent coefficient of variation on the level of unemployment when the unemployment rate is 6 percent. In addition, North Carolina joined the group of direct-use States. In 1989, variable coefficient time-series models for monthly estimation of State employment and unemployment were introduced for 39 States and the District of Columbia. Further improvement was effected with the implementation of signal-plus-noise models in 1994. These models rely heavily on monthly CPS data, as well as current wage and salary employment and unemployment insurance statistics. State labor force estimation for the direct-use States was based the time series modeling approach beginning in January 1996.

Improvements introduced with the redesign in January 2005 ensured that State estimates add to the national estimates of employment and unemployment each month, through real-time benchmarking. In doing so, the benchmark changed from annual State-level estimates of employment and unemployment to monthly national estimates of these measures. In this way, economic shocks are reflected in the State estimates on a real-time basis, and end-of-year revisions are significantly smaller.

Historical benchmarking is part of the annual processing activities performed on the models. The first two steps, revision of inputs and model re-estimation, are the same for both the not-seasonally-adjusted (NSA) series and the seasonally-adjusted series. The final step, benchmarking to historical control totals, differs by series. The NSA estimates are benchmarked to monthly Division model controls which have been controlled to monthly national CPS estimates. This ensures that the monthly State NSA estimates sum to the national CPS estimates. The annual average of the NSA estimates is used to control the monthly seasonally-adjusted model estimates. This process preserves the underlying smoothness in the model estimates that would be lost by applying the monthly benchmarking procedure.

However, the current procedure had an unanticipated impact on the historical benchmarking for the seasonally-adjusted estimates during 2008. Unemployment rose steeply in the nation and all States during 2008. The benchmark methodology that required the use of the annual average as the historical control total for the seasonally-adjusted estimates meant that unemployment rates were adjusted

downward during the latter months of 2008. This impacted comparisons with January 2009 unemployment estimates that continued to reflect the steep economic decline. In addition to issues with historical benchmarking, the monthly real-time benchmarking procedure introduces volatility into the current seasonally-adjusted estimates, producing estimates with spurious turning points that are difficult to explain to data users.

II. Current Action

To address these serious issues, the BLS proposes modifying the procedures for the seasonally-adjusted estimates and implementing a smoothing methodology for both current and historical seasonally-adjusted series. Smoothing the current series will reduce the number of spurious turning points in the estimates. For historical estimates, the first two steps in annual processing: revising model inputs and re-estimating the series, are unchanged. The last step, benchmarking to control totals, will be revised for the seasonally-adjusted estimates. The use of the annual average of the NSA series as the control total will be dropped. Instead, as in current monthly estimation, the historical seasonally-adjusted series will be adjusted by the same pro-rata factor used in adjusting the NSA estimates to the national control totals. Since the pro-rata factors fluctuate from month-to-month, this procedure will introduce additional variability into the historical series, which could dominate the monthly change in the benchmarked series. Smoothing the series following the application of the pro-rata adjustment will reduce the volatility added. The smoother selected is the Henderson Trend Filter (H13).

Detailed descriptions of the current and proposed approaches are available from the office listed above.

III. Desired Focus of Comments

Comments and recommendations are requested from the public on the use of the Henderson Trend Filter (H13) to smooth the LAUS current and historical seasonally adjusted estimates.

Signed at Washington, DC, this 17th day of November 2009.

Kimberley Hill,

Acting Chief, Division of Management Systems, Bureau of Labor Statistics.

[FR Doc. E9-27930 Filed 11-19-09; 8:45 am]

BILLING CODE 4510-24-P

DEPARTMENT OF LABOR

Employment and Training Administration

Request for Certification of Compliance—Rural Industrialization Loan and Grant Program

AGENCY: Employment and Training Administration, Labor.

ACTION: Notice.

SUMMARY: The Employment and Training Administration is issuing this notice to announce the receipt of a "Certification of Non-Relocation and Market and Capacity Information Report" (Form 4279-2) for the following:

Applicant/Location: Legacy Senior Care Group, LLC/Richfield, Ohio.

Principal Product/Purpose: The loan, guarantee, or grant application is to enable a new business venture to construct and manage an assisted living facility that also offers memory care. The NAICS industry code for this enterprise is: 623311 Continuing Care Retirement Communities.

DATES: All interested parties may submit comments in writing no later than December 4, 2009. Copies of adverse comments received will be forwarded to the applicant noted above.

ADDRESSES: Address all comments concerning this notice to Anthony D. Dais, U.S. Department of Labor, Employment and Training Administration, 200 Constitution Avenue, NW., Room S-4231, Washington, DC 20210; or e-mail Dais.Anthony@dol.gov; or transmit via fax (202) 693-3015 (this is not a toll-free number).

FOR FURTHER INFORMATION CONTACT: Anthony D. Dais, at telephone number (202) 693-2784 (this is not a toll-free number).

SUPPLEMENTARY INFORMATION: Section 188 of the Consolidated Farm and Rural Development Act of 1972, as established under 29 CFR Part 75, authorizes the United States Department of Agriculture to make or guarantee loans or grants to finance industrial and business activities in rural areas. The Secretary of Labor must review the application for financial assistance for the purpose of certifying to the Secretary of Agriculture that the assistance is not calculated, or likely, to result in: (a) A transfer of any employment or business activity from one area to another by the loan applicant's business operation; or, (b) An increase in the production of goods, materials, services, or facilities in an area where there is not sufficient

demand to employ the efficient capacity of existing competitive enterprises unless the financial assistance will not have an adverse impact on existing competitive enterprises in the area. The Employment and Training Administration within the Department of Labor is responsible for the review and certification process. Comments should address the two bases for certification and, if possible, provide data to assist in the analysis of these issues.

Signed at Washington, DC this 10th of November, 2009.

Jane Oates,

Assistant Secretary for Employment and Training.

[FR Doc. E9-27871 Filed 11-19-09; 8:45 am]

BILLING CODE 4510-FN-P

NATIONAL SCIENCE FOUNDATION

Agency Information Collection Activities: Proposed Collection, Comment Request

AGENCY: National Science Foundation.

ACTION: Notice.

SUMMARY: The National Science Foundation (NSF) is announcing plans to request clearance for this collection. In accordance with the requirement of Section 3506 (c)(2)(A) of the Paperwork Reduction Act of 1995, we are providing opportunity for public comment on this action. After obtaining and considering public comment, NSF will prepare the submission requesting OMB clearance of this collection for no longer than three years.

Comments are invited on (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the Agency, including whether the information shall have practical utility; (b) the accuracy of the Agency's estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information on respondents, including through the use of automated collection techniques or other forms of information technology; and (d) ways to minimize the burden of the collection of information of respondents, including through the use of automated collection techniques or other forms of information technology.

DATES: Written comments should be received by January 19, 2010, to be assured of consideration. Comments received after that date will be considered to the extent practicable.

ADDRESSES: Written comments regarding the information collection and

requests for copies of the proposed information collection request should be addressed to Suzanne Plimpton, Reports Clearance Officer, National Science Foundation, 4201 Wilson Boulevard, Room 295, Arlington, VA 22230, or by e-mail to splimpton@nsf.gov.

FOR FURTHER INFORMATION CONTACT:

Suzanne Plimpton on (703) 292-7556 or send e-mail to splimpton@nsf.gov.

Individuals who use a telecommunications device for the deaf (TDD) may call the Federal Information Relay Service (FIRS) at 1-800-877-8339 between 8 a.m. and 8 p.m., Eastern time, Monday through Friday.

SUPPLEMENTARY INFORMATION:

Title of Collection: Quantitative Evaluation of the ADVANCE Program.
OMB Control No.: 3145-NEW.
Expiration Date of Approval: Not applicable.

Abstract: The ADVANCE Program was established by the National Science Foundation in 2001 to address the underrepresentation and inadequate advancement of women on STEM (Science, Technology, Engineering, and Mathematics) faculties at postsecondary institutions. The evaluation being conducted by Westat focuses on the outcomes of two ADVANCE program components: (a) The first two (2001 and 2003) cohorts of Institutional Transformation (IT) awardees, and (b) both (2002 and 2004) cohorts of individuals receiving ADVANCE Fellows awards. The study will rely on a thorough review of project documents and relevant literature; a survey (facilitated online via WebEx) and an outcome indicator data form (distributed and completed electronically) for the 19 IT awardee institutions; and, a mail survey, with telephone followup as needed, of all 59 former Fellows. In addition, the study will use data from the 2001 and 2008 administrations of the Survey of Doctorate Recipients (SDR) for comparison purposes.

The evaluation of the IT component has two primary goals: to compare selected gender equity outcomes for STEM faculty at the 19 IT Cohorts 1 and 2 institutions and at other similar U.S. four-year colleges and universities that have not subsequently received ADVANCE IT awards, and to develop innovative institutional-level measures of changes in gender equity climate and practices that can be applied to evaluating the outcomes of the IT award. The primary goal of the Fellows evaluation is to compare the career trajectories of ADVANCE Fellows with those of similar individuals who were not awarded these fellowships.

Respondents: Faculty and staff at institutions of higher education and individuals holding doctoral degrees in STEM fields awarded an NSF ADVANCE Fellowship.

Estimated Number of Annual Respondents: 139.

Burden on the Public: 1859 hours.

Dated: November 17, 2009.

Suzanne H. Plimpton,

Reports Clearance Officer, National Science Foundation.

[FR Doc. E9-27928 Filed 11-19-09; 8:45 am]

BILLING CODE 7555-01-P

NUCLEAR REGULATORY COMMISSION

[Docket No. 50-282 and 50-306; NRC-2009-0507]

Northern States Power Company; Prairie Island Nuclear Generating Plant, Units 1 and 2; Notice of Availability of the Draft Supplement 39 to the Generic Environmental Impact Statement for License Renewal of Nuclear Plants, the License Renewal of Prairie Island Nuclear Generating Plants, Units 1 and 2

Notice is hereby given that the U.S. Nuclear Regulatory Commission (NRC or Commission) has published a draft plant-specific supplement to the Generic Environmental Impact Statement for License Renewal of Nuclear Plants (GEIS), NUREG-1437, regarding the renewal of operating licenses DPR-42 and DPR-60 for an additional 20 years of operation for Prairie Island Nuclear Generating Plant, Unit 1 and Unit 2 (PINGP 1 and 2). PINGP 1 and 2 are located in Red Wing, Minnesota, on the west bank of the Mississippi River in Goodhue County. Possible alternatives to the proposed action (license renewal) include no action and reasonable alternative energy sources.

The draft Supplement 39 to the GEIS is publicly available at the NRC Public Document Room (PDR), located at One White Flint North, 11555 Rockville Pike, Rockville, Maryland, 20852, or from the NRC's Agencywide Documents Access and Management System (ADAMS). The ADAMS Public Electronic Reading Room is accessible at <http://adamswebsearch.nrc.gov/dologin.htm>. The Accession Number for the draft Supplement 39 to the GEIS is ML093170484. Persons who do not have access to ADAMS, or who encounter problems in accessing the documents located in ADAMS, should contact the NRC's PDR reference staff by telephone at 1-800-397-4209, or 301-415-4737,

or by e-mail at pdr.resource@nrc.gov. In addition, a copy of the draft supplement to the GEIS is available to local residents near the site at the Red Wing Public Library, 225 East Avenue, Red Wing, Minnesota 55066.

Any interested party may submit comments on the draft supplement to the GEIS for consideration by the NRC staff. To be considered, comments on the draft supplement to the GEIS and the proposed action must be received by January 30, 2010; the NRC staff is able to ensure consideration only for comments received on or before this date. Comments received after the due date will be considered only if it is practical to do so. Written comments on the draft supplement to the GEIS should be sent to: Chief, Rulemaking and Directives Branch, Division of Administrative Services, Office of Administration, Mailstop TWB 5B01, U.S. Nuclear Regulatory Commission, Washington, DC, 20555-0001.

Electronic comments may be submitted to the NRC by e-mail at PrairieIslandEIS@nrc.gov. All comments received by the Commission, including those made by Federal, State, local agencies, Native American Tribes, or other interested persons, will be made available electronically at the Commission's PDR in Rockville, Maryland, and through ADAMS.

The NRC staff will hold public meetings prior to the close of the public comment period to present an overview of the draft plant-specific supplement to the GEIS and to accept public comments on the document. The times, date, and location of the meetings will be announced in a meeting notice at a later date. The meetings will be transcribed and will include: (1) A presentation of the contents of the draft plant-specific supplement to the GEIS, and (2) the opportunity for interested government agencies, organizations, and individuals to provide comments on the draft report. Additionally, the NRC staff will host informal discussions one hour prior to the start of each session at the same location. No comments on the draft supplement to the GEIS will be accepted during the informal discussions. To be considered, comments must be provided either at the transcribed public meeting or in writing. Persons may pre-register to attend or present oral comments at the meeting by contacting Ms. Elaine Keegan, the NRC Environmental Project Manager at 1-800-368-5642, extension 8517, or by e-mail at Elaine.Keegan@nrc.gov, no later than Friday, December 4, 2009. Members of the public may also register to provide oral comments within 15 minutes of the start of each session. Individual, oral

comments may be limited by the time available, depending on the number of persons who register. If special equipment or accommodations are needed to attend or present information at the public meeting, the need should be brought to Ms Keegan's attention no later than December 1, 2009, to provide the NRC staff adequate notice to determine whether the request can be accommodated.

FOR FURTHER INFORMATION CONTACT: Ms. Elaine Keegan, Projects Branch 2, Division of License Renewal, Office of Nuclear Reactor Regulation, U.S. Nuclear Regulatory Commission, Mail Stop O-11F1, Washington, DC, 20555-0001. Ms. Keegan may be contacted at the aforementioned telephone number or e-mail address.

Dated at Rockville, Maryland, this 16th day of November 2009.

For the Nuclear Regulatory Commission.

David J. Wrona,

Chief, Projects Branch 2, Division of License Renewal, Office of Nuclear Reactor Regulation.

[FR Doc. E9-27911 Filed 11-19-09; 8:45 am]

BILLING CODE 7590-01-P

NUCLEAR REGULATORY COMMISSION

[Docket No. 50-005; NRC-2009-0495]

Pennsylvania State University: Penn State Breazeale Reactor; Environmental Assessment and Finding of No Significant Impact; Correction

AGENCY: Nuclear Regulatory Commission.

ACTION: Notice of Environmental Assessment and Finding of No Significant Impact; Correction.

SUMMARY: This document corrects a notice appearing in the *Federal Register* on November 12, 2009 (74 FR 58319), that considers issuance of a renewal of Facility Operating License No. R-2, to be held by Pennsylvania State University. This action is necessary to correct a typographical error where the International System of Units symbol for the prefix "micro" (μ) was incorrectly displayed as \pm and $\pm m$.

FOR FURTHER INFORMATION CONTACT: Linh Tran, Senior Project Manager, Research and Test Reactors Branch A, Division of Policy and Rulemaking, Office of Nuclear Reactor Regulation, U.S. Nuclear Regulatory Commission, telephone (301) 415-4103, e-mail Linh.Tran@nrc.gov.

SUPPLEMENTARY INFORMATION:

In FR doc. E9-27282, published on November 12, 2009, make the following corrections:

1. On page 58320, in the second column, under the headings I. Radiological Impact Environmental Effects of Reactor Operations, the fifth sentence is corrected to read as follows:

Licensee calculations, based on those measurements, indicate that annual Argon-41 releases result in an offsite concentration of $3.2E-10$ microCuries per milliliter ($\mu\text{Ci/ml}$), which is below the limit of $1.0E-8$ $\mu\text{Ci/ml}$ specified in 10 CFR Part 20, Appendix B for air effluent releases.

2. On page 58320, in the third column, first complete paragraph, the seventh and eighth sentences are corrected to read as follows:

According to the licensee, the leakage resulted in the release of approximately 1.3 milliCuries of tritium, at a concentration of $2.8E-5$ $\mu\text{Ci/ml}$. This concentration is a fraction of the limit of $1E-3$ $\mu\text{Ci/ml}$ specified in 10 CFR Part 20, Appendix B for liquid effluents.

Dated at Rockville, Maryland, this 16th day of November 2009.

For the Nuclear Regulatory Commission.

Michael T. Lesar,

Liaison Officer, Nuclear Regulatory Commission.

[FR Doc. E9-27912 Filed 11-19-09; 8:45 am]

BILLING CODE 7590-01-P

POSTAL REGULATORY COMMISSION

[Docket No. CP2010-10; Order No. 341]

New Postal Product

AGENCY: Postal Regulatory Commission.
ACTION: Notice.

SUMMARY: The Commission is noticing a recently-filed Postal Service request to add a Global Expedited Package Services 2 contract to the Competitive Product List. The Postal Service has also filed a related contract. This notice addresses procedural steps associated with these filings.

DATES: Comments are due: November 23, 2009.

ADDRESSES: Submit comments electronically via the Commission's Filing Online system at <http://www.prc.gov>. Commenters who cannot submit their views electronically should contact the person identified in **FOR FURTHER INFORMATION CONTACT** by telephone for advice on alternatives to electronic filing.

FOR FURTHER INFORMATION CONTACT: Stephen L. Sharfman, General Counsel, 202-789-6820 or stephen.sharfman@prc.gov.

SUPPLEMENTARY INFORMATION:

- I. Introduction
- II. Notice of Filing
- III. Ordering Paragraphs

I. Introduction

On November 12, 2009, the Postal Service filed a notice announcing that it has entered into an additional Global Expedited Package Services 2 (GEPS 2) contract.¹

GEPS 2 provides volume-based incentives for mailers that send large volumes of Express Mail International (EMI) and/or Priority Mail International (PMI). The Postal Service believes the instant contract is functionally equivalent to the previously submitted GEPS 2 contracts and is supported by the Governors' Decision filed in Docket No. CP2008-4.² *Id.* at 1.

The instant contract. The Postal Service filed the instant contract pursuant to 39 CFR 3015.5. In addition, the Postal Service contends that the contract is in accordance with Order No. 290.³ In Order No. 328, the Commission approved an extension of the current contract with this customer in order for the Postal Service to complete its internal procedures and file the instant contract for regulatory review.⁴ The term of the instant contract is 1 year from the completion of the regulatory review. Notice at 2-3.

In support of its Notice, the Postal Service filed four attachments as follows:

1. Attachment 1—an application for non-public treatment of materials to maintain the contract and supporting documents under seal;
2. Attachment 2—a redacted copy of Governors' Decision No. 08-7 which establishes prices and classifications for GEPS contracts, a description of applicable GEPS contracts, formulas for prices, an analysis and certification of the formulas and certification of the Governors' vote;
3. Attachment 3—a redacted copy of the contract and applicable annexes; and

¹ Notice of United States Postal Service Filing of Functionally Equivalent Global Expedited Package Services 2 Negotiated Service Agreement and Application for Non-Public Treatment of Materials Filed Under Seal, November 12, 2009 (Notice).

² See Docket No. CP2008-4, Notice of United States Postal Service of Governors' Decision Establishing Prices and Classifications for Global Expedited Package Services Contracts, May 20, 2008.

³ See Docket No. CP2009-50, Order Granting Clarification and Adding Global Expedited Package Services 2 to the Competitive Product List, August 28, 2009 (Order No. 290).

⁴ See Docket No. CP2008-23, Order Granting Motion for Temporary Relief, October 29, 2009 (Order No. 328).

4. Attachment 4—a certified statement required by 39 CFR 3015.5(c)(2).

Functional equivalency. The Postal Service asserts that the instant contract is functionally equivalent to the contract in Docket No. CP2009-50 and prior GEPS 2 contracts. *Id.* at 3-4. It also contends that the instant contract meets the requirements of Governors' Decision No. 08-7 for rates for GEPS contracts. *Id.* at 3. The Postal Service states that the basic difference between the contract in Docket No. CP2009-50 and the instant contract is customer-specific information including the customer's name, address, representative to receive notices, identity of the signatory, and provisions clarifying tender locations, minimum revenue and/or volume requirements, and liquidated damages. *Id.* at 3-4. It asserts that the instant contract and all GEPS 2 contracts have similar cost and market characteristics and is functionally equivalent in all relevant aspects. *Id.* at 4. The Postal Service concludes that this contract is in compliance with 39 U.S.C. 3633, and requests that this contract be included within the GEPS 2 product. *Id.*

II. Notice of Filing

The Commission establishes Docket No. CP2010-10 for consideration of matters related to the contract identified in the Postal Service's Notice.

Interested persons may submit comments on whether the Postal Service's contract is consistent with the policies of 39 U.S.C. 3632, 3633 or 3642. Comments are due no later than November 23, 2009. The public portions of these filings can be accessed via the Commission's Web site (<http://www.prc.gov>).

The Commission appoints Paul L. Harrington to serve as Public Representative in this proceeding.

III. Ordering Paragraphs

It is ordered:

1. The Commission establishes Docket No. CP2010-10 for consideration of the issues raised in this docket.
2. Comments by interested persons in these proceedings are due no later than November 23, 2009.
3. Pursuant to 39 U.S.C. 505, Paul L. Harrington is appointed to serve as officer of the Commission (Public Representative) to represent the interests of the general public in this proceeding.

4. The Secretary shall arrange for publication of this order in the **Federal Register**.

By the Commission.

Judith M. Grady,

Acting Secretary.

[FR Doc. E9-27936 Filed 11-19-09; 8:45 am]

BILLING CODE 7710-FW-S

RECOVERY ACCOUNTABILITY AND TRANSPARENCY BOARD**Privacy Act of 1974; System of Records**

AGENCY: Recovery Accountability and Transparency Board

ACTION: Notice of new Privacy Act systems of records.

SUMMARY: The Recovery Accountability and Transparency Board (Board or RATB) proposes two new systems of records subject to the Privacy Act of 1974, as amended (Privacy Act or the Act), entitled "RATB Investigative Files" and "RATB Fraud Hotline Program Files."

ADDRESSES: Comments may be submitted:

By Mail or Hand Delivery: Jennifer Dure, Office of General Counsel, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006;

By Fax: (202) 254-7970; or

By E-mail to the Board:

comments@ratb.gov.

All comments on the proposed new systems of records should be clearly identified as such.

FOR FURTHER INFORMATION CONTACT:

Jennifer Dure, General Counsel, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006, (202) 254-7900.

SUPPLEMENTARY INFORMATION: Title 5 U.S.C. 552a(e)(4) and (11) provide that the public be given a 30-day period in which to comment on any new routine use of a system of records. The Office of Management and Budget (OMB), which has oversight responsibilities under the Act, requires a 40-day period in which to conclude its review of the new systems. Therefore, please submit any comments by December 30, 2009.

In accordance with 5 U.S.C. 552a(r), the Board has provided a report to OMB and the Congress on the proposed systems of records.

Recovery Accountability and Transparency Board**Table of Contents**

RATB-11—RATB Investigative Files.

RATB-12—RATB Fraud Hotline Program Files.

RATB—11**SYSTEM NAME:**

RATB Investigative Files.

SECURITY CLASSIFICATION:

The majority of the information in the system is Sensitive but Unclassified.

SYSTEM LOCATION:

Recovery Accountability and Transparency Board, located at 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006.

CATEGORIES OF INDIVIDUALS COVERED BY THE SYSTEM:

In connection with its investigative duties, the RATB maintains records on the following categories of individuals:

(a) Individuals or entities who are or have been the subject of investigations identified by the RATB;

(b) Individuals who are or have been witnesses, complainants, or informants in investigations identified by the RATB; and

(c) Individuals or entities that have been identified as potential subjects or parties to an RATB investigation.

CATEGORIES OF RECORDS IN THE SYSTEM:

Information relating to investigation, including:

(a) Letters, memoranda, and other documents describing complaints or alleged criminal, civil, or administrative misconduct.

(b) Investigative files which include general intelligence and relevant data, leads for Inspectors General (or other applicable oversight and law enforcement entities), reports of investigations and related exhibits, statements, affidavits, and records obtained during an investigation.

AUTHORITY FOR MAINTENANCE OF SYSTEM:

The RATB's enabling legislation, the American Recovery and Reinvestment Act of 2009 (Pub. L. 111-5) (Recovery Act), sections 1521 *et seq.*

PURPOSE(S):

The purpose of this system of records is to enable the RATB to carry out its responsibilities under its enabling legislation, the Recovery Act. The RATB is statutorily directed to coordinate and conduct oversight of covered funds to prevent fraud, waste, and abuse.

ROUTINE USES OF RECORDS MAINTAINED IN THE SYSTEM, INCLUDING CATEGORIES OF USERS AND THE PURPOSES OF SUCH USES:

In addition to those disclosures generally permitted under 5 U.S.C. 552a(b) of the Privacy Act, these records or information contained therein may specifically be disclosed outside the RATB as a routine use pursuant to 5 U.S.C. 552a(b)(3) as follows:

A. To the appropriate federal, state, local, or tribal agency responsible for investigating, prosecuting, enforcing, or implementing a statute, rule, regulation, or order, if the information is relevant to a violation or potential violation of civil or criminal law or regulation within the jurisdiction of the receiving entity.

B. To any individual or entity when necessary to elicit information that will assist an RATB review or audit.

C. To appropriate officials and employees of a federal agency or entity which requires information relevant to a decision concerning the hiring, appointment, or retention of an individual; the issuance, renewal, suspension, or revocation of a security clearance; the execution of a security or suitability investigation; the letting of a contract; or the issuance or revocation of a grant or other benefit.

D. To a Member of Congress or staff acting upon the Member's behalf when the Member or staff requests the information on behalf of, and at the request of, the individual who is the subject of the record.

E. Information may be disclosed to the Department of Justice (DOJ), or in a proceeding before a court, adjudicative body, or other administrative body before which the RATB is authorized to appear, when:

1. The RATB, or any component thereof; or

2. Any employee of the RATB in his or her official capacity; or

3. Any employee of the RATB in his or her individual capacity where the DOJ or the RATB has agreed to represent the employee; or

4. The United States, if the RATB determines that litigation is likely to affect the RATB or any of its components, is a party to litigation or has an interest in such litigation, and the use of such records by the DOJ or the RATB is deemed by the RATB to be relevant and necessary to the litigation, provided, however, that in each case it has been determined that the disclosure is compatible with the purpose for which the records were collected.

F. Information may be disclosed to the National Archives and Records Administration in records management inspections.

G. Information may be disclosed to contractors, grantees, consultants, or volunteers performing or working on a contract, service, grant, cooperative agreement, job, or other activity for the RATB and who have a need to have access to the information in the performance of their duties or activities for the RATB.

DISCLOSURE TO CONSUMER REPORTING AGENCIES:

Not applicable.

POLICIES AND PRACTICES FOR STORING, RETRIEVING, ACCESSING, RETAINING, AND DISPOSING OF RECORDS IN THE SYSTEM: STORAGE:

Information in this system is stored electronically in office automation equipment.

RETRIEVABILITY:

Each lead or investigation is assigned a file number and all records relating to a particular investigation are filed and retrieved by that number. Records may also be retrievable by the names of subjects.

SAFEGUARDS:

Computer records are maintained in a secure, password-protected computer system. Paper records are maintained in lockable file cabinets. All records are maintained in secure, access-controlled areas.

RETENTION AND DISPOSAL:

Records will be retained and disposed of in accordance with RATB Records Control Schedules approved by the National Archives and Records Administration.

SYSTEM MANAGER AND ADDRESS:

Assistant Director, Investigations, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006.

NOTIFICATION PROCEDURE:

Address inquiries to the System Manager listed above.

RECORD ACCESS PROCEDURES:

The major part of this system is exempt from this requirement pursuant to 5 U.S.C. 552a(j)(2) and (k)(2). To the extent that this system is not subject to exemption, it is subject to access. A determination as to exemption shall be made at the time a request for access is received. A request for access to records contained in this system shall be made in writing, with the envelope and the letter clearly marked "Privacy Access Request." Include in the request the full name of the individual involved, his or her current address, date and place of birth, notarized signature (or submitted with date and signature under penalty of perjury), and any other identifying number or information which may be of assistance in locating the record. The requester shall also provide a return address for transmitting the information. Access requests shall be directed to the System Manager listed above.

CONTESTING RECORDS PROCEDURES:

Requesters shall direct their request to the System Manager listed above, stating clearly and concisely what information is being contested, the reason for contesting it, and the proposed amendment to the information.

RECORD SOURCE CATEGORIES:

The subjects of investigations; individuals with whom the subjects of investigations are associated; federal, state, local, and foreign law enforcement and non-law enforcement agencies; private citizens; witnesses; informants; and public source materials.

RATB—12**SYSTEM NAME:**

RATB Fraud Hotline Program Files

SECURITY CLASSIFICATION:

The majority of the information in the system is Sensitive but Unclassified.

SYSTEM LOCATION:

Recovery Accountability and Transparency Board, located at 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006.

CATEGORIES OF INDIVIDUALS COVERED BY THE SYSTEM:

Persons who report information to the RATB concerning the possible existence of activities constituting a violation of law, rules, or regulations, fraud, abuse, mismanagement, or gross waste of funds, and the subject of the complaints.

CATEGORIES OF RECORDS IN THE SYSTEM:

Letters, memoranda, other communications and documents describing complaints of alleged criminal, civil, or administrative misconduct relating to covered funds.

AUTHORITY FOR MAINTENANCE OF SYSTEM:

The RATB's enabling legislation, the American Recovery and Reinvestment Act of 2009 (Pub. L. 111-5) (Recovery Act), sections 1521 *et seq.*

PURPOSE(S):

The purpose of this system of records is to enable the RATB to carry out its responsibilities under its enabling legislation, the Recovery Act. The RATB is statutorily directed to coordinate and conduct oversight of covered funds to prevent fraud, waste, and abuse.

ROUTINE USES OF RECORDS MAINTAINED IN THE SYSTEM, INCLUDING CATEGORIES OF USERS AND THE PURPOSES OF SUCH USES:

In addition to those disclosures generally permitted under 5 U.S.C. 552a(b) of the Privacy Act, these records or information contained therein may specifically be disclosed outside the

RATB as a routine use pursuant to 5 U.S.C. 552a(b)(3) as follows:

A. To the appropriate federal, state, local, or tribal agency responsible for investigating, prosecuting, enforcing, or implementing a statute, rule, regulation, or order, if the information is relevant to a violation or potential violation of civil or criminal law or regulation within the jurisdiction of the receiving entity.

B. To any individual or entity when necessary to elicit information that will assist an RATB review or audit.

C. To appropriate officials and employees of a federal agency or entity which requires information relevant to a decision concerning the hiring, appointment, or retention of an individual; the issuance, renewal, suspension, or revocation of a security clearance; the execution of a security or suitability investigation; the letting of a contract; or the issuance or revocation of a grant or other benefit.

D. To a Member of Congress or staff acting upon the Member's behalf when the Member or staff requests the information on behalf of, and at the request of, the individual who is the subject of the record.

E. Information may be disclosed to the Department of Justice (DOJ), or in a proceeding before a court, adjudicative body, or other administrative body before which the RATB is authorized to appear, when:

1. The RATB, or any component thereof; or
2. Any employee of the RATB in his or her official capacity; or
3. Any employee of the RATB in his or her individual capacity where the DOJ or the RATB has agreed to represent the employee; or
4. The United States, if the RATB determines that litigation is likely to affect the RATB or any of its components, is a party to litigation or has an interest in such litigation, and the use of such records by the DOJ or the RATB is deemed by the RATB to be relevant and necessary to the litigation, provided, however, that in each case it has been determined that the disclosure is compatible with the purpose for which the records were collected.

F. Information may be disclosed to the National Archives and Records Administration in records management inspections.

G. Information may be disclosed to contractors, grantees, consultants, or volunteers performing or working on a contract, service, grant, cooperative agreement, job, or other activity for the RATB and who have a need to have access to the information in the

performance of their duties or activities for the RATB.

POLICIES AND PRACTICES FOR STORING, RETRIEVING, ACCESSING, RETAINING, AND DISPOSING OF RECORDS IN THE SYSTEM:**STORAGE:**

Hard copy files and a computer database.

RETRIEVABILITY:

Records are retrievable by case number or subject name.

SAFEGUARDS:

Computer records are maintained in a secure, password protected computer system. Paper records are maintained in lockable file cabinets. All records are maintained in secure, access-controlled areas or buildings.

RETENTION AND DISPOSAL:

Records will be retained and disposed of in accordance with RATB Records Control Schedules approved by the National Archives and Records Administration.

SYSTEM MANAGER AND ADDRESS:

Assistant Director, Investigations, Recovery Accountability and Transparency Board, 1717 Pennsylvania Avenue, NW., Suite 700, Washington, DC 20006.

NOTIFICATION PROCEDURE:

Address inquiries to the System Manager listed above.

RECORD ACCESS PROCEDURES:

The major part of this system is exempt from this requirement pursuant to 5 U.S.C. 552a(k)(2) and (k)(5). To the extent that this system is not subject to exemption, it is subject to access. A determination as to exemption shall be made at the time a request for access is received. A request for access to records contained in this system shall be made in writing, with the envelope and the letter clearly marked "Privacy Access Request." Include in the request the full name of the individual involved, his or her current address, date and place of birth, notarized signature (or submitted with date and signature under penalty of perjury), and any other identifying number or information which may be of assistance in locating the record. The requester shall also provide a return address for transmitting the information. Access requests shall be directed to the System Manager listed above.

CONTESTING RECORDS PROCEDURES:

Requesters shall direct their request to the System Manager listed above, stating clearly and concisely what information is being contested, the reason for

contesting it, and the proposed amendment to the information.

RECORD SOURCE CATEGORIES:

Complainants who are employees of federal, state, and local agencies, and private citizens. Records in the system come from complainants through the telephone, mail, electronic mail, facsimile, and the web site <http://www.recovery.gov>.

Ivan J. Flores,

Paralegal Specialist, Recovery Accountability and Transparency Board.

[FR Doc. E9-27899 Filed 11-19-09; 8:45 am]

BILLING CODE 6820-GA-P

DEPARTMENT OF STATE

[Public Notice 6815]

Notice of Information Collection Under Emergency Review: Forms DS-2053, DS-2054; Medical Examination for Immigrant or Refugee Applicant; DS-3030, Chest X-Ray and Classification Worksheet; OMB Control Number 1405-0113

AGENCY: Department of State.

ACTION: Notice of request for emergency OMB approval.

SUMMARY: The Department of State has submitted the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the emergency review procedures of the Paperwork Reduction Act of 1995.

- *Title of Information Collection:* Medical Examination for Immigrant or Refugee Applicant.
- *OMB Control Number:* 1405-0113.
- *Type of Request:* Emergency Review.
- *Originating Office:* Bureau of Consular Affairs, Office of Visa Services (CA/VO)
- *Form Number:* DS-2053, DS-2054, DS-3030.
- *Respondents:* Immigrant visa and refugee applicants.
- *Estimated Number of Respondents:* 630,000 per year.
- *Estimated Number of Responses:* 630,000 per year.
- *Average Hours Per Response:* 1 hour.
- *Total Estimated Burden:* 630,000 hours annually.
- *Frequency:* Once per application.
- *Obligation To Respond:* Required to Obtain Benefit.

The proposed information collection is published to obtain comments from the public and affected agencies. Emergency review and approval of this

collection has been requested from OMB by January 4, 2010. If granted, the emergency approval is only valid for 180 days. Direct comments to the Department of State Desk Officer in the Office of Information and Regulatory Affairs at the Office of Management and Budget (OMB).

You may submit comments by the following methods:

- *E-mail:* oir_submission@omb.eop.gov. You must include the DS form number, information collection title, and OMB control number in the subject line of your message.
- *Fax:* 202-395-5806. *Attention:* Desk Officer for Department of State.

During the first 60 days of the emergency approval period, a regular review of this information collection is also being undertaken. The submitting agency requests written comments and suggestions from the public and affected agencies concerning the proposed collection of information. Comments will be accepted until 60 days from the date that this notice is published in the **Federal Register**.

You may submit comments by any of the following methods:

- *E-mail:* VisaRegs@state.gov (Subject line must read OMB 1405-0113 Reauthorization).

- *Mail (paper, disk, or CD-ROM submissions):* Chief, Legislation and Regulations Division, Visa Services—OMB 1405-0113 Reauthorization, 2401 E Street, NW., Washington, DC 20520-30106.

- *Fax:* (202) 663-3898.

You must include the DS form number (if applicable), information collection title, and OMB control number in any correspondence.

FOR FURTHER INFORMATION CONTACT:

Direct requests for additional information regarding the collection listed in this notice, including requests for copies of the proposed information collection and supporting documents, to Lauren Prosnik of the Office of Visa Services, U.S. Department of State, 2401 E Street, NW., L-603, Washington, DC 20522, who may be reached at (202) 663-2951.

SUPPLEMENTARY INFORMATION: *We are soliciting public comments to permit the Department to:*

- Evaluate whether the proposed information collection is necessary for the proper performance of our functions.
- Evaluate the accuracy of our estimate of the burden of the proposed collection, including the validity of the methodology and assumptions used.

- Enhance the quality, utility, and clarity of the information to be collected.

- Minimize the reporting burden on those who are to respond, including the use of automated collection techniques or other forms of technology.

Abstract of proposed collection:

INA Section 221(d) requires that prior to the issuance of an immigrant visa the applicant undergo a physical and mental examination. INA Section 412(b)(4)(B) requires that the United States Government “provide for the identification of refugees who have been determined to have medical conditions affecting the public health and requiring treatment.” Form DS-2053, Medical Examination for Immigrant or Refugee Applicant (1991 Technical Instructions); Form DS-2054, Medical Examination for Immigrant or Refugee Applicant (2007 Technical Instructions); and DS-3030, Chest X-Ray and Classification Worksheet (2007 Technical Instructions) are designed to record the results of the medical examination. A panel physician performs the medical examination of the applicant and completes the forms.

A final rule was published by the Department of Health and Human Services (HHS) in the **Federal Register** on November 2, 2009 that amends HHS regulations to remove the Human Immunodeficiency Virus (HIV) from the definition of *communicable disease of public health significance* and to remove references to HIV from medical examinations for aliens. In order to comply with this rule, effective January 4, 2010, an emergency approval review has been requested from OMB.

The Department must make available to panel physicians conducting medical exams for visa applicants revised medical exam forms. Forms DS-2053, DS-2054 and DS-3030 are being revised to remove references to HIV.

Methodology:

The medical forms are sent to the applicant in the applicant’s package. The applicant takes the forms to the panel physician to use during the medical examination. The panel physician completes the medical examination and fills out the forms. The forms are then submitted in hard copy to the consular officer for processing.

November 16, 2009.

Edward Ramotowski,

Acting Deputy Assistant Secretary, Bureau of Consular Affairs, Department of State.

[FR Doc. E9-27967 Filed 11-19-09; 8:45 am]

BILLING CODE 4710-06-P

DEPARTMENT OF TRANSPORTATION**Surface Transportation Board****[STB Finance Docket No. 35315]****Standard Railroad Corporation—
Acquisition and Operation
Exemption—General Railway
Corporation d/b/a Iowa Northwestern
Railroad**

Standard Railroad Corporation (SRC), a noncarrier, has filed a verified notice of exemption under 49 CFR 1150.31 to acquire from General Railway Corporation d/b/a Iowa Northwestern Railroad (IANW), and to operate a rail line approximately 0.4 miles long. The line extends between milepost 215.0, west of Superior, and milepost 215.4, west of Mackenzie Junction, in Dickinson County, IA.

The transaction is expected to be consummated on or after December 4, 2009 (30 days after the verified notice was filed).

SRC certifies that its projected annual revenues as a result of this transaction will not result in SRC becoming a Class II or Class I rail carrier. SRC further certifies that its projected annual revenues upon becoming a Class III carrier will not exceed \$5 million.

Pursuant to the Consolidated Appropriations Act, 2008, Public Law 110-161, § 193, 121 Stat. 1844 (2007), nothing in this decision authorizes the following activities at any solid waste rail transfer facility: collecting, storing, or transferring solid waste outside of its original shipping container; or separating or processing solid waste (including baling, crushing, compacting, and shredding). The term "solid waste" is defined in section 1004 of the Solid Waste Disposal Act, 42 U.S.C. 6903.

If SRC's verified notice contains false or misleading information, the exemption is void *ab initio*. Petitions to revoke the exemption under 49 U.S.C. 10502(d) may be filed at any time. The filing of a petition to revoke will not automatically stay the effectiveness of the exemption. Stay petitions must be filed no later than November 27, 2009 (at least 7 days before the exemption becomes effective).

An original and 10 copies of all pleadings, referring to STB Finance Docket No. 35315, must be filed with the Surface Transportation Board, 395 E Street, SW., Washington, DC 20423-0001. In addition, one copy of each pleading must be served on Fritz R. Kahn, Fritz R. Kahn P.C., 1920 N Street, NW, (8th floor), Washington, DC 20036.

Board decisions and notices are available on our Web site at <http://www.stb.dot.gov>.

Decided: November 17, 2009.

By the Board, Rachel D. Campbell,
Director, Office of Proceedings.

Kulunie L. Cannon,

Clearance Clerk.

[FR Doc. E9-27886 Filed 11-19-09; 8:45 am]

BILLING CODE 4915-01-P

DEPARTMENT OF TRANSPORTATION**Federal Transit Administration****[FTA Docket No. FTA-2009-0055]****Notice of Request for the Approval of
Information Collection**

AGENCY: Federal Transit Administration (FTA), DOT.

ACTION: Notice of request for comments.

SUMMARY: In accordance with the Paperwork Reduction Act of 1995, this notice announces the intention of the Federal Transit Administration (FTA) to request the Office of Management and Budget (OMB) to approve the following information collection:

49 U.S.C. 5308—Clean Fuels Grant Program.

DATES: Comments must be submitted before January 19, 2010.

ADDRESSES: To ensure that your comments are not entered more than once into the docket, submit comments identified by the docket number by only one of the following methods:

1. *Web site:* <http://www.regulations.gov>. Follow the instructions for submitting comments on the U.S. Government electronic docket site. (**Note:** The U.S. Department of Transportation's (DOT's) electronic docket is no longer accepting electronic comments.) All electronic submissions must be made to the U.S. Government electronic docket site at <http://www.regulations.gov>. Commenters should follow the directions below for mailed and hand-delivered comments.

2. *Fax:* 202-366-7951.

3. *Mail:* U.S. Department of Transportation, 1200 New Jersey Avenue, SE., Docket Operations, M-30, West Building, Ground Floor, Room W12-140, Washington, DC 20590-0001.

4. *Hand Delivery:* U.S. Department of Transportation, 1200 New Jersey Avenue, SE., Docket Operations, M-30, West Building, Ground Floor, Room W12-140, Washington, DC 20590-0001 between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

Instructions: You must include the agency name and docket number for this notice at the beginning of your comments. Submit two copies of your comments if you submit them by mail.

For confirmation that FTA has received your comments, include a self-addressed stamped postcard. Note that all comments received, including any personal information, will be posted and will be available to Internet users, without change, to <http://www.regulations.gov>. You may review DOT's complete Privacy Act Statement in the **Federal Register** published April 11, 2000, (65 FR 19477), or you may visit <http://www.regulations.gov>.

Docket: For access to the docket to read background documents and comments received, go to <http://www.regulations.gov> at any time. Background documents and comments received may also be viewed at the U.S. Department of Transportation, 1200 New Jersey Avenue, SE., Docket Operations, M-30, West Building, Ground Floor, Room W12-140, Washington, DC 20590-0001 between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

FOR FURTHER INFORMATION CONTACT: Kimberly Sledge, FTA Office of Program Management (202) 366-2053, or *e-mail:* kimberly.sledge@dot.gov.

SUPPLEMENTARY INFORMATION: Interested parties are invited to send comments regarding any aspect of this information collection, including: (1) The necessity and utility of the information collection for the proper performance of the functions of the FTA; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the collected information; and (4) ways to minimize the collection burden without reducing the quality of the collected information. Comments submitted in response to this notice will be summarized and/or included in the request for OMB approval of this information collection.

Title: 49 U.S.C. Section 5308—Clean Fuels Grant Program (*OMB Number:* 2132-NEW).

Background: The Section 5308 Clean Fuels Grant Program was initiated as a formula program under the Transportation Equity Act for the 21st Century (TEA-21) in June 1998. The program was reauthorized in August 2005 under the Safe, Accountable, Flexible, Efficient Transportation Equity Act: A Legacy for Users (SAFETEA-LU) as a grant program. The program supports the development and deployment of clean fuel and advanced propulsion technologies for transit buses by providing funds for clean fuel vehicles and facilities. To meet program oversight responsibilities, FTA needs information on the operations and performance of clean fuel technology buses to help assess the reliability,

benefits and costs of these technologies compared to conventional vehicle technologies.

Respondents: State and local government and public transportation authorities located in areas designated as non-attainment or maintenance for ozone or carbon monoxide.

Estimated Annual Burden on Respondents: 32 hours for each respondent.

Estimated Total Annual Burden: 512.
Frequency: Semi-annual.

Issued: November 16, 2009.

Ann M. Linnertz,

Associate Administrator for Administration.

[FR Doc. E9-27897 Filed 11-19-09; 8:45 am]

BILLING CODE 4910-57-P

DEPARTMENT OF TRANSPORTATION

Surface Transportation Board

[STB Finance Docket No. 35237]

City of Davenport, IA—Construction and Operation Exemption—in Scott County, IA

AGENCY: Surface Transportation Board, DOT.

ACTION: Correction to Notice of Availability of the Environmental Assessment.

SUMMARY: This document contains a correction to the title of the Notice of Availability of the Environmental Assessment served and published in the *Federal Register* on Monday, October 26, 2009 (74 FR 55085) by the Board's Section of Environmental Analysis. That notice, published in this docket, was titled "Eastern Iowa Industrial Center Rail Project—Construction and Operation Exemption—City of Davenport, Iowa." The correct title should read, "City of Davenport, IA—Construction and Operation Exemption—in Scott County, IA."

FOR FURTHER INFORMATION CONTACT:

Christa Dean, (202) 245-0299.

[Assistance for the hearing impaired is available through the Federal Information Relay Service (FIRS) at: (800) 877-8339.]

SUPPLEMENTARY INFORMATION: On October 26, 2009, the Board served a Notice of Availability of the Environmental Assessment in this docket. The notice is related to a petition filed on July 21, 2009, by the City of Davenport, IA, seeking an exemption under 49 U.S.C. 10502 from the prior approval requirements¹ of 49

U.S.C. 10901 to construct approximately 2.8 miles of rail line in Scott County, IA. The Board instituted a proceeding in this matter under 49 U.S.C. 10502(b) by decision served October 19, 2009. This notice corrects the title of the Notice of Availability of the Environmental Assessment.

Decided: November 16, 2009.

By the Board, Rachel D. Campbell, Director, Office of Proceedings.

Kulunie L. Cannon,

Clearance Clerk.

[FR Doc. E9-27884 Filed 11-19-09; 8:45 am]

BILLING CODE 4915-01-P

DEPARTMENT OF TRANSPORTATION

Surface Transportation Board

[STB Docket No. AB-331 (Sub-No. 1X)]

Bi-State Development Agency of the Missouri-Illinois Metropolitan District—Discontinuance of Service Exemption—in the City of St. Louis, MO

On November 2, 2009, Bi-State Development Agency of the Missouri-Illinois Metropolitan District (Bi-State) filed with the Surface Transportation Board a petition under 49 U.S.C. 10502 for exemption from the provisions of 49 U.S.C. 10903 to discontinue service over a 1.43-mile line of railroad extending from milepost 1.8 in St. Louis, MO, to milepost 3.23 in St. Louis, MO. The line traverses U.S. Postal Service Zip Codes 63110 and 63108.¹

According to Bi-State, the line does not contain federally granted rights-of-way. Any documentation in Bi-State's possession will be made available promptly to those requesting it.

The interest of railroad employees will be protected by the conditions set forth in *Oregon Short Line R. Co.—Abandonment—Goshen*, 360 I.C.C. 91 (1979).

By issuance of this notice, the Board is instituting an exemption proceeding pursuant to 49 U.S.C. 10502(b). A final decision will be issued by February 19, 2010.

¹ Bi-State initially filed this petition for exemption on October 1, 2009, but supplemented it on November 2, 2009, to comply with the newspaper publication requirements of 49 CFR 1105.12. Under that provision, a petitioner must notify the public by publishing a notice of the proposed action in a newspaper of general circulation in each county that the line traverses and must certify to the Board that it has done so by the date its petition is filed. On November 2, 2009, Bi-State certified to the Board that it has satisfied the newspaper publication requirement. Therefore, November 2, 2009, will be considered the official filing date of the petition for exemption.

Any offer of financial assistance (OFA) under 49 CFR 1152.27(b)(2) to subsidize continued rail service will be due no later than 10 days after service of a decision granting the petition for exemption. Each offer must be accompanied by a \$1,500 filing fee. See 49 CFR 1002.2(f)(25).²

All filings in response to this notice must refer to STB Docket No. AB-331 (Sub-No. 1X), and must be sent to: (1) Surface Transportation Board, 395 E Street, SW., Washington, DC 20423-0001; and (2) Theodore J. Williams, Jr., Williams Venker & Sanders LLC, 100 North Broadway, Suite 2100, St. Louis, MO 63102. Replies to the Bi-State petition are due on or before December 10, 2009.

Persons seeking further information concerning discontinuance procedures may contact the Board's Office of Public Assistance, Governmental Affairs, and Compliance at (202) 245-0230 or refer to the full abandonment and discontinuance regulations at 49 CFR part 1152. Questions concerning environmental issues may be directed to the Board's Section of Environmental Analysis (SEA) at (202) 245-0305. Assistance for the hearing impaired is available through the Federal Information Relay Service (FIRS) at 1-800-877-8339.

Board decisions and notices are available on our Web site at: <http://www.stb.dot.gov>.

Decided: November 13, 2009.

By the Board, Rachel D. Campbell, Director, Office of Proceedings.

Anne K. Quinlan,

Acting Secretary.

[FR Doc. E9-27794 Filed 11-19-09; 8:45 am]

BILLING CODE 4915-01-P

DEPARTMENT OF TRANSPORTATION

Federal Transit Administration

Intent To Prepare an Environmental Impact Statement for Proposed Intermodal Transit Improvements in Hercules, CA

AGENCY: Federal Transit Administration (FTA), U.S. Department of Transportation.

ACTION: Notice of Intent (NOI) to prepare an Environmental Impact Statement (EIS).

² Because this is a discontinuance proceeding and not an abandonment, trail use/rail banking and public use conditions are not appropriate. Similarly, no environmental or historic documentation is required under 49 CFR 1105.6(c)(2) and 1105.8.

¹ In an amendment filed on September 8, 2009, the City clarified that it also seeks operation authority.

SUMMARY: The Federal Transit Administration (FTA) in cooperation with the City of Hercules, CA (City) is planning to prepare a joint Environmental Impact Statement/ Environmental Impact Report (EIS/EIR) for the construction of a proposed intermodal transit center project which includes a new passenger train station on the existing Capitol Corridor line, a transit bus terminal, access roadways, and parking facilities, located in Hercules, California. The project would serve commuters, visitors and recreational users who desire an alternative way to travel to and from the City of Hercules and the San Francisco Bay area and the Sacramento area to access employment, entertainment, and recreational destinations. This EIS/EIR will not study a ferry terminal as part of the proposed project. Any future ferry terminal will be evaluated under a separate environmental document. However, the potential for a future ferry terminal will be considered in the cumulative impact analysis for this project. The purpose of this Notice of Intent (NOI) is to alert interested parties on the preparation of an EIS/EIR, to provide information on the proposed transit project, to invite participation in the EIS/EIR process, including comments on the scope of the EIS/EIR, and to announce the public scoping meeting that will be conducted.

DATES: The City of Hercules has already initiated coordination with Federal, State and local agencies. On November 18th, 2009, the City of Hercules participated in an interagency meeting hosted at the U.S. Army Corps of Engineers office in San Francisco, California and presented an overview of the project and invited agency comment on the proposed project. Through the development of a public and agency coordination plan, coordination with responsible and cooperating agencies will continue throughout the review of the EIR/EIS and through permit coordination.

A scoping meeting will be held on December 8th, 2009 at 5:30 p.m. at the Hercules Library, Large Conference Room, 109 Civic Drive, Hercules, CA. Written comments on the scope of the EIS/EIR including the project's purpose and need, the alternatives to be considered, the impacts to be evaluated, and the methodologies to be used in the evaluations should be sent to Lisa Hammon, Assistant City Manager, City of Hercules, 111 Civic Drive, Hercules CA 94547 by December 23, 2009. Comments may also be offered at the scoping meeting.

The general public and agency representatives with an interest in the proposed project are encouraged to attend this public meeting. The project's purpose and need and the description of alternatives for the proposed project will be presented at this meeting. Representatives of Native American Tribal governments and all Federal, State, regional and local agencies that may have an interest in any aspect of the project will be invited to be participating or cooperating agencies, as appropriate.

ADDRESSES: The scoping meeting will be held on December 8th, 2009 at 5:30 p.m. at the Hercules Library, Large Conference Room, 109 Civic Drive, Hercules, CA. The meeting facilities will be accessible to persons with disabilities. If special translation or signing services or other special accommodations are needed, please contact Lisa Hammon, Assistant City Manager, at (510) 799-8251, or by e-mail at: LHammon@ci.hercules.ca.us at least 48 hours before the scoping meeting. Paper copies of scoping materials may be obtained from Lisa Hammon. Also, scoping materials will be available at the meetings and on the City of Hercules Web site [<http://www.ci.hercules.ca.us>].

Written comments on proposed project should be sent to Lisa Hammon, Assistant City Manager, City of Hercules, 111 Civic Drive, Hercules, CA 94547 by December 23, 2009.

Further Information: For further information contact Paul Page, Federal Transit Administration, San Francisco Regional Office at (415) 744-3133.

SUPPLEMENTARY INFORMATION: Scoping: The FTA and the City of Hercules invite all interested individuals, and organizations, public agencies and Native American Tribes to comment on the scope of the EIS, including the project's purpose and need, the alternatives to be studied, the impacts to be evaluated and the evaluation methods to be used. Comments should focus on: alternatives that may be less costly or have fewer environmental or community impacts while achieving similar transportation objectives, and the identification of any significant social, economic or environmental issues related to alternatives. This is not considered a transit project of unusual complexity. Therefore, in line with CEQ 1502.7 (page limits) FTA is setting a limit of 250 pages (exclusive of technical appendices) for this EIS. The document should emphasize graphics, maps and visual simulations, minimize technical jargon and be accessible to

members of the public with limited technical expertise.

Description of Study Area and Proposed Project: The project site is located in Hercules, California, on the shoreline of San Pablo Bay (a part of San Francisco Bay), approximately 1 mile northwest of Interstate 80 (I-80). This is the City's Waterfront District which has been planned for transit oriented development. Project components would include: (1) Grade separation and realignment of a portion of the existing Union Pacific railroad tracks, including the construction of a rail platform, retaining walls and the replacement of a bridge crossing Refugio Creek; (2) construction of a station building; (3) extension of John Muir Parkway, including the construction of Bayfront Boulevard over Refugio Creek, a new Transit Loop Drive, Civic Plaza and surface parking; (4) realignment and restoration of a portion of Refugio Creek from San Pablo Bay upstream approximately 1000 feet to the existing restored segment; (5) construction of segments of the East Bay Park Regional District's recreational trail along the shoreline from Pinole trail to Victoria by the Bay; and, (6) a pedestrian walkway over the railroad tracks to provide a connection to the Hercules Point open space area. The project will also require the relocation of existing utility and gas lines and an outfall to Refugio Creek.

Purpose and Need for the Project: Residents of the San Francisco Bay Area depend heavily on region wide and transbay commuting. Despite the use of existing public transit services, particularly rail and buses, traffic congestion continues to increase, affecting hundreds of thousands of Bay Area residents and creating both economic and environmental costs. The severity of congestion will increase in the future as population and employment in the Bay Area increase. The purpose of the proposed project is to increase local and regional mobility and transportation options by providing new and expanded transit services with intermodal connections that will encourage use of public transit. The project would provide bus-to-train connections, in addition to providing car commuters with access to new transit options that would divert traffic from I-80, the most congested corridor in the Bay Area for the past six years. An expanded and more convenient transit system with new train connections to existing bus services would provide commuters with more options and would primarily decrease car usage and its associated impacts, rather than divert riders from existing

buses, BART, or Capitol Corridor. Key project objectives are to:

1. Reduce vehicle trips on Interstate 80, the most congested freeway in the Bay Area, by providing alternatives to commuting in single occupant vehicles.

2. Provide coordinated, intermodal transit connections by bus, train, and a potential future ferry and human-powered connections by bicycling and walking for transport to/from jobs, recreational uses, educational opportunities, etc.

3. Improve emergency response by having rail and (ultimately) ferry service available in case of a natural or man-made disaster that disables the Bay Bridge or other highways/roadways. Ferry and rail service could also deliver goods and services in an emergency.

4. Support transit-oriented development and "new urbanist" standards by providing the transportation links within the 43-acre waterfront development which also includes housing (including affordable housing), retail, office, and commercial space.

5. Improve safety along the railroad corridor by providing completely grade-separated access to the railroad tracks from the adjacent development by constructing a series of retaining walls and fences for approximately one mile along the waterfront and by constructing over-crossings to Hercules Point and the future ferry terminal.

6. Implement the Goals, Policies and Programs in the City of Hercules General Plan to:

- Develop transportation facilities to provide access to the region, particularly public transit systems (buses, ridesharing, rail transit, as well as potential over-water transit) (Land Use Policy 3A, Circulation Policy e).
- Establish trail linkage between Pinole and Rodeo as part of the regional bay access trail system (Land Use Program 14A.2 and Open Space/Conservation Policy 1b) and continue to improve and protect Refugio Creek as a major environmental amenity (Program 14.A.3).

7. Improve Refugio Creek to allow adequate flows into the Bay without resulting in flooding.

8. Implement the City of Hercules Waterfront Master Plan Initiative and its directive to construct an intermodal transit center on Block I.

Alternatives: The EIS/EIR will include a Build alternative. Included in the Build analysis will be design alternatives to the proposed project that will meet both NEPA and CEQA requirements and are intended to reduce potential environmental effects, including impacts to sensitive biological

habitat. A No Action (No Build) alternative will also be evaluated in the EIS/EIR which would continue with the existing bus services without the construction of a train station and a new bus terminal at the same location. This alternative serves as the baseline against which the environmental effects of the proposed project and other alternatives will be evaluated.

Traffic congestion is an ongoing and steadily increasing problem in the Bay Area, regardless of economic conditions. Alternatives to reduce traffic congestion have been explored in numerous previous studies. According to the Metropolitan Transportation Commission (MTC), the Bay Bridge approach corridor along Interstate 80 (I-80) from State Route 4 (SR-4) in Hercules to the Bay Bridge experiences the worst congestion in the Bay Area. Caltrans' Bay Area monitoring program found that between 1992 and 2005, traffic delay in the region as a whole more than doubled from 64,100 hours to 135,700 hours. According to Caltrans' 2006 report, between 2001 and 2005, traffic delay on the I-80 segment from SR-4 to the Bay Bridge metering lights increased by 16 percent, from 9,410 hours to 10,930 hours (MTC 2007). This segment includes the stretch of I-80 that passes near the proposed HITC project. MTC projects that traffic congestion will continue to worsen; by 2020, MTC expects that Bay Bridge traffic will increase by 50 percent and be "at capacity" for nearly five hours a day during the morning and afternoon peak hours. MTC also predicts that many more Bay Area workers, due to high housing costs, will be living far from their jobs, demanding that they spend more time commuting and polluting on roadways. Even during an economic downturn, BART runs at capacity through the Transbay Tube during peak hours. Improvements in commuter bus service are dependent upon traffic flow, and are limited by the need for more road capacity and more dedicated High Occupancy Vehicle (HOV) lanes for significant expansion. Increased train and transit services would provide expanded commute capacity while avoiding corresponding increases in traffic congestion.

Additionally, the San Francisco Bay Area Water Transit Authority (WTA), now the Water Emergency Transportation Authority (WETA), is a regional agency authorized by the State of California to operate a comprehensive San Francisco Bay Area public water transit system. In 2003, the WTA's plan, "A Strategy to Improve Public Transit with an Environmentally Friendly Ferry System" (the Plan) was approved by

statute (Senate Bill 915, Ch. 714, stats of 2003). The Plan drew on extensive technical studies that examined ridership demand, cost effectiveness, vessel design, environmental impacts, safety, and operations. A Hercules-San Francisco route was identified in the Plan as a potential future ferry route. The potential environmental effects of proposed new ferry service on San Francisco Bay under the WTA Plan were studied at a program planning level in the Program Environmental Impact Report (Program EIR) prepared in 2003. The 2003 Program EIR included analysis of a Hercules/Rodeo location and seven other potential new ferry service locations around the Bay and Delta. While ferry service is anticipated for the city of Hercules, current planning for the ferry is still in development and is considered a future project. The current project proposes only to construct a rail and bus transit facility.

As part of the General Plan for the City of Hercules, the proposed project is intended to be the central element of a transit-oriented development (TOD) project that will include residential and commercial development clustered around transit facilities to enable local residents to use public transit and reduce the need for automobile use. The planned TOD, known as Hercules Bayfront, is not part of the project considered in this EIR/EIS, and will be the subject of a separate environmental review.

Probable Effects/Potential Impacts for Analysis: The purpose of the EIS process is to explore in a public setting potentially significant effects of implementing the proposed project and alternatives on the physical, human, and natural environment.

Implementation of the project components will result in direct effects to the physical environment and may include the loss of special aquatic sites such as tidal wetlands, mudflats, and riparian areas. Mitigation will be incorporated into the project design by first avoiding and minimizing impacts to resources. Compensatory mitigation will be provided for unavoidable impacts. Based on preliminary investigations for special status species the project may affect, but is unlikely to adversely affect, any species listed as threatened or endangered under the State or Federal endangered species acts.

The proposed project would extend the John Muir Parkway, provide parking for short-term and long-term parking, and develop a new access point to commuter rail. Each of these activities may encourage automobile traffic in the

area that could adversely affect levels of service at nearby intersections.

FTA Procedures: Regulations implementing NEPA, as well as provisions of SAFETEA-LU, call for public involvement in the EIS process. Section 6002 of SAFETEA-LU requires that FTA do the following: (1) Extend an invitation to other Federal and non-Federal agencies and Indian Tribes that may have an interest in the proposed project to become participating agencies; (2) provide an opportunity for involvement by participating agencies and the public in helping to define the purpose and need for a proposed project, as well as the range of alternatives for consideration; and (3) establish a plan for coordinating public and agency participation in and comment on the environmental review process. An invitation to become a participating agency will be extended to other Federal and non-Federal agencies and Indian Tribes that may have an interest in the proposed project. It is possible that we may not be able to identify all Federal and non-Federal agencies and Indian Tribes that may have such an interest. Any Federal or non-Federal agency or Indian Tribe interested in the proposed project that does not receive an invitation to become a participating agency should notify, at the earliest opportunity, the City at the **ADDRESSES** or phone number above.

A comprehensive public involvement program has been developed. A technical advisory committee called the Project Development Team, consisting of representatives of State, regional and local agencies, is in place. The program also includes a public scoping process, a public review/comment period and public hearing on the Draft Environmental Impact Statement, development and distribution of project newsletters and posting of information on the project Web site. The purposes of and need for the proposed project have been preliminarily identified in this notice. We invite the public and participating agencies to consider the preliminary statement of purposes of and need for the proposed project, as well as potential alternatives, and the public is welcome to use the public scoping process to further define the issues of concern among all parties interested in the project. Comments on potential significant environmental impacts that may be associated with the proposed project are also welcomed. All comments and suggestions will be given serious consideration. Comments on potentially significant environmental impacts that may be associated with the proposed project are also welcomed. There will be additional opportunities

to participate in the scoping process at the public meetings announced in this notice. In accordance with 23 CFR 771.105(a) and 771.133, FTA will comply with all Federal environmental laws, regulations and executive orders applicable to the proposed project during the environmental review process to the maximum extent practicable. These requirements include, but are not limited to, the regulations of the Council on Environmental Quality implementing NEPA (40 CFR parts 1500–1508 and 23 CFR part 771), the project-level air quality conformity regulation of the U.S. Environmental Protection Agency (EPA) (40 CFR part 93), section 404(b)(1) guidelines of EPA (40 CFR part 230), Executive Orders 11988, 11990 and 12898 regarding floodplains, wetlands, and environmental justice, respectively; Section 106 of the National Historic Preservation Act (36 CFR Part 800); Section 7 of the Endangered Species Act (50 CFR part 402); and Section 4(f) of the Department of Transportation Act (23 CFR 774). The EIR portion of the document will be prepared in accordance with the California Environmental Quality Act (CEQA) and the California Code of Regulations, Title 14, section 15000 *et seq.*

Dated: November 13, 2009.

Leslie T. Rogers,

Federal Transit Administration.

[FR Doc. E9–27896 Filed 11–19–09; 8:45 am]

BILLING CODE 4910–57–P

DEPARTMENT OF TRANSPORTATION

Maritime Administration

[USCG–2006–24644]

TORP Terminal LP, Bienville Offshore Energy Terminal Liquefied Natural Gas Deepwater Port License Application; Preparation of Supplemental Environmental Impact Statement

AGENCY: Maritime Administration, DOT.

ACTION: Notice of availability; Notice of public meeting; Request for comments.

SUMMARY: The Maritime Administration, in cooperation with the U.S. Coast Guard announces the availability of the Draft Supplemental Environmental Impact Statement (DSEIS) for the TORP Terminal LP, Bienville Offshore Energy Terminal (BOET) Liquefied Natural Gas (LNG) Deepwater Port license amended application. The amended application describes a project that would be located in the Gulf of Mexico, in Main Pass block MP 258, approximately 63 miles south of Mobile Point, Alabama.

Publication of this notice begins a 45 day comment period and provides information on how to participate in the process.

DATES: The public meeting will be held in Mobile, Alabama on December 9, 2009, from 6 p.m. to 8 p.m., and will be preceded by an open house from 5 p.m. to 6 p.m. The public meeting may end later than the stated time, depending on the number of persons wishing to speak. Material submitted in response to this request for comments must reach the Docket Management Facility by January 4, 2010.

ADDRESSES: The open house and public meeting will be held at the: Mobile Convention Center, One South Water Street, Mobile, Alabama 36602; *telephone:* 251–208–2100.

The amended application, comments and associated documentation are available for viewing at the Federal Docket Management System (FDMS) Web site: <http://www.regulations.gov> under docket number USCG–2006–24644.

Docket submissions for USCG–2006–24644 should be addressed to: Department of Transportation, Docket Management Facility, West Building, Ground Floor, Room W12–140, 1200 New Jersey Avenue, SE., Washington, DC 20590–0001.

The Docket Management Facility accepts hand-delivered submissions, and makes docket contents available for public inspection and copying at the above address between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays. The Facility telephone number is 202–366–9329, the fax number is 202–493–2251, and the Web site for electronic submissions or for electronic access to docket contents is <http://www.regulations.gov>.

FOR FURTHER INFORMATION CONTACT: Mr. Patrick Marchman, Maritime Administration, telephone: 202–366–8805, e-mail:

Patrick.Marchman@dot.gov; or Mr.

Linden Houston, Maritime Administration, telephone: 202–366–4839, e-mail: Linden.Houston@dot.gov.

If you have questions on viewing the docket, call Renee V. Wright, Program Manager, Docket Operations, telephone 202–493–0402.

SUPPLEMENTARY INFORMATION:

Public Meeting and Open House

We invite you to learn about the proposed deepwater port at an informational open house and to comment at a public meeting immediately afterwards on the proposed action and the evaluation contained in the DSEIS. In order to allow everyone a

chance to speak at the public meeting, we may limit speaker time, or extend the meeting hours, or both. You must identify yourself, and any organization you represent, by name. Your remarks will be recorded or transcribed for inclusion in the public docket.

You may submit written material at the public meeting, either in place of or in addition to speaking. Written material must include your name and address and will be included in the public docket.

Public docket materials will be made available to the public on the Federal Docket Management Facility (*see* Request for Comments).

Our public meeting location is wheelchair-accessible. If you plan to attend the open house or public meeting and need special assistance such as sign language interpretation or other reasonable accommodation, please notify the Maritime Administration (*see* **FOR FURTHER INFORMATION CONTACT**) at least 3 business days in advance. Include your contact information as well as information about your specific needs.

Request for Comments

We request public comments or other relevant information on the DSEIS. The public meeting is not the only opportunity you have to comment. In addition to or in place of attending this meeting, you may submit comments to the Docket Management Facility during the public comment period (*see* **DATES**). We will consider all comments and material received during the comment period.

Submissions should include:

- Docket number USCG–2006–24644.
- Your name and address.

Submit comments or material using only one of the following methods:

- Electronic submission to FDMS, <http://www.regulations.gov>.
- Fax, mail, or hand delivery to the Docket Management Facility (*see* **ADDRESSES**). Faxed or hand delivered submissions must be unbound, no larger than 8½ by 11 inches, and suitable for copying and electronic scanning. If you mail your submission and want to know when it reaches the Facility, include a stamped, self-addressed postcard or envelope.

Regardless of the method used for submitting comments or material, all submissions will be posted, without change, to the FDMS Web site (<http://www.regulations.gov>) and will include any personal information you provide. Therefore, submitting this information makes it public. You may wish to read the Privacy and Use Notice that is available on the FDMS Web site and the

Department of Transportation Privacy Act Notice that appeared in the **Federal Register** on April 11, 2000 (65 FR 19477) (*see* Privacy Act). You may view docket submissions at the Department of Transportation Docket Management Facility or electronically on the FDMS Web site (*see* **ADDRESSES**).

Background

The Notice of Intent to Prepare a Supplemental EIS for the proposed action was published in the **Federal Register** at 74 FR 39136, August 5, 2009. The DSEIS, application materials and associated comments are available on the docket. Information from the “Summary of the Application” from previous **Federal Register** notices is included below for your convenience.

Proposed Action and Alternatives

The proposed action requiring environmental review is the Federal licensing of the proposed deepwater port described in the “Summary of the Application” below. The alternatives to approving and licensing the proposed port are: (1) Approving and licensing with conditions (including conditions designed to mitigate environmental impact), or (2) denying the application, which for purposes of environmental review is the “no-action” alternative. These alternatives are more fully discussed in the DSEIS. The Maritime Administration and the U.S. Coast Guard are the lead Federal agencies for the preparation of the Supplemental EIS. You can address any questions about the proposed action or the DSEIS to the Maritime Administration project managers identified in **FOR FURTHER INFORMATION CONTACT**.

Summary of the Application

TORP Terminal LP proposes to own, construct, and operate a deepwater port, the Bienville Offshore Energy Terminal (BOET), in the Federal waters of the Outer Continental Shelf on Main Pass Block MP 258, approximately 63 miles south of Mobile Point, Alabama, in a water depth of approximately 425 feet. The proposed BOET deepwater port would be capable of mooring a single LNG carrier (LNGC) of up to approximately 265,000 cubic meters (m³) (8.8 million cubic feet [ft³]) in capacity.

The LNGC would be off-loaded using a HiLoad LNG off-loading and regasification unit (HiLoad), which is proprietary technology consisting of a remotely operated floating LNG transfer and regasification unit that connects to the hull of the LNGC. The HiLoad unit would regasify the LNG and deliver the gas via flexible gas pipes to the floating

regasification unit (FRU) located approximately 250 ft (150 m) from the HiLoad unit. Ambient air vaporizers (AAVs) with methanol as an intermediate fluid (IF) would be located aboard the FRU and would provide the heat required to regasify the LNG, all in a closed-loop vaporization system design.

At the FRU, the gas would be metered and sent out via interconnect pipelines to four existing offshore pipelines (Dauphin Natural Gas Pipeline, Williams Natural Gas Pipeline, Destin Natural Gas Pipeline, and Viosca Knoll Gathering System [VKGS] Gas Pipeline) that connect to the onshore natural gas transmission pipeline system. The natural gas would be delivered to customers through existing facilities. BOET would have an average throughput capacity of 1.2 billion standard cubic feet of gas per day (Bscfd) (33.9 million cubic meters of gas per day [m³/day]).

BOET’s major components would include a turret mooring system (TMS), a FRU, a HiLoad unit, two mooring lines that connect the HiLoad to the FRU, two high pressure (HP) flexible gas pipes, two floating IF hoses, two umbilicals, and 22.7 mi (36 km) of new subsea pipelines.

No new onshore pipelines or LNG storage facilities are proposed with this action. A shore based facility will be used to facilitate movement of personnel, equipment, supplies, and disposable materials between the terminal and shore.

BOET will also require permits from the Environmental Protection Agency (EPA) pursuant to the provisions of the Clean Air Act, as amended, and the Clean Water Act, as amended.

Should a license be issued, construction of the deepwater port would be expected to take 30 months with startup of commercial operations anticipated in 2014. The deepwater port, if licensed, would be designed, constructed, and operated in accordance with applicable codes and standards and would have an expected operating life of approximately 25 years.

Privacy Act

Electronic copies of all comments received into the Federal Docket Management System can be searched by the name of the individual submitting the comment (or signing the comment, if submitted on behalf of an association, business, labor union, *etc.*). The DOT Privacy Act Statement can be viewed in the **Federal Register** published on April 11, 2000 (Volume 65, Number 70, pages 19477–78) or you may visit <http://www.regulations.gov>.

Authority: 49 CFR 1.66.

Dated: November 16, 2009.

By order of the Maritime Administrator.

Christine Gurland,

Secretary, Maritime Administration.

[FR Doc. E9-27975 Filed 11-19-09; 8:45 am]

BILLING CODE 4910-81-P

DEPARTMENT OF TRANSPORTATION

Federal Motor Carrier Safety Administration

[Docket No. FMCSA-2009-0271]

Identification of Interstate Motor Vehicles: New York City, Cook County and New Jersey Tax Identification Requirements; Petition for Determination

AGENCY: Federal Motor Carrier Safety Administration (FMCSA), Department of Transportation (DOT).

ACTION: Notice of Petition for Determination; extension of comment period.

SUMMARY: The Federal Motor Carrier Safety Administration (FMCSA) extends until December 3, 2009 the comment period for the Petition for Determination that was published on October 19, 2009.

DATES: Comments are due on or before December 3, 2009.

ADDRESSES: You may submit comments identified by the Federal Docket Management System Number in the heading of this document by any of the following methods—Internet, facsimile, regular mail, or hand-delivery. Do not submit the same comments by more than one method. However, to allow effective public participation before the comment period deadline, the Agency encourages use of the Web site that is listed first. It will provide the most efficient and timely method of receiving and processing your comments.

- *Federal eRulemaking Portal:* Go to <http://www.regulations.gov>. Follow the online instructions for submitting comments.

- *Fax:* 1-202-493-2251.

- *Mail:* Docket Management Facility; U.S. Department of Transportation, Room W12-140, 1200 New Jersey Avenue, SE., Washington, DC 20590-0001.

- *Hand Delivery:* Ground floor, Room W12-140, 1200 New Jersey Avenue, SE., Washington, DC, between 9 a.m. and 5 p.m., ET., Monday through Friday, except Federal holidays.

Instructions: All submissions must include the Agency name and docket number for this action. Note that all comments received will be posted

without change to <http://www.regulations.gov>, including any personal information provided. Refer to the Privacy Act heading on <http://www.regulations.gov> for further information.

Public Participation: The regulations.gov system is generally available 24 hours each day, 365 days each year. You can find electronic submission and retrieval help and guidelines under the “Help” section of the Web site. For notification that FMCSA received the comments, please include a self-addressed, stamped envelope or postcard, or print the acknowledgement page that appears after submitting comments on line. Copies or abstracts of all documents referenced in this notice are in the docket: FMCSA-2009-0271. For access to the docket to read background documents or comments received, go to <http://www.regulations.gov> at any time or to Room W12-140, 1200 New Jersey Avenue, SE., Washington, DC, between 9 a.m. and 5 p.m., ET., Monday through Friday, except Federal holidays. All comments received before the close of business on the comment closing date indicated above will be considered and will be available for examination in the docket at the above address. Comments received after the closing date will be considered to the extent practicable. FMCSA may, however, issue a final determination at any time after the close of the comment period. In addition to late comments, FMCSA will also continue to file in the public docket relevant information that becomes available after the comment closing date. Interested persons should monitor the public docket for new material.

FOR FURTHER INFORMATION CONTACT:

Genevieve D. Sapir, Office of the Chief Counsel, Federal Motor Carrier Safety Administration, 1200 New Jersey Avenue SE., Washington, DC 20590, (202) 366-7056; e-mail Genevieve.Sapir@dot.gov.

SUPPLEMENTARY INFORMATION: On October 19, 2009, FMCSA published a Notice of Petition for Determination inviting public comment on three petitions submitted by the American Trucking Associations (ATA) requesting determinations that the Commercial Motor Vehicle (CMV) identification requirements imposed by the State of New Jersey, New York City, and Cook County, Illinois, are preempted by Federal law (74 FR 53578). The Agency provided the public with a 30-day comment period scheduled to expire on November 18, 2009. Because of the level of interest this notice has generated,

FMCSA extends the comment period until December 3, 2009.

Issued on: November 16, 2009.

David K. Tochen,

Acting Chief Counsel.

[FR Doc. E9-27855 Filed 11-17-09; 11:15 am]

BILLING CODE 4910-EX-P

DEPARTMENT OF THE TREASURY

Office of Thrift Supervision

[AC-31; OTS Nos. 02186 and H4650]

Versailles Savings and Loan Company, Versailles, OH; Approval of Conversion Application

Notice is hereby given that on November 12, 2009, the Office of Thrift Supervision approved the application of Versailles Savings and Loan Company, Versailles, Ohio, to convert to the stock form of organization. Copies of the application are available for inspection by appointment (*phone number:* (202) 906-5922 or *e-mail:* public.info@ots.treas.gov) at the Public Reading Room, 1700 G Street, NW., Washington, DC 20552, and the OTS Central Regional Office, 1 South Wacker Drive, Suite 2000, Chicago, Illinois 60606.

Dated: November 16, 2009.

By the Office of Thrift Supervision.

Sandra E. Evans,

Federal Register Liaison.

[FR Doc. E9-27995 Filed 11-19-09; 8:45 am]

BILLING CODE 6720-01-P

DEPARTMENT OF THE TREASURY

Fiscal Service

Surety Companies Acceptable on Federal Bonds—Change In Business Address; American Home Assurance Company; Granite State Insurance Company; Insurance Company of the State of Pennsylvania (The); National Union Fire Insurance Company of Pittsburgh, PA; New Hampshire Insurance Company

AGENCY: Financial Management Service, Fiscal Service, Department of the Treasury.

ACTION: Notice.

SUMMARY: This is Supplement No. 5 to the Treasury Department Circular 570, 2009 Revision, published July 1, 2009, at 74 FR 31536.

FOR FURTHER INFORMATION CONTACT: Surety Bond Branch at (202) 874-6850.

SUPPLEMENTARY INFORMATION: Notice is hereby given by the Treasury that the above-named companies formally changed their "BUSINESS ADDRESS" to "175 WATER STREET, 18TH FLOOR, NEW YORK, NY 10038" effective June 30, 2009.

Federal bond-approving officials should annotate their reference copies

of the Treasury Department Circular 570 ("Circular"), 2009 Revision, to reflect this change.

The Circular may be viewed and downloaded through the Internet at <http://www.fms.treas.gov/c570>.

Questions concerning this notice may be directed to the U.S. Department of the Treasury, Financial Management

Service, Financial Accounting and Services Division, Surety Bond Branch, 3700 East-West Highway, Room 6F01, Hyattsville, MD 20782.

Laura Carrico,

Acting Director, Financial Accounting and Services Division.

[FR Doc. E9-27861 Filed 11-19-09; 8:45 am]

BILLING CODE 4810-35-M



Federal Register

**Friday,
November 20, 2009**

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Part II

Department of Health and Human Services

Centers for Medicare & Medicaid Services

**42 CFR Parts 410, 416, and 419
Medicare Program: Changes to the
Hospital Outpatient Prospective Payment
System and CY 2010 Payment Rates;
Changes to the Ambulatory Surgical
Center Payment System and CY 2010
Payment Rates; Final Rule**

DEPARTMENT OF HEALTH AND HUMAN SERVICES**Centers for Medicare & Medicaid Services****42 CFR Parts 410, 416, and 419**

[CMS-1414-FC]

RIN 0938-AP41

Medicare Program: Changes to the Hospital Outpatient Prospective Payment System and CY 2010 Payment Rates; Changes to the Ambulatory Surgical Center Payment System and CY 2010 Payment Rates**AGENCY:** Centers for Medicare & Medicaid Services (CMS), HHS.**ACTION:** Final rule with comment period.

SUMMARY: This final rule with comment period revises the Medicare hospital outpatient prospective payment system (OPPS) to implement applicable statutory requirements and changes arising from our continuing experience with this system. In this final rule with comment period, we describe the changes to the amounts and factors used to determine the payment rates for Medicare hospital outpatient services paid under the prospective payment system. These changes are applicable to services furnished on or after January 1, 2010.

In addition, this final rule with comment period updates the revised Medicare ambulatory surgical center (ASC) payment system to implement applicable statutory requirements and changes arising from our continuing experience with this system. In this final rule with comment period, we set forth the applicable relative payment weights and amounts for services furnished in ASCs, specific HCPCS codes to which these changes will apply, and other pertinent ratesetting information for the CY 2010 ASC payment system. These changes are applicable to services furnished on or after January 1, 2010.

DATES: *Effective Date:* The provisions of this rule are effective January 1, 2010.

Comment Period: We will consider comments on the subject areas listed in the **SUPPLEMENTARY INFORMATION** section of this rule that are received at one of the addresses provided in the **ADDRESSES** section of this rule no later than 5 p.m. EST on December 29, 2009.

Application Deadline for New Class of New Technology Intraocular Lenses: Request for review of applications for a new class of new technology intraocular lenses must be received by 5 p.m. EST on March 8, 2010.

ADDRESSES: In commenting, please refer to file code CMS-1414-FC. Because of staff and resource limitations, we cannot accept comments by facsimile (FAX) transmission.

You may submit comments in one of four ways (no duplicates, please):

1. *Electronically.* You may submit electronic comments on this regulation to <http://www.regulations.gov>. Follow the instructions for "Comment or Submission" and enter the file code to find the document accepting comments.

2. *By regular mail.* You may mail written comments (one original and two copies) to the following address ONLY: Centers for Medicare & Medicaid Services, Department of Health and Human Services, *Attention:* CMS-1414-FC, P.O. Box 8013, Baltimore, MD 21244-1850.

Please allow sufficient time for mailed comments to be received before the close of the comment period.

3. *By express or overnight mail.* You may send written comments (one original and two copies) to the following address ONLY: Centers for Medicare & Medicaid Services, Department of Health and Human Services, *Attention:* CMS-1414-FC, Mail Stop C4-26-05, 7500 Security Boulevard, Baltimore, MD 21244-1850.

4. *By hand or courier.* If you prefer, you may deliver (by hand or courier) your written comments (one original and two copies) before the close of the comment period to one of the following addresses:

a. Room 445-G, Hubert H. Humphrey Building, 200 Independence Avenue, SW., Washington, DC 20201.

(Because access to the interior of the HHH Building is not readily available to persons without Federal Government identification, commenters are encouraged to leave their comments in the CMS drop slots located in the main lobby of the building. A stamp-in clock is available for persons wishing to retain a proof of filing by stamping in and retaining an extra copy of the comments being filed.)

b. 7500 Security Boulevard, Baltimore, MD 21244-1850.

If you intend to deliver your comments to the Baltimore address, please call the telephone number (410) 786-9994 in advance to schedule your arrival with one of our staff members.

Comments mailed to the addresses indicated as appropriate for hand or courier delivery may be delayed and received after the comment period.

For information on viewing public comments, see the beginning of the **SUPPLEMENTARY INFORMATION** section.

Applications for a new class of new technology intraocular lenses: Requests

for review of applications for a new class of new technology intraocular lenses must be sent by regular mail to ASC/NTOL, Division of Outpatient Care, Mailstop C4-05-17, Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244-1850.

FOR FURTHER INFORMATION CONTACT:

Alberta Dwivedi, (410) 786-0378, Hospital outpatient prospective payment issues.

Dana Burley, (410) 786-0378,

Ambulatory surgical center issues.

Michele Franklin, (410) 786-4533, and

Jana Lindquist, (410) 786-4533,

Partial hospitalization and community mental health center issues.

James Poyer, (410) 786-2261, Reporting of quality data issues.

SUPPLEMENTARY INFORMATION:

Comment Subject Areas: We will consider comments on the following subject areas discussed in this final rule with comment period that are received by the date and time indicated in the **DATES** section of this final rule with comment period:

(1) The payment classifications assigned to HCPCS codes identified in Addenda B, AA, and BB to this final rule with comment period with the "NI" comment indicator;

(2) Recognition of plasma protein fraction as a blood product or a biological for OPPS payment, as discussed in section II.A.1.d.(2) of this final rule with comment period;

(3) Potential alternative coding schemes for reporting hospital clinic visits for new and established patients, as discussed in section IX.B.1. of this final rule with comment period;

(4) The possibility of extending the direct supervision requirements for hospital-based partial hospitalization program services to those same services in community mental health centers, as discussed in section XII.D.3. of this final rule with comment period; and

(5) The possibility of establishing direct physician supervision requirements for ASC services, as discussed in section XV.A.3. of this final rule with comment period.

Inspection of Public Comments: All comments received before the close of the comment period are available for viewing by the public, including any personally identifiable or confidential business information that is included in a comment. We post all comments received before the close of the comment period on the following Web site as soon as possible after they have been received: <http://www.regulations.gov>. Follow the search

instructions on that Web site to view public comments.

Comments received timely will also be available for public inspection as they are received, generally beginning approximately 3 weeks after publication of a document, at the headquarters of the Centers for Medicare & Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244, on Monday through Friday of each week from 8:30 a.m. to 4 p.m. EST. To schedule an appointment to view public comments, phone 1-800-743-3951.

Electronic Access

This **Federal Register** document is also available from the **Federal Register** online database through *GPO Access*, a service of the U.S. Government Printing Office. Free public access is available on a Wide Area Information Server (WAIS) through the Internet and via asynchronous dial-in. Internet users can access the database by using the World Wide Web; the Superintendent of Documents' home page address is <http://www.gpoaccess.gov/index.html>, by using local WAIS client software, or by telnet to *swais.access.gpo.gov*, then login as guest (no password required). Dial-in users should use communications software and modem to call (202) 512-1661; type swais, then login as guest (no password required).

Alphabetical List of Acronyms Appearing in This Final Rule

ACEP American College of Emergency Physicians	CPT [Physicians'] Current Procedural Terminology, Fourth Edition, 2009, copyrighted by the American Medical Association	OPD [Hospital] Outpatient department
AHA American Hospital Association	CR Cardiac rehabilitation	OPPS [Hospital] Outpatient prospective payment system
AHIMA American Health Information Management Association	CRNA Certified registered nurse anesthetist	PBD Provider-based department
AMA American Medical Association	CY Calendar year	PHP Partial hospitalization program
AMP Average manufacturer price	DMEPOS Durable medical equipment, prosthetics, orthotics, and supplies	PM Program memorandum
AOA American Osteopathic Association	DMERC Durable medical equipment regional carrier	PPI Producer Price Index
APC Ambulatory payment classification	DRA Deficit Reduction Act of 2005, Public Law 109-171	PPS Prospective payment system
ASC Ambulatory Surgical Center	DSH Disproportionate share hospital	PR Pulmonary rehabilitation
ASP Average sales price	EACH Essential Access Community Hospital	PRA Paperwork Reduction Act
AWP Average wholesale price	E/M Evaluation and management	QAPI Quality Assessment and Performance Improvement
BBA Balanced Budget Act of 1997, Public Law 105-33	EPO Erythropoietin	QIO Quality Improvement Organization
BBRA Medicare, Medicaid, and SCHIP [State Children's Health Insurance Program] Balanced Budget Refinement Act of 1999, Public Law 106-113	ESRD End-stage renal disease	RAC Recovery Audit Contractor
BCA Blue Cross Association	FACA Federal Advisory Committee Act, Public Law 92-463	RFA Regulatory Flexibility Act
BCBSA Blue Cross and Blue Shield Association	FAR Federal Acquisition Regulations	RHQDAPU Reporting Hospital Quality Data for Annual Payment Update [Program]
BIPA Medicare, Medicaid, and SCHIP Benefits Improvement and Protection Act of 2000, Public Law 106-554	FDA Food and Drug Administration	RHHI Regional home health intermediary
CAH Critical access hospital	FFS Fee-for-service	SBA Small Business Administration
CAP Competitive Acquisition Program	FSS Federal Supply Schedule	SCH Sole community hospital
CBSA Core-Based Statistical Area	FTE Full-time equivalent	SDP Single Drug Pricer
CCR Cost-to-charge ratio	FY Federal fiscal year	SI Status indicator
CERT Comprehensive Error Rate Testing	GAO Government Accountability Office	TEFRA Tax Equity and Fiscal Responsibility Act of 1982, Public Law 97-248
CKD Chronic kidney disease	GME Graduate medical education	TOPS Transitional outpatient payments
CMHC Community mental health center	HCPCS Healthcare Common Procedure Coding System	USPDI United States Pharmacopoeia Drug Information
CMS Centers for Medicare & Medicaid Services	HCRIS Hospital Cost Report Information System	WAC Wholesale acquisition cost
CORF Comprehensive outpatient rehabilitation facility	HHA Home health agency	
	HIPAA Health Insurance Portability and Accountability Act of 1996, Public Law 104-191	
	HOPD Hospital outpatient department	
	HOPQDRP Hospital Outpatient Quality Data Reporting Program	
	ICD-9-CM International Classification of Diseases, Ninth Edition, Clinical Modification	
	ICR Intensive cardiac rehabilitation	
	IDE Investigational device exemption	
	IME Indirect medical education	
	I/OCE Integrated Outpatient Code Editor	
	IOL Intraocular lens	
	IPPS [Hospital] Inpatient prospective payment system	
	IVIG Intravenous immune globulin	
	KDE Kidney disease education	
	MAC Medicare Administrative Contractor	
	MedPAC Medicare Payment Advisory Commission	
	MDH Medicare-dependent, small rural hospital	
	MIEA-TRHCA Medicare Improvements and Extension Act Under Division B, Title I of the Tax Relief Health Care Act of 2006, Public Law 109-432	
	MIPPA Medicare Improvements for Patients and Providers Act of 2008, Public Law 110-275	
	MMA Medicare Prescription Drug, Improvement, and Modernization Act of 2003, Public Law 108-173	
	MMSEA Medicare, Medicaid, and SCHIP Extension Act of 2007, Public Law 110-173	
	MPFS Medicare Physician Fee Schedule	
	MSA Metropolitan Statistical Area	
	NCCI National Correct Coding Initiative	
	NCD National Coverage Determination	
	NTIOL New technology intraocular lens	
	OIG [HHS] Office of the Inspector General	
	OMB Office of Management and Budget	

In this document, we address two payment systems under the Medicare program: the hospital outpatient prospective payment system (OPPS) and the revised ambulatory surgical center (ASC) payment system. The provisions relating to the OPPS are included in sections I. through XIV., and XVI. through XXI. of this final rule with comment period and in Addenda A, B, C (Addendum C is available on the Internet only; we refer readers to section XVIII.A. of this final rule with comment period), D1, D2, E, L, and M to this final rule with comment period. The provisions related to the revised ASC payment system are included in sections XV., XVI., and XVIII. through XXI. of this final rule with comment period and in Addenda AA, BB, DD1, DD2, and EE to this final rule with comment period. (Addendum EE is available on the Internet only; we refer readers to section XVIII.B. of this final rule with comment period.)

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- I. Background and Summary of the CY 2010 OPSS/ASC Final Rule With Comment Period**
- A. Legislative and Regulatory Authority for the Hospital Outpatient Prospective Payment System*
- When Title XVIII of the Social Security Act (the Act) was enacted, Medicare payment for hospital outpatient services was based on hospital-specific costs. In an effort to ensure that Medicare and its beneficiaries pay appropriately for services and to encourage more efficient delivery of care, the Congress mandated replacement of the reasonable cost-based payment methodology with a prospective payment system (PPS). The Balanced Budget Act (BBA) of 1997 (Pub. L. 105-33) added section 1833(t) to the Act authorizing implementation of a PPS for hospital outpatient services. The OPSS was first implemented for services furnished on or after August 1, 2000. Implementing regulations for the OPSS are located at 42 CFR part 419.
- The Medicare, Medicaid, and SCHIP Balanced Budget Refinement Act (BBRA) of 1999 (Pub. L. 106-113) made major changes in the hospital outpatient prospective payment system (OPSS). The following Acts made additional changes to the OPSS: the Medicare, Medicaid, and SCHIP Benefits Improvement and Protection Act (BIPA) of 2000 (Pub. L. 106-554); the Medicare Prescription Drug, Improvement, and Modernization Act (MMA) of 2003 (Pub. L. 108-173); the Deficit Reduction Act (DRA) of 2005 (Pub. L. 109-171), enacted on February 8, 2006; the Medicare Improvements and Extension Act under Division B of Title I of the Tax Relief and Health Care Act (MIEA-TRHCA) of 2006 (Pub. L. 109-432), enacted on December 20, 2006; the Medicare, Medicaid, and SCHIP Extension Act (MMSEA) of 2007 (Pub. L. 110-173), enacted on December 29, 2007; and the Medicare Improvements for Patients and Providers Act (MIPPA) of 2008 (Pub. L. 110-275), enacted on July 15, 2008.
- Under the OPSS, we pay for hospital outpatient services on a rate-per-service basis that varies according to the ambulatory payment classification (APC) group to which the service is assigned. We use the Healthcare Common Procedure Coding System (HCPCS) codes (which include certain Current Procedural Terminology (CPT) codes) and descriptors to identify and

group the services within each APC group. The OPPS includes payment for most hospital outpatient services, except those identified in section I.B. of this final rule with comment period. Section 1833(t)(1)(B)(ii) of the Act provides for payment under the OPPS for hospital outpatient services designated by the Secretary (which includes partial hospitalization services furnished by community mental health centers (CMHCs) and hospital outpatient services that are furnished to inpatients who have exhausted their Part A benefits, or who are otherwise not in a covered Part A stay. Section 611 of Public Law 108–173 added provisions for Medicare coverage for an initial preventive physical examination, subject to the applicable deductible and coinsurance, as an outpatient department service, payable under the OPPS.

The OPPS rate is an unadjusted national payment amount that includes the Medicare payment and the beneficiary copayment. This rate is divided into a labor-related amount and a nonlabor-related amount. The labor-related amount is adjusted for area wage differences using the hospital inpatient wage index value for the locality in which the hospital or CMHC is located.

All services and items within an APC group are comparable clinically and with respect to resource use (section 1833(t)(2)(B) of the Act). In accordance with section 1833(t)(2) of the Act, subject to certain exceptions, services and items within an APC group cannot be considered comparable with respect to the use of resources if the highest median (or mean cost, if elected by the Secretary) for an item or service in the APC group is more than 2 times greater than the lowest median cost for an item or service within the same APC group (referred to as the “2 times rule”). In implementing this provision, we generally use the median cost of the item or service assigned to an APC group.

For new technology items and services, special payments under the OPPS may be made in one of two ways. Section 1833(t)(6) of the Act provides for temporary additional payments, which we refer to as “transitional pass-through payments,” for at least 2 but not more than 3 years for certain drugs, biological agents, brachytherapy devices used for the treatment of cancer, and categories of other medical devices. For new technology services that are not eligible for transitional pass-through payments, and for which we lack sufficient data to appropriately assign them to a clinical APC group, we have established special APC groups based

on costs, which we refer to as New Technology APCs. These New Technology APCs are designated by cost bands which allow us to provide appropriate and consistent payment for designated new procedures that are not yet reflected in our claims data. Similar to pass-through payments, an assignment to a New Technology APC is temporary; that is, we retain a service within a New Technology APC until we acquire sufficient data to assign it to a clinically appropriate APC group.

B. Excluded OPPS Services and Hospitals

Section 1833(t)(1)(B)(i) of the Act authorizes the Secretary to designate the hospital outpatient services that are paid under the OPPS. While most hospital outpatient services are payable under the OPPS, section 1833(t)(1)(B)(iv) of the Act excludes payment for ambulance, physical and occupational therapy, and speech-language pathology services, for which payment is made under a fee schedule. Section 614 of Public Law 108–173 amended section 1833(t)(1)(B)(iv) of the Act to exclude payment for screening and diagnostic mammography services from the OPPS. The Secretary exercised the authority granted under the statute to also exclude from the OPPS those services that are paid under fee schedules or other payment systems. Such excluded services include, for example, the professional services of physicians and nonphysician practitioners paid under the Medicare Physician Fee Schedule (MPFS); laboratory services paid under the clinical diagnostic laboratory fee schedule (CLFS); services for beneficiaries with end-stage renal disease (ESRD) that are paid under the ESRD composite rate; and services and procedures that require an inpatient stay that are paid under the hospital inpatient prospective payment system (IPPS). We set forth the services that are excluded from payment under the OPPS in § 419.22 of the regulations.

Under § 419.20(b) of the regulations, we specify the types of hospitals and entities that are excluded from payment under the OPPS. These excluded entities include: Maryland hospitals, but only for services that are paid under a cost containment waiver in accordance with section 1814(b)(3) of the Act; critical access hospitals (CAHs); hospitals located outside of the 50 States, the District of Columbia, and Puerto Rico; and Indian Health Service hospitals.

C. Prior Rulemaking

On April 7, 2000, we published in the **Federal Register** a final rule with comment period (65 FR 18434) to implement a prospective payment system for hospital outpatient services. The hospital OPPS was first implemented for services furnished on or after August 1, 2000. Section 1833(t)(9) of the Act requires the Secretary to review certain components of the OPPS, not less often than annually, and to revise the groups, relative payment weights, and other adjustments that take into account changes in medical practices, changes in technologies, and the addition of new services, new cost data, and other relevant information and factors.

Since initially implementing the OPPS, we have published final rules in the **Federal Register** annually to implement statutory requirements and changes arising from our continuing experience with this system. These rules can be viewed on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/>. We published in the **Federal Register** on November 18, 2008 the CY 2009 OPPS/ASC final rule with comment period (73 FR 68502). In that final rule with comment period, we revised the OPPS to update the payment weights and conversion factor for services payable under the CY 2009 OPPS on the basis of claims data from January 1, 2007, through December 31, 2007, and to implement certain provisions of Public Law 110–173 and Public Law 110–275. In addition, we responded to public comments received on the provisions of the November 27, 2007 final rule with comment period (72 FR 66580) pertaining to the APC assignment of HCPCS codes identified in Addendum B to that rule with the new interim (“NI”) comment indicator, and public comments received on the July 18, 2008 OPPS/ASC proposed rule for CY 2009 (73 FR 41416).

Subsequent to publication of the CY 2009 OPPS/ASC final rule with comment period, we published in the **Federal Register** on January 26, 2009, a correction notice (74 FR 4343 through 4344) to correct certain technical errors in the CY 2009 OPPS/ASC final rule with comment period.

On July 20, 2009, we issued in the **Federal Register** (74 FR 35232) a proposed rule for the CY 2010 OPPS/ASC payment system to implement statutory requirements and changes arising from our continuing experience with both systems.

D. Advisory Panel on Ambulatory Payment Classification (APC) Groups

1. Authority of the APC Panel

Section 1833(t)(9)(A) of the Act, as amended by section 201(h) of Public Law 106–113, and redesignated by section 202(a)(2) of Public Law 106–113, requires that we consult with an outside panel of experts to review the clinical integrity of the payment groups and their weights under the OPSS. The Act further specifies that the panel will act in an advisory capacity. The Advisory Panel on Ambulatory Payment Classification (APC) Groups (the APC Panel), discussed under section I.D.2. of this final rule with comment period, fulfills these requirements. The APC Panel is not restricted to using data compiled by CMS, and it may use data collected or developed by organizations outside the Department in conducting its review.

2. Establishment of the APC Panel

On November 21, 2000, the Secretary signed the initial charter establishing the APC Panel. This expert panel, which may be composed of up to 15 representatives of providers (currently employed full-time, not as consultants, in their respective areas of expertise) subject to the OPSS, reviews clinical data and advises CMS about the clinical integrity of the APC groups and their payment weights. The APC Panel is technical in nature, and it is governed by the provisions of the Federal Advisory Committee Act (FACA). Since its initial chartering, the Secretary has renewed the APC Panel's charter four times: on November 1, 2002; on November 1, 2004; on November 21, 2006; and on November 2, 2008. The current charter specifies, among other requirements, that: the APC Panel continues to be technical in nature; is governed by the provisions of the FACA; may convene up to three meetings per year; has a Designated Federal Officer (DFO); and is chaired by a Federal official designated by the Secretary.

The current APC Panel membership and other information pertaining to the APC Panel, including its charter, **Federal Register** notices, membership, meeting dates, agenda topics, and meeting reports, can be viewed on the CMS Web site at: http://www.cms.hhs.gov/FACA/05_AdvisoryPanelonAmbulatoryPaymentClassificationGroups.asp#TopOfPage.

3. APC Panel Meetings and Organizational Structure

The APC Panel first met on February 27 through March 1, 2001. Since the

initial meeting, the APC Panel has held 16 meetings, with the last meeting taking place on August 5 and 6, 2009. Prior to each meeting, we publish a notice in the **Federal Register** to announce the meeting and, when necessary, to solicit nominations for APC Panel membership and to announce new members.

The APC Panel has established an operational structure that, in part, includes the use of three subcommittees to facilitate its required APC review process. The three current subcommittees are the Data Subcommittee, the Visits and Observation Subcommittee, and the Packaging Subcommittee. The Data Subcommittee is responsible for studying the data issues confronting the APC Panel and for recommending options for resolving them. The Visits and Observation Subcommittee reviews and makes recommendations to the APC Panel on all technical issues pertaining to observation services and hospital outpatient visits paid under the OPSS (for example, APC configurations and APC payment weights). The Packaging Subcommittee studies and makes recommendations on issues pertaining to services that are not separately payable under the OPSS, but whose payments are bundled or packaged into APC payments. Each of these subcommittees was established by a majority vote from the full APC Panel during a scheduled APC Panel meeting, and their continuation as subcommittees was last approved at the August 2009 APC Panel meeting. At that meeting, the APC Panel recommended that the work of these three subcommittees continue, and we accept those recommendations of the APC Panel. All subcommittee recommendations are discussed and voted upon by the full APC Panel.

Discussions of the other recommendations made by the APC Panel at the August 2009 meeting are included in the sections of this final rule with comment period that are specific to each recommendation. For discussions of earlier APC Panel meetings and recommendations, we refer readers to previously published hospital OPSS/ASC proposed and final rules, the CMS Web site mentioned earlier in this section, and the FACA database at: <http://fido.gov/facadata/base/public.asp>.

Comment: Several commenters requested that CMS include ASC representation on the APC Panel. Because the revised ASC payment system is based upon the same APC groups and relative payment weights as the OPSS, the commenters believed that

ASC representation on the APC Panel would ensure input from representatives of all care settings that provide surgical services whose payment groups and payment weights are affected by the OPSS. Further, the commenters urged CMS to revise the APC Panel's charter to reflect the current alignment of the OPSS and the revised ASC payment system by including representation from the ASC industry on the APC Panel, as the commenters believed is permitted by the statute.

Response: We acknowledge that the revised ASC payment system provides Medicare payments to ASCs for surgical procedures that are based, in most cases, on the relative payment weights of the OPSS. However, CMS is statutorily required to have an appropriate selection of representatives of "providers" as members of the APC Panel. The current APC Panel charter requires that "Each Panel member must be employed full-time by a hospital, hospital system, or other Medicare provider subject to payment under the OPSS," which does not include ASCs because ASCs are not providers. We refer readers to section 1833(t)(9)(A) of the Act and § 400.202 of our regulations for specific requirements and definitions. ASCs are suppliers, not providers. The charter must comply with the statute, which does not include representatives of suppliers on the APC Panel. Therefore, although we understand the concerns of the commenters regarding ASC input on the APC Panel now that the ASC payment system is based on the OPSS relative payment weights, we cannot revise the charter to include ASC representation.

E. Background and Summary of the CY 2010 OPSS/ASC Proposed Rule

A proposed rule appeared in the July 20, 2009 **Federal Register** (74 FR 35232) that set forth proposed changes to the Medicare hospital OPSS for CY 2010 to implement statutory requirements and changes arising from our continuing experience with the system. In addition, we set forth proposed changes to the revised Medicare ASC payment system for CY 2010, including updated payment weights, covered surgical procedures, and covered ancillary items and services based on the proposed OPSS update. Finally, we set forth proposed quality measures for the Hospital Outpatient Quality Data Reporting Program (HOP QDRP) for reporting quality data for annual payment rate updates for CY 2011 and subsequent calendar years, the requirements for data collection and submission for the annual payment

update, and a proposed reduction in the OPSS payment for hospitals that fail to meet the HOP QDRP requirements for the CY 2010 payment update, in accordance with the statutory requirement. The following is a summary of the major proposed changes included in the CY 2010 OPSS/ASC proposed rule:

1. Updates Affecting OPSS Payments

In section II. of the proposed rule, we set forth—

- The methodology used to recalibrate the APC relative payment weights.
- The proposed changes to packaged services.
- The proposed update to the conversion factor used to determine payment rates under the OPSS. In this section, we set forth proposed changes in the amounts and factors for calculating the full annual update increase to the conversion factor.
- The proposed retention of our current policy to use the IPPS wage indices to adjust, for geographic wage differences, the portion of the OPSS payment rate and the copayment standardized amount attributable to labor-related cost.
- The proposed update of statewide average default CCRs.
- The proposed application of hold harmless transitional outpatient payments (TOPs) for certain small rural hospitals.
- The proposed payment adjustment for rural SCHs.
- The proposed calculation of the hospital outpatient outlier payment.
- The calculation of the proposed national unadjusted Medicare OPSS payment.
- The proposed beneficiary copayments for OPSS services.

2. OPSS Ambulatory Payment Classification (APC) Group Policies

In section III. of the proposed rule, we discussed—

- The proposed additions of new HCPCS codes to APCs.
- The proposed establishment of a number of new APCs.
- Our analyses of Medicare claims data and certain recommendations of the APC Panel.
- The application of the 2 times rule and proposed exceptions to it.
- The proposed changes to specific APCs.
- The proposed movement of procedures from New Technology APCs to clinical APCs.

3. OPSS Payment for Devices

In section IV. of the proposed rule, we discussed the proposed pass-through

payment for specific categories of devices and the proposed adjustment for devices furnished at no cost or with partial or full credit.

4. OPSS Payment Changes for Drugs, Biologicals, and Radiopharmaceuticals

In section V. of the proposed rule, we discussed the proposed CY 2010 OPSS payment for drugs, biologicals, and radiopharmaceuticals, including the proposed payment for drugs, biologicals, and radiopharmaceuticals with and without pass-through status.

5. Estimate of OPSS Transitional Pass-Through Spending for Drugs, Biologicals, Radiopharmaceuticals, and Devices

In section VI. of the proposed rule, we discussed the estimate of CY 2010 OPSS transitional pass-through spending for drugs, biologicals, and devices.

6. OPSS Payment for Brachytherapy Sources

In section VII. of the proposed rule, we discussed payment for brachytherapy sources.

7. OPSS Payment for Drug Administration Services

In section VIII. of the proposed rule, we set forth our proposed policy concerning coding and payment for drug administration services.

8. OPSS Payment for Hospital Outpatient Visits

In section IX. of the proposed rule, we set forth our proposed policies for the payment of clinic and emergency department visits and critical care services based on claims data.

9. Payment for Partial Hospitalization Services

In section X. of the proposed rule, we set forth the proposed payment for partial hospitalization services, including the proposed separate threshold for outlier payments for CMHCs.

10. Procedures That Will Be Paid Only as Inpatient Procedures

In section XI. of the proposed rule, we discussed the procedures that we proposed to remove from the inpatient list and assign to APCs for payment under the OPSS.

11. OPSS Nonrecurring Technical and Policy Changes and Clarifications

In section XII. of the proposed rule, we discussed nonrecurring technical issues, proposed policy changes, and provided policy clarifications.

12. OPSS Payment Status and Comment Indicators

In section XIII. of the proposed rule, we discussed our proposed changes to the definitions of status indicators assigned to APCs and presented our proposed comment indicators for the final rule with comment period.

13. OPSS Policy and Payment Recommendations

In section XIV. of the proposed rule, we addressed recommendations made by the Medicare Payment Advisory Commission (MedPAC) in its March 2009 report to Congress, by the Office of Inspector General (OIG), and by the APC Panel regarding the OPSS for CY 2010.

14. Updates to the Ambulatory Surgical Center (ASC) Payment System

In section XV. of the proposed rule, we discussed the proposed updates of the revised ASC payment system and payment rates for CY 2010.

15. Reporting Quality Data for Annual Payment Rate Updates

In section XVI. of the proposed rule, we discussed the proposed quality measures for reporting hospital outpatient (HOP) quality data for the annual payment update factor for CY 2011 and subsequent calendar years; set forth the requirements for data collection and submission for the annual payment update; and discussed the reduction in the OPSS payment for hospitals that fail to meet the HOP Quality Data Reporting Program (QDRP) requirements for CY 2010.

16. Healthcare-Associated Conditions

In section XVII. of the proposed rule, we discussed public responses to a December 2008 CMS public listening session addressing the potential extension of the principle of Medicare not paying more under the IPPS for the care of preventable hospital-acquired conditions experienced by a Medicare beneficiary during a hospital inpatient stay to medical care in other settings that are paid under other Medicare payment systems, including the OPSS, for those healthcare-associated conditions that occur or result from care in those other settings.

17. Regulatory Impact Analysis

In section XXI. of the proposed rule, we set forth an analysis of the impact the proposed changes would have on affected entities and beneficiaries.

F. Public Comments Received in Response to the CY 2010 OPSS/ASC Proposed Rule

We received approximately 1,527 timely pieces of correspondence containing multiple comments on the CY 2010 OPSS/ASC proposed rule. We note that we received some public comments that were outside of the scope of the CY 2010 OPSS/ASC proposed rule. These out-of-scope public comments are not addressed in this final rule with comment period.

New (and substantially revised) CY 2010 HCPCS codes are designated with comment indicator "NI" in Addenda B, AA, and BB of this final rule with comment period to signify that their CY 2010 interim OPSS and/or ASC treatment are open to public comment on this final rule with comment period. Summaries of the public comments that are within the scope of the CY 2010 proposals and our responses to those comments are set forth in the various sections of this final rule with comment period under the appropriate headings.

G. Public Comments Received in Response to the November 18, 2008 OPSS/ASC Final Rule With Comment Period

We received approximately 41 timely pieces of correspondence on the CY 2009 OPSS/ASC final rule with comment period, some of which contained multiple comments on the interim APC assignments and/or status indicators of HCPCS codes identified with comment indicator "NI" in Addendum B of that final rule with comment period. Summaries of those public comments on topics open to comment in the CY 2009 OPSS/ASC final rule with comment period and our responses to them are set forth in the various sections of this final rule with comment period under the appropriate headings.

II. Updates Affecting OPSS Payments

A. Recalibration of APC Relative Weights

1. Database Construction

a. Database Source and Methodology

Section 1833(t)(9)(A) of the Act requires that the Secretary review and revise the relative payment weights for APCs at least annually. In the April 7, 2000 OPSS final rule with comment period (65 FR 18482), we explained in detail how we calculated the relative payment weights that were implemented on August 1, 2000 for each APC group.

For CY 2010, we proposed to use the same basic methodology that we

described in the April 7, 2000 OPSS final rule with comment period to recalibrate the APC relative payment weights for services furnished on or after January 1, 2010, and before January 1, 2011 (CY 2010). That is, we proposed to recalibrate the relative payment weights for each APC based on claims and cost report data for hospital outpatient department (HOPD) services. We proposed to use the most recent available data to construct the database for calculating APC group weights. Therefore, for the purpose of recalibrating the APC relative payment weights for CY 2010, we used approximately 141 million final action claims for hospital outpatient department services furnished on or after January 1, 2008, and before January 1, 2009. (For exact counts of claims used, we refer readers to the claims accounting narrative under supporting documentation for this final rule with comment period on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/HORD/>.)

Of the 141 million final action claims for services provided in hospital outpatient settings used to calculate the CY 2010 OPSS payment rates for this final rule with comment period, approximately 107 million claims were the type of bill potentially appropriate for use in setting rates for OPSS services (but did not necessarily contain services payable under the OPSS). Of the 107 million claims, approximately 50 million claims were not for services paid under the OPSS or were excluded as not appropriate for use (for example, erroneous cost-to-charge ratios (CCRs) or no HCPCS codes reported on the claim). From the remaining 58 million claims, we created approximately 99 million single records, of which approximately 68 million were "pseudo" single or "single session" claims (created from 26 million multiple procedure claims using the process we discuss later in this section). Approximately 657,000 claims were trimmed out on cost or units in excess of ± 3 standard deviations from the geometric mean, yielding approximately 99 million single bills for median setting. As described in section II.A.2. of this final rule with comment period, our data development process is designed with the goal of using appropriate cost information in setting the APC relative weights. The bypass process is described in section II.A.1.b. of this final rule with comment period. This section discusses how we develop "pseudo" single claims, with the intention of using more appropriate data from the available claims. In some cases, the bypass process allows us to use

some portion of the submitted claim for cost estimation purposes, while the remaining information on the claim continues to be unusable. Consistent with the goal of using appropriate information in our data development process, we only use claims (or portions of each claim) that are appropriate for ratesetting purposes. Ultimately, we were able to use for CY 2010 ratesetting some portion of 95 percent of the CY 2008 claims containing services payable under the OPSS.

As proposed, the APC relative weights and payments for CY 2010 in Addenda A and B to this final rule with comment period were calculated using claims from CY 2008 that were processed before January 1, 2009 and continue to be based on the median hospital costs for services in the APC groups. We selected claims for services paid under the OPSS and matched these claims to the most recent cost report filed by the individual hospitals represented in our claims data. We continue to believe that it is appropriate to use the most current full calendar year claims data and the most recently submitted cost reports to calculate the median costs underpinning the APC relative payment weights and the CY 2010 payment rates.

We did not receive any public comments on our proposal to base the CY 2010 APC relative weights on the most currently available cost reports and on claims for services furnished in CY 2008. Therefore, for the reasons noted above in this section, we are finalizing our data source for the recalibration of the CY 2010 APC relative payment weights as proposed, without modification, as described in this section of this final rule with comment period.

b. Use of Single and Multiple Procedure Claims

For CY 2010, in general, we proposed to continue to use single procedure claims to set the medians on which the APC relative payment weights would be based, with some exceptions as discussed below in this section. We generally use single procedure claims to set the median costs for APCs because we believe that the OPSS relative weights on which payment rates are based should be derived from the costs of furnishing one procedure and because, in many circumstances, we are unable to ensure that packaged costs can be appropriately allocated across multiple procedures performed on the same date of service.

We agree that, optimally, it is desirable to use the data from as many claims as possible to recalibrate the APC relative payment weights, including

those claims for multiple procedures. As we have for several years, we continued to use date of service stratification and a list of codes to be bypassed to convert multiple procedure claims to “pseudo” single procedure claims. Through bypassing specified codes that we believe do not have significant packaged costs, we are able to use more data from multiple procedure claims. In many cases, this enables us to create multiple “pseudo” single claims from claims that were submitted as multiple procedure claims spanning multiple dates of service, or claims that contained numerous separately paid procedures reported on the same date on one claim. We refer to these newly created single procedure claims as “pseudo” single claims. The history of our use of a bypass list to generate “pseudo” single claims is well documented, most recently in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68512 through 68519). In addition, for CY 2008, we increased packaging and created the first composite APCs. This also increased the number of bills that we were able to use for median calculation by enabling us to use claims that contained multiple major procedures that previously would not have been usable. Further, for CY 2009, we expanded the composite APC model to one additional clinical area, multiple imaging services (73 FR 68559 through 68569), which also increased the number of bills we were able to use to calculate APC median costs. We refer readers to section II.A.2.e. of this final rule with comment period for discussion of the use of claims to establish median costs for composite APCs.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35239 through 35241), we proposed to continue to apply these processes to enable us to use as much claims data as possible for ratesetting for the CY 2010 OPSS. This process enabled us to create, for this final rule with comment period, approximately 68 million “pseudo” single claims, including multiple imaging composite “single session” bills (we refer readers to section II.A.2.e.(5) of this final rule with comment period for further discussion), to add to the approximately 32 million “natural” single bills. For this final rule with comment period, “pseudo” single and “single session” procedure bills represent 68 percent of all single bills used to calculate median costs.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35239 through 35241), we proposed to bypass 438 HCPCS codes for CY 2010. Since the inception of the bypass list, we have calculated the

percent of “natural” single bills that contained packaging for each HCPCS code and the amount of packaging on each “natural” single bill for each code. Each year, we generally retain the codes on the previous year’s bypass list and use the update year’s data (for CY 2010, data available for the February 2009 APC Panel meeting from CY 2008 claims processed through September 30, 2008 and CY 2007 claims data processed through June 30, 2008 used to model the final payment rates for CY 2009) to determine whether it would be appropriate to propose to add additional codes to the previous year’s bypass list. For CY 2010, we proposed to continue to bypass all of the HCPCS codes on the CY 2009 OPSS bypass list. We also proposed to add to the bypass list for CY 2010 all HCPCS codes not on the CY 2009 bypass list that, using both CY 2009 final rule and February 2009 APC Panel data, met the same previously established empirical criteria for the bypass list that are summarized below. Because we must make some assumptions about packaging in the multiple procedure claims in order to assess a HCPCS code for addition to the bypass list, we assume that the representation of packaging on “natural” single claims for any given code is comparable to packaging for that code in the multiple claims. The proposed criteria for the bypass list were:

- There are 100 or more “natural” single claims for the code. This number of single claims ensures that observed outcomes are sufficiently representative of packaging that might occur in the multiple claims.

- Five percent or fewer of the “natural” single claims for the code have packaged costs on that single claim for the code. This criterion results in limiting the amount of packaging being redistributed to the separately payable procedures remaining on the claim after the bypass code is removed and ensures that the costs associated with the bypass code represent the cost of the bypassed service.

- The median cost of packaging observed in the “natural” single claims is equal to or less than \$50. This criterion also limits the amount of error in redistributed costs. Throughout the bypass process, we do not know the dollar value of the packaged cost that should be appropriately attributed to the other procedures on the claim. Ensuring that redistributed costs associated with a bypass code are small in amount and volume protects the validity of cost estimates for low cost services billed with the bypassed service.

- The code is not a code for an unlisted service.

In addition, we proposed to continue to include on the bypass list HCPCS codes that CMS medical advisors believe have minimal associated packaging based on their clinical assessment of the complete CY 2010 OPSS proposal. Some of these codes were identified by CMS medical advisors and some were identified in prior years by commenters with specialized knowledge of the packaging associated with specific services, especially on a multiple procedure claim. We also proposed to continue to include on the bypass list certain HCPCS codes in order to purposefully direct the assignment of packaged costs to a companion code where services always appear together and where there would otherwise be few single claims available for ratesetting. For example, we have previously discussed our reasoning for adding HCPCS code G0390 (Trauma response team associated with hospital critical care service) and the CPT codes for additional hours of drug administration to the bypass list (73 FR 68513 and 71 FR 68117 through 68118).

As a result of the multiple imaging composite APCs that we established in CY 2009, we note that the program logic for creating “pseudo” singles from bypassed codes that are also members of multiple imaging composite APCs changed. When creating the set of “pseudo” single claims, claims that contain “overlap bypass codes,” that is, those HCPCS codes that are both on the bypass list and are members of the multiple imaging composite APCs, were identified first. These HCPCS codes were then processed to create multiple imaging composite “single session” bills, that is, claims containing HCPCS codes from only one imaging family, thus suppressing the initial use of these codes as bypass codes. However, these “overlap bypass codes” were retained on the bypass list because, at the end of the “pseudo” single processing logic, we reassessed the claims without suppression of the “overlap bypass codes” under our longstanding “pseudo” single process to determine whether we could convert additional claims to “pseudo” single claims. (We refer readers to section II.A.2.b. of this final rule with comment period for further discussion of the treatment of “overlap bypass codes.”) This process also created multiple imaging composite “single session” bills that could be used for calculating composite APC median costs. “Overlap bypass codes” that would be members of the proposed multiple imaging composite APCs were

identified by asterisks (*) in Table 1 of the CY 2010 OPPS/ASC proposed rule (74 FR 35242 through 35252).

At the February 2009 APC Panel Meeting, the APC Panel recommended that CMS place CPT code 76098 (Radiological examination, surgical specimen) on the bypass list and reassign the code to APC 0260 (Level I Plain Film Except Teeth) in response to a public presentation requesting that CMS make these changes. Although CPT code 76098 would not be eligible for addition to the bypass list because the frequency and magnitude of packaged costs in its "natural" single claims exceed the empirical criteria, the presenter suggested that the "natural" single claims represented aberrant billing with inappropriate packaged services and pointed out that the packaged services support the surgical procedures that commonly are also reported on claims for CPT code 76098. The presenter suggested that bypassing CPT code 76098 would properly allocate packaged costs to surgical procedures on these claims, and would increase the number of single claims available for ratesetting for both CPT code 76098 and the associated surgical breast procedures. The APC Panel indicated that the issues raised by the presenter appeared to be consistent with clinical practice and subsequently made the recommendation to bypass CPT code 76098 and reassign the code to APC 0260 based on the code's revised cost.

Based on the APC Panel's specific recommendation for CPT code 76098, we studied the billing patterns for the code in the "natural" single and multiple major claims in the CY 2008 claims data available for the February 2009 APC Panel. The presenter asserted that CPT code 76098 is commonly billed with surgical breast procedures and our claims data from the multiple procedure claims confirm this observation. However, as noted above, there are also a significant number of "natural" single bills in those data (1,303), and these "natural" single claims include costly packaged services, such as CPT code 19290 (Preoperative placement of needle localization wire, breast) and CPT 77032 code (Mammographic guidance for needle placement, breast (eg, for wire localization or for injection), each lesion, radiological supervision and interpretation). We have received anecdotal information indicating that hospitals may place guidance wires prior to surgery in the hospital's radiology department and then examine the surgical specimen in the radiology department after its surgical removal. This information,

along with the number of observed "natural" single claims, suggests that the packaged costs might appropriately be associated with the radiological examination of the breast specimen. Although bypassing CPT code 76098 would allow for the creation of more "pseudo" single claims for ratesetting, it would also require the assumption that all packaging on the claim would be correctly assigned to the remaining major procedure where it exists and that on "natural" single bills no packaging would be appropriately associated with CPT code 76098. Given the number of "natural" single bills for CPT code 76098 and the significant packaged costs on these claims, we are not confident that placement of this code on the bypass list is appropriate.

While we did not propose to place CPT code 76098 on the bypass list, we wanted to continue to provide separate payment for this procedure when appropriate. We believe that CPT code 76098 is generally ancillary and supportive to surgical breast procedures. In CY 2008 we established a group of conditionally packaged codes, called "T-packaged codes," whose payment is packaged when one or more separately paid surgical procedures with status indicator "T" are provided during a hospital encounter. In order to provide separate payment for CPT code 76098 when not provided with a separately payable surgical procedure and also to recognize its ancillary and supportive nature when it accompanies separately payable procedures, we proposed to conditionally package CPT code 76098 as a "T-packaged code" for CY 2010, identified with status indicator "Q2" in Addendum B to the CY 2010 OPPS/ASC proposed rule. As a "T-packaged code," CPT code 76098 would receive separate payment except where it appears with a surgical procedure, in which case its payment would be packaged. Designating CPT 76098 in this way allows the separate payment to appropriately account for the packaged costs that appear on the code's "natural" single bills, while also allowing us to use more multiple procedure claims that include both a surgical procedure and CPT code 76098 to set the payment rates for the related surgical procedures. The CPT code-specific median cost of CPT code 76098 in the CY 2008 claims data available for the February 2009 APC Panel meeting was approximately \$346, consistent with its CY 2009 assignment to APC 0317 (Level II Miscellaneous Radiology Procedures), which had an observed APC median cost in those data of approximately \$339. In contrast, the

median cost of APC 0260, the APC reassignment recommended by the APC Panel, was much lower in the APC Panel data, approximately \$46. Therefore, we did not accept the APC Panel's recommendation to reassign CPT code 76098. Instead, we proposed to continue its assignment to APC 0317 for CY 2010 in those cases where CPT code 76098 is separately paid.

Comment: Several commenters requested that CMS add CPT code 76098 to the bypass list and reassign it to APC 0260. The commenters believed that CPT 76098 is similar with respect to resource use to the other codes assigned to APC 0260. The commenters also claimed that including CPT code 76098 on the bypass list would appropriately make more claims available for ratesetting purposes for the CPT code itself and the surgical breast procedures that appear with CPT code 76098 in the multiple major procedure claims. Another commenter supported the proposal to not include CPT code 76098 on the CY 2010 bypass list.

Response: The hospital claims data show that there is significant packaging associated with CPT code 76098. Therefore, we believe CPT code 76098 is not appropriate for inclusion on the bypass list.

In examining the billing patterns for CPT 76098, we noted its failure to meet the empirical criteria for inclusion on the bypass list. The significant number of "natural" single claims suggests that these claims are an accurate representation of hospital billing practices in certain clinical situations. Further, we believe the packaging on these claims is properly associated with the code. Anecdotal information on the placement of wires prior to surgery suggests that the packaging on the "natural" single claims reflects appropriate billing in some clinical scenarios, such as when hospitals place guidance wires prior to surgery in the hospital's radiology department and then examine the surgical specimen in the radiology department after its surgical removal. This example illustrates appropriate billing on "natural" single claims for CPT code 76098 because the hospital has accurately reported all services that the hospital provided to the patient on the claim. In this case, the hospital did not provide the associated surgical breast procedure; therefore, all packaging would be appropriately associated with CPT code 76098, which is the separately payable service that the hospital provided to the patient. This scenario contradicts the commenter's belief that the significant packaging on the "natural" single claims for CPT code

76098 would represent aberrant hospital billing. As a result, for the CY 2010 proposed rule, we did not propose to add CPT code 76098 to the bypass list. However, based on our examination of the claims data for the proposed rule, we agreed that CPT 76098 is generally ancillary and supportive to surgical breast procedures. In order to provide appropriate separate payment for CPT code 76098 when the service is not furnished with a separately payable surgical procedure, and also to recognize its ancillary and supportive nature when it accompanies separately payable procedures, we proposed to conditionally package CPT code 76098 as a "T-packaged code" for CY 2010, identified with status indicator "Q2" in Addendum B to the proposed rule. Designating CPT code 76098 as a "T-packaged code" allows the separate payment to appropriately account for the packaged costs that appear on the code's "natural" single bills, while also allowing us to use more multiple procedure claims that include a surgical procedure and CPT code 76098 to set the payment rates for the related surgical procedures. In turn, we are able to use more data from the multiple procedure claims with CPT code 76098 to set payment rates for the surgical breast procedures on those claims. We continue to believe that classifying CPT code 76098 as a conditionally packaged code with status indicator "Q2" is the proper policy to both provide appropriate payment when the service is billed by itself and appropriate payment for the associated surgical breast procedures that it supports.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to assign status indicator "Q2" to CPT code 76098. When the service is furnished with a separately payable surgical procedure with status indicator "T" on the same day, payment for CPT code 76098 is packaged. Otherwise, payment for CPT code 76098 is made separately through APC 0317, which has a final APC median cost of approximately \$374. We are not adding CPT code 76098 to the bypass list for CY 2010.

Comment: Many commenters supported the current methodology of bypassing HCPCS codes and the goal of using more data from the multiple major claims. A few commenters noted that some of the HCPCS codes on the proposed CY 2010 bypass list do not meet the empirical criteria described above and observed that many codes that meet the empirical criteria were not included on the proposed bypass list. The commenters highlighted findings

from supporting data analysis to illustrate their points. Several commenters also raised concerns about the transparency of the bypass process. The commenters suggested that the empirical criteria were not explained clearly and were applied inconsistently. Other commenters believed that there is a lack of transparency regarding the addition of codes to the bypass list and the bypass process in general.

The commenters requested detailed explanations about which codes are included on the bypass list, asking that CMS identify any codes on the bypass list that do not meet the empirical criteria and the reason for their inclusion. Several commenters believed that modifying the specific empirical criteria that the median packaged cost be less than \$50 on less than 5 percent of "natural" single bills would increase the number of potential bypass codes and "pseudo" single claims. Some commenters suggested adopting a different threshold of some low percentage of total packaged costs on the code's single claims as a percent of total costs on all single claims. They believed that a percentage approach could provide more stability in the ratesetting process. One commenter also suggested that more generous empirical thresholds could be appropriate for a select set of HCPCS codes by subtracting the average packaged cost of the bypass code from other costs on the date of service where the code appears and is used as a bypass code, specifically to increase the number of claims available for setting payment rates for APCs for low dose rate brachytherapy services. A few commenters recommended that the median packaged cost threshold of \$50 on less than 5 percent of "natural" single bills be updated as CMS has not updated the threshold since its introduction, and one commenter claimed the packaged cost threshold was arbitrary. Several commenters also indicated that the HCPCS codes CMS proposed to add to the CY 2010 OPPS bypass list were not actually incorporated into CMS' ratesetting process.

Response: As discussed above in this section, we only apply the empirical criteria to the "natural" single claims. The bypass list is intended to consist of services that have minimal or no associated packaging, and in recent years, also includes codes for services that we wish to explicitly treat as not having packaged costs for purposes of OPPS payment. We refer readers to our previous discussions regarding the inclusion of additional hours of drug administration services (73 FR 68513) and HCPCS G0390 (71 FR 68117

through 68118) on the bypass list for further detail. Extracting "pseudo" single bills or unique estimates of a single service's total resource cost from claims containing multiple procedures requires making some assumptions about the amount of packaging associated with every service. As reflected in the CY 2005 proposed rule (69 FR 50474 through 50475), our empirical criteria of 100 "natural" single claims, 5 percent or fewer "natural" single claims with packaging, and median packaged cost less than \$50 are intended to be conservative, that is, to limit the amount and impact of redistributed packaging from expanding the bypass list. These criteria ensure that the packaged costs associated with bypass codes are limited, based on the best information that we have in the "natural" single procedure claims. Bypassing codes with significant associated packaging would inappropriately redistribute these packaged costs to major procedures billed with the bypass codes in the multiple procedure claims, when the individual line-items for the bypass codes are removed to create "pseudo" single claims. Because we recognize that the "natural" single claims are not always good representations of the code when it is reported on multiple major claims, for example, a service with only 20 "natural" single claims, we also judiciously include procedures on the bypass list that both CMS' medical advisors and public commenters identify as not including significant packaging and for which our own data analyses do not suggest that inclusion on the bypass list would result in an inappropriate redistribution of packaged costs. Finally, our general policy each year has been to retain codes from the previous year's bypass list without reevaluation of these codes in the context of the empirical criteria based upon updated data. We listed and discussed these empirical criteria most recently in the CY 2010 OPPS/ASC proposed rule (74 FR 35240 through 35241). The empirical criteria have remained unchanged since first implemented because it has been our experience that they effectively limit the inappropriate redistribution of packaged costs when we create "pseudo" single procedure claims.

In examining the empirical data provided by commenters supporting their requests for additions to the bypass list, we believe that the research supporting these public comments applied the empirical criteria to all single claims rather than only to the "natural" single claims. We note that

this application of the empirical criteria is inconsistent with our methodology of generalizing about packaging in the multiple procedure claims from the “natural” single procedure claims. We do not believe that it would be appropriate to expand the bypass list by assuming that our packaging redistribution after application of the current bypass list should be used to identify additional candidates for the bypass list. Clearly comparing all single bills, not just “natural” single bills, would lead to the conclusion that many more codes are eligible for inclusion on the bypass list but could also compound any inappropriate cost redistribution created by the current “pseudo” single claim development process. The OPSS pays for individual items and services and some APCs do not contain many services and some of these services are low cost. Further, some payment rates are based on a small sample of single procedure claims. Because redistributing even a small amount of packaging could have a potentially large impact on median costs for small volume or low cost APCs, we believe our current empirical criteria and reliance on “natural” single procedure claims provide the most appropriate bypass policy.

Some commenters indicated that a packaged cost threshold based on a percentage of low packaged costs out of total costs for all single bills would be more appropriate. We believe that using a percentage could allow some significant packaged costs to be redistributed. Specifically, implementing this change to the empirical criteria could redistribute a low percentage of packaged cost out of total cost for all single bills to a very inexpensive service, leading to potential distortions in the APC relative weights. This would be contrary to one primary purpose of the empirical criteria, which is to limit the inappropriate redistribution of packaged costs in the bypass process. We also do not understand how adopting this policy would introduce greater stability. If the policy increased the size of the bypass list, it could introduce greater instability by inappropriately redistributing more variable packaged costs from year to year. With regard to the suggestion that we subtract an average packaged cost for the bypass code from each multiple procedure claim, we believe that this would inappropriately remove cost information from the claims used for ratesetting and assume that the removal of that average cost is appropriate in most cases.

While we are not adopting the commenters' suggested revisions to the

empirical criteria for the CY 2010 OPSS bypass list, we acknowledge that the \$50 median packaged cost threshold has not been updated for several years and that the real value of this packaged cost threshold criterion has declined due to inflation. Consequently, we will consider whether it would be appropriate to update the \$50 dollar packaged cost threshold for inflation when identifying potential bypass codes in future rulemaking.

The bypass list we used to calculate payment rates for this final rule with comment period omits 11 of the 14 HCPCS codes that we newly proposed to add to the bypass list for the CY 2010 OPSS. Although these 14 proposed codes met the empirical criteria for inclusion on the bypass list for CY 2010 and although we listed them in Table 1 of the proposed rule (74 FR 35242 through 35352), we inadvertently omitted them from the bypass list that we used to calculate the median costs and payment rates that we proposed for CY 2010. To ensure consistency between the proposed rule and the final rule with comment period, we began our modeling for this final rule with comment period using the same list of bypass codes that we used to create the median costs and payment rates that we proposed for CY 2010. Three proposed radiation oncology code additions are an exception to this approach. In this final rule with comment period, we are including these three proposed bypass codes both because they meet the empirical criteria and because commenters on the CY 2010 OPSS/ASC proposed rule specifically requested that we add them to the CY 2010 bypass list. These three codes are: CPT code 77300 (Basic radiation dosimetry, central axis depth dose calculation, TDF, NSD, gap calculation, off axis factor, tissue inhomogeneity factors, calculation of non-ionizing radiation surface and depth dose, as required during course of treatment, only when prescribed by the treating physician); CPT code 77331 (Special dosimetry (e.g., TLD, microdosimetry)(specify), only when prescribed by the treating physician); and CPT code 77370 (Special medical radiation physics consultation).

Thus, the bypass list that we used to calculate the payment rates in this final rule with comment period does not include 11 of the 14 codes proposed for inclusion on the CY 2010 bypass list. These 11 HCPCS codes are identified in Table 1 of this final rule with comment period. In response to commenters' requests that we document additions to the bypass list, we have included a column in the list of bypass codes in

Table 2 to identify additions for the CY 2010 update year, and we will continue to identify new additions in future rulemaking.

Comment: A few commenters noted that CMS removed radiation oncology HCPCS codes that did not meet the empirical criteria from the bypass list for the CY 2009 OPSS/ASC final rule with comment period. Observing that this action had an adverse effect on the median costs for those codes and services frequently billed with those codes, the commenters requested that a number of the radiation oncology CPT codes be added to the bypass list, including CPT codes 77295 (Therapeutic radiology simulation-aided field setting, 3-dimensional); 77299 (Unlisted procedure, therapeutic radiology clinical treatment planning); 77300 (Basic radiation dosimetry calculation, central axis depth dose calculation, TDF, NSD, gap calculation, off axis factor, tissue inhomogeneity factors, calculation of non-ionizing radiation surface and depth dose, as required during course of treatment, only when prescribed by treating physician); 77301 (Intensity modulated radiotherapy plan, including dose-volume histograms for target and critical structure partial tolerance specifications); 77310 (Teletherapy, isodose plan (whether hand or computer calculated); intermediate (three or more treatment ports directed to a single area of interest)); 77315 (Teletherapy, Isodose plan (whether hand or computer calculated); complex (mantle or inverted Y, tangential ports, the use of wedges, compensators, complex blocking, rotational beam, or special beam considerations)); 77327 (Brachytherapy isodose plan; intermediate (multiplane dosage calculations, application involving 5 to 10 sources/ribbons, remote afterloading brachytherapy, 9 to 12 sources)); 77328 (Brachytherapy isodose plan; complex (multiplane isodose plan, volume implant calculations, over 10 sources/ribbons used, special spatial reconstruction, remote afterloading brachytherapy, over 12 sources)); 77331 (Special dosimetry (e.g., TLD, microdosimetry) (specify), only when prescribed by the treating physician); 77336 (Continuing medical physics consultation, including assessment of treatment parameters, quality assurance of dose delivery, and review of patient treatment documentation in support of the radiation oncologist, reporter per week of therapy); 77370 (Special medical radiation physics consultation); 77371 (Radiation treatment delivery, stereotactic radiosurgery (SRS),

complete course of treatment of cranial lesion(s) consisting of 1 session; multi-source Cobalt 60 based); 77401 (Radiation treatment delivery, superficial and/or ortho voltage); 77470 (Special treatment procedure (e.g., total body irradiation, hemibody radiation, per oral, endocavitary or intraoperative cone irradiation)); 77600 (Hyperthermia, externally generated; superficial (i.e., heating to a depth of 4 cm or less)); 77783 (Remote afterloading high intensity brachytherapy; 9–12 source positions or catheters); and 77789 (Surface application of radiation source).

Response: Some of the HCPCS codes that commenters suggested that we add to the bypass list are already included on the bypass list for this final rule with comment period, including CPT codes 77301, 77315, 77336, and 77401. These codes met the empirical criteria in earlier years and, because of our policy to retain codes once they have been added to the bypass list, these codes continue on the bypass list. However, many of the codes that commenters requested for addition the CY 2010 bypass list do not meet the empirical criteria because the percentage of “natural” single procedure claims with packaging exceeds 5 percent and, for some, the low volume of “natural” single claims prevents us from making an accurate assessment about packaging in the multiple procedure claims. Most of these codes have a low packaged median cost in the “natural” single procedure claims.

We examined the billing patterns for these HCPCS codes in the multiple major claims to better understand the potential impact that adding the recommended codes that do not meet the empirical criteria to the bypass list might have on the redistribution of packaged costs. We specifically analyzed the amount of packaged cost on the same date of service as the suggested bypass codes and other codes in the same clinical series as the recommended bypass codes in the multiple procedure claims, as well as the number of other procedures appearing on the same date of service, the APCs associated with these procedures, and whether any of these other procedures were already included on the bypass list. For three codes, specifically CPT codes 77600 (Hyperthermia, externally generated; superficial (i.e. heating to a depth of 4cm or less)); 77605 (Hyperthermia, externally generated; deep (i.e. heating to depths greater than 4 cm)); and 77610 (Hyperthermia generated by interstitial probe(s); 5 or fewer interstitial applicators), we did not observe a

significant amount of additional packaging on the multiple procedure claims or many other services, so we believe that including these codes on the bypass list would result in a limited amount of redistributed packaged cost. Therefore, we added these three codes to the CY 2010 bypass list. We also observed packaged costs associated with CPT code 77327, but the amount was proportionally limited relative to the procedure costs on the same date of service, and we believe that we can appropriately add this code to the CY 2010 bypass list.

As discussed above in this section, we also are adding the radiation oncology codes that we proposed to include on the CY 2010 bypass list, specifically CPT codes 77300, 77331, and 77370, because these codes meet the empirical criteria, they were proposed for addition to the bypass list, and several commenters specifically requested these codes be included on the bypass list. However, several codes in the commenters’ suggested additions to the bypass list not only failed the empirical criteria in the “natural” single procedure claims, but also were associated with significant packaged costs proportional to the costs of the other procedures appearing on the same date of service and the presence of many other separately paid procedures. Most of this packaged cost on claims for the candidate bypass codes was reported as revenue code charges without HCPCS codes, and we could not ascertain whether some of the packaging should be associated with the suggested bypass code or with one of the many other procedures appearing on the same date of service in the multiple claims. Because we would be unable to allocate the packaged cost among services or to determine that it was not associated with the candidate bypass list code, we believe it would be inappropriate to add these HCPCS codes to the bypass list. Although previous commenters have suggested that packaging of radiation guidance services in CY 2008 reduced the number of claims available for setting payment rates for radiation oncology services, it is notable that only a small portion of the packaged costs on the claims for radiation oncology services could be attributed to the radiation guidance services. In summary, we are not adding CPT codes 77295, 77299, 77310, 77328, 77371, 77470, 77783, and 77789 to the final CY 2010 bypass list.

We always appreciate the empirical information that commenters submitted regarding their suggested additions to the bypass list. However, we note that, due to the redistributive properties of

the bypass list and our process for creating “pseudo” single procedure claims, we always must examine the redistributive impact of additions to the bypass list on all HCPCS code and APC median costs. Future recommendations from the public for additions to the bypass list should consider the global impact on APCs and HCPCS codes of changes to the bypass list in order to facilitate our evaluation of codes suggested for inclusion on the bypass list in the future.

Comment: Some commenters supported the inclusion of the HCPCS codes for additional hours of drug administration on the bypass list. In addition, several commenters requested that CPT 90768 (Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); concurrent infusion (List separately in addition to code for primary procedure)) be made separately payable and added to the bypass list to ensure consistent treatment of codes for additional hours of drug administration under the bypass list.

Response: We appreciate the commenters’ support and have continued to include the separately payable codes for additional hours of drug administration on the CY 2010 bypass list. Bypassing these drug administration codes, and associating all the packaging with the code for the initial hour of drug administration, enables us to use many correctly coded claims for initial drug administration services that would otherwise not be available for ratesetting. We did not include CPT 90768 on the CY 2010 bypass list because we proposed to unconditionally package its successor code (CPT code 96368 (Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); concurrent infusion (List separately in addition to code for primary procedure))) in CY 2010 and, therefore, CPT code 90768 is not a candidate for the bypass list. Our final CY 2010 policy to package payment for CPT code 96368 is discussed in section VIII.B. of this final rule with comment period.

As discussed above, the bypass list consists of separately paid services with no or minimal packaging or separately paid services that CMS knowingly prices without including packaged costs and associates any packaging with the other service(s) billed on the same date of service. The purpose of the bypass list is to help develop better estimates of total resource costs for a given separately payable procedure through creating “pseudo” single procedure claims from the multiple procedure claims by removing line-items without

packaging from each claim's date of service. Including packaged codes on the bypass list would remove valid packaging from a multiple procedure claim and would not allow CMS to derive more estimates of a service's total resource costs from multiple procedure claims. We have previously discussed our reasons for packaging CPT code 90768 in the CY 2009 OPPS/ASC final rule with final period (73 FR 68674).

Comment: Several commenters supported the inclusion of HCPCS code G0340 (Image-guided robotic linear accelerator-based stereotactic radiosurgery, delivery including collimator changes and custom plugging, fractionated treatment, all lesion, per session, second through fifth session, maximum) on the bypass list.

Response: We appreciate the commenters' support and have continued to include HCPCS code G0340 on the CY 2010 bypass list.

Comment: One commenter requested that CMS examine whether changes to the bypass list or other edits included in CMS' ratesetting processes negatively affected the proposed CY 2010 payment

rates for APC 0651 (Complex Interstitial Radiation Source Application) and composite APC 8001 (LDR Prostate Brachytherapy Composite).

Response: In analyzing the impact of the final CY 2010 bypass list changes on APCs 0651 and 8001, we noted modest changes in both single procedure claim frequency and median costs. In the case of composite APC 8001, bypass list changes increased the single procedure claims available for ratesetting purposes and reduced the median cost by roughly 2 percent. APC 0651 experienced a modest increase of 3 percent in the single procedure claims available for ratesetting and its median cost also increased by about 3 percent. Neither APC 0651 nor composite APC 8001 experienced significant fluctuations in median cost or single procedure claim frequency due to the line-item trim discussed in section II.A.2.(a) of this final rule with comment period.

After consideration of the public comments received, we are adopting, as final, our proposed methodology to use a bypass list to create "pseudo" single claims. To ensure consistency between

the CY 2010 proposed and final rules, we began our consideration of comments using the same list of bypass codes for this final rule with comment period that we used to calculate the median costs and payment rates that we proposed for CY 2010, which was the CY 2009 final rule bypass list. We added HCPCS codes to the CY 2010 bypass list based on whether they met the empirical criteria and, if they did not, whether we believe that the amount of redistributed packaged cost that their inclusion on the bypass list would generate would be appropriate. We ultimately added seven codes to the CY 2010 bypass list. The list of CY 2010 bypass code additions that we proposed in the CY 2010 OPPS/ASC proposed rule but did not implement in this final rule with comment period appears in Table 1. Table 2 below is the final list of bypass codes for CY 2010. "Overlap bypass codes" that are members of the multiple imaging composite APCs are identified by asterisks (*) in Table 2. HCPCS codes that have been added for CY 2010 are also identified by asterisks (*) in Table 2.

TABLE 1—PROPOSED CY 2010 BYPASS CODE ADDITIONS EXCLUDED FROM FINAL CY 2010 BYPASS LIST

CY 2010 HCPCS Code	CY 2010 Short descriptor
57452	Exam of cervix w/scope.
76120	Cine/video x-rays.
76813	Ob us nuchal meas, 1 gest.
88314	Histochemical stain.
88367	Insitu hybridization, auto.
92700	Ent procedure/service.
94660	Pos airway pressure, CPAP.
95971	Analyze neurostim, simple.
99406	Behav chng smoking 3–10 min.
99407	Behav chng smoking >10 min.
G0249	Provide INR test mater/equip.

**TABLE 2.—FINAL CY 2010 BYPASS CODES FOR CREATING
“PSEUDO” SINGLE CLAIMS FOR CALCULATING MEDIAN COSTS**

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
11056	Trim skin lesions, 2 to 4		
11057	Trim skin lesions, over 4		
11300	Shave skin lesion		
11301	Shave skin lesion		
11719	Trim nail(s)		
11720	Debride nail, 1-5		
11721	Debride nail, 6 or more		
11954	Therapy for contour defects		
17000	Destruct premalg lesion		
17003	Destruct premalg les, 2-14		
29220	Strapping of low back		
31231	Nasal endoscopy, dx		
31579	Diagnostic laryngoscopy		
51798	Us urine capacity measure		
53661	Dilation of urethra		
54240	Penis study		
56820	Exam of vulva w/scope		
57150	Treat vagina infection		
67820	Revise eyelashes		
69210	Remove impacted ear wax		
69220	Clean out mastoid cavity		
70030	X-ray eye for foreign body		
70100	X-ray exam of jaw		
70110	X-ray exam of jaw		
70120	X-ray exam of mastoids		
70130	X-ray exam of mastoids		
70140	X-ray exam of facial bones		
70150	X-ray exam of facial bones		
70160	X-ray exam of nasal bones		
70200	X-ray exam of eye sockets		
70210	X-ray exam of sinuses		
70220	X-ray exam of sinuses		
70250	X-ray exam of skull		
70260	X-ray exam of skull		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
70328	X-ray exam of jaw joint		
70330	X-ray exam of jaw joints		
70336	Magnetic image, jaw joint	*	
70355	Panoramic x-ray of jaws		
70360	X-ray exam of neck		
70370	Throat x-ray & fluoroscopy		
70371	Speech evaluation, complex		
70450	Ct head/brain w/o dye	*	
70480	Ct orbit/ear/fossa w/o dye	*	
70486	Ct maxillofacial w/o dye	*	
70490	Ct soft tissue neck w/o dye	*	
70544	Mr angiography head w/o dye	*	
70551	Mri brain w/o dye	*	
71010	Chest x-ray		
71015	Chest x-ray		
71020	Chest x-ray		
71021	Chest x-ray		
71022	Chest x-ray		
71023	Chest x-ray and fluoroscopy		
71030	Chest x-ray		
71034	Chest x-ray and fluoroscopy		
71035	Chest x-ray		
71100	X-ray exam of ribs		
71101	X-ray exam of ribs/chest		
71110	X-ray exam of ribs		
71111	X-ray exam of ribs/chest		
71120	X-ray exam of breastbone		
71130	X-ray exam of breastbone		
71250	Ct thorax w/o dye	*	
72010	X-ray exam of spine		
72020	X-ray exam of spine		
72040	X-ray exam of neck spine		
72050	X-ray exam of neck spine		
72052	X-ray exam of neck spine		
72069	X-ray exam of trunk spine		
72070	X-ray exam of thoracic spine		
72072	X-ray exam of thoracic spine		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
72074	X-ray exam of thoracic spine		
72080	X-ray exam of trunk spine		
72090	X-ray exam of trunk spine		
72100	X-ray exam of lower spine		
72110	X-ray exam of lower spine		
72114	X-ray exam of lower spine		
72120	X-ray exam of lower spine		
72125	Ct neck spine w/o dye	*	
72128	Ct chest spine w/o dye	*	
72131	Ct lumbar spine w/o dye	*	
72141	Mri neck spine w/o dye	*	
72146	Mri chest spine w/o dye	*	
72148	Mri lumbar spine w/o dye	*	
72170	X-ray exam of pelvis		
72190	X-ray exam of pelvis		
72192	Ct pelvis w/o dye	*	
72202	X-ray exam sacroiliac joints		
72220	X-ray exam of tailbone		
73000	X-ray exam of collar bone		
73010	X-ray exam of shoulder blade		
73020	X-ray exam of shoulder		
73030	X-ray exam of shoulder		
73050	X-ray exam of shoulders		
73060	X-ray exam of humerus		
73070	X-ray exam of elbow		
73080	X-ray exam of elbow		
73090	X-ray exam of forearm		
73100	X-ray exam of wrist		
73110	X-ray exam of wrist		
73120	X-ray exam of hand		
73130	X-ray exam of hand		
73140	X-ray exam of finger(s)		
73200	Ct upper extremity w/o dye	*	
73218	Mri upper extremity w/o dye	*	
73221	Mri joint upr extrem w/o dye	*	
73510	X-ray exam of hip		
73520	X-ray exam of hips		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
73540	X-ray exam of pelvis & hips		
73550	X-ray exam of thigh		
73560	X-ray exam of knee, 1 or 2		
73562	X-ray exam of knee, 3		
73564	X-ray exam, knee, 4 or more		
73565	X-ray exam of knees		
73590	X-ray exam of lower leg		
73600	X-ray exam of ankle		
73610	X-ray exam of ankle		
73620	X-ray exam of foot		
73630	X-ray exam of foot		
73650	X-ray exam of heel		
73660	X-ray exam of toe(s)		
73700	Ct lower extremity w/o dye	*	
73718	Mri lower extremity w/o dye	*	
73721	Mri jnt of lwr extre w/o dye	*	
74000	X-ray exam of abdomen		
74010	X-ray exam of abdomen		
74020	X-ray exam of abdomen		
74022	X-ray exam series, abdomen		
74150	Ct abdomen w/o dye	*	
74210	Contrst x-ray exam of throat		
74220	Contrast x-ray, esophagus		
74230	Cine/vid x-ray, throat/esoph		
74246	Contrst x-ray uppr gi tract		
74247	Contrst x-ray uppr gi tract		
74249	Contrst x-ray uppr gi tract		
76100	X-ray exam of body section		
76510	Ophth us, b & quant a		
76511	Ophth us, quant a only		
76512	Ophth us, b w/non-quant a		
76513	Echo exam of eye, water bath		
76514	Echo exam of eye, thickness		
76516	Echo exam of eye		
76519	Echo exam of eye		
76536	Us exam of head and neck		
76645	Us exam, breast(s)		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
76700	Us exam, abdom, complete	*	
76705	Echo exam of abdomen	*	
76770	Us exam abdo back wall, comp	*	
76775	Us exam abdo back wall, lim	*	
76776	Us exam k transpl w/doppler	*	
76801	Ob us < 14 wks, single fetus		
76805	Ob us >= 14 wks, sngl fetus		
76811	Ob us, detailed, sngl fetus		
76816	Ob us, follow-up, per fetus		
76817	Transvaginal us, obstetric		
76830	Transvaginal us, non-ob		
76856	Us exam, pelvic, complete	*	
76857	Us exam, pelvic, limited	*	
76870	Us exam, scrotum	*	
76880	Us exam, extremity		
76970	Ultrasound exam follow-up		
76977	Us bone density measure		
77072	X-rays for bone age		
77073	X-rays, bone length studies		
77074	X-rays, bone survey, limited		
77075	X-rays, bone survey complete		
77076	X-rays, bone survey, infant		
77077	Joint survey, single view		
77078	Ct bone density, axial		
77079	Ct bone density, peripheral		
77080	Dxa bone density, axial		
77081	Dxa bone density/peripheral		
77082	Dxa bone density, vert fx		
77083	Radiographic absorptiometry		
77084	Magnetic image, bone marrow		
77300	Radiation therapy dose plan		*
77301	Radiotherapy dose plan, imrt		
77315	Teletx isodose plan complex		
77327	Brachytx isodose calc interm		*
77331	Special radiation dosimetry		*
77336	Radiation physics consult		
77370	Radiation physics consult		*

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
77401	Radiation treatment delivery		
77600	Hyperthermia treatment		*
77605	Hyperthermia treatment		*
77610	Hyperthermia treatment		*
80500	Lab pathology consultation		
80502	Lab pathology consultation		
85097	Bone marrow interpretation		
86510	Histoplasmosis skin test		
86850	RBC antibody screen		
86870	RBC antibody identification		
86880	Coombs test, direct		
86885	Coombs test, indirect, qual		
86886	Coombs test, indirect, titer		
86890	Autologous blood process		
86900	Blood typing, ABO		
86901	Blood typing, Rh (D)		
86903	Blood typing, antigen screen		
86904	Blood typing, patient serum		
86905	Blood typing, RBC antigens		
86906	Blood typing, Rh phenotype		
86930	Frozen blood prep		
86970	RBC pretreatment		
86977	RBC pretreatment, serum		
88104	Cytopath fl nongyn, smears		
88106	Cytopath fl nongyn, filter		
88107	Cytopath fl nongyn, sm/fltr		
88108	Cytopath, concentrate tech		
88112	Cytopath, cell enhance tech		
88160	Cytopath smear, other source		
88161	Cytopath smear, other source		
88162	Cytopath smear, other source		
88172	Cytopathology eval of fna		
88173	Cytopath eval, fna, report		
88182	Cell marker study		
88184	Flowcytometry/ tc, 1 marker		
88185	Flowcytometry/tc, add-on		
88300	Surgical path, gross		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
88302	Tissue exam by pathologist		
88304	Tissue exam by pathologist		
88305	Tissue exam by pathologist		
88307	Tissue exam by pathologist		
88311	Decalcify tissue		
88312	Special stains		
88313	Special stains		
88321	Microslide consultation		
88323	Microslide consultation		
88325	Comprehensive review of data		
88331	Path consult intraop, 1 bloc		
88342	Immunohistochemistry		
88346	Immunofluorescent study		
88347	Immunofluorescent study		
88348	Electron microscopy		
88358	Analysis, tumor		
88360	Tumor immunohistochem/manual		
88361	Tumor immunohistochem/comput		
88365	Insitu hybridization (fish)		
88368	Insitu hybridization, manual		
89049	Chct for mal hyperthermia		
89230	Collect sweat for test		
89240	Pathology lab procedure		
90472	Immunization admin, each add		
90474	Immune admin oral/nasal addl		
90761	Hydrate iv infusion, add-on		
90766	Ther/proph/dg iv inf, add-on		
90767	Tx/proph/dg addl seq iv inf		
90770	Sc ther infusion, addl hr		
90771	Sc ther infusion, reset pump		
90775	Tx/pro/dx inj new drug addon		
90801	Psy dx interview		
90802	Intac psy dx interview		
90804	Psytx, office, 20-30 min		
90805	Psytx, off, 20-30 min w/e&m		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
90806	Psytx, off, 45-50 min		
90807	Psytx, off, 45-50 min w/e&m		
90808	Psytx, office, 75-80 min		
90809	Psytx, off, 75-80, w/e&m		
90810	Intac psytx, off, 20-30 min		
90811	Intac psytx, 20-30, w/e&m		
90812	Intac psytx, off, 45-50 min		
90816	Psytx, hosp, 20-30 min		
90818	Psytx, hosp, 45-50 min		
90826	Intac psytx, hosp, 45-50 min		
90845	Psychoanalysis		
90846	Family psytx w/o patient		
90847	Family psytx w/patient		
90853	Group psychotherapy		
90857	Intac group psytx		
90862	Medication management		
92002	Eye exam, new patient		
92004	Eye exam, new patient		
92012	Eye exam established pat		
92014	Eye exam & treatment		
92020	Special eye evaluation		
92025	Corneal topography		
92081	Visual field examination(s)		
92082	Visual field examination(s)		
92083	Visual field examination(s)		
92135	Ophth dx imaging post seg		
92136	Ophthalmic biometry		
92225	Special eye exam, initial		
92226	Special eye exam, subsequent		
92230	Eye exam with photos		
92240	Icg angiography		
92250	Eye exam with photos		
92275	Electroretinography		
92285	Eye photography		
92286	Internal eye photography		
92520	Laryngeal function studies		
92541	Spontaneous nystagmus test		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
92546	Sinusoidal rotational test		
92548	Posturography		
92552	Pure tone audiometry, air		
92553	Audiometry, air & bone		
92555	Speech threshold audiometry		
92556	Speech audiometry, complete		
92557	Comprehensive hearing test		
92567	Tympanometry		
92582	Conditioning play audiometry		
92585	Auditor evoke potent, compre		
92603	Cochlear implt f/up exam 7 >		
92604	Reprogram cochlear implt 7 >		
92626	Eval aud rehab status		
93005	Electrocardiogram, tracing		
93017	Cardiovascular stress test		
93225	ECG monitor/record, 24 hrs		
93226	ECG monitor/report, 24 hrs		
93231	ECG monitor/record, 24 hrs		
93232	ECG monitor/report, 24 hrs		
93236	ECG monitor/report, 24 hrs		
93270	ECG recording		
93271	ECG/monitoring and analysis		
93278	ECG/signal-averaged		
93727	Analyze ilr system		
93731	Analyze pacemaker system		
93732	Analyze pacemaker system		
93733	Telephone analy, pacemaker		
93734	Analyze pacemaker system		
93735	Analyze pacemaker system		
93736	Telephonic analy, pacemaker		
93741	Analyze ht pace device sngl		
93742	Analyze ht pace device sngl		
93743	Analyze ht pace device dual		
93744	Analyze ht pace device dual		
93786	Ambulatory BP recording		
93788	Ambulatory BP analysis		
93797	Cardiac rehab		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
93798	Cardiac rehab/monitor		
93875	Extracranial study		
93880	Extracranial study		
93882	Extracranial study		
93886	Intracranial study		
93888	Intracranial study		
93922	Extremity study		
93923	Extremity study		
93924	Extremity study		
93925	Lower extremity study		
93926	Lower extremity study		
93930	Upper extremity study		
93931	Upper extremity study		
93965	Extremity study		
93970	Extremity study		
93971	Extremity study		
93975	Vascular study		
93976	Vascular study		
93978	Vascular study		
93979	Vascular study		
93990	Doppler flow testing		
94015	Patient recorded spirometry		
94690	Exhaled air analysis		
95115	Immunotherapy, one injection		
95117	Immunotherapy injections		
95165	Antigen therapy services		
95250	Glucose monitoring, cont		
95805	Multiple sleep latency test		
95806	Sleep study, unattended		
95807	Sleep study, attended		
95808	Polysomnography, 1-3		
95812	Eeg, 41-60 minutes		
95813	Eeg, over 1 hour		
95816	Eeg, awake and drowsy		
95819	Eeg, awake and asleep		
95822	Eeg, coma or sleep only		
95869	Muscle test, thor paraspinal		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
95872	Muscle test, one fiber		
95900	Motor nerve conduction test		
95921	Autonomic nerv function test		
95925	Somatosensory testing		
95926	Somatosensory testing		
95930	Visual evoked potential test		
95950	Ambulatory eeg monitoring		
95953	EEG monitoring/computer		
95970	Analyze neurostim, no prog		
95972	Analyze neurostim, complex		
95974	Cranial neurostim, complex		
95978	Analyze neurostim brain/1h		
96000	Motion analysis, video/3d		
96101	Psycho testing by psych/phys		
96111	Developmental test, extend		
96116	Neurobehavioral status exam		
96118	Neuropsych tst by psych/phys		
96119	Neuropsych testing by tec		
96150	Assess hlth/behave, init		
96151	Assess hlth/behave, subseq		
96152	Intervene hlth/behave, indiv		
96153	Intervene hlth/behave, group		
96402	Chemo hormon antineopl sq/im		
96411	Chemo, iv push, addl drug		
96415	Chemo, iv infusion, addl hr		
96417	Chemo iv infus each addl seq		
96423	Chemo ia infuse each addl hr		
96900	Ultraviolet light therapy		
96910	Photochemotherapy with UV-B		
96912	Photochemotherapy with UV-A		
96913	Photochemotherapy, UV-A or B		
96920	Laser tx, skin < 250 sq cm		
98925	Osteopathic manipulation		
98926	Osteopathic manipulation		
98927	Osteopathic manipulation		
98940	Chiropractic manipulation		
98941	Chiropractic manipulation		

CY 2010 HCPCS Code	CY 2010 Short Descriptor	"Overlap Bypass Codes"	CY 2010 additions
98942	Chiropractic manipulation		
99204	Office/outpatient visit, new		
99212	Office/outpatient visit, est		
99213	Office/outpatient visit, est		
99214	Office/outpatient visit, est		
99241	Office consultation		
99242	Office consultation		
99243	Office consultation		
99244	Office consultation		
99245	Office consultation		
0144T	CT heart wo dye; qual calc		
G0008	Admin influenza virus vac		
G0101	CA screen;pelvic/breast exam		
G0127	Trim nail(s)		
G0130	Single energy x-ray study		
G0166	Extrnl counterpulse, per tx		
G0175	OPPS Service,sched team conf		
G0340	Robt lin-radsurg fractx 2-5		
G0344	Initial preventive exam		
G0365	Vessel mapping hemo access		
G0367	EKG tracing for initial prev		
G0376	Smoke/tobacco counseling >10		
G0389	Ultrasound exam AAA screen		
G0390	Trauma Respons w/hosp criti		
M0064	Visit for drug monitoring		
Q0091	Obtaining screen pap smear		

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c. Calculation of CCRs

(1) Development of the CCRs

We calculated hospital-specific overall ancillary CCRs and hospital-specific departmental CCRs for each hospital for which we had CY 2008 claims data from the most recent available hospital cost reports, in most cases, cost reports beginning in CY 2007. For the CY 2010 OPPS ratesetting, we used the set of claims processed during CY 2008. We applied the hospital-specific CCR to the hospital's charges at the most detailed level possible, based on a revenue code-to-cost center crosswalk that contains a hierarchy of CCRs used to estimate costs from charges for each revenue code. That crosswalk is available for review

and continuous comment on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/03_crosswalk.asp#TopOfPage. We calculated CCRs for the standard and nonstandard cost centers accepted by the electronic cost report database. In general, the most detailed level at which we calculated CCRs was the hospital-specific departmental level. For a discussion of the hospital-specific overall ancillary CCR calculation, we refer readers to the CY 2007 OPPS/ASC final rule with comment period (71 FR 67983 through 67985).

In the CY 2010 OPPS/ASC proposed rule (74 FR 35253), we proposed to continue using the hospital-specific overall ancillary and departmental CCRs to convert charges on the claims reported under specific revenue codes

to estimated costs through application of a revenue code-to-cost center crosswalk for CY 2010.

We did not receive any public comments on this proposal. Therefore, we are finalizing our proposal for CY 2010, without modification, to calculate hospital-specific overall and departmental CCRs as described above in this section.

(2) Charge Compression

Since the implementation of the OPPS, some commenters have raised concerns about potential bias in the OPPS cost-based weights due to "charge compression," which is the practice of applying a lower charge markup to higher-cost services and a higher charge markup to lower-cost services. (We discuss our CCR calculation in section

II.A.1.c. of this final rule with comment period and how we use these CCRs to estimate cost on hospital outpatient claims in detail in section II.A.2.a. of this final rule with comment period). As a result, the cost-based weights incorporate aggregation bias, undervaluing high cost items and overvaluing low cost items when an estimate of average markup, embodied in a single CCR, is applied to items of widely varying costs in the same cost center. Commenters on previous rules have expressed increased concern about the impact of charge compression when CMS began setting the relative weights for payment under the IPPS based on the costs of inpatient hospital services, rather than the charges for the services.

To explore this issue, in August 2006 we awarded a contract to RTI International (RTI) to study the effects of charge compression in calculating the IPPS relative weights, particularly with regard to the impact on inpatient diagnosis-related group (DRG) payments, and to consider methods to capture better the variation in cost and charges for individual services when calculating costs for the IPPS relative weights across services in the same cost center. Of specific note was RTI's analysis of a regression-based methodology estimating an average adjustment for CCR by type of revenue code from an observed relationship between provider cost center CCRs and proportional billing of high and low cost services in the revenue codes associated with the cost center in the claims data. RTI issued a report in March 2007 with its findings on charge compression. The report is available on the CMS Web site at: <http://www.cms.hhs.gov/reports/downloads/Dalton.pdf>. Although this report was focused largely on charge compression in the context of the IPPS cost-based relative weights, several of the findings were relevant to the OPSS. Therefore, we discussed the findings and our responses to that report in the CY 2008 OPSS/ASC proposed rule (72 FR 42641 through 42643) and reiterated them in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66599 through 66602).

RTI noted in its 2007 report that its research was limited to IPPS DRG cost-based weights and that it did not examine potential areas of charge compression specific to hospital outpatient services. We were concerned that the analysis was too limited in scope because typically hospital cost report CCRs encompass both inpatient and outpatient services for each cost center. Further, because both the IPPS and OPSS rely on cost-based weights, we preferred to introduce any

methodological adjustments to both payment systems at the same time. We believe that because charge compression affects the cost estimates for services paid under both IPPS and OPSS in the same way, it is appropriate that we would use the same or, at least, similar approaches to address the issue. Finally, we noted that we wished to assess the educational activities being undertaken by the hospital community to improve cost reporting accuracy in response to RTI's findings, either as an adjunct to or in lieu of regression-based adjustments to CCRs.

We expanded RTI's analysis of charge compression to incorporate outpatient services. In August 2007, we again contracted with RTI. Under this contract, we asked RTI to evaluate the cost estimation process for the OPSS relative weights. This research included a reassessment of the regression-based CCR models using hospital outpatient and inpatient charge data, as well as a detailed review of the OPSS revenue code-to-cost center crosswalk and the OPSS' hospital-specific CCR methodology. In evaluating cost-based estimation, in general, the results of RTI's analyses impact both the OPSS APC relative weights and the IPPS MS-DRG (Medicare severity) relative weights. The RTI final report can be found on RTI's Web site at: http://www.rti.org/reports/cms/HHSM-500-2005-00291/PDF/Refining_Cost_to_Charge_Ratios_200807_Final.pdf. For a complete discussion of the RTI recommendations, public comments, and our responses, we refer readers to the CY 2009 OPSS/ASC final rule with comment period (73 FR 68519 through 68527).

In the FY 2009 IPPS final rule, we finalized our proposal for both the OPSS and IPPS to add one cost center to the cost report so that, in general, the costs and charges for relatively inexpensive medical supplies would be reported separately from the costs and charges for more expensive implantable devices (such as pacemakers and other implantable devices). Specifically, we created one cost center for "Medical Supplies Charged to Patients" and one cost center for "Implantable Devices Charged to Patients." This change split the CCR for "Medical Supplies and Equipment" into one CCR for medical supplies and another CCR for implantable devices. In response to the majority of commenters on the proposal set forth in the FY 2009 IPPS proposed rule, we finalized a definition of the "Implantable Devices Charged to Patients" cost center as capturing the costs and charges billed with the following UB-04 revenue codes: 0275

(Pacemaker), 0276 (Intraocular lens), 0278 (Other implants), and 0624 (FDA investigational devices). We made this change to the cost report form for cost reporting periods beginning in the spring of 2009. Because there is generally a 3-year lag between the availability of cost report data for IPPS and OPSS ratesetting purposes in a given calendar year, we believe we will be able to use data from the revised cost report form to estimate costs from charges associated with UB-04 revenue codes 0275, 0276, 0278, and 0624 for implantable devices in order to more accurately estimate the costs of device-related procedures for the CY 2013 OPSS relative weights. For a complete discussion of the proposal, public comments, and our responses, we refer readers to the FY 2009 IPPS final rule (73 FR 48458 through 45467).

For the CY 2009 OPSS/ASC proposed rule, we made a similar proposal for drugs, proposing to split the "Drugs Charged to Patients" cost center into two cost centers: one for drugs with high pharmacy overhead costs and one for drugs with low pharmacy overhead costs (73 FR 41492). We noted that we expected that CCRs from the proposed new cost centers would be available in 2 to 3 years to refine OPSS drug cost estimates by accounting for differential hospital markup practices for drugs with high and low pharmacy overhead costs. However, after consideration of the public comments received and the APC Panel recommendations, we did not finalize our proposal to split the single standard "Drugs Charged to Patients" cost center into two cost centers, and instead indicated in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68659) that we would continue to explore other potential approaches to improve our drug cost estimation methodology. Unlike implantable devices, we do not currently have a policy to address charge compression in our cost estimation for expensive drugs and biologicals. In section V.B.3. of the CY 2010 OPSS/ASC proposed rule (74 FR 35326 through 35333), we proposed an adjustment to our cost estimation methodology for drugs and biologicals to address charge compression by proposing to shift a portion of the pharmacy overhead cost associated with packaged drugs and biologicals from those packaged drugs and biologicals to separately payable drugs and biologicals; proposing payment for separately payable drugs and biologicals at ASP+4 percent; and proposing a proportional reduction in the total amount of pharmacy overhead cost

associated with packaged drugs and biologicals prior to our estimating the total resource costs of individual OPSS services.

Finally, in the CY 2009 OPSS/ASC final rule with comment period, we indicated that we would be making some OPSS-specific changes in response to the RTI report recommendations. With regard to modifying the cost reporting preparation software in order to impose fixed descriptions for nonstandard cost centers, we indicated that the change would be made for the next release of the cost report software. We anticipate that these changes will be made to the cost reporting software in CY 2010 and will act as a quality check for hospitals to review their choice of nonstandard cost center code to ensure that the reporting of nonstandard cost centers is accurate, while not significantly increasing provider burden. In addition to improving the reporting mechanism for the nonstandard cost centers, we indicated in the CY 2009 OPSS/ASC final rule with comment period that we also planned to add the new nonstandard cost centers for Cardiac Rehabilitation, Hyperbaric Oxygen Therapy, and Lithotripsy. We expect that changes to add these nonstandard cost centers also will be made for cost reports beginning in CY 2010. Furthermore, we noted in the FY 2010 IPPS final rule (74 FR 43781 through 43782) that we are updating the cost report form to eliminate outdated requirements, in conjunction with the Paperwork Reduction Act (PRA), and that we had proposed actual changes to the cost reporting form, the attending cost reporting software, and the cost report instructions in Chapters 36 and 40 of the PRM-II. The comment period for this proposal (74 FR 31738) ended on August 31, 2009. We believe that improved cost report software, the incorporation of new nonstandard cost centers, and elimination of outdated requirements will improve the accuracy of the cost data contained in the electronic cost report data files and, therefore, the accuracy of our cost estimation processes for the OPSS relative weights. As has been described above, CMS has taken steps to address charge compression in the IPPS and OPSS, and continues to examine ways in which it can improve the accuracy of its cost estimation process.

Comment: Several commenters expressed support for the policy adopted in the FY 2009 IPPS final rule, with application to both the OPSS and IPPS, to create one cost center for "Medical Supplies Charged to Patients" and one cost center for "Implantable

Devices Charged to Patients." Some commenters recommended that CMS verify the accuracy of the CCRs derived from the new cost centers by comparing CCRs calculated from the new cost center against regression-based CCRs or by undertaking other activities to ensure that data reported in these revised cost centers are consistent and accurate.

One commenter stated that hospitals are reluctant to bill for devices that do not remain in the patient upon discharge, specifically cryoablation probes, under revenue code 0278 (*Medical/Surgical Supplies: Other Implants*). The commenter requested that CMS work with hospitals to revise the common hospital practice of billing for cryoablation probes under revenue code 0272 (*Medical/Surgical Supplies: Sterile Supplies*) rather than revenue code 0278. The commenter asserted that billing cryoablation probes under revenue code 0272 would result in estimating costs from charges using a CCR derived from the revised cost center for "Medical Supplies Charged to Patients," rather than one derived from the "Implantable Devices Charged to Patients," even though cryoablation probes are high cost implantable devices. The commenter believed that, without a change in the revenue code under which many hospitals report cryoablation probes, the recent cost center changes for medical supplies would negatively bias the estimated cost of cryoablation probes and the accuracy of the APC payment rates for cryoablation procedures.

Some commenters suggested that CMS engage in outreach and educational activities to hospitals on the changes to the cost report and the reporting of charges with respect to the medical device and medical supply cost centers so that hospitals can appropriately report data. The commenters recommended that the outreach activities go beyond the "distribution of bulletins that are used to inform providers about changes to the Medicare program."

Response: We appreciate the commenters' support for our CY 2009 policy to split the "Medical Supplies Charged to Patients" into one cost center for "Medical Supplies Charged to Patients" and one cost center for "Implantable Devices Charged to Patients". In the FY 2009 IPPS final rule (73 FR 48458 through 48467), we explained in detail the reasoning behind the development of the cost center split and our decision to ultimately have hospitals use the American Hospital Association's National Uniform Billing Committee (NUBC) revenue codes to determine what would be reported in

the "Medical Supplies Charged to Patients" and the "Implantable Devices Charged to Patients" cost centers. In that discussion, we noted that while we require that the device broadly be considered implantable to have its costs and charges included in the new "Implantable Devices Charged to Patients" cost center, our final policy did not require the device to remain in the patient at discharge (73 FR 48462 through 48463). We typically do not specify a revenue code-to-cost center crosswalk that hospitals must adopt to prepare their cost report, recognizing hospitals' need to interpret the NUBC definitions and cost reporting requirements within the context of their own financial systems. In response to comments on our proposal to create the new cost center in the FY 2009 IPPS final rule, we did define the new "Implantable Devices Charged to Patients" cost center by the revenue codes that we believe would map to this cost center to facilitate ease of reporting by hospitals. We note that revenue code definitions are established by the NUBC, and we fully expect hospitals to follow existing guidelines regarding revenue code use. Specifically with regard to reporting cryoablation probes, we do not believe that the current NUBC definition of revenue code 0278 (*Medical/Surgical Supplies and Devices* (also see 062x, an extension of 027x); *Other implants* (a)) precludes reporting hospital charges for cryoablation probes under this revenue code. Therefore, we believe hospitals can report charges for cryoablation probes under the revenue code 0278 using the definitions in the official UB 04 Data Specifications Manual.

As discussed in the FY 2010 IPPS final rule (74 FR 43780), we reiterated that we had not proposed any policy changes with respect to the use of revenue codes or alternative ways of identifying high-cost devices. We refer readers to the discussion in the FY 2009 IPPS final rule concerning our current policy on these matters (73 FR 48462). Hospitals were able to report costs and charges for the new "Implantable Devices Charged to Patients" cost center for cost reporting periods beginning on or after May 1, 2009 as line 55.30 on Form 2552-96 and, at the time of development of this final rule with comment period, we anticipate that hospitals will be able to report costs and charges for the new cost center as line 69 on the revised draft Medicare hospital cost report form CMS-2552-10 beginning February 1, 2010.

In the FY 2009 IPPS final rule (73 FR 48463), we agreed that once the data reflecting the cost center changes become available for ratesetting, we

would evaluate the CCRs that we derive from the new “Medical Supplies Charged to Patients” and “Implantable Devices Charged to Patients” cost centers and that we would continue to analyze cost report data. In the FY 2010 IPPS final rule (74 FR 43782), we indicated that we might consider the results of regression analyses as one way to evaluate costs and charges reported in the new cost center. However, we point out that we do not believe it is appropriate to “pick and choose” between CCRs; rather, the determining factor should be payment accuracy, regardless of whether one method increases or decreases payment for devices (73 FR 48463). That is, the validity of the CCRs resulting from the newly implemented cost center cannot be determined to be accurate simply because they will result in higher overall cost estimates for procedures that rely on implantable devices and, therefore, higher APC payment rates.

As discussed in the FY 2010 IPPS rule, we believe it is early to plan specific outreach activities on the revised cost report form CMS-2552-10 and the new “Implantable Devices Charged to Patients” cost center, given that the comment period for the revised cost reporting forms closed on August 31, 2009. We agree that such educational activities are important, and we have been considering various options for educating the provider community that would involve fiscal intermediaries, Medicare administrative contractors, and cost report vendors. We look forward to working with the provider community on these initiatives.

Comment: A few commenters noted that two revenue codes became effective for reporting radiopharmaceuticals, specifically 0343 (Nuclear Medicine; Diagnostic Radiopharmaceuticals) for diagnostic preparations and 0344 (Nuclear Medicine; Therapeutic Radiopharmaceuticals) for therapeutic preparations in October 2004; and that this more specific revenue code reporting should help capture the unique costs and charges of radiopharmaceuticals. The commenters also pointed out that the costs and charges associated with these revenue codes likely would be reported by hospitals under the broader radiology cost center on the Medicare hospital cost report. They expressed concern that, because the CCR used to estimate charges for these revenue codes encompasses a large volume of many different services, the specificity of charge information in the claims data gained through use of the new revenue codes would not translate into better

cost estimation for diagnostic and therapeutic radiopharmaceuticals under the OPSS. The commenters suggested that CMS require hospitals to report costs and charges for these two revenue codes as unique cost centers on the cost report.

Response: We agree with the commenters that the broader the range and volume of services included in a given cost center, the more the resulting CCR calculated from the costs and charges for that cost center represents a weighted average of included services. To the extent that the revenue codes implemented in October 2004, specifically 0343 (Nuclear Medicine; Diagnostic Radiopharmaceuticals) for diagnostic preparations and 0344 (Nuclear Medicine; Therapeutic Radiopharmaceuticals) for therapeutic preparations, have no specific associated cost center in which to capture their unique costs and charges and to the extent hospitals report these costs and charges in cost center 4100 “Radiology—Diagnostic” or 4200 “Radiology—Therapeutic,” the CCRs for cost centers 4100 and 4200 that CMS uses to estimate costs from charges on claims for specific radiopharmaceuticals will reflect the average cost and markup associated with all diagnostic and therapeutic radiology procedures. However, our policy for establishing new cost centers requires a public review process that allows commenters the opportunity to provide input on any changes, and many commenters historically have not been interested in adding cost centers to the cost report because of the associated hospital administrative burden.

As we have noted above, we have recently undertaken regulatory comment and response on our effort to update the cost report. The proposed draft hospital cost report Form CMS-2552-10 went on **Federal Register** public display at the Office of the Federal Register on July 2, 2009, for a 60-day review and comment period, which ended on August 31, 2009. As we stated in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68525 through 68526), that notice and comment procedure is the process by which we are considering public comments requesting additional cost centers. We will consider all comments for new cost centers submitted through that process as we work to improve and modify the hospital cost report. We also note that we make the revenue code-to-cost center crosswalk that we use to match Medicare hospital cost report information with claims data continually available for inspection and comment on the CMS Web site: [\[www.cms.hhs.gov/HospitalOutpatientPPS\]\(http://www.cms.hhs.gov/HospitalOutpatientPPS\).](http://</p></div><div data-bbox=)

Comment: One commenter believed that the proposed drug cost center split discussed in the CY 2009 OPSS/ASC proposed rule would represent an unnecessary burden for hospitals.

Response: While we welcome comments regarding OPSS policy, we note that the drug cost center proposal was a CY 2009 proposal which was not finalized (73 FR 68654 through 68657). We have not proposed a policy to split the drug cost center for CY 2010.

Comment: One commenter requested that CMS issue clarifying instructions for reporting computed tomography (CT) and magnetic resonance imaging (MRI) equipment and supported the creation of new cost centers to capture the unique costs and charges of CT scanning, MRI, and other radiology procedures.

Response: We did not propose to implement separate standard radiology cost centers for CT Scanning, MRI, and other radiology procedures due to the significant number of comments we received in response to our general request in the CY 2009 OPSS/ASC proposed rule for comments and reactions to RTI’s recommendations. The commenters on the CY 2009 OPSS/ASC proposed rule were generally in favor of these cost centers in theory, but suggested that the allocation of capital cost across these cost centers was not consistent or consistently accurate across hospitals and that smaller hospitals might not have sufficiently sophisticated accounting systems to accurately allocate costs (73 FR 68526). In that discussion, we expressed our preference for establishing these cost centers as standard cost centers because standard cost centers constitute the minimum set of cost centers that a hospital is required to report, assuming that the hospital maintains separate departments for those services and reports the costs and charges for these departments in separate accounts within its own internal accounting systems. We believe this step would improve the accuracy of radiology payment by encouraging greater and more consistent reporting of the costs and charges specifically associated with advanced imaging services. However, we also noted that nonstandard cost centers already are available for CT Scanning and MRI and that hospitals that provide these services and maintain a separate account for each of these services in their internal accounting records to capture the costs and charges are currently required, in accordance with § 413.53(a)(1), to report these cost centers on the cost report, even if CMS

does not identify a nonstandard cost center code for the department(s).

As we stated in the CY 2009 OPPS/ASC final rule with comment (73 FR 68525 through 68526) and in response to an earlier comment in this section, we will consider public comments requesting additional cost centers in response to the PRA **Federal Register** notice for the proposed draft cost report form CMS–2552–10. The comment period for this proposal ended August 31, 2009.

Comment: A few commenters expressed concern about the timing for implementing the nonstandard cost center for cardiac rehabilitation, suggesting that a delay could limit beneficiary access to cardiac rehabilitation services because the proposed CY 2010 payment was too low. The commenters noted that the new CCRs would not be available for setting OPPS payment rates until CY 2013.

Response: While we understand the commenters' concern regarding the timing of implementing the cardiac rehabilitation nonstandard cost center, in our CY 2009 OPPS/ASC final rule with comment period discussion (73 FR 68524), we explained our preference for improving the accuracy of the APC relative weights through long-term changes to the cost report rather than implementing short-term statistical adjustments, in order to ensure that actual hospital data are used to set payment rates. As discussed above, we currently anticipate we will implement new nonstandard cost centers for Cardiac Rehabilitation, Hyperbaric Oxygen Therapy, and Lithotripsy with the revised Medicare hospital cost report form in CY 2010.

We have approximately 2.5 million CY 2008 claims from almost 2,000 hospitals for cardiac rehabilitation sessions available for setting the CY 2010 payment rates for these services. Given that the OPPS payment for the services has been highly stable for the past several years, we have no reason to believe that Medicare beneficiaries' access to cardiac rehabilitation will be limited in CY 2010 based on the final OPPS payment rates for the services. Further discussion of CY 2010 payment for traditional and intensive cardiac rehabilitation services is included in section XII.B. of this final rule with comment period.

Comment: One commenter believed that CMS continues to expand and complicate the antiquated Medicare cost report rather than to design a helpful tool. The commenter believes that the current "piecemeal" approach to revising the cost report is costly and

burdensome. Based on that impression, the commenter recommended that CMS partner with the hospital industry to consider more comprehensive changes to the cost report.

Response: In the FY 2009 IPPS proposed and final rules (73 FR 23546 and 73 FR 48461) and CY 2009 OPPS/ASC proposed rule and final rule with comment period (73 FR 41431 and 73 FR 68526), we stated that we began a comprehensive review of the Medicare hospital cost report, and splitting the current cost center for "Medical Supplies Charged to Patients" into one line for "Medical Supplies Charged to Patients" and another line for "Implantable Devices Charged to Patients" is part of that initiative to update and revise the cost report. We also explained that in the context of the effort to update the cost report and eliminate outdated requirements, we would make changes to the cost report form and cost report instructions that would be available to the public for comment. Thus, the public would have an opportunity to suggest the more comprehensive reforms that one commenter on the CY 2010 OPPS/ASC proposed rule advocates. Similarly, the public would be able to offer suggestions for ensuring that these reforms are made in a manner that is not disruptive to hospitals' billing and accounting systems, and within the guidelines of General Accepted Accounting Principles (GAAP), which are consistent with the Medicare principles of reimbursement and sound accounting practices. The proposed draft hospital cost report Form CMS–2552–10 went on **Federal Register** public display at the Office of the Federal Register on July 2, 2009, for a 60-day review and comment period, which ended on August 31, 2009. We will consider comments from the public as we work to improve and modify the hospital cost report. The cost center for "Implantable Devices Charged to Patients" is available for use for cost reporting periods beginning on or after May 1, 2009. The revised hospital cost report Form CMS–2552–10 will be effective for cost reporting periods beginning on or after February 1, 2010 (74 FR 43781 through 43782).

2. Data Development Process and Calculation of Median Costs

In this section of this final rule with comment period, we discuss the use of claims to calculate final OPPS payment rates for CY 2010. The hospital OPPS page on the CMS Web site on which this final rule with comment period is posted provides an accounting of claims used in the development of the final

payment rates at: <http://www.cms.hhs.gov/HospitalOutpatientPPS>. The accounting of claims used in the development of this final rule with comment period is included on the CMS Web site under supplemental materials for the CY 2010 OPPS/ASC final rule with comment period. That accounting provides additional detail regarding the number of claims derived at each stage of the process. In addition, below in this section we discuss the file of claims that comprise the data set that is available for purchase under a CMS data use agreement. Our CMS Web site, <http://www.cms.hhs.gov/HospitalOutpatientPPS>, includes information about purchasing the "OPPS Limited Data Set," which now includes the additional variables previously available only in the OPPS Identifiable Data Set, including ICD–9–CM diagnosis codes and revenue code payment amounts. This file is derived from the CY 2008 claims that were used to calculate the final payment rates for the CY 2010 OPPS.

As proposed, we used the methodology described in sections II.A.2.b. through e. of this final rule with comment period to establish the relative weights used in calculating the final OPPS payment rates for CY 2010 shown in Addenda A and B to this final rule with comment period.

a. Claims Preparation

For the CY 2010 OPPS/ASC proposed rule, we used the CY 2008 hospital outpatient claims processed before January 1, 2009 to calculate the median costs of APCs, which in turn are used to set the proposed relative weights for CY 2010. To begin the calculation of the relative weights for CY 2010, we pulled all claims for outpatient services furnished in CY 2008 from the national claims history file. This is not the population of claims paid under the OPPS, but all outpatient claims (including, for example, critical access hospital (CAH) claims and hospital claims for clinical laboratory services for persons who are neither inpatients nor outpatients of the hospital). In the discussion that follows, we have updated the information to reflect the claims available for this final rule with comment period, specifically CY 2008 claims processed through June 30, 2009.

We then excluded claims with condition codes 04, 20, 21, and 77. These are claims that providers submitted to Medicare knowing that no payment would be made. For example, providers submit claims with a condition code 21 to elicit an official denial notice from Medicare and

document that a service is not covered. We then excluded claims for services furnished in Maryland, Guam, the U.S. Virgin Islands, American Samoa, and the Northern Mariana Islands because hospitals in those geographic areas are not paid under the OPSS.

We divided the remaining claims into the three groups shown below. Groups 2 and 3 comprise the 107 million claims that contain hospital bill types paid under the OPSS.

1. Claims that were not bill types 12X, 13X (hospital bill types), 14X (laboratory specimen bill types), or 76X (CMHC bill types). Other bill types are not paid under the OPSS and, therefore, these claims were not used to set OPSS payment.

2. Claims that were bill types 12X, 13X or 14X. Claims with bill types 12X and 13X are hospital outpatient claims. Claims with bill type 14X are laboratory specimen claims, of which we use a subset for the limited number of services in these claims that are paid under the OPSS.

3. Claims that were bill type 76X (CMHC). (These claims are later combined with any claims in item 2 above with a condition code 41 to set the per diem partial hospitalization rates determined through a separate process.)

To convert charges on the claims to estimated cost, we needed to multiply those charges by the CCR associated with each revenue code as discussed in section II.A.1.c.(1) of this final rule with comment period. For the CCR calculation process, we used the same general approach that we used in developing the final APC rates for CY 2007, using the revised CCR calculation that excluded the costs of paramedical education programs and weighted the outpatient charges by the volume of outpatient services furnished by the hospital. We refer readers to the CY 2007 OPSS/ASC final rule with comment period for more information (71 FR 67983 through 67985). We first limited the population of cost reports to only those for hospitals that filed outpatient claims in CY 2008 before determining whether the CCRs for such hospitals were valid.

We then calculated the CCRs for each cost center and the overall ancillary CCR for each hospital for which we had claims data. We did this using hospital-specific data from the Hospital Cost Report Information System. We used the most recent available cost report data, in most cases, cost reports with cost reporting periods beginning in CY 2007. As proposed, for this final rule with comment period, we used the most recently submitted cost reports to

calculate the CCRs to be used to calculate median costs for the final CY 2010 OPSS payment rates. If the most recent available cost report was submitted but not settled, we looked at the last settled cost report to determine the ratio of submitted to settled cost using the overall ancillary CCR, and we then adjusted the most recent available submitted but not settled cost report using that ratio. We then calculated both an overall ancillary CCR and cost center-specific CCRs for each hospital. We used the overall ancillary CCR referenced in section II.A.1.c.(1) of this final rule with comment period for all purposes that require use of an overall ancillary CCR.

We then flagged CAH claims, which are not paid under the OPSS, and claims from hospitals with invalid CCRs. The latter included claims from hospitals without a CCR; those from hospitals paid an all-inclusive rate; those from hospitals with obviously erroneous CCRs (greater than 90 or less than .0001); and those from hospitals with overall ancillary CCRs that were identified as outliers (3 standard deviations from the geometric mean after removing error CCRs). In addition, we trimmed the CCRs at the cost center (that is, departmental) level by removing the CCRs for each cost center as outliers if they exceeded ± 3 standard deviations from the geometric mean. We used a four-tiered hierarchy of cost center CCRs, which is the revenue code-to-cost center crosswalk, to match a cost center to every possible revenue code appearing in the outpatient claims that is relevant to OPSS services, with the top tier being the most common cost center and the last tier being the default CCR. If a hospital's cost center CCR was deleted by trimming, we set the CCR for that cost center to "missing" so that another cost center CCR in the revenue center hierarchy could apply. If no other cost center CCR could apply to the revenue code on the claim, we used the hospital's overall ancillary CCR for the revenue code in question. For example, if a visit was reported under the clinic revenue code but the hospital did not have a clinic cost center, we mapped the hospital-specific overall ancillary CCR to the clinic revenue code. The revenue code-to-cost center crosswalk is available for inspection and comment on the CMS Web site: <http://www.cms.hhs.gov/HospitalOutpatientPPS>. Revenue codes not used to set medians or to model impacts are identified with an "N" in the revenue code-to-cost center crosswalk.

As we proposed, we updated the revenue code-to-cost center crosswalk to

more accurately reflect the current use of revenue codes. We indicated in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68531) that we intended to assess the NUBC revenue codes to determine whether any changes to the list of packaged revenue codes should be proposed for the CY 2010 OPSS. We expanded this evaluation to review all revenue codes in the revenue code-to-cost center crosswalk that we have used for OPSS ratesetting purposes in recent years against the CY 2008 NUBC definitions of revenue codes in place for CY 2008. As a result of that review, we proposed to revise the revenue code-to-cost center crosswalk as described in Table 2 of the CY 2010 OPSS/ASC proposed rule (74 FR 35256 through 35261).

Comment: Two commenters specifically addressed the proposed OPSS treatment of a number of revenue codes for CY 2010 and submitted identical, detailed recommendations. In general, the commenters agreed with the proposed treatment of revenue codes 0253 (Pharmacy; Take Home Drugs); 0290 (Durable Medical Equipment (other than renal); General Classification); 0291 (Durable Medical Equipment; Rental); 0292 (Durable Medical Equipment; Purchase of New DME); 0293 (Durable Medical Equipment; Purchase of Used DME); 0294 (Durable Medical Equipment; Supplies/Drugs for DME); 052x (Free-Standing Clinic; All Classifications); 066X (Respite Care; All Classifications); 0749, 0759, 0779, 0799, and 0910 (All Reserved); and 0948 (Other Therapeutic Services—Pulmonary Rehabilitation). The commenters disagreed with the proposed treatment of the revenue codes as displayed in Table 3 below, which provides the commenters' perspective on each revenue code.

Response: Specifically, our revenue proposal addressed: (1) Acknowledging that costs estimated from charges are associated with specific revenue codes when calculating OPSS payment rates; (2) identifying the appropriate cost center CCR that should be used to estimate costs for certain revenue codes; and (3) packaging of revenue center costs into the costs of separately paid procedures when revenue charges are reported without a HCPCS code. The commenters addressed some revenue codes that were explicitly identified and discussed in the CY 2010 OPSS/ASC proposed rule (74 FR 35256 through 35266), as well as some additional revenue codes. Table 3 below displays our response to each area where the commenters disagreed with our proposed treatment of the revenue code.

We note that we continually make our review and comment, and we welcome revenue code-to-cost center crosswalk available on the CMS Web site for

further comments on the crosswalk from the public at any time.

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TABLE 3.--COMMENTS AND RESPONSES ON CMS TREATMENT OF REVENUE CODES TO WHICH COMMENTERS OBJECTED

Revenue Code	Public Comment	CMS Response
0273 (Medical/Surgical Supplies and Devices; Take Home Supplies)	The commenters disagreed that estimated costs associated with 0273 should be included in OPPS ratesetting and, more specifically, disagreed that CMS should package the costs for revenue code 0273 when it appears without a HCPCS code because to do so would not be consistent with the exclusion and absence of packaging for revenue code 0253 (Pharmacy - Take Home Drugs).	We agree and will exclude the costs estimated from charges reported under revenue code 0273. We have removed revenue code 0273 from the final CY 2010 packaged revenue code list.
0274 (Medical/Surgical Supplies and Devices; Prosthetic/Orthotic Devices)	The commenters indicated that nonimplantable prosthetics/orthotics are not covered under the OPPS and that if CMS excludes charges reported under revenue codes 0290 (Durable Medical Equipment) and 0292 (Purchase of new DME), CMS should also exclude charges reported under revenue code 0274.	We disagree. The definition of revenue code 0274 for CY 2008 (the year on which the CY 2010 claims data are based) is not limited to nonimplantable prosthetics/orthotics. Therefore, it is possible that a hospital would report charges for implantable prosthetic devices under this revenue code and it would be appropriate to include those charges in the calculation of payments under the OPPS.

Revenue Code	Public Comment	CMS Response
0299 (Durable Medical Equipment; Other Equipment)	The commenters claimed that costs estimated from charges reported under revenue code 0299 should not be considered in OPSS ratesetting and agreed that estimated costs associated with revenue code 0299 should be excluded from the packaged revenue code list because the commenter asserted that all costs estimated from charges reported with DME revenue codes should be excluded from OPSS ratesetting in general.	We disagree. We will continue to recognize costs estimated from charges reported under revenue code 0299 because we believe that hospitals may infrequently report charges for DME used to care for hospital outpatients under this revenue code. We note that for CY 2007, hospitals reported approximately \$1,700 in total charges under this revenue code.
030X (Laboratory; All Classifications)	The commenters objected to the inclusion of costs estimated from charges reported under revenue code 030X in the OPSS ratesetting methodology because they believe that these charges are attributable to services paid under the clinical laboratory fee schedule.	We disagree and will continue to recognize costs estimated from charges reported with revenue code 030X because clinical pathology services that are paid under the OPSS are sometimes reported under this revenue code. To disallow charges under this revenue code would exclude valid cost estimates for services that are paid under the OPSS.
0500 (Outpatient Services; General Classification)	The commenters disagreed with the current exclusion of revenue code 0500 because the commenters believe that exclusion of charges under this revenue code may exclude charges for Part B inpatients that are covered under the OPSS when those patients have exhausted their Part A benefit.	We disagree and will exclude this revenue code from the OPSS because Medicare's longstanding policy is not to recognize this revenue code for Medicare payment (Medicare Claims Processing Manual 100-04, Chapter 25).

Revenue Code	Public Comment	CMS Response
0509 (Outpatient Services; Other Outpatient)	The commenters disagreed with the exclusion of revenue code 0509 because the commenters believe that exclusion of charges under this revenue code may exclude charges for Part B inpatients that are covered under the OPSS when those patients have exhausted their Part A benefit.	We disagree that charges for revenue code 0509 would be for services for inpatients receiving covered part B services. This revenue code is for services provided to outpatients, and inpatients are not outpatients. Without additional rationale, we are unwilling to include estimated costs associated with this revenue code.
056X (Home Health (hh) Medical Social Services; All Classifications)	The commenters agreed that these codes should not be allowed for the OPSS and recommended for that reason they should be excluded from the packaged revenue code list.	We agree and will remove them from the packaged revenue code list. We note that because we excluded charges reported under these revenue codes from OPSS, there were no costs to package, regardless of whether they were on the packaged revenue code file.

Revenue Code	Public Comment	CMS Response
0623 (Medical/surgical Supplies – Extension of 027X; Surgical Dressings)	The commenters disagreed with the proposal to include costs estimated from charges for revenue code 0623 in OPSS ratesetting and the proposal to add this revenue code to the list of packaged revenue codes because the commenters believe that Medicare instructions require items billed under this revenue code to be paid based on the DMEPOS fee schedule. The commenter stated that if CMS excludes DME charges under revenue codes 0290 and 0292, then for consistency, CMS should also exclude charges for surgical supplies under revenue code 0623.	We do not believe that this revenue code contains only implantable DME items, unlike revenue codes 0290 and 0292. In particular, the instructions in the NUBC Data Specifications Manual and the Medicare Claims Processing Manual (100-04; Chapter 25) identify revenue codes in the 062X series as being an extension of the revenue code 027X series (medical supplies). Further, we are unable to identify instructions that require CMS to pay for items charged under revenue code 0623 through the DMEPOS fee schedule. Therefore, we will recognize costs estimated from charges associated with revenue code 0623 for ratesetting and include revenue code 0623 on the packaged revenue code list for the CY 2010 OPSS.
0709 (Cast Room; Reserved) 0719 (Recovery Room; Reserved)	The commenters agreed that these codes should not be allowed for the OPSS and recommended for that reason they should be excluded from the packaged revenue code file.	We agree and will remove them from the packaged revenue code list. We note that because we excluded charges reported under these revenue codes from OPSS, there were no costs to package, regardless of whether they were on the packaged revenue code file.

Revenue Code	Public Comment	CMS Response
<p>0722 (Labor Room/Delivery – Delivery Room)</p> <p>0723 (Labor Room/Delivery; Circumcision)</p> <p>0724 (Labor Room/Delivery; Birthing Center)</p> <p>0729 (Labor Room/Delivery; Other Labor Room/Delivery)</p>	<p>The commenters requested that revenue codes 0722, 0723, 0724, and 0729 be included on the list of packaged revenue codes and that CMS package any associated estimated costs appearing without a HCPCS code into the major separately paid code with which they appear.</p>	<p>We disagree. We believe that these revenue codes are largely associated with major separately paid procedures and that hospitals should report these revenue codes together with the HCPCS codes for the procedures. We do not believe that hospitals should report uncoded charges in these procedure-specific revenue codes, and we do not recognize these revenue codes as containing unspecified charges in the I/OCE. We note that for CY 2007, hospitals reported only \$3,400 in charges without HCPCS codes across all of these revenue codes.</p>
<p>0931 (Medical Rehabilitation Day Program; Half Day)</p> <p>0932 (Medical Rehabilitation Day Program; Full Day)</p>	<p>The commenters agreed that these codes should not be allowed for OPPS and recommend for that reason they should be excluded from the packaged revenue code list and should not have a cost center assignment.</p>	<p>We agree and will remove the cost center assignment. We note that because we excluded charges reported under these revenue codes from OPPS, there were no allowed charges to which to apply the cost center 6000.</p>

Revenue Code	Public Comment	CMS Response
0942 (Other Therapeutic Services; Education/Training)	The commenters stated that education and training occur in many areas of the hospital and that CMS should use the hospital overall ancillary CCR, rather than the clinic CCR, to estimate costs from charges reported under revenue code 0942.	We do not disagree with the commenter on the potential varied hospital locations of education and training services, but we do not believe this is a reason to use the overall ancillary CCR. We believe that the charges and costs associated with delivery of education and training would more likely resemble the costs associated with clinic services, such as visits, than the costs and charges of surgical and radiology services that dominate the overall ancillary CCR. In addition, our claims data document that hospitals rarely use this revenue code. For CY 2007, total charges reported under revenue code 0942 were approximately \$700.

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Comment: One commenter suggested that CMS include dates in the revenue code-to-cost center crosswalk document to allow hospitals and CMS to easily track the effective dates for each change.

Response: We appreciate the desire to track changes to the revenue code-to-cost center crosswalk. However, rather than document changes to individual revenue codes in the crosswalk, we will provide the public with the current and past copies of the same revenue code-to-cost center crosswalk that we directly

incorporate into our modeling of the OPPS payment rates each year.

Table 4 below shows the update to the revenue codes for which estimated costs on each claim for this final rule with comment period are based and incorporates the costs for those revenue codes into APC median cost estimates. Column A of Table 4 provides the 2008 revenue code and description. Column B indicates whether the charges reported with the revenue code will be converted to cost and incorporated into median cost estimates for CY 2010.

Column C indicates whether the charges reported with the revenue code were converted to cost and incorporated into median cost estimates for the CY 2009 OPPS. In both columns, a "Y" indicates that the charges will be converted to cost in CY 2010 (or were converted for CY 2009), and an "N" indicates that charges reported under the revenue code will not be converted to cost and incorporated into median cost estimates. Finally, Column D provides our rationale for the CY 2010 final change.

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**TABLE 4.—CHANGES TO CY 2010 OPPTS REVENUE CODES
INCLUDED IN THE REVENUE CODE-TO-COST CENTER CROSSWALK**

A	B	C	D
2008 Revenue Code and Description	CY 2010 Inclusion in Median Cost Estimates	CY 2009 Inclusion in Median Cost Estimates	Rationale for CY 2010 Change
0290 -- Durable Medical Equipment (Other than Renal); General classification 0292 -- Durable Medical Equipment (Other than Renal); Purchase of New DME	N	Y	We are not considering charges reported under revenue codes 0290 and 0292 for OPPTS ratesetting because we believe that these charges are not for items for which payment may be made under the OPPTS. Only implantable DME is paid under the OPPTS and we believe that implantable DME is reported as a supply or implant under revenue code 0270, 0278, or 0279.
0392 -- Administration, Processing and Storage for Blood and Blood Components; Processing and Storage	Y	Missing	We are adding revenue 0392, which was previously omitted from the crosswalk, and considering these charges for OPPTS ratesetting because we believe that hospitals may correctly choose to report charges for blood processing and storage under this revenue code.
0500 -- Outpatient Services; General Classification 0509 -- Outpatient Services; Other Outpatient	N	Missing	We are adding previously omitted revenue codes 0500 and 0509 to the crosswalk because they are valid revenue codes, but we are not considering charges reported under them for OPPTS ratesetting because Medicare historically has not recognized revenue code 0500 for payment. Furthermore, we believe that hospitals primarily use revenue code 0509 to report charges that are paid under methodologies other than the OPPTS.

A	B	C	D
2008 Revenue Code and Description	CY 2010 Inclusion in Median Cost Estimates	CY 2009 Inclusion in Median Cost Estimates	Rationale for CY 2010 Change
<p>0520 -- Free-Standing Clinic; General Classification</p> <p>0523 -- Free-Standing Clinic: Family Practice Clinic</p> <p>0524 -- Free-Standing Clinic ; Visit by RHC/FQHC practitioner to a Member in a Covered Part A Stay at SNF</p> <p>0525 -- Free-Standing Clinic; Visit by RHC/FQHC practitioner to a Member in a SNF (not in a Covered Part A Stay) or NF, or ICF MR or Other Residential Facility</p> <p>0527 -- Free-Standing Clinic; Visiting Nurse Service(s) to a Member's Home when in a Home Health Shortage Area</p> <p>0528 -- Free-Standing Clinic; RHC/FQHC Practitioner to Other non RHC/FQHC site</p> <p>0529 -- Other Free-Standing Clinic</p>	N	<p>Y for 0520, 0523, 0526 and 0529;</p> <p>Missing for 0524, 0525, 0527, 0528</p>	<p>We are not considering charges reported under revenue codes 0520, 0523, 0524, 0525, 0527, 0528 and 0529 for purposes of OPSS ratesetting because we do not believe that services that would be reported under these revenue codes would be paid under the OPSS. To be paid under the OPSS, therapeutic services must be furnished directly by a hospital or under arrangements with the hospital, and all must be furnished in the hospital or a provider-based department of the hospital. A freestanding clinic or RHC is not a hospital or a provider-based department of a hospital. An FQHC may, under rare circumstances, be a provider-based department of a hospital if it meets the requirements in §413.65(n), but covered FQHC services furnished by an FQHC that is a provider-based department of a hospital are not paid under the OPSS.</p> <p>We also are adding revenue codes 0524, 0525, 0527, and 0528 which are now omitted from the crosswalk, to the crosswalk because they are valid revenue codes. We believe the crosswalk should reflect the existence of these revenue codes in the data, but we are not considering their charges for OPSS ratesetting because, as noted above, we do not believe that services that would be reported under these revenue codes would be paid under the OPSS.</p>

A	B	C	D
2008 Revenue Code and Description	CY 2010 Inclusion in Median Cost Estimates	CY 2009 Inclusion in Median Cost Estimates	Rationale for CY 2010 Change
<p>0560 -- Home Health (HH)-Medical Social Services; General Classification</p> <p>0561 -- Home Health (HH) Medical Social Services; Visit Charge</p> <p>0562 -- Home Health (HH) Medical Social Services; Hourly Charge</p> <p>0569 -- Home Health (HH) Medical Social Services; Other Medical Social Service</p>	N	Y	<p>We are not considering charges reported under revenue codes 0560, 0561, 0562 and 0569 because to be paid under the OPSS, therapeutic services must be furnished directly by a hospital or under arrangements with the hospital, and all must be furnished in the hospital or a provider-based department of the hospital. Home health care is furnished in a home and, therefore, does not meet the criteria for payment under the OPSS.</p>
<p>0623 -- Medical Surgical Supplies -- Extension of 027X; Surgical Dressings</p>	Y	N	<p>We are considering charges reported under revenue code 0623 because we believe that these charges may be associated with surgical dressings applied during procedures for which payment is made under the OPSS and should be allowed for purposes of ratesetting.</p>
<p>0660 -- Respite Care; General Classification</p> <p>0661 -- Respite Care; Hourly Charge -Nursing</p> <p>0662 -- Respite Care; Hourly Charge/Aide/Homemaker/Companion</p> <p>0663 -- Respite Care; Daily Respite Charge</p> <p>0669 -- Respite Care; Other Respite Care</p>	N	Missing	<p>We are adding previously omitted revenue codes 0660, 0661, 0662, 0663, and 0669 to the crosswalk, but not considering charges reported under these revenue codes for OPSS ratesetting. We do not believe that respite care services would meet the requirements for payment under the OPSS. We are adding these revenue codes to the crosswalk to reflect the existence of these codes in the data. However, we will not consider charges reported under these codes for ratesetting because we do not believe that services reported under these revenue codes would be paid under the OPSS and, therefore, we believe the charges would be inappropriate for use in OPSS ratesetting.</p>

A	B	C	D
2008 Revenue Code and Description	CY 2010 Inclusion in Median Cost Estimates	CY 2009 Inclusion in Median Cost Estimates	Rationale for CY 2010 Change
0709 – Cast Room; RESERVED 0719 -- Recovery Room; RESERVED 0749 – EEG (Electroencephalogram); RESERVED 0759 – Gastro-Intestinal (GI) Services; RESERVED 0779 – Preventive Care Services; RESERVED 0799 – Extra-Corporeal Shock Wave Therapy (Formerly Lithotripsy); RESERVED 0910 – Behavioral Health Treatments/Services – Extension of 090X; RESERVED (Use 090 for General Classification)	N	Y	We are not considering charges under revenue codes 0709, 0719, 0749, 0759, 0779, 0799, and 0910 for OPSS ratesetting because no charges should be reported under a revenue code that is reserved.
0931 -- Medical Rehabilitation Day Program; Half Day 0932 -- Medical Rehabilitation Day Program; Full Day	N	Missing	We are adding previously omitted revenue codes 0931 and 0932 to the crosswalk to reflect their existence in the NUBC dataset. However, we will not consider charges reported using these revenue codes for ratesetting because the NUBC rules prohibit hospitals from reporting charges under these revenue codes.
0948 -- Other Therapeutic Services (also see 095x, an extension of 094x); Pulmonary Rehabilitation	Y	Missing	We are considering charges reported under revenue code 0948 for purposes of OPSS ratesetting. Through our assessment of the NUBC revenue code definitions, we believe that hospitals report charges for services paid under the OPSS under revenue code 0948.

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Also, as a result of our comprehensive review of the revenue codes included in the revenue code-to-cost center crosswalk, as we proposed, we are adding revenue codes to the hierarchy of primary, secondary, and tertiary hospital cost report cost centers that

result in the departmental CCRs that we use to estimate cost from charges for some revenue codes or to revise the applicable cost centers associated with a given revenue code. Table 5 below lists the revenue codes for which we made changes to the revenue code-to-cost center crosswalk for CY 2010

ratesetting and our rationale for each change. With the exception of revenue code 0942 (Other Therapeutic Services; Education/Training), the revenue codes for which we made changes to the designated departmental CCRs are those identified in our comprehensive review that are also listed above in Table 4.

TABLE 5—CHANGES TO CY 2010 OPPTS HIERARCHY OF COST CENTERS IN THE REVENUE CODE-TO-COST CENTER CROSSWALK

2008 Revenue code and description	Rationale for CY 2010 change
0392—Administration, Processing and Storage for Blood and Blood Components; Processing and Storage.	We crosswalked charges under revenue code 0392 to cost center 4700 (Blood Storing, Processing, & Transfusing) because we believe that cost center 4700 is the most likely departmental cost center to which hospitals would assign the costs of blood processing and storage. We made no secondary or tertiary cost centers because we believe that no other departmental cost centers are appropriate.
0623—Medical Surgical Supplies—Extension of 027X; Surgical Dressings.	We crosswalked the charges reported under revenue code 0623 to cost center 5500 (Medical Supplies Charged to Patients) as the primary cost center because we believe that the costs associated with the charges for surgical dressings are most likely to be assigned by hospitals to cost center 5500. We made no secondary or tertiary cost centers because we believe that no other departmental cost centers are appropriate.
0942—Other Therapeutic Services (also see 095x, an extension of 094x); Educ/Training.	We crosswalked the charges under revenue code 0942 to cost center 6000 (Clinic) as the primary cost center. Previously, the charges under revenue code 0942 were crosswalked to the overall ancillary CCR. As discussed above, we believe that cost center 6000 is a more appropriate primary cost center. We made no secondary or tertiary cost centers because we believe that no other departmental cost centers are appropriate.
0948—Other Therapeutic Services (also see 095x, an extension of 094x); Pulmonary Rehabilitation.	We crosswalked the charges under revenue code 0948 to cost center 4900 (Respiratory Therapy) as primary and to cost center 6000 (Clinic) as secondary because we believe that hospitals are most likely to assign the costs of these services to these cost centers. We are not establishing a tertiary cost center.

Having revised the revenue code-to-cost center crosswalk, we then converted the charges to costs on each claim by applying the CCR that we believed was best suited to the revenue code indicated on the line with the charge. One exception to this general methodology for converting charges to costs on each claim is the calculation of median blood costs, as discussed in section II.A.2.d.(2) of the proposed rule and this final rule with comment period.

Thus, we applied CCRs as described above to claims with bill type 12X, 13X, or 14X, excluding all claims from CAHs and hospitals in Maryland, Guam, the U.S. Virgin Islands, American Samoa, and the Northern Mariana Islands and claims from all hospitals for which CCRs were flagged as invalid.

We identified claims with condition code 41 as partial hospitalization services of hospitals and moved them to another file. These claims were combined with the 76X claims identified previously to calculate the partial hospitalization per diem rates. We note that the separate file containing partial hospitalization claims is included in the files that are available for purchase as discussed above.

We then excluded claims without an HCPCS code. We moved to another file claims that contained nothing but influenza and pneumococcal pneumonia (PPV) vaccines. Influenza and PPV vaccines are paid at reasonable cost and, therefore, these claims are not used to set OPPTS rates.

We next copied line-item costs for drugs, blood, and brachytherapy sources (the lines stay on the claim, but are copied onto another file) to a separate

file. No claims were deleted when we copied these lines onto another file. These line-items are used to calculate a per unit mean and median cost and a per day mean and median cost for drugs, therapeutic radiopharmaceutical agents, and brachytherapy sources, as well as other information used to set payment rates, such as a unit-to-day ratio for drugs.

To implement our policy to redistribute some portion of total cost for packaged drugs and biologicals to the separately payable drugs and biologicals as acquisition and pharmacy overhead and handling costs discussed in section V.B.3. of this final rule with comment period, we used the line-item cost data for drugs and biologicals for which we had an HCPCS code with ASP pricing information to calculate the ASP+X values first for all drugs and biologicals, and then for separately payable drugs and biologicals and for packaged drugs and biologicals, respectively, by taking the ratio of total claim cost for each group relative to total ASP dollars (per unit of each drug or biological HCPCS code's July 2009 ASP amount multiplied by total units for each drug or biological in the CY 2008 claims data). These values are ASP+11 percent, ASP-3 percent, and ASP+259 percent, respectively. As we discuss in greater detail in section V.B.3. of this final rule with comment period, we are finalizing a policy to redistribute \$150 million of the total cost in our claims data for packaged drugs and biologicals that have an associated ASP from packaged drugs with an ASP to separately payable drugs and biologicals. The \$150 million is, roughly, one-third of the difference of

\$445 million between the total cost of packaged drugs and biologicals with an associated ASP in our CY 2008 claims data (\$616 million) and ASP for the same drugs and biologicals (\$171 million). In response to comments that CMS excluded valid overhead and handling costs associated with drugs lacking ASP information, largely costs estimated from uncoded charges reported under pharmacy revenue codes, we also are finalizing a policy to redistribute an additional \$50 million of the total cost in our claims data for drugs and biologicals lacking an ASP, largely for estimated costs associated with uncoded charges billed under pharmacy revenue code series 025X (Pharmacy (also see 063X, an extension of 025X)), 026X (IV Therapy), and 063X (Pharmacy—Extension of 025X). As we state in section V.B.3. of this final rule with comment period, because we do not know ASP for this subset of drug costs, we do not know the amount of associated pharmacy overhead. We observe about \$656 million for drugs lacking an ASP in our CY 2008 claims data. This total excludes the cost of diagnostic and therapeutic radiopharmaceuticals because they are not reported under pharmacy revenue codes or under the pharmacy cost center on the cost report.

Removing a total of \$150 million in pharmacy overhead cost from packaged drugs and biologicals reduces the \$616 million to \$466 million, a 24 percent reduction. Removing \$50 million from the cost of drugs lacking an ASP reduces the \$656 million to \$606 million, an 8 percent reduction. To implement our final CY 2010 policy to redistribute \$150 million in claim cost from

packaged drugs and biologicals with an ASP to separately payable drugs and biologicals and \$50 million in claim cost from packaged drugs and biologicals lacking an ASP, including uncoded pharmacy revenue code charges, we multiplied the cost of each packaged drug or biological with an HCPCS code and ASP pricing information in our CY 2008 claims data by 0.76, and we multiplied all other packaged drug costs in our CY 2008 claims data, excluding those for diagnostic radiopharmaceuticals, by 0.92. We also added the redistributed \$200 million to the total cost of separately payable drugs and biologicals in our CY 2008 claims data, which increased the relationship between the total cost for separately payable drugs and biologicals and ASP dollars for the same drugs and biologicals to ASP+4 percent.

For CY 2010, we added an additional trim in our claims preparation to remove line-items that were not paid during claim processing, presumably for a line-item rejection or denial. The number of edits for valid OPSS payment in the Integrated Outpatient Code Editor (I/OCE) and elsewhere has grown significantly in the past few years, especially with the implementation of the full spectrum of National Correct Coding Initiative (NCCI) edits. To ensure that we are using valid claims that represent the cost of payable services to set payment rates, we removed line-items with an OPSS status indicator for the claim year (CY 2008) and a status indicator of "S," "T," "V," or "X" when separately paid under the final CY 2010 payment system. This logic preserves charges for services that would not have been paid in the claim year but for which some estimate of cost is needed for the prospective year, such as services newly proposed to come off the inpatient list for CY 2010 which were assigned status indicator "C" in the claim year.

Using February 2009 APC Panel data, we estimate that the impact of removing line-items with valid status indicators that received no CY 2008 payment was limited to approximately 1.4 percent of all line-items for separately paid services. This additional trim reduced the number of single bills available for ratesetting by 1.5 percent. For approximately 92 percent of procedural APCs, we observed a change in the APC median cost of less than 1 percent. A handful of APCs experienced greater changes in median cost. For example, APC 0618 (Trauma Response with Critical Care) experienced declines in both the number of single bills used to set the median cost and the estimated

median cost itself. This occurred because the I/OCE has an edit to ensure that HCPCS code G0390 (Trauma response team activation associated with hospital critical care service), which is assigned to APC 0618, receives payment only when one unit of G0390 appears with both a revenue code in the 68x series and CPT code 99291 (Critical care, evaluation and management of the critically ill or critically injured patient; first 30–74 minutes) on the claim for the same date of service, as described in the CY 2007 OPSS/ASC final rule with comment period (71 FR 68134). If the I/OCE criteria are not met, HCPCS code G0390 is not separately paid, and we found that a number of CY 2008 claims including HCPCS code G0390 did not meet the criteria for payment. On the other hand, a few APCs had greater estimated median costs and greater numbers of single bills as a result of this additional trim, presumably because removing lines from the claim allowed us to identify more single bills. We believe that removing lines with valid status indicators that were edited and not paid during claims processing increases the accuracy of the single bills used to determine the APC median costs for ratesetting.

Comment: One commenter claimed that the removal of charges and costs from denied lines was in contrast to longstanding policy for hospital inpatient services. A few commenters expressed concern about APC 0312 (Radioelement Applications), noting that there has been significant fluctuation in the payment rates for this APC in the past. They believe that implementing the proposed line-item trim, which removed a significant number of single claims, may have contributed to that instability. The commenters suggested that historical data would not indicate any reason for significant line-items to be trimmed. One commenter believed that the payment rates for low dose rate prostate brachytherapy were arbitrary and unfair. Based on the commenters' impression that the purpose of the line-item trim was to act as a quality check, the commenters requested that the line-item trim be suppressed for APC 0312.

Response: While payment systems such as the IPPS do not remove charge and cost data, this is largely due to the differences in the fundamental structures of the two payment systems. The IPPS is a system based on DRGs that relies on significant bundling of services under common clinical scenarios, while the OPSS is largely based on payment for a specific individual service. These differences in payment approach under each system

are reflected in the way that data are used to establish the payment weights, from the CCRs used to reduce charges to cost to the structure of how charge and cost information is classified. One byproduct of the differences between the IPPS and the OPSS is the level of editing in each system to ensure that a correct payment is made. Similarly, there are many NCCI edits to ensure that payment is made to hospitals for outpatient services only when there is correct coding because there are hundreds of APCs that may contribute to inappropriate unbundling of services when those services are reported for a hospital outpatient encounter. In the CY 2010 OPSS/ASC proposed rule (74 FR 35262), we indicated that removing lines with valid status indicators that were edited and not paid during claims processing increases the accuracy of the single bills used for ratesetting. Doing so allows the single bills used for ratesetting purposes to be representative of those services as they would be paid in the prospective year.

In studying the billing patterns for HCPCS codes that are assigned to APC 0312, we noted that the line-item trim removes a number of unpaid single bills for this APC, as the commenters had suggested. However, we also observed a general decline in the reporting of services assigned to this APC that was unrelated to the line-item trim, suggesting that a portion of the observed decline in the number of single bills available for ratesetting is due to an actual reduction in the frequency that the services assigned to APC 0312 are furnished. While we understand the commenters' concern regarding the reduction of single bills used in ratesetting for APC 0312, the data suggest that the reduction is due in part to a decline in the billing of individual services assigned to the APC. Further, we believe that removing these line-items which have likely been rejected or denied is appropriate in light of the goal of using accurate single procedure claims for ratesetting under the OPSS.

After consideration of the public comments received relating to our CY 2010 proposal for claims preparation, we are adopting it as final, with modification to the treatment of certain revenue codes as described in Table 4 in this section.

b. Splitting Claims and Creation of "Pseudo" Single Claims

(1) Splitting Claims

We then split the remaining claims into five groups: single majors, multiple majors, single minors, multiple minors, and other claims. (Specific definitions

of these groups follow below.) In the CY 2010 OPPS/ASC proposed rule (74 FR 35262), we proposed to continue our current policy of defining major procedures as any HCPCS code having a status indicator of “S,” “T,” “V,” or “X;” defining minor procedures as any code having a status indicator of “F,” “G,” “H,” “K,” “L,” “R,” “U,” or “N,” and classifying “other” procedures as any code having a status indicator other than one that we have classified as major or minor. For CY 2010, we proposed to continue assigning status indicator “R” to blood and blood products; status indicator “U” to brachytherapy sources; status indicator “Q1” to all “STVX-packaged codes;” status indicator “Q2” to all “T-packaged codes;” and status indicator “Q3” to all codes that may be paid through a composite APC based on composite-specific criteria or paid separately through single code APCs when the criteria are not met. As discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68709), we established status indicators “Q1,” “Q2,” and “Q3” to facilitate identification of the different categories of codes. We proposed to treat these codes in the same manner for data purposes for CY 2010 as we have treated them since CY 2008. Specifically, we proposed to continue to evaluate whether the criteria for separate payment of codes with status indicator “Q1” or “Q2” are met in determining whether they are treated as major or minor codes. As discussed earlier in this section, because we proposed to treat CPT code 76098 as conditionally packaged, this logic now includes the addition of CPT code 76098 as a “Q2” code. Codes with status indicator “Q1” or “Q2” are carried through the data either with status indicator “N” as packaged or, if they meet the criteria for separate payment, they are given the status indicator of the APC to which they are assigned and are considered as “pseudo” single major codes. Codes assigned status indicator “Q3” are paid under individual APCs unless they occur in the combinations that qualify for payment as composite APCs and, therefore, they carry the status indicator of the individual APC to which they are assigned through the data process and are treated as major codes during both the split and “pseudo” single creation process. The calculation of the median costs for composite APCs from multiple major claims is discussed in section II.A.2.e. of this final rule with comment period.

Specifically, we divided the remaining claims into the following five groups:

1. *Single Major Claims:* Claims with a single separately payable procedure (that is, status indicator “S,” “T,” “V,” or “X,” which includes codes with status indicator “Q3”); claims with one unit of a status indicator “Q1” code (“STVX-packaged”) where there was no code with status indicator “S,” “T,” “V,” or “X” on the same claim on the same date; or claims with one unit of a status indicator “Q2” code (“T-packaged”) where there was no code with a status indicator “T” on the same claim on the same date.

2. *Multiple Major Claims:* Claims with more than one separately payable procedure (that is, status indicator “S,” “T,” “V,” or “X,” which includes codes with status indicator “Q3”), or multiple units of one payable procedure. These claims include those codes with a status indicator “Q2” code (“T-packaged”) where there was no procedure with a status indicator “T” on the same claim on the same date of service but where there was another separately paid procedure on the same claim with the same date of service (that is, another code with status indicator “S,” “V,” or “X”). We also include in this set, claims that contained one unit of one code when the bilateral modifier was appended to the code and the code was conditionally or independently bilateral. In these cases, the claims represented more than one unit of the service described by the code, notwithstanding that only one unit was billed.

3. *Single Minor Claims:* Claims with a single HCPCS code that was assigned status indicator “F,” “G,” “H,” “K,” “L,” “R,” “U,” or “N” and not status indicator “Q1” (“STVX-packaged”) or status indicator “Q2” (“T-packaged”) code.

4. *Multiple Minor Claims:* Claims with multiple HCPCS codes that are assigned status indicator “F,” “G,” “H,” “K,” “L,” “R,” “U,” or “N;” claims that contain more than one code with status indicator “Q1” (“STVX-packaged”) or more than one unit of a code with status indicator “Q1” but no codes with status indicator “S,” “T,” “V,” or “X” on the same date of service; or claims that contain more than one code with status indicator “Q2” (T-packaged), or “Q2” and “Q1,” or more than one unit of a code with status indicator “Q2” but no code with status indicator “T” on the same date of service.

5. *Non-OPPS Claims:* Claims that contain no services payable under the OPPS (that is, all status indicators other than those listed for major or minor

status). These claims were excluded from the files used for the OPPS. Non-OPPS claims have codes paid under other fee schedules, for example, durable medical equipment or clinical laboratory tests, and do not contain a code for a separately payable or packaged OPPS service. Non-OPPS claims include claims for therapy services paid sometimes under the OPPS but billed, in these non-OPPS cases, with revenue codes indicating that the therapy services would be paid under the Medicare Physician Fee Schedule (MPFS).

The claims listed in numbers 1, 2, 3, and 4 above are included in the data file that can be purchased as described above. Claims that contain codes to which we have assigned status indicators “Q1” (“STVX-packaged”) and “Q2” (“T-packaged”) appear in the data for the single major file, the multiple major file, and the multiple minor file used in this final rule with comment period. Claims that contain codes to which we have assigned status indicator “Q3” (composite APC members) appear in both the data of the single and multiple major files used in this final rule with comment period, depending on the specific composite calculation.

Because we did not receive any public comments on our proposed process of organizing claims by type, we are finalizing our CY 2010 proposal without modification.

(2) Creation of “Pseudo” Single Claims

As we proposed, to develop “pseudo” single claims for this final rule with comment period, we examined both the multiple major claims and the multiple minor claims. We first examined the multiple major claims for dates of service to determine if we could break them into “pseudo” single procedure claims using the dates of service for all lines on the claim. If we could create claims with single major procedures by using dates of service, we created a single procedure claim record for each separately payable procedure on a different date of service (that is, a “pseudo” single).

We also used the bypass codes listed earlier in Table 1 and discussed in section II.A.1.b. of this final rule with comment period to remove separately payable procedures that we determined contained limited or no packaged costs or that were otherwise suitable for inclusion on the bypass list from a multiple procedure bill. As discussed above, we ignore the “overlap bypass codes,” that is, those HCPCS codes that are both on the bypass list and are members of the multiple imaging

composite APCs, in this initial assessment for “pseudo” single claims. The CY 2010 “overlap bypass codes” are listed in Table 1 in section II.A.1.b. of this final rule with comment period. When one of the two separately payable procedures on a multiple procedure claim was on the bypass list, we split the claim into two “pseudo” single procedure claim records. The single procedure claim record that contained the bypass code did not retain packaged services. The single procedure claim record that contained the other separately payable procedure (but no bypass code) retained the packaged revenue code charges and the packaged HCPCS code charges. We also removed lines that contained multiple units of codes on the bypass list and treated them as “pseudo” single claims by dividing the cost for the multiple units by the number of units on the line. Where one unit of a single, separately payable procedure code remained on the claim after removal of the multiple units of the bypass code, we created a “pseudo” single claim from that residual claim record, which retained the costs of packaged revenue codes and packaged HCPCS codes. This enabled us to use claims that would otherwise be multiple procedure claims and could not be used.

We then assessed the claims to determine if the criteria for the multiple imaging composite APCs, discussed in section II.A.2.e.(5) of this final rule with comment period, were met. Where the criteria for the imaging composite APCs were met, we created a “single session” claim for the applicable imaging composite service and determined whether we could use the claim in ratesetting. For HCPCS codes that are both conditionally packaged and are members of a multiple imaging composite APC, we first assessed whether the code would be packaged and if so, the code ceased to be available for further assessment as part of the composite APC. Because the packaged code would not be a separately payable procedure, we considered it to be unavailable for use in setting the composite APC median cost. Having identified “single session” claims for the imaging composite APCs, we reassessed the claim to determine if, after removal of all lines for bypass codes, including the “overlap bypass codes,” a single unit of a single separately payable code remained on the claim. If so, we attributed the packaged costs on the claim to the single unit of the single remaining separately payable code other than the bypass code to create a “pseudo” single

claim. We also identified line-items of overlap bypass codes as a “pseudo” single claim. This allowed us to use more claims data for ratesetting purposes.

We also examined the multiple minor claims to determine whether we could create “pseudo” single procedure claims. Specifically, where the claim contained multiple codes with status indicator “Q1” (“STVX-packaged”) on the same date of service or contained multiple units of a single code with status indicator “Q1,” we selected the status indicator “Q1” HCPCS code that had the highest CY 2008 relative weight, set the units to one on that HCPCS code to reflect our policy of paying only one unit of a code with a status indicator of “Q1.” We then packaged all costs for the following into a single cost for the “Q1” HCPCS code that had the highest CY 2008 relative weight to create a “pseudo” single claim for that code: additional units of the status indicator “Q1” HCPCS code with the highest CY 2008 relative weight; other codes with status indicator “Q1;” and all other packaged HCPCS codes and packaged revenue code costs. We changed the status indicator for selected codes from the data status indicator of “N” to the status indicator of the APC to which the selected procedure was assigned for further data processing and considered this claim as a major procedure claim. We used this claim in the calculation of the APC median cost for the status indicator “Q1” HCPCS code.

Similarly, where a multiple minor claim contained multiple codes with status indicator “Q2” (“T-packaged”) or multiple units of a single code with status indicator “Q2,” we selected the status indicator “Q2” HCPCS code that had the highest CY 2008 relative weight, set the units to one on that HCPCS code to reflect our policy of paying only one unit of a code with a status indicator of “Q2.” We then packaged all costs for the following into a single cost for the “Q2” HCPCS code that had the highest CY 2008 relative weight to create a “pseudo” single claim for that code: additional units of the status indicator “Q2” HCPCS code with the highest CY 2008 relative weight; other codes with status indicator “Q2;” and other packaged HCPCS codes and packaged revenue code costs. We changed the status indicator for the selected code from a data status indicator of “N” to the status indicator of the APC to which the selected code was assigned, and we considered this claim as a major procedure claim.

Lastly, where a multiple minor claim contained multiple codes with status indicator “Q2” (“T-packaged”) and

status indicator “Q1” (“STVX-packaged”), we selected the status indicator “Q2” HCPCS code (“T-packaged”) that had the highest relative weight for CY 2008 and set the units to one on that HCPCS code to reflect our policy of paying only one unit of a code with a status indicator of “Q2.” We then packaged all costs for the following into a single cost for the selected (“T-packaged”) HCPCS code to create a “pseudo” single claim for that code: additional units of the status indicator “Q2” HCPCS code with the highest CY 2008 relative weight; other codes with status indicator “Q2;” codes with status indicator “Q1” (“STVX-packaged”); and other packaged HCPCS codes and packaged revenue code costs. We favor status indicator “Q2” over “Q1” HCPCS codes because “Q2” HCPCS codes have higher CY 2008 relative weights. If a status indicator “Q1” HCPCS code had a higher CY 2008 relative weight, it would become the primary code for the simulated single bill process. We changed the status indicator for the selected status indicator “Q2” (“T-packaged”) code from a data status indicator of “N” to the status indicator of the APC to which the selected code was assigned and we considered this claim as a major procedure claim.

We excluded those claims that we were not able to convert to single claims even after applying all of the techniques for creation of “pseudo” singles to multiple major and to multiple minor claims. As has been our practice in recent years, we also excluded claims that contained codes that were viewed as independently or conditionally bilateral and that contained the bilateral modifier (Modifier 50 (Bilateral procedure)) because the line-item cost for the code represented the cost of two units of the procedure, notwithstanding that the code appeared with a unit of one.

Comment: One commenter noted that the bilateral procedure logic did not appear to appropriately exclude claims with bilateral codes from the single major claims, having observed bilateral procedure codes in that claims subset. Also, the commenter suggested that the conditional packaging of the status indicator “Q2” (“T-packaged”) codes did not appear to be treated consistently with the policy we proposed, which was that a “Q2” procedure with the highest scaled weight would be paid separately when there is no status indicator “T” procedure on the claim and that the costs of any other “Q2” codes on the claim would be packaged.

Response: In seeking to address the commenter’s observations, we discovered that the bilateral logic was

not processed correctly as we proposed. Similarly, inaccurate program logic in the weight comparison for status indicator “Q2” (“T-packaged”) codes caused the packaging to be assigned based on order of precedence rather than by weight.

For this final rule with comment period, we accurately applied the bilateral and status indicator “Q2” (“T-packaged”) weight comparison packaging logic, consistent with the proposed and final policy. The national unadjusted payments for CY 2010 accurately reflect the policy that we proposed to continue for CY 2010 OPPS and that we are finalizing in this final rule with comment period.

After consideration of the public comment received, we are finalizing our CY 2010 proposal, without modification, for the process by which we develop “pseudo” single procedure claims.

c. Completion of Claim Records and Median Cost Calculations

We then packaged the costs of packaged HCPCS codes (codes with status indicator “N” listed in Addendum B to this final rule with comment period and the costs of those lines for codes with status indicator “Q1” or “Q2” when they are not separately paid), and the costs of packaged revenue codes into the cost of the single major procedure remaining on the claim. For CY 2010, this packaging also included the redistributed packaged pharmacy overhead cost relative to the units of separately payable drugs on each single procedure claim.

As noted in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66606), for the CY 2008 OPPS, we adopted an APC Panel recommendation that requires CMS to review the final list of packaged revenue codes for consistency with OPPS policy and ensure that future versions of the I/OCE edit accordingly. We compared the packaged revenue codes in the I/OCE to the final list of packaged revenue codes for the CY 2009 OPPS (73 FR 68531 through 68532) that we used for packaging costs in median calculation. As a result of that analysis, we proposed to use the packaged revenue codes for CY 2010 that were displayed in Table 4 of the CY 2010 OPPS/ASC proposed rule (74 FR 35265 through 35266).

As noted in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68531), we replaced the NUBC standard abbreviations for the revenue codes listed in Table 2 of the CY 2009 OPPS/ASC proposed rule with the most current NUBC descriptions of the

revenue code categories and subcategories to better articulate the meanings of the revenue codes without actually changing the proposed list of revenue codes. In the course of making the changes in labeling for the revenue codes in Table 2 of the CY 2009 OPPS/ASC final rule with comment period, we noticed some changes to revenue categories and subcategories that we believed warranted further review for future OPPS updates. Although we finalized the list of packaged revenue codes in Table 2 for CY 2009, we indicated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68531) that we intended to assess the NUBC revenue codes to determine whether any changes to the list of packaged revenue codes should be proposed for the CY 2010 OPPS. We specifically requested public input and discussion on this issue during the comment period of the CY 2009 OPPS/ASC final rule with comment period. We did not receive any public comments on this issue. As we discuss in section II.A.2.a. of this final rule with comment period, we have completed that analysis for all revenue codes in the revenue code-to-cost center crosswalk. As discussed in the CY 2010 OPPS/ASC proposed rule (74 FR 35264 through 35265), as a result, we proposed to add several revenue codes to the list of packaged revenue codes for the CY 2010 OPPS. Specifically, we believe that the costs derived from charges reported under revenue codes 0261 (IV Therapy; Infusion Pump); 0392 (Administration, Processing and Storage for Blood and Blood Components; Processing and Storage); 0623 (Medical Supplies—Extension of 027X, Surgical Dressings); 0943 (Other Therapeutic Services (also see 095X, an extension of 094X), Cardiac Rehabilitation); and 0948 (Other Therapeutic Services (also see 095X, an extension of 094X), Pulmonary Rehabilitation) are appropriately packaged into payment for other OPPS services when charges appear on lines with these revenue codes but no HCPCS code appears on the line. Revenue codes that we proposed to add to the CY 2010 packaged revenue code list were identified by asterisks (*) in Table 4 of the CY 2010 OPPS/ASC proposed rule.

The public comments that we received that resulted in our changing the list of packaged revenue codes for CY 2010 are discussed in section II.A.2.a. of this final rule with comment period. Thus, we are finalizing the proposed packaged revenue codes for CY 2010, with modification. The final CY 2010 packaged revenue codes are listed in Table 6 below. Revenue codes

that we are adding to the CY 2010 packaged revenue code list are identified by asterisks (*) in Table 6.

TABLE 6—FINAL CY 2010 PACKAGED REVENUE CODES

Revenue code	Description
0250	Pharmacy; General Classification.
0251	Pharmacy; Generic Drugs.
0252	Pharmacy; Non-Generic Drugs.
0254	Pharmacy; Drugs Incident to Other Diagnostic Services.
0255	Pharmacy; Drugs Incident to Radiology.
0257	Pharmacy; Non-Prescription.
0258	Pharmacy; IV Solutions.
0259	Pharmacy; Other Pharmacy.
0260	IV Therapy; General Classification.
*0261	IV Therapy; Infusion Pump.
0262	IV Therapy; IV Therapy/Pharmacy Svcs.
0263	IV Therapy; IV Therapy/Drug/Supply Delivery.
0264	IV Therapy; IV Therapy/Supplies.
0269	IV Therapy; Other IV Therapy.
0270	Medical/Surgical Supplies and Devices; General Classification.
0271	Medical/Surgical Supplies and Devices; Non-sterile Supply.
0272	Medical/Surgical Supplies and Devices; Sterile Supply.
0275	Medical/Surgical Supplies and Devices; Pacemaker.
0276	Medical/Surgical Supplies and Devices; Intraocular Lens.
0278	Medical/Surgical Supplies and Devices; Other Implants.
0279	Medical/Surgical Supplies and Devices; Other Supplies/Devices.
0280	Oncology; General Classification.
0289	Oncology; Other Oncology.
0343	Nuclear Medicine; Diagnostic Radiopharmaceuticals.
0344	Nuclear Medicine; Therapeutic Radiopharmaceuticals.
0370	Anesthesia; General Classification.
0371	Anesthesia; Anesthesia Incident to Radiology.
0372	Anesthesia; Anesthesia Incident to Other DX Services.
0379	Anesthesia; Other Anesthesia.
0390	Administration, Processing and Storage for Blood and Blood Components; General Classification.

TABLE 6—FINAL CY 2010 PACKAGED REVENUE CODES—Continued

Revenue code	Description
*0392	Administration, Processing and Storage for Blood and Blood Components; Processing and Storage.
0399	Administration, Processing and Storage for Blood and Blood Components; Other Blood Handling.
0621	Medical Surgical Supplies—Extension of 027X; Supplies Incident to Radiology.
0622	Medical Surgical Supplies—Extension of 027X; Supplies Incident to Other DX Services.
*0623	Medical Supplies—Extension of 027X, Surgical Dressings.
0624	Medical Surgical Supplies—Extension of 027X; FDA Investigational Devices.
0630	Pharmacy—Extension of 025X; Reserved.
0631	Pharmacy—Extension of 025X; Single Source Drug.
0632	Pharmacy—Extension of 025X; Multiple Source Drug.
0633	Pharmacy—Extension of 025X; Restrictive Prescription.
0681	Trauma Response; Level I Trauma.
0682	Trauma Response; Level II Trauma.
0683	Trauma Response; Level III Trauma.
0684	Trauma Response; Level IV Trauma.
0689	Trauma Response; Other.
0700	Cast Room; General Classification.
0710	Recovery Room; General Classification.
0720	Labor Room/Delivery; General Classification.
0721	Labor Room/Delivery; Labor.
0732	EKG/ECG (Electrocardiogram); Telemetry.
0762	Specialty Room—Treatment/Observation Room; Observation Room.
0801	Inpatient Renal Dialysis; Inpatient Hemodialysis.
0802	Inpatient Renal Dialysis; Inpatient Peritoneal Dialysis (Non-CAPD).
0803	Inpatient Renal Dialysis; Inpatient Continuous Ambulatory Peritoneal Dialysis (CAPD).
0804	Inpatient Renal Dialysis; Inpatient Continuous Cycling Peritoneal Dialysis (CCPD).
0809	Inpatient Renal Dialysis; Other Inpatient Dialysis.

TABLE 6—FINAL CY 2010 PACKAGED REVENUE CODES—Continued

Revenue code	Description
0810	Acquisition of Body Components; General Classification.
0819	Inpatient Renal Dialysis; Other Donor.
0821	Hemodialysis-Outpatient or Home; Hemodialysis Composite or Other Rate.
0824	Hemodialysis-Outpatient or Home; Maintenance—100%.
0825	Hemodialysis-Outpatient or Home; Support Services.
0829	Hemodialysis-Outpatient or Home; Other OP Hemodialysis.
0942	Other Therapeutic Services (also see 095X, an extension of 094x); Education/Training.
*0943	Other Therapeutic Services (also see 095X, an extension of 094X), Cardiac Rehabilitation.
*0948	Other Therapeutic Services (also see 095X, an extension of 094X), Pulmonary Rehabilitation.

In addition, we excluded: (1) claims that had zero costs after summing all costs on the claim, and (2) claims containing packaging flag number 3. Effective for services furnished on or after July 1, 2004, the I/OCE assigned packaging flag number 3 to claims on which hospitals submitted token charges for a service with status indicator “S” or “T” (a major separately payable service under the OPSS) for which the fiscal intermediary or MAC was required to allocate the sum of charges for services with a status indicator equaling “S” or “T” based on the relative weight of the APC to which each code was assigned. We do not believe that these charges, which were token charges as submitted by the hospital, are valid reflections of hospital resources. Therefore, we deleted these claims. We also deleted claims for which the charges equaled the revenue center payment (that is, the Medicare payment) on the assumption that where the charge equaled the payment, to apply a CCR to the charge would not yield a valid estimate of relative provider cost.

For the remaining claims, we then standardized 60 percent of the costs of the claim (which we have previously determined to be the labor-related portion) for geographic differences in labor input costs. We made this adjustment by determining the wage index that applied to the hospital that

furnished the service and dividing the cost for the separately paid HCPCS code furnished by the hospital by that wage index. As has been our policy since the inception of the OPSS, we proposed to use the pre-reclassified wage indices for standardization because we believe that they better reflect the true costs of items and services in the area in which the hospital is located than the post-reclassification wage indices and, therefore, would result in the most accurate unadjusted median costs.

We also excluded claims that were outside 3 standard deviations from the geometric mean of units for each HCPCS code on the bypass list (because, as discussed above, we used claims that contain multiple units of the bypass codes).

After removing claims for hospitals with error CCRs, claims without HCPCS codes, claims for immunizations not covered under the OPSS, and claims for services not paid under the OPSS, approximately 58 million claims were left. Using these 58 million claims, we created approximately 99 million single and “pseudo” single claims, of which we used 99 million single bills (after trimming out approximately 657,000 claims as discussed above in this section) in the CY 2010 median development and ratesetting.

We used these claims to calculate the CY 2010 median costs for each separately payable HCPCS code and each APC. The comparison of HCPCS code-specific and APC medians determines the applicability of the 2 times rule. Section 1833(t)(2) of the Act provides that, subject to certain exceptions, the items and services within an APC group cannot be considered comparable with respect to the use of resources if the highest median (or mean cost, if elected by the Secretary) for an item or service in the group is more than 2 times greater than the lowest median cost for an item or service within the same group (the 2 times rule). Finally, we reviewed the median costs for this final rule with comment period and reassigned HCPCS codes to different APCs where we believed that it was appropriate. Section III. of this final rule with comment period includes a discussion of certain HCPCS code assignment changes that resulted from examination of the median costs, review of the public comments, and for other reasons. The APC medians were recalculated after we reassigned the affected HCPCS codes. Both the HCPCS code-specific medians and the APC medians were weighted to account for the inclusion of multiple units of the bypass codes in the creation of “pseudo” single bills.

Comment: Several commenters objected to the volatility of the OPPS rates from year to year. The commenters asserted that the absence of stability in the OPPS rates creates budgeting, planning, and operating problems for hospitals, and that as more care is provided on an outpatient, rather than inpatient basis, the need for stable payment rates from one year to the next becomes more important to hospitals. Some commenters suggested that CMS limit reductions in APC payments to a set percentage, with one commenter noting that CMS dampened payment decreases for blood and blood products to mitigate large payment fluctuations in order to limit provider losses. One commenter suggested that the median costs from claims be adjusted to limit changes from year to year. Another commenter suggested that CMS perform a thorough examination of the payment rates and examine billed charges, costs, median and mean costs, and CCRs to isolate the source of the fluctuations as well as mandate a review of all APCs that fluctuate above a certain percentage, similar to the 2 times rule.

Response: There are a number of factors pertinent to the OPPS that may cause median costs to change from one year to the next. Some of these are a reflection of hospital behavior, and some of them are a reflection of fundamental characteristics of the OPPS as defined in statute. For example, the OPPS payment rates are based on hospital cost report and claims data. However, hospital costs and charges change each year and this results in both changes to the CCRs taken from the most currently available cost reports and also differences in the charges on the claims that are the basis of the calculation of the median costs on which OPPS rates are based. Similarly, hospitals adjust their mix of services from year to year by offering new services and ceasing to furnish services and changing the proportion of the various services they furnish, which have an impact on the CCRs that we derive from their cost reports. CMS cannot stabilize these hospital-driven fundamental inputs to the calculation of OPPS payment rates.

Moreover, there are other essential elements of the OPPS which contribute to the changes in relative weights each year. These include, but are not limited to, reassignments of HCPCS codes to APCs to rectify 2 times violations as required by the law, to address the costs of new services, to address differences in hospitals' costs that may result from changes in medical practice, and to respond to public comments. Our efforts to improve payment accuracy may also

contribute to payment volatility in the short run, as may be the case when we are eventually able to use more specific CCRs to estimate the costs of implantable devices, based on the final policy that we adopted to disaggregate the single cost center for medical supplies into two more specific cost centers, as described in the FY 2009 IPPS final rule (73 FR 48458 through 48467). Moreover, for some services, we cannot avoid using small numbers of claims, either because the volume of services is naturally low or because the claims data do not facilitate the calculation of a median cost for a single service. Where there are small numbers of claims that are used in median calculation, there is more volatility in the median cost from one year to the next. Lastly, changes to OPPS payment policy (for example, changes to packaging) also contribute to some extent to the fluctuations in the OPPS payment rates for the same services from year to year.

We cannot avoid the naturally occurring volatility in the cost report and claims data that hospitals submit and on which the payment rates are based. Moreover (with limited exceptions), we reassign HCPCS codes to APCs where it is necessary to avoid 2 times violations. However, we have made other changes to resolve some of the other potential reasons for instability from year to year. Specifically, we continue to seek ways to use more claims data so that we have fewer APCs for which there are small numbers of single bills used to set the APC median costs. Moreover, we have tried to eliminate APCs with very small numbers of single bills where we could do so. We recognize that changes to payment policies, such as the packaging of payment for ancillary and supportive services and the implementation of composite APCs, may contribute to volatility in payment rates in the short term, but we believe that larger payment packages and bundles should help to stabilize payments in future years by enabling us to use more claims data and by establishing payments for larger groups of services.

While we recognize the reasoning behind a policy that would dampen both increases and decreases in the weights or payment rates of the OPPS, this would not be as simple or beneficial as commenters have implied. Implementing such a dampening policy would require the assumption that payment policy is static from year to year. Based on the commenters' own acknowledgement, and the data used to develop the OPPS, we know that this is not true. Further, in seeking to mitigate

fluctuations in the OPPS, implementing such a system would make payments less reflective of the true service costs. Dampening payments across all APCs in this way could unfairly harm those hospitals whose true cost for a service increases significantly, while inappropriately benefiting those hospitals whose true cost for a service decreases significantly. While one commenter requested that CMS adopt a policy to investigate any APCs that fluctuate above a certain threshold, this mandate would be unnecessary since we already examine all APCs that experience significant median cost fluctuations, as described in the CY 2010 OPPS/ASC proposed rule (74 FR 35626 through 35627).

Comment: Some commenters asked that CMS provide an adjustment for medical education costs under the OPPS because many of the costs of teaching services are now incurred in the HOPD as services previously furnished only in the inpatient setting are now being furnished in the HOPD. They also noted that the OPPS did not have a teaching adjustment while many of the other Medicare payment systems, such as inpatient, psychiatric, and rehabilitation facilities, already include one. These commenters stated that CMS indicated that it would study the costs and payment differential among different classes of providers in the April 7, 2000 OPPS final rule but has not done so. They recommended that CMS study whether the hospital outpatient costs of teaching hospitals are higher than the costs of other hospitals for purposes of determining whether there should be a teaching hospital adjustment. The commenters explained that analysis of 2007 Medicare cost reports showed that the average outpatient margins were -30.4 for major teaching hospitals, -13.8 for other teaching hospitals, and -14.4 for nonteaching hospitals. They believed that these findings demonstrated that the hospital outpatient costs of major teaching hospitals are significantly greater than the costs of other hospitals. The commenters requested that CMS conduct its own analysis and that if that analysis showed a difference due to the unique missions of teaching hospitals, CMS should add a teaching adjustment to the OPPS.

Response: Unlike payment under the IPPS, the law does not provide for payment for indirect medical education costs to be made under the OPPS. Section 1833(t)(2)(E) of the Act states that the Secretary shall establish, in a budget neutral manner " * * * other adjustments as determined to be necessary to ensure equitable payments,

such as adjustments for certain classes of hospitals.” We have not found such an adjustment to be necessary to ensure equitable payments to teaching hospitals and, therefore, have not developed such an adjustment. Furthermore, in this final rule with comment period, we have developed payment weights that we believe provide appropriate and adequate payment for the complex medical services, such as new technology services and device-dependent procedures, which we understand are furnished largely by teaching hospitals. We note that teaching hospitals benefit from the recalibration of the APCs in this final rule with comment period. The final CY 2010 impacts by class of hospital are displayed in Table 73 in section XXI.B. of this final rule with comment period.

After consideration of the public comments we received, we are finalizing our proposed CY 2010 methodology for calculating the median costs upon which the CY 2010 OPSS payment rates are based, with modifications as discussed throughout this section.

In some cases, APC median costs are calculated using variations of the process outlined above. Section II.A.2.d. of this final rule with comment period that follows addresses the calculation of single APC criteria-based median costs. Section II.A.2.e. of this final rule with comment period discusses the calculation of composite APC criteria-based median costs. Section X.B. of this final rule with comment period addresses the methodology for calculating the median cost for partial hospitalization services.

At the February 2009 APC Panel Meeting, the APC Panel recommended that CMS study the claims data for any APC in which the calculated payment reduction would be greater than 10 percent. The APC Panel also recommended that CMS provide a list of APCs to the APC Panel at the next meeting with a proposed payment rate change of greater than 10 percent. While we recognize the concerns the APC Panel expressed with regards to cost variability in the system, we already engage in a standard review process for all APCs that experience significant changes in median costs. We study all significant changes in estimated cost to determine the effect that proposed and final payment policies have on the APC payment rates and ensure that these policies are appropriate and that the intended cost estimation methodologies have been correctly applied. We note that there are a number of factors that cause APC median costs to change from

one year to the next. Some of these are a reflection of hospital behavior, and some of them are a reflection of fundamental characteristics of the OPSS as defined in the statute. With limited exceptions, we are required by law to reassign HCPCS codes to APCs where it is necessary to avoid 2 times violations. Thus, there are various mechanisms already in place to ensure that we assess changes in cost and adjust APC weights accordingly or justify why we have not made adjustments. We plan to continue our examination of all APCs that experience changes of greater than 10 percent. In the CY 2010 OPSS/ASC proposed rule (74 FR 35267), we indicated that we would provide the APC Panel with a list of the APCs with proposed changes in costs of more than 10 percent for CY 2010 at the next CY 2009 APC Panel meeting. Accordingly, we accepted this recommendation of the APC Panel in full.

At the August 2009 meeting of the APC Panel, we provided the APC Panel a list of all APCs fluctuating by more than 10 percent when comparing the CY 2010 proposed rule APC median costs to those based on CY 2009 final rule data. We found that the median costs for 7 APCs decreased by 10 percent or more and the median costs for 63 APCs increased by 10 percent or more. These changes occurred due to some of the reasons described earlier, including reassignment of HCPCS codes from one APC to another to resolve 2 times violations, modeling changes such as the removal of lines for codes that were not payable in CY 2008 under the OPSS payment rules, low volumes of services influencing the claims used to determine APC median costs, and updated cost and charge information from hospital claims and cost reports. We noted that the median costs for 63 APCs increased by 10 percent or more and that the reasons for the increases were similar to the reasons for the decreases of more than 10 percent but, in general, we found nothing that raised concern regarding the data process we used to calculate the proposed median costs. The APC Panel discussed the different APCs on the list but did not express any significant concern with the fluctuations. As a result, they did not make any further recommendations related to the list of APCs with median costs fluctuating by greater than 10 percent.

At the February 2009 APC Panel meeting, we reviewed and examined the data process in preparation for the CY 2010 rulemaking cycle. At this meeting, the APC Panel recommended that the Data Subcommittee continue its work and we accepted that recommendation.

The APC Panel further recommended at the August 2009 meeting that the Data Subcommittee continue its work. We are accepting this most recent recommendation, and we will continue to work closely with the APC Panel's Data Subcommittee to prepare and review data and analyses relevant to the APC configurations and OPSS payment policies for hospital outpatient items and services.

d. Calculation of Single Procedure APC Criteria-Based Median Costs

(1) Device-Dependent APCs

Device-dependent APCs are populated by HCPCS codes that usually, but not always, require that a device be implanted or used to perform the procedure. For a full history of how we have calculated payment rates for device-dependent APCs in previous years and a detailed discussion of how we developed the standard device-dependent APC ratesetting methodology, we refer readers to the CY 2008 OPSS/ASC final rule with comment period (72 FR 66739 through 66742). Overviews of the procedure-to-device edits and device-to-procedure edits used in ratesetting for device-dependent APCs are available in the CY 2005 OPSS final rule with comment period (69 FR 65761 through 65763) and the CY 2007 OPSS/ASC final rule with comment period (71 FR 68070 through 68071).

In the CY 2010 OPSS/ASC proposed rule (74 FR 35267), we proposed to revise our standard methodology for calculating median costs for device-dependent APCs, which utilizes claims data that generally represent the full cost of the required device, to exclude claims that contain the “FC” modifier. Specifically, we proposed to calculate the median costs for device-dependent APCs for CY 2010 using only the subset of single procedure claims from CY 2008 claims data that pass the procedure-to-device and device-to-procedure edits; do not contain token charges (less than \$1.01) for devices; do not contain the “FB” modifier signifying that the device was furnished without cost to the provider, supplier, or practitioner, or where a full credit was received; and do not contain the “FC” modifier signifying that the hospital received partial credit for the device. The “FC” modifier became effective January 1, 2008, and is present for the first time on claims that would be used in OPSS ratesetting for CY 2010. We stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35267) that we believe the standard methodology for calculating median costs for device-

dependent APCs, further refined to exclude claims with the "FC" modifier, gives us the most appropriate median costs for device-dependent APCs in which the hospital incurs the full cost of the device.

The median costs for the majority of device-dependent APCs that were calculated using the CY 2010 proposed rule claims data were generally stable, with most median costs increasing moderately compared to the median costs upon which the CY 2009 OPPS payment rates were based. However, the median costs for APC 0225 (Implantation of Neurostimulator Electrodes, Cranial Nerve) and APC 0418 (Insertion of Left Ventricular Pacing Electrode) demonstrated significant fluctuation. Specifically, the proposed CY 2010 median cost for APC 0225 increased approximately 49 percent compared to the final CY 2009 median cost, although this APC median cost had declined by approximately the same proportion from CY 2008 to CY 2009. The proposed CY 2010 median cost for APC 0418, which had decreased approximately 45 percent from CY 2008 to CY 2009, showed an increase of approximately 56 percent based on the claims data available for the CY 2010 proposed rule. As indicated in the CY 2010 OPPS/ASC proposed rule (74 FR 35267), we believe the fluctuations in median costs for these two APCs are a consequence of the small number of single bills upon which the median costs are based and the small number of providers of these services. As we have stated in the past, some fluctuation in relative costs from year to year is to be expected in a prospective payment system for low volume device-dependent APCs, particularly where there are small numbers of single bills from a small number of providers.

At the February 2009 meeting of the APC Panel, one presenter stated that the assignment of the single-array cranial neurostimulator pulse generator implantation procedure described by CPT code 61885 (Insertion or replacement of cranial neurostimulator pulse generator or receiver, direct or inductive coupling; with connection to a single electrode array) to APC 0039 (Level I Implantation of Neurostimulator Generator), along with the peripheral/gastric neurostimulator pulse generator implantation procedure described by CPT code 64590 (Insertion or replacement of peripheral or gastric neurostimulator pulse generator or receiver, direct or inductive coupling) is not appropriate, given the clinical and cost differences between the two procedures. According to the presenter, the cranial procedure described by CPT

code 61885 is more similar clinically and in terms of resource utilization to the spinal neurostimulator pulse generator implantation procedure described by CPT code 63685 (Insertion or replacement of spinal neurostimulator pulse generator or receiver, direct or inductive coupling), which is the only CPT code assigned to APC 0222 (Level II Implantation of Neurostimulator) for CY 2009. The presenter requested that the APC Panel recommend that CMS restructure the existing configuration of neurostimulator pulse generator implantation APCs for CY 2010 by splitting APC 0039, so that procedures involving peripheral/gastric neurostimulators and cranial neurostimulators would be in distinct APCs, or by reassigning the cranial neurostimulator pulse generator implantation procedure described by CPT code 61885 from APC 0039 to APC 0222. In response to this request, the APC Panel recommended that CMS combine APC 0039 and APC 0222 for CY 2010, given the overall similarity in median costs among the cranial, peripheral/gastric, and spinal neurostimulator pulse generator implantation procedures assigned to these two APCs. The APC Panel also recommended that CMS maintain the configuration of APC 0315 (Level III Implantation of Neurostimulator Generator) as it currently exists in CY 2009 for CY 2010. The dual-array cranial neurostimulator pulse generator implantation procedure described by CPT code 61886 (Insertion or replacement of cranial neurostimulator pulse generator or receiver, direct or inductive coupling; with connection to two or more electrode arrays) is currently the only procedure assigned to APC 0315.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35267 through 35268), we stated that we agree with the APC Panel that the median costs of the procedures described by CPT codes 61885, 63685, and 64590 are sufficiently similar to warrant placement of the CPT codes into a single APC, rather than two APCs. We accepted the APC Panel's recommendation and, therefore, proposed to reassign CPT code 63685 to APC 0039, to delete APC 0222, and to maintain the current configuration of APC 0315 for CY 2010. We also proposed to change the title of APC 0315 to "Level II Implantation of Neurostimulator Generator" to reflect the proposed two-level, rather than three-level, structure of the neurostimulator pulse generator implantation APCs.

In reviewing the APC Panel recommendation for consolidating APC 0039 and APC 0222, we observed that the median costs of the procedures assigned to APC 0425 (Level II Arthroplasty or Implantation with Prosthesis) and APC 0681 (Knee Arthroplasty) also are sufficiently similar to warrant combining these two APCs into one APC. The proposed median cost for the only procedure currently assigned to APC 0681, described by CPT code 27446 (Arthroplasty, knee, condyle and plateau; medial OR lateral compartment), was approximately \$7,464 based on the claims data available for the CY 2010 OPPS/ASC proposed rule. This proposed median cost was very similar to the proposed median cost of approximately \$7,852 calculated for APC 0425, which included other procedures involving the implantation of prosthetic devices into bone, similar to the procedure described by CPT code 27446. Given the shared resource and clinical characteristics of the procedures included in APC 0425 and the only procedure assigned to APC 0681 for CY 2009, in the CY 2010 OPPS/ASC proposed rule, we proposed to consolidate these two APCs by reassigning CPT code 27446 to APC 0425, and deleting APC 0681. We also noted that, over the past several years, the median cost for CPT code 27446 has fluctuated due to a low volume of services being performed by a small number of providers, and to a single provider performing the majority of services (73 FR 68535). We indicated in the CY 2010 OPPS/ASC proposed rule (74 FR 35268) that we believe by reassigning CPT code 27446 to APC 0425 and deleting APC 0681, we can maintain greater stability from year to year in the payment rate for this knee arthroplasty service, while also paying appropriately for the service.

At its August 2009 meeting, the APC Panel heard a joint presentation from neurostimulator manufacturers who asserted that CMS' proposal to consolidate spinal, peripheral/gastric, and single-array cranial neurostimulator pulse generator implantation procedures into a single APC does not adequately capture facility resources associated with the different types of neurostimulator pulse generators involved in these procedures and would undermine access to rechargeable neurostimulators. The neurostimulator manufacturers asked the APC Panel to recommend to CMS a revised, three-level APC configuration for neurostimulator pulse generator implantation procedures that would

differentiate payment for procedures involving rechargeable and nonrechargeable neurostimulators. Following discussion of this request, the APC Panel recommended that CMS adopt the two-level neurostimulator pulse generator implantation APC configuration proposed by CMS for CY 2010.

Comment: Many commenters supported CMS' proposal to continue using the standard methodology for calculating median costs for device-dependent APCs, revised to exclude claims that contain the "FC" modifier. The commenters stated that the exclusion of partial credit claims would result in APC median costs that more appropriately reflect true hospital costs. Some commenters also supported the mandatory reporting of all HCPCS device C-codes to encourage hospitals to remain vigilant in reporting the costs of performing services involving devices. The commenters urged CMS to continue educating hospitals on the importance of accurate coding for devices, supplies, and other technologies to help ensure these items are more appropriately reflected in future years' payment rates for outpatient services.

Some commenters recommended CMS continue examining and refining the ratesetting methodology for procedures involving devices in order to encourage the continued development and proliferation of new technology. The commenters also encouraged CMS to develop mechanisms for capturing the costs of devices included on multiple procedure claims.

Response: We appreciate the commenters' support of the standard device-dependent APC ratesetting methodology, including our proposal to refine the methodology to exclude claims that contain the "FC" modifier. As we have stated in the past (73 FR 68535 through 68536), we agree that accurate reporting of device, supply, and technology charges will help to ensure that these items are appropriately accounted for in future years' OPSS payment rates. We encourage stakeholders to carefully review HCPCS code descriptors, as well as any guidance CMS may have provided for specific HCPCS codes. In addition, we have provided further instructions on the billing of medical and surgical supplies in the October 2008 OPSS update (Transmittal 1599, Change Request 6196, dated September 19, 2008) and the April 2009 OPSS update (Transmittal 1702, Change Request 6416, dated March 13, 2009). For HCPCS codes that are paid under the OPSS, providers may also submit inquiries to the AHA Central Office on

HCPCS, which serves as a clearinghouse on the proper use of Level I HCPCS codes for hospitals and certain Level II HCPCS codes for hospitals, physicians, and other health professionals. Inquiries must be submitted using the approved form, which may be downloaded from the AHA Web site (<http://www.ahacentraloffice.org>) and either faxed to 312-422-4583 or mailed directly to the AHA Central Office: Central Office on HCPCS, American Hospital Association, One North Franklin, Floor 29, Chicago, IL 60606.

We agree with the commenters that we should continue to encourage the development and proliferation of new technology under the OPSS. We have special mechanisms to provide payment for new technologies and services under the OPSS, including new technology APCs and transitional pass-through payments for certain devices. We refer readers to sections III.C. and IV.A., respectively, of this final rule with comment period for more information on these payment methodologies. For all OPSS services, we continue our efforts to use the data from as many multiple procedure claims as possible, through approaches such as use of the bypass list and date splitting of claims as described further in section II.A. of this final rule with comment period, and through methodologies such as increased packaging and composite APCs. We refer readers to section II.A.2.e. of this final rule with comment period for a detailed summary of the public comments related to the establishment of a composite payment methodology for procedures involving cardiac resynchronization therapy defibrillators and pacemakers and our responses.

Comment: Many commenters responded to CMS' proposal to revise the APC configuration for neurostimulator pulse generator implantation procedures from a three-level structure to a two-level structure. While one commenter supported the proposal to combine the single-array cranial neurostimulator pulse generator implantation procedure, described by CPT code 61885 and used for vagus nerve stimulation, with the spinal neurostimulator pulse generator implantation procedure, described by CPT code 63685, many commenters argued that the proposed two-level configuration for neurostimulator pulse generator implantation procedures would threaten patient access to rechargeable spinal neurostimulators. These commenters asserted that hospitals may be unable to offer rechargeable spinal neurostimulator pulse generators at the proposed CY

2010 payment rate for APC 0039, which, according to the commenters, is substantially less than the cost of the device and the CY 2009 payment rate for the procedure. Some commenters presented an analysis of CY 2008 OPSS claims data available for the CY 2010 OPSS/ASC proposed rule that demonstrated a \$4,132 difference in costs for spinal neurostimulator pulse generator implantation procedures involving rechargeable devices compared to the same procedures involving nonrechargeable devices. According to these commenters, this difference in cost warrants a separate APC for rechargeable spinal neurostimulator pulse generator procedures. They argued that while the cost difference does not violate the 2 times rule, it is large enough to influence hospitals to choose the lower cost nonrechargeable spinal neurostimulator pulse generators instead of the rechargeable devices if hospitals receive the same payment for the implantation procedure, regardless of the type of technology that is used. Several commenters noted that the threat to patient access to rechargeable spinal neurostimulators should be of particular concern to CMS, given the Agency's past recognition of the technology's ability to reduce the need for device replacements and the associated surgical risks, thereby reducing costs while providing optimal therapy.

Some commenters also stated that the consolidation of APC 0039 and APC 0222 would result in a disproportionately small number of single claims for procedures involving spinal neurostimulator pulse generators being used in ratesetting compared to the number of single claims for other types of neurostimulator pulse generator implantation procedures (specifically, peripheral/gastric and single-array cranial), further reducing the payment for these procedures relative to their costs. The commenters pointed out that, because spinal neurostimulator pulse generator implantation procedures are almost always performed with permanent lead placement procedures, rather than being staged as is common with other neurostimulator implantation procedures, they are typically not captured in the single claims used to calculate the median cost for consolidated APC 0039, upon which payment for that APC would be based. Many commenters argued that the proposed policy would be inconsistent with CMS' rationale in the CY 2008 OPSS/ASC final rule with comment period for implementing the current

APC configuration for neurostimulator pulse generator implantation procedures, which places the spinal neurostimulator pulse generator implantation procedure in its own APC. According to the commenters, CMS implemented a separate APC for this procedure because, unlike other neurostimulator pulse generator implantation procedures that involve only the less costly nonrechargeable devices, spinal neurostimulator pulse generator implantation procedures utilize either the more costly rechargeable device or the less costly nonrechargeable device. The commenters summarized CMS' assessment in the CY 2008 OPPTS/ASC final rule with comment period that the placement of the procedure described by CPT code 63685 as the only procedure in APC 0222 would enable CMS to calculate payment rates for spinal neurostimulator implantation procedures that reflect changes in surgical practice based on clinical, rather than financial, considerations.

Many commenters asserted that CMS' proposed APC configuration for neurostimulator pulse generator implantation procedures would result in APC 0039 being overly broad and clinically heterogeneous. The commenters stated that the spinal, peripheral/gastric, and single-array cranial neurostimulator pulse generator implantation procedures proposed for assignment to APC 0039 are clinically disparate and involve widely diverse neurostimulator technologies (including vagus nerve stimulators for epilepsy, sacral nerve stimulators for urinary incontinence, gastric pacemakers for chronic nausea and vomiting, and spinal neurostimulators for chronic neuropathic pain). One commenter requested that the CY 2010 proposal for neurostimulator pulse generator implantation procedures be reviewed by a pain management physician and a certified coder working in pain management.

According to the commenters, in order to address these concerns, CMS should differentiate payment for procedures involving rechargeable and nonrechargeable neurostimulators by revising the current (CY 2009) three-level APC payment structure for neurostimulator pulse generator implantation procedures. The commenters stated that their recommended configuration would group peripheral/gastric and spinal neurostimulator pulse generator implantation procedures (described by CPT codes 64590 and 63685, respectively) involving nonrechargeable devices in Level 1; single-array cranial

neurostimulator pulse generator implantation procedures (described by CPT code 61885) involving nonrechargeable devices in Level 2; and dual-array cranial neurostimulator pulse generator implantation procedures (described by CPT code 61886) and any neurostimulator pulse generator implantation procedure involving rechargeable devices in Level 3. According to the commenters, this APC configuration for neurostimulator pulse generator implantation procedures could be implemented by assigning APCs based on the presence of HCPCS device C-codes present on claims or through the creation of new Level II HCPCS G-codes that would distinguish procedures performed to implant nonrechargeable neurostimulator pulse generators from those performed to implant rechargeable neurostimulator pulse generators. The commenters asserted that CMS has shown a willingness to use alternative mapping schemes in the past to differentiate resource costs for procedures involving technologies such as drug-eluting coronary stents, implantable cardioverter defibrillators (ICDs), and linear accelerator-based stereotactic radiosurgery (LINAC-SRS), when there are important technology and facility resource cost differences that cannot be identified through the use of existing CPT codes.

The commenters urged CMS to maintain the current neurostimulator pulse generator implantation APC configuration as adopted in CY 2008 if the Agency decides not to implement their recommended three-level technology-specific APC configuration, or to create a four-level APC configuration in which the existing APC 0039 is split, with one APC for single-array cranial neurostimulator pulse generator implantation procedures and a separate APC for peripheral/gastric neurostimulator pulse generator implantation procedures. According to the commenters, either approach would yield more accurate payment rates than CMS' proposal for CY 2010.

Response: We do not agree with the commenters who argued that we should not implement our CY 2010 proposal to revise the APC configuration of neurostimulator pulse generator implantation procedures from a three-level structure to a two-level structure. We are finalizing our CY 2010 proposal to reassign CPT code 63685 to APC 0039, to delete APC 0222, and to maintain the current configuration of APC 0315. We believe that the final CY 2010 median costs for the neurostimulator pulse generator implantation procedures, described by

CPT codes 61885, 63685, and 64590, are sufficiently similar to warrant their placement in a single APC, as demonstrated in Table 7 below. The difference between the procedure with the highest median cost in APC 0039, described by CPT code 63685, and the procedure with the lowest median cost in APC 0039, described by CPT code 64590, is approximately \$3,000. Even if we were to consider the difference in costs between spinal neurostimulator pulse generator implantation procedures described by CPT code 63685 when they are performed with a rechargeable device compared to when they are performed with a nonrechargeable device, estimated by the commenters to be approximately \$4,000, the grouping of these procedures in the same APC would not violate the 2 times rule. We also point out that, as demonstrated in Table 7, we use a similar number of single claims with each of the CPT codes assigned to APC 0039 to calculate the median cost upon which the final CY 2010 payment rate for APC 0039 is based.

We do not agree with the commenters that these modest differences in costs, either among the various types of neurostimulator pulse generator implantation procedures assigned to APC 0039 or among the same types of procedures involving rechargeable versus nonrechargeable devices, are sufficiently substantial to result in hospitals denying access to the limited subset of patients for whom the more expensive rechargeable technology is clinically indicated. We note that payment based on a measure of central tendency is a principle of any prospective payment system. As we have stated in the past (73 FR 68562), in some individual cases, payment exceeds the average cost, and in other cases, payment is less than the average cost. On balance, however, payment should approximate the relative cost of the average case, recognizing that, as a prospective payment system, the OPPTS is a system of averages.

In addition to being similar in terms of resource utilization, we believe the procedures described by CPT codes 61885, 63685, and 64590 are comparable from a clinical perspective because they all involve the subcutaneous placement of a neurostimulator pulse generator. We do not agree with the commenters who argued that these procedures should be considered clinically disparate because they use widely diverse technologies for very different clinical indications. It is not uncommon under the OPPTS to group procedures described by relatively general HCPCS codes that

may utilize a wide variety of technologies and may be performed to treat different patient populations in the same APC. Furthermore, as stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 66537), the standard device-dependent APC ratesetting methodology does not take into consideration patient diagnoses. In response to the commenter who requested that the CY 2010 proposal for neurostimulator pulse generator implantation procedures be reviewed by a pain management physician and a certified coder working in pain management, we note that the CMS staff involved in reviewing the clinical characteristics of the APC groups include medical advisors from a variety of specialties as well as certified coders.

We also do not agree that we should not implement the two-level APC configuration for neurostimulator pulse generator implantation procedures as proposed for CY 2010 because, as argued by some commenters, it would be inconsistent with our rationale in the CY 2008 OPPS/ASC final rule with comment period to maintain a separate APC solely for spinal neurostimulator pulse generator implantation procedures. It is our standard process under the OPPS to reassess the composition of APCs, including reviewing the median costs of individual HCPCS codes, annually when we have the most current claims and Medicare cost report data, and to propose through our annual rulemaking cycle changes that we believe are necessary to maintain and improve the clinical and resource homogeneity of APCs based on the updated data. In CY 2008, the median costs for the single-array cranial and peripheral/gastric neurostimulator pulse generator implantation procedures described by CPT codes 61885 and 64590 of \$12,799 and \$10,954, respectively, were more divergent from the median cost calculated for the spinal neurostimulator pulse generator implantation procedure of \$15,150 using the CY 2006 claims and cost report data available at that time, compared to the median costs for these procedures calculated from the CY 2008 claims and cost report data available for this CY 2010 OPPS/ASC final rule with comment period, as demonstrated in Table 7 below.

Finally, we do not agree with the commenters that we should differentiate payment for neurostimulator pulse generator implantation procedures based on the type of technology that is implanted (that is, rechargeable or nonrechargeable), nor do we agree with the commenters that past CMS policy to

use alternative mapping schemes to differentiate resource costs for certain procedures, such as those involving drug-eluting stents, ICDs, and LINAC-SRS, serves as a precedent to do so. As we have stated in the past (72 FR 66715 through 66716 and 73 FR 68538), a policy to provide different payments for the same procedures according to the types of devices implanted would not be consistent with our overall strategy under the OPPS to encourage hospitals to use resources more efficiently by increasing the size of the payment bundles. The circumstances surrounding the payment policies and coding configurations for drug-eluting stents (67 FR 66732 through 66734), ICDs (72 FR 66702 through 66703), and LINAC-SRS (72 FR 66734 through 66737) are markedly different from the circumstances surrounding neurostimulator pulse generator implantation procedures. We developed HCPCS G-codes to distinguish payment for procedures involving drug-eluting stents from procedures involving non-drug-eluting stents because drug-eluting stents did not meet the criteria for transitional pass-through payment or for payment under a New Technology APC. Unlike drug-eluting stents, rechargeable spinal neurostimulators were granted pass-through status under the OPPS in CY 2006, which lasted until December 31, 2007. In the case of ICDs, we created HCPCS G-codes to gather cost data on single and dual chamber ICDs, but we did not differentiate payment for ICD insertion based on the type of technology that was used (72 FR 66703). Finally, our policy to utilize HCPCS G-codes rather than CPT codes for payment under the OPPS for LINAC-SRS treatment delivery services recognizes the vastly different capital equipment costs required for various LINAC-SRS services, rather than differences in the costs of single-use devices implanted in patients during the same procedure.

Comment: Some commenters disagreed with CMS' presentation at the August 2009 APC Panel meeting of the proposed CY 2010 line-item median costs for the two device HCPCS C-codes that describe neurostimulator pulse generators, specifically HCPCS code C1767 (Generator, neurostimulator (implantable), nonrechargeable) and HCPCS code C1820 (Generator, neurostimulator (implantable), with rechargeable battery and charging system). The commenters disputed the accuracy of the data presented by CMS, specifically that the line-item median costs for HCPCS codes C1767 and C1820 are \$9,606 and \$9,636, respectively,

based on CY 2008 claims available for the CY 2010 OPPS/ASC proposed rule. According to the commenters, these line-item median costs are inconsistent with the commenters' analyses of CY 2010 OPPS/ASC proposed rule data, which indicated that the line-item median costs for HCPCS codes C1767 and C1820 are \$10,580 and \$13,587, respectively. One commenter urged CMS to reanalyze the data and to disregard the APC Panel's support of the proposed CY 2010 APC configuration for neurostimulator pulse generator implantation procedures if the data were found to be erroneous. Another commenter characterized CMS' presentation of the line-item median costs for HCPCS codes C1767 and C1820 as incomplete because OPPS payment rates are based upon median costs that include all packaged items and services associated with providing a procedure as they appear on single claims, and not the line-item median costs for individual devices. The commenter asked CMS to ensure that all data presented to the APC Panel in the future is full and appropriate information for decisionmaking.

Response: In response to the commenters' concerns, we reassessed our methodology for calculating the proposed CY 2010 line-item median costs for HCPCS codes C1767 and C1820 and verified that the information presented to the APC Panel is accurate based on the CY 2008 claims data available for the CY 2010 OPPS/ASC proposed rule. The line-item statistics for these HCPCS codes, along with all other HCPCS codes recognized under the OPPS, are released to the public as part of the OPPS limited data set. We do not agree with the commenters that the presentation of these data was incomplete or inappropriate. We frequently consider line-item median costs for devices and other packaged items and services as one data element among several when we evaluate the clinical and resource homogeneity of APCs, particularly when stakeholders voice concerns that the costs of different items are driving procedure costs or influencing hospitals' decisions to provide certain services. An advantage of the line-item median costs is that they represent data from all OPPS claims, and not just the single claims that we are able to use in ratesetting for procedures. Therefore, we believe that a comparison of line-item costs is particularly appropriate for different types of neurostimulator pulse generators because one of the commenters' concerns was that there are relatively few single claims available

for ratesetting for the implantation of spinal neurostimulator pulse generators. We would expect the device costs on multiple procedure claims to be reflective of the hospital costs of these neurostimulator pulse generators, because commenters stated that multiple procedure claims resulted from the most typical spinal neurostimulator

implantation procedures. Furthermore, we would not expect there to be significant packaged costs associated with the neurostimulator pulse generators described by these device HCPCS codes. Therefore, we would expect the line-item median costs to accurately reflect the differential costs of non-rechargeable and rechargeable

neurostimulator technology. We note that the APC Panel members are well-acquainted with the OPPS ratesetting methodology, including the use of single procedure claims and not line-item median costs for individual items, to calculate the median costs upon which OPPS payment rates are based.

TABLE 7—CY 2010 APC CONFIGURATION FOR PAYMENT OF NEUROSTIMULATOR PULSE GENERATOR IMPLANTATION PROCEDURES

CY 2010 APC	Revised APC Title for CY 2010	CY 2010 CPT Code	CY 2010 CPT Code Descriptor	CY 2010 CPT Code Median Cost	CY 2010 CPT Code Single Claims	CY 2010 APC Median Cost
0039	Level I Implantation of Neurostimulator Generator.	61885	Insertion or replacement of cranial neurostimulator pulse generator or receiver, direct or inductive coupling; with connection to a single electrode array.	\$14,141	1,260	\$13,766
		63685	Insertion or replacement of spinal neurostimulator pulse generator or receiver, direct or inductive coupling.	15,802	1,262	13,766
		64590	Insertion or replacement of peripheral or gastric neurostimulator pulse generator or receiver, direct or inductive coupling.	12,726	1,978	13,766
0315	Level II Implantation of Neurostimulator Generator.	61886	Insertion or replacement of cranial neurostimulator pulse generator or receiver, direct or inductive coupling; with connection to two or more electrode arrays.	18,350	1,004	18,350

Comment: Several commenters expressed support for the proposed CY 2010 payment rate for the implantation of auditory osseointegrated devices, described by CPT codes 69714 (Implantation, osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; without mastoidectomy); 69715 (Implantation, osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; with mastoidectomy); 69717 (Replacement (including removal of existing device), osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; without mastoidectomy); and 69718 (Replacement (including removal of existing device), osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; with mastoidectomy) and assigned to APC 0425. Other commenters, however, stated that the proposed payment rate for APC 0425 is less than hospitals' device and service-related costs associated with the procedures described by these CPT codes and urged CMS to consider a slight increase in the payment for APC 0425.

Response: We agree with the commenters that the payment rate for APC 0425, calculated from the standard device-dependent APC ratesetting methodology, appropriately reflects hospitals' relative costs for providing the procedures assigned to APC 0425 as reported to us in the claims and cost report data. We used 1,410 single claims from CY 2008 to calculate the median cost upon which the final CY 2010 payment rate for APC 0425 is based. The final CY 2010 median cost for APC 0425 is approximately \$7,932, slightly higher than the final CY 2009 median cost for APC 0425 of \$7,863. We note that we were able to use significantly more single claims in ratesetting for APC 0425 for CY 2010 compared to CY 2009 (1,410 single claims from CY 2008 compared to 668 single claims from CY 2007). We disagree with the commenters who requested an additional increase in the payment rate for APC 0425, because this would artificially and inaccurately inflate payment rates. A fundamental principle of the OPPS is that it is based on relative weights, and as we have stated in the past (73 FR 68541), it is the relativity of the costs to one another, rather than absolute cost, that is important in setting payment rates. To deviate from our standard OPPS ratesetting methodology and increase the payment rates for certain procedures

beyond their relative costs as derived from claims and cost report data would skew this relativity.

Comment: Some commenters supported CMS' proposal to reassign CPT code 27446 to APC 0425 and to delete APC 0681. Other commenters, however, opposed the consolidation of these two APCs, arguing that the procedure described by CPT code 27446 is clinically dissimilar from the arthroplasty procedures currently assigned to APC 0425. The commenters recommended that CMS continue to maintain APC 0681 for CY 2010 and to add other total knee arthroplasty procedures to this APC, along with the procedure described by CPT code 27446.

Response: We disagree with the commenters who argued that it is necessary to maintain APC 0681 specifically for knee arthroplasty procedures because we do not believe it is appropriate to maintain an APC that is not necessary to classify services into groups that are similar clinically and in terms of resource utilization. We continue to believe that CPT code 27446 is most appropriately assigned to APC 0425 for CY 2010, as we proposed, based on consideration of the procedure's clinical and resource characteristics. As described in section XI.B. of this final rule with comment

period, we are not removing any total knee arthroplasty procedures from the inpatient list.

Comment: Several commenters supported the proposed payment rate for the implantation of cochlear implants, described by CPT code 69930 (Cochlear device implantation, with or without mastoidectomy) and assigned to APC 0259 (Level VII ENT Procedures). These commenters stated that while hospitals' device and service-related costs for these procedures likely still exceed the proposed payment rate for APC 0259, they represent an improvement in payment relative to CY 2009 that may lead to better access to care for Medicare beneficiaries.

Response: We appreciate the commenters' support of the proposed payment rate for APC 0259. We believe that the standard device-dependent APC ratesetting methodology results in a payment rate that reflects hospitals' relative costs for providing the procedure assigned to this APC as reported to us in the claims and cost report data.

Comment: One commenter concurred with CMS' proposal that APC 0385 (Level I Prosthetic Urological Procedures) and APC 0386 (Level II Prosthetic Urological Procedures) continue to be recognized as device-

dependent APCs. The commenter supported CMS' continued application of procedure-to-device edits for procedures assigned to these APCs.

Response: We appreciate the commenter's support of the continued recognition of APC 0385 and 0386 as device-dependent APCs. We agree that claims processing edits for devices that are integral to the performance of procedures assigned to device-dependent APCs are an important element of the standard device-dependent APC ratesetting methodology.

Comment: One commenter urged CMS not to reduce the payment for the procedure described by CPT code 62361 (Implantation or replacement of device for intrathecal or epidural drug infusion; nonprogrammable pump), which is assigned to APC 0227 (Implantation of Drug Infusion Device). The commenter stated that patient access to this procedure is limited due to recent payment cuts.

Response: The final CY 2010 median cost for APC 0227 of approximately \$13,268 is approximately 10 percent higher than the median cost of \$12,006, upon which the final CY 2009 payment rate was based, and approximately 13 percent higher than the median cost of \$11,569, upon which the final CY 2008

payment rate was based. We believe that the final CY 2010 median cost for APC 0227 of \$13,268, which is calculated using the standard device-dependent APC methodology, results in a final CY 2010 payment rate that accurately and appropriately reflects hospitals' costs for providing the service described by CPT code 62361 and will not result in any barriers to patient care.

In summary, after consideration of the public comments we received, we are finalizing our proposed CY 2010 payment policies for device-dependent APCs, without modification. The CY 2010 OPPS payment rates for device-dependent APCs are based on their median costs calculated from CY 2008 claims and the most recent cost report data, using only claims that pass the device edits, do not contain token charges for devices, and do not have a modifier signifying that the device was furnished without cost or with full or partial credit. We continue to believe that the median costs calculated from the single claims that meet these criteria represent the most valid estimated relative costs of these services to hospitals when they incur the full cost of the devices required to perform the procedures. The CY 2010 device-dependent APCs are listed in Table 8 below.

TABLE 8—CY 2010 DEVICE-DEPENDENT APCs

CY 2010 APC	CY 2010 Status indicator	CY 2010 APC Title
0039	S	Level I Implantation of Neurostimulator Generator
0040	S	Percutaneous Implantation of Neurostimulator Electrodes
0061	S	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electrodes
0082	T	Coronary or Non-Coronary Atherectomy
0083	T	Coronary or Non-Coronary Angioplasty and Percutaneous Valvuloplasty
0084	S	Level I Electrophysiologic Procedures
0085	T	Level II Electrophysiologic Procedures
0086	T	Level III Electrophysiologic Procedures
0089	T	Insertion/Replacement of Permanent Pacemaker and Electrodes
0090	T	Insertion/Replacement of Pacemaker Pulse Generator
0104	T	Transcatheter Placement of Intracoronary Stents
0106	T	Insertion/Replacement of Pacemaker Leads and/or Electrodes
0107	T	Insertion of Cardioverter-Defibrillator
0108	T	Insertion/Replacement/Repair of Cardioverter-Defibrillator Leads
0115	T	Cannula/Access Device Procedures
0202	T	Level VII Female Reproductive Procedures
0225	S	Implantation of Neurostimulator Electrodes, Cranial Nerve
0227	T	Implantation of Drug Infusion Device
0229	T	Transcatheter Placement of Intravascular Shunts
0259	T	Level VII ENT Procedures
0293	T	Level V Anterior Segment Eye Procedures
0315	S	Level II Implantation of Neurostimulator Generator
0384	T	GI Procedures with Stents
0385	S	Level I Prosthetic Urological Procedures
0386	S	Level II Prosthetic Urological Procedures
0418	T	Insertion of Left Ventricular Pacing Electrode
0425	T	Level II Arthroplasty or Implantation with Prosthesis
0427	T	Level II Tube or Catheter Changes or Repositioning
0622	T	Level II Vascular Access Procedures
0623	T	Level III Vascular Access Procedures
0648	T	Level IV Breast Surgery

TABLE 8—CY 2010 DEVICE-DEPENDENT APCs—Continued

CY 2010 APC	CY 2010 Status indicator	CY 2010 APC Title
0652	T	Insertion of Intraperitoneal and Pleural Catheters
0653	T	Vascular Reconstruction/Fistula Repair with Device
0654	T	Insertion/Replacement of a Permanent Dual Chamber Pacemaker
0655	T	Insertion/Replacement/Conversion of a Permanent Dual Chamber Pacemaker
0656	T	Transcatheter Placement of Intracoronary Drug-Eluting Stents
0674	T	Prostate Cryoablation
0680	S	Insertion of Patient Activated Event Recorders

(2) Blood and Blood Products

Since the implementation of the OPSS in August 2000, we have made separate payments for blood and blood products through APCs rather than packaging payment for them into payments for the procedures with which they are administered. Hospital payments for the costs of blood and blood products, as well as for the costs of collecting, processing, and storing blood and blood products, are made through the OPSS payments for specific blood product APCs.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35269), we proposed to continue to establish payment rates for blood and blood products using our blood-specific CCR methodology, which utilizes actual or simulated CCRs from the most recently available hospital cost reports to convert hospital charges for blood and blood products to costs. This methodology has been our standard ratesetting methodology for blood and blood products since CY 2005. It was developed in response to data analysis indicating that there was a significant difference in CCRs for those hospitals with and without blood-specific cost centers, and past comments indicating that the former OPSS policy of defaulting to the overall hospital CCR for hospitals not reporting a blood-specific cost center often resulted in an underestimation of the true hospital costs for blood and blood products. Specifically, in order to address the differences in CCRs and to better reflect hospitals' costs, we proposed to continue to simulate blood CCRs for each hospital that does not report a blood cost center by calculating the ratio of the blood-specific CCRs to hospitals' overall CCRs for those hospitals that do report costs and charges for blood cost centers. We would then apply this mean ratio to the overall CCRs of hospitals not reporting costs and charges for blood cost centers on their cost reports in order to simulate blood-specific CCRs for those hospitals. We calculated the median costs upon which the proposed CY 2010 payment rates for blood and blood products were based using the

actual blood-specific CCR for hospitals that reported costs and charges for a blood cost center and a hospital-specific simulated blood-specific CCR for hospitals that did not report costs and charges for a blood cost center.

We stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35269) that we continue to believe the hospital-specific, blood-specific CCR methodology better responds to the absence of a blood-specific CCR for a hospital than alternative methodologies, such as defaulting to the overall hospital CCR or applying an average blood-specific CCR across hospitals. Because this methodology takes into account the unique charging and cost accounting structure of each provider, we believe that it yields more accurate estimated costs for these products. We indicated that we believe continuing with this methodology in CY 2010 would result in median costs for blood and blood products that appropriately reflect the relative estimated costs of these products for hospitals without blood cost centers and, therefore, for these blood products in general.

Comment: One commenter expressed appreciation for CMS' recognition of the complexities of calculating payment rates for blood and blood products and the accommodations CMS has made through the blood and blood product ratesetting methodology to ensure the calculated rates are as fair as possible. However, several commenters stated that the proposed payment rates for many blood and blood products are less than the costs hospitals incur acquiring, managing, and processing them, and that the claims-based cost data for blood and blood products are error-prone and subject to significant and unexplained fluctuations. They noted that the payment decreases for several blood and blood products seem inexplicable because prices for blood have been increasing due to new technologies and tests required to ensure the continued safety of the blood supply and increasingly expensive donor recruitment and retention efforts. According to the commenters, a

comparison of the proposed APC payment changes for blood and blood products to the producer price index (PPI) for blood and organ banks, which increased 3.1 percent from July 2008 to July 2009, indicates that the blood product payment rates in the CY 2010 OPSS/ASC proposed rule do not reflect overall pricing trends in the blood banking industry. The commenters asked CMS to adjust the CY 2010 payment rates for blood and blood products by increasing all of the CY 2009 payment rates by 3.1 percent, or by implementing a 3.1 percent payment floor for CY 2010 payment rates compared to CY 2009 payment rates for blood and blood products. One commenter particularly urged CMS to apply a 3 percent minimum increase in payment for the highest volume blood product, described by HCPCS code P9016 (Red blood cells, leukocytes reduced, each unit). The commenters asserted that the use of the PPI for blood and organ banks in calculating hospital payment is not unprecedented. They stated that in the CY 2005 OPSS final rule, CMS used the PPI for blood and derivatives for human use in calculating the payment rates for low-volume blood products. They also pointed out that CMS recognized the value of the PPI for blood and organ banks by using it to update blood and blood product prices in the market basket under the IPPS for CY 2010.

Response: We continue to believe that using blood-specific CCRs applied to hospital claims data results in payments that appropriately reflect hospitals' relative costs of providing blood and blood products as reported to us by hospitals. We do not believe it is necessary or appropriate to use the PPI for blood and organ banks as a benchmark for updating the payment rates for blood and blood products from year to year, because it is not our standard process under the OPSS for any item or service to update payment rates by implementing across-the-board, product-specific inflation updates to the payment rates that were in place the year before. Rather, we annually update

payment groups and payment weights using the most recently available hospital claims and cost report data. This process allows us to recalibrate the payment groups and payment weights in response to changes in hospitals' costs from year to year. A fundamental principle of the OPPS is that it is based on relative weights, and as we have stated in the past (73 FR 68541), it is the relativity of the costs to one another, rather than absolute cost, that is important in setting payment rates. To deviate from our standard OPPS ratesetting methodology and update the payment rates for blood and blood products by the PPI would skew this relativity.

We also note that, as discussed in section II.B. of this final rule with comment period, we are required by law to update the conversion factor used to determine payment rates under the OPPS. For CY 2010, the update is equal to the hospital inpatient market basket increase. The PPI for blood and organ banks is one of several price proxies used to calculate the hospital inpatient market basket (74 FR 43847), which represents the change in price over time of the same mix (quantity and intensity) of goods and services purchased to provide hospital services. In this way, the PPI for blood and blood products is already incorporated in the CY 2010 payment rates for blood and blood products.

Comment: One commenter noted that the proposed CY 2010 median costs for several blood and blood products fluctuated significantly relative to CY 2009. The commenter expressed concern about potentially large payment decreases and noted that, in the past, CMS dampened payment decreases for blood and blood products to limit product losses. The commenter requested that CMS disclose the source of the fluctuations in CY 2010 median costs for blood and blood products and implement a dampening policy to mitigate significant payment fluctuations, not only for blood and blood products but for all other services.

Response: As stated previously, we continue to believe that using blood-specific CCRs applied to hospital claims data results in payments that appropriately reflect hospitals' relative costs of providing blood and blood products as reported to us by hospitals. We do not believe it is necessary or appropriate to implement a dampening policy to mitigate significant payment fluctuations, for blood and blood products or for any other items and services payable under the OPPS, as described in section II.A.2.c. of this final rule with comment period. As we have

stated in the past (73 FR 68541), it is our common practice to review significant changes in median costs from year to year and from the proposed rule to the final rule for a given calendar year. The final CY 2010 median costs for more than two-thirds of all blood and blood products changed by a margin of less than 10 percent compared to the CY 2009 median costs. Of the remaining blood and blood products, 8 demonstrated decreases in median costs of greater than 10 percent, and 5 demonstrated increases in median costs of greater than 10 percent. We determined that the fluctuations in median costs for these 13 blood and blood products were due to contributions of additional claims, the addition or removal of individual hospitals furnishing particular blood and blood products, and revised cost report data. For all APCs whose payment rates are based upon relative payment weights, we note that the quality and accuracy of reported units and charges significantly influence the median costs that are the basis for our payment rates, especially for low volume items and services. Beyond our standard OPPS trimming methodology (described in section II.A.2. of this final rule with comment period) that we apply to those claims that have passed various types of claims processing edits, it is not our general policy to judge the accuracy of hospital coding and charging for purposes of ratesetting.

Comment: One commenter recommended that CMS recognize plasma protein fraction (PPF) products as drugs under the OPPS and assign status indicator "K" (Nonpass-Through Drugs and Nonimplantable Biologicals, Including Therapeutic Radiopharmaceuticals) to HCPCS codes P9043 (Infusion, plasma protein fraction (human), 5%, 50 ml) and P9048 (Infusion, plasma protein fraction (human), 5%, 250 ml), rather than assigning them status indicator "R" (Blood and Blood Products). The commenter also requested that CMS instruct providers to use the appropriate infusion CPT codes for administration of PPF, rather than blood transfusion codes. According to the commenter, PPF is similar clinically to albumin in terms of how it is derived and the patients for whom it is indicated. The commenter also stated that, according to the AABB, both albumin and PPF are blood derivatives that should be billed with pharmacy revenue codes. According to the commenter, the AABB also indicates that the administration of blood derivatives, including PPF, should be billed with injection or infusion CPT

codes rather than blood transfusion CPT codes.

Response: We did not propose to change the status indicators for the PPF products described by HCPCS codes P9043 and P9048 from "R" to "K" for CY 2010. Because changing the status indicators for these products as the commenter recommended could have significant payment implications, we believe we should not consider such a potential change in policy without seeking input from all interested stakeholders through our annual rulemaking cycle. Specifically, changing the status indicator from "R" to "K" would require us to calculate the payment rates for PPF using mean unit cost from hospital claims, as we currently do for albumin products, rather than using our standard blood-specific CCR methodology for blood and blood products.

We last addressed the issue of whether plasma-derived therapies and their recombinant analogs should be considered blood and blood products for purposes of payment under the OPPS in the CY 2003 OPPS final rule with comment period (67 FR 66774) and the CY 2004 OPPS final rule with comment period (68 FR 63455). We stated that, because these products are highly processed and not manufactured by local blood banks, they do not have the same access and safety concerns as other blood and blood products. Therefore, we did not consider any plasma-derived products and their recombinant analogs, including albumin and immune globulins, to fall under the category of blood and blood products (67 FR 66774).

We are requesting comments on this final rule with comment period that address whether PPF should be recognized as a blood and blood product, designated with status indicator "R," or as a nonpass-through drug and biological, designated with status indicator "K." Specifically, we are interested in how PPF is derived and manufactured, and whether the same access and safety concerns that apply to the blood and blood products recognized under the OPPS for payment purposes also apply to PPF. Finally, we are interested in the relationship between albumin and PPF, from clinical, manufacturing, and safety perspectives, and whether there would be a rationale for treating these products similarly for payment purposes under the OPPS. We will consider these comments as we prepare for the CY 2011 annual rulemaking cycle.

Comment: One commenter asked if the product "prepoled cryoprecipitate"

would be added to the list of blood and blood products.

Response: The existing HCPCS code that describes cryoprecipitate products, P9012 (Cryoprecipitate, each unit), is recognized under the OPPS for payment purposes as a blood and blood product. We note there is an established process in place for requesting a revision to the Level II HCPCS codes if stakeholders believe the current codes cannot adequately address all clinical circumstances. The Level II HCPCS coding system is a comprehensive and standardized system that classifies similar products that are medical in nature into categories for the purpose of efficient claims processing. The process and criteria for revising Level II HCPCS codes is available on the CMS Web site at: http://www.cms.hhs.gov/MedHCPCSGenInfo/02_HCPCSCODING_PROCESS.asp#TopOfPage.

After consideration of the public comments we received, we are finalizing, without modification, our proposal to calculate the median costs upon which the CY 2010 payment rates for blood and blood products are based using the blood-specific CCR methodology that we have utilized since CY 2005. We believe that continuing this methodology in CY 2010 results in median costs for blood and blood products that appropriately reflect the relative estimated costs of these products for hospitals without blood cost centers and, therefore, for these products in general.

We refer readers to Addendum B to this final rule with comment period for the final CY 2010 payment rates for blood and blood products, which are identified with status indicator "R." For more detailed discussion of the blood-specific CCR methodology, we refer readers to the CY 2005 OPPS proposed rule (69 FR 50524 through 50525). For a full history of OPPS payment for blood and blood products, we refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66807 through 66810).

(3) Single Allergy Tests

In the CY 2010 OPPS/ASC proposed rule (74 FR 35269), we proposed to continue with our methodology of differentiating single allergy tests ("per test") from multiple allergy tests ("per visit") by assigning these services to two different APCs to provide accurate payments for these tests in CY 2010. Multiple allergy tests are currently assigned to APC 0370 (Allergy Tests), with a median cost calculated based on the standard OPPS methodology. We provided billing guidance in CY 2006 in Transmittal 804 (issued on January 3,

2006) specifically clarifying that hospitals should report charges for the CPT codes that describe single allergy tests to reflect charges "per test" rather than "per visit" and should bill the appropriate number of units of these CPT codes to describe all of the tests provided. However, as noted in the CY 2010 OPPS/ASC proposed rule (74 FR 35269), our CY 2008 claims data available for that proposed rule for APC 0381 did not reflect improved and more consistent hospital billing practices of "per test" for single allergy tests. The median cost of APC 0381, calculated for the proposed rule according to the standard single claims OPPS methodology, was approximately \$55, significantly higher than the CY 2009 median cost of APC 0381 of approximately \$23 calculated according to the "per unit" methodology, and greater than we would expect for these procedures that are to be reported "per test" with the appropriate number of units. Some claims for single allergy tests still appear to provide charges that represent a "per visit" charge, rather than a "per test" charge. Therefore, consistent with our payment policy for single allergy tests since CY 2006, we proposed to calculate a "per unit" median cost for APC 0381, based upon 530 claims containing multiple units or multiple occurrences of a single CPT code. The proposed CY 2010 median cost for APC 0381 using the "per unit" methodology was approximately \$29. For a full discussion of this methodology, we refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66737).

We did not receive any public comments on our CY 2010 proposal for payment of single allergy tests. Therefore, we are finalizing our CY 2010 proposal, without modification, to calculate a "per unit" median cost for APC 0381 as described above in this section. The final CY 2010 median cost of APC 0381 is approximately \$29.

(4) Echocardiography Services

In CY 2008, we implemented a policy whereby payment for all contrast agents is packaged into the payment for the associated imaging procedure, regardless of whether the contrast agent met the OPPS drug packaging threshold. Section 1833(t)(2)(G) of the Act requires us to create additional APC groups of services for procedures that use contrast agents to classify them separately from those procedures that do not utilize contrast agents. To reconcile this statutory provision with our final policy of packaging all contrast agents, for CY 2008, we calculated HCPCS code-specific median costs for all separately

payable echocardiography procedures that may be performed with contrast agents by isolating single and "pseudo" single echocardiography claims with the following CPT codes where a contrast agent was also billed on the claim:

- 93303 (Transthoracic echocardiography for congenital cardiac anomalies; complete);
- 93304 (Transthoracic echocardiography for congenital cardiac anomalies; follow-up or limited study);
- 93307 (Echocardiography, transthoracic, real-time with image documentation (2D) with or without M-mode recording; complete);
- 93308 (Echocardiography, transthoracic, real-time with image documentation (2D) with or without M-mode recording; follow-up or limited study);
- 93312 (Echocardiography, transesophageal, real time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report);
- 93315 (Transesophageal echocardiography for congenital cardiac anomalies; including probe placement, image acquisition, interpretation and report);
- 93318 (Echocardiography, transesophageal (TEE) for monitoring purposes, including probe placement, real time 2-dimensional image acquisition and interpretation leading to ongoing (continuous) assessment of (dynamically changing) cardiac pumping function and to therapeutic measures on an immediate time basis); and
- 93350 (Echocardiography, transthoracic, real-time with image documentation (2D), with or without M-mode recording, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report).

After reviewing HCPCS code-specific median costs, we determined that all echocardiography procedures that may be performed with contrast agents are reasonably similar both clinically and in terms of resource use. In CY 2008, we created APC 0128 (Echocardiogram with Contrast) to provide payment for echocardiography procedures that are performed with a contrast agent. We refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66643 through 66646) for more information on this methodology.

In order for hospitals to identify and receive appropriate payment for echocardiography procedures performed with contrast beginning in CY 2008, we created eight new HCPCS codes (C8921

through C8928) that corresponded to the related CPT echocardiography codes and assigned them to the newly created APC 0128. We instructed hospitals to report the CPT codes when performing echocardiography procedures without contrast and to report the new HCPCS C-codes when performing echocardiography procedures with contrast, or without contrast followed by with contrast. As is our standard policy with regard to new codes, the APC assignment of these codes was then open to comment in that final rule.

We used the same process to calculate median costs for these codes for CY 2009 as we used for CY 2008 to separately identify echocardiography services provided with contrast and those provided without contrast because the data reported under these new codes were not yet available for CY 2009 ratesetting.

In addition, for CY 2009, the American Medical Association (AMA) revised several CPT codes in the 93000 series to more specifically describe particular services provided during echocardiography procedures. The CY 2009 descriptor for new CPT code 93306 (Echocardiography, transthoracic real-time with image documentation (2D), includes M-mode recording, when performed, complete, with spectral Doppler echocardiography, and with color flow Doppler echocardiography) includes the services described in CY 2008 by three CPT codes: 93307; 93320 (Doppler echocardiography, pulsed wave and/or continuous wave with spectral display; complete); and 93325 (Doppler echocardiography color flow velocity mapping). Therefore, the service described in CY 2009 by new CPT code 93306 was reported in the CY 2008 data with three CPT codes, specifically CPT codes 93307, 93320, and 93325. In CY 2008, the hospital received separate payment for CPT code 93307 through APC 0269 (Level II Echocardiogram without Contrast Except Transesophageal), into which payment for the other two services was packaged. The revised CY 2009 descriptor of CPT code 93307 explicitly excludes services described by CPT codes 93320 and 93325.

To estimate the hospital costs of CPT codes 93306 and 93307 based on their CY 2009 descriptors and the corresponding HCPCS codes C8929 and C8923 for CY 2009, we used claims data from CY 2007. As described in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68542 through 68544), we manipulated our CY 2007 single and "pseudo" single claims data to simulate the new CY 2009 definitions of these services. Specifically, we

selected claims for CPT code 93307 on which CPT codes 93320 and 93325 were also present and we treated the summed costs on these claims as if they were a single procedure claim for CPT code 93306. Similarly, we selected single claims for CPT code 93307 to reflect the newly revised descriptor for CY 2009; that is, we included those claims where CPT code 93307 was not billed with packaged CPT code 93320 or CPT code 93325 on the same claim. We then applied our CY 2009 methodology for calculating HCPCS code-specific median costs for these echocardiography procedures with and without contrast by dividing the new set of claims for CPT codes 93306 and 93307 into those billed with and without contrast agents. We assigned the costs for simulated CPT codes 93306 and 93307 reported without contrast to those CPT codes. We then assigned the costs for simulated CPT codes 93306 and 93307 reported with contrast to new HCPCS code C8929 (Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, complete, with spectral Doppler echocardiography, and with color flow Doppler echocardiography) and revised HCPCS code C8923 (Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, complete, without spectral or color Doppler echocardiography), respectively. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68542 through 68544), we assigned these CPT and HCPCS codes to APCs for CY 2009 based on their simulated median costs and clinical characteristics. New CY 2009 CPT code 93306 and HCPCS code C8929 were assigned comment indicator "NI" in that final rule with comment period, to signify that they were new codes whose interim final OPPS treatment was open to comment on that final rule with comment period.

The CY 2010 OPPS/ASC proposed rule was the first opportunity to have claims data available from hospitals for echocardiography services performed with contrast (or without contrast followed by with contrast) and reported with HCPCS codes C8921 through C8928. With the exception of HCPCS code C8923, which had a significant change in its code descriptor for CY 2009, in the CY 2010 OPPS/ASC proposed rule (74 FR 35271), we proposed to use our standard

methodology to set the CY 2010 OPPS payment rates for these echocardiography services performed with contrast, taking into consideration their HCPCS code-specific median costs from CY 2008 claims.

For CY 2010 ratesetting, we proposed to employ an alternative ratesetting methodology for CPT codes 93306 and 93307 and HCPCS codes C8929 and C8923 that is similar to the approach we used for CY 2009 in order to account for the new codes and revised code descriptors for which CY 2008 data are unavailable. However, in the case of the proposed CY 2010 cost estimation, our CY 2008 claims for CPT code 93307 were only for services performed without contrast, and we have CY 2008 claims for HCPCS C8923 for the comparable services performed with contrast. Specifically, we selected claims for CPT code 93307 on which CPT codes 93320 and 93325 were also present and we treated the summed costs on these claims as if they were a single procedure claim for CPT code 93306 in order to simulate the median cost for CPT code 93306, for which CY 2008 claims data are not available. We then selected single claims for CPT code 93307 to reflect the newly revised descriptor for CY 2009; that is, we included those claims where CPT code 93307 was not billed with either packaged CPT code 93320 or CPT code 93325 on the same claim in order to simulate an appropriate CY 2010 proposed median cost for CPT code 93307. We assigned the costs of HCPCS code C8923 when reported with CPT codes 93320 and 93325 to HCPCS code C8929 and the costs of HCPCS code C8923 when reported without CPT code 93320 or 93325 to HCPCS code C8923.

Following publication of the CY 2009 OPPS/ASC final rule with comment period, several stakeholders brought a number of concerns to our attention, including the interim APC assignment of new CPT code 93351 (Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision) and the corresponding new HCPCS code C8930 (Transthoracic echocardiography, with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using

treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision). These stakeholders noted that new CY 2009 CPT code 93351 was created to include the services reported previously by CPT codes 93015 (Cardiovascular stress test using maximal or submaximal treadmill or bicycle exercise, continuous electrocardiographic monitoring, and/or pharmacological stress; with physician supervision, with interpretation and report) and 93350 (Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report). Because new CY 2009 CPT code 93351 was meant to include the services previously reported with both the CPT codes for a transthoracic echocardiogram during rest and stress (CPT code 93350 is recognized under the OPPS) and a cardiovascular stress test (CPT code 93017 is recognized under the OPPS, rather than CPT code 93015), these stakeholders disagreed with our assignments of both CPT codes 93350 and 93351 to APC 0269 for CY 2009.

Upon review of these concerns and our CY 2008 data, in the CY 2010 OPPS/ASC proposed rule (74 FR 35271), we proposed for CY 2010 to use an alternative methodology to simulate median costs for CPT code 93351 and corresponding HCPCS code C8930, for which CY 2008 claims data are unavailable, and for CPT code 93350 and corresponding HCPCS code C8928 (Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report). That is, we proposed to use claims that contain both CPT codes 93350 and 93017 (Cardiovascular stress test using maximal or submaximal treadmill or bicycle exercise, continuous electrocardiographic monitoring, and/or pharmacological stress; tracing only, without interpretation and report) to simulate the median cost for CPT code 93351. We also proposed to use the remaining claims that contain CPT code 93350 but that do not contain CPT code 93017 to develop the proposed CY 2010 median cost for CPT code 93350. For

our proposed rule analysis, we identified over 74,000 CY 2008 claims with both CPT code 93350 and CPT code 93017 on the same date of service and no other separately paid services appearing on the same date after applying our bypass processing logic, discussed in section II.A.1.b. of the proposed rule (74 FR 35240 through 35241). We treated these modified claims containing both CPT codes 93350 and 93017 as a single service and we calculated a proposed median cost of approximately \$604. Therefore, for CY 2010, we proposed to reassign CPT code 93351 to revised APC 0270 (Level III Echocardiogram without Contrast), which had a proposed APC median cost of approximately \$596. We proposed to continue to assign CPT code 93350 to APC 0269, which had a proposed APC median cost of approximately \$456, based on its proposed HCPCS code-specific median cost of approximately \$406 based on approximately 11,000 single claims. Furthermore, we proposed to use claims for HCPCS code C8928 that are reported with CPT code 93017 on the same claim to simulate the CY 2010 median cost for HCPCS code C8930. We identified over 4,000 claims in the proposed rule data with both HCPCS code C8930 and CPT code 93017 on the same date of service and no other separately paid services appearing on the same date after applying our bypass processing logic, discussed in section II.A.1.b. of the proposed rule (74 FR 35240 through 35241), that we modified to treat HCPCS code C8930 and CPT code 93017 as a single service. We calculated a HCPCS code-specific proposed median cost of approximately \$706. Therefore, we proposed to continue to assign HCPCS code C8930 to APC 0128 with a proposed APC median cost of approximately \$660. We also proposed to continue to assign HCPCS code C8928 to APC 0128, based on its HCPCS code-specific proposed median cost of approximately \$595 based on approximately 1,000 single claims.

Comment: One commenter on the CY 2009 OPPS/ASC final rule with comment period addressed the interim final treatment of new CPT code 93306 for CY 2009. The commenter requested that CMS not recognize CPT code 93306 under the OPPS because this code represents the combination of three services already described by existing CPT codes 93307, 93320, and 93325. Alternatively, the commenter recommended that CMS could instruct hospitals to continue billing CPT codes 93320 and 93325 in association with CPT code 93306 in order to encourage

consistent reporting of services described by CPT codes 93320 and 93325 when they are furnished with any echocardiography service. The commenter believed that requiring the use of CPT code 93306 may confuse hospitals, as other echocardiography services require the separate reporting of CPT codes 93320 and 93325 when these additional procedures are performed. Because there are already existing codes for the services described by CPT code 93306 and hospitals could inappropriately stop reporting CPT codes 93320 and 93325 in association with other echocardiography services, the commenter requested that CMS not recognize CPT code 93306 for payment under the OPPS. According to the commenter, under all circumstances, hospitals would continue to report CPT code 93320 or CPT code 93325 when they are performed with any echocardiography procedure, a practice preferred by the commenter. Similarly, the commenter recommended that CMS not recognize the corresponding HCPCS code C8929 that describes CPT code 93306 when furnished with contrast because the contrast echocardiography procedure could also be reported using existing HCPCS code C8921 and CPT codes 93320 and 93325.

Response: As is our standard methodology, we review new CPT codes annually and assign status indicators to all new codes and provide APC assignments, if applicable, for codes that describe services that may be performed in the hospital outpatient department (which includes provider-based clinics located on and off campus). We consider CPT code 93306 to be part of the standard CPT code set hospitals use for reporting services under the OPPS, and the service described by the code is one that we believe could be furnished to a hospital outpatient and potentially covered and, therefore, paid by Medicare under the OPPS. We incorporated CPT code 93306 in the CY 2009 OPPS/ASC final rule with comment period, assigning it a separately payable status indicator and APC, consistent with our belief that the service described by this code could be appropriately reported by hospitals when they furnish the service in the HOPD. Furthermore, as described in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68543), we used a special cost estimation methodology to estimate the expected cost of CPT code 93306 based on hospital claims data for the individual predecessor codes in order to inform our interim final assignment of CPT code 93306 to a clinical APC for CY 2009.

Regarding the commenter's alternative suggestion that we instruct hospitals to continue to report CPT codes 93320 and 93325 when performed in association with the procedure described by CPT code 93306, we will not instruct hospitals to continue to report CPT codes 93320 and 93325 when billing for CPT code 93306 because CPT code 93306 incorporates the services described by CPT codes 93320 and 93325 in its code descriptor. Billing separately for these services when reporting CPT 93306 would not be consistent with correct coding principles and could create greater confusion and unnecessary burden for hospitals. Whenever possible, hospitals have repeatedly encouraged us to follow standard coding guidelines in order to reduce their administrative burden in reporting services differently for Medicare, and our recognition of CPT code 93306 for payment under the OPSS is consistent with hospitals' general request to us.

Finally, as we are continuing to instruct hospitals to use CPT code 93306 for CY 2010, it continues to be appropriate for hospitals to bill using HCPCS code C8929 when furnishing the service described by CPT code 93306 with contrast. In the case of CPT code 93306 and other CPT codes for echocardiography services, we have developed parallel HCPCS C-codes to report each procedure when furnished with contrast in order to provide payment through separate APCs for those echocardiography services furnished with and without contrast.

Comment: Several commenters on the CY 2010 OPSS/ASC proposed rule expressed support for the revisions to the echocardiography APCs included in the CY 2010 OPSS/ASC proposed rule. One commenter noted appreciation for the proposed reassignment of CPT code 93351 from APC 0269 to APC 0270. However, one commenter on the CY 2009 OPSS/ASC final rule with comment period suggested that the new CY 2009 CPT code 93351 should not be recognized for payment under the OPSS. The commenter reasoned that the comprehensive service described by CPT code 93351 is comprised of two services previously reported with CPT codes 93350 and 93015: CPT code 93015 includes physician supervision and interpretation, which are not hospital outpatient services; and CPT code 93015 is reported by nonhospital practitioners and is not recognized for payment under the OPSS.

In addition, a commenter on the CY 2010 OPSS/ASC proposed rule stated that a more appropriate treatment of CPT code 93351 under the OPSS would

be to not recognize this code for payment under the OPSS but, rather, to continue to recognize for payment several existing CPT codes which, when reported in combination, would describe the service that would otherwise be reported with CPT code 93351 alone. The commenter believed that CPT code 93351 was created specifically for services performed in nonfacility settings and that the intent of the CPT Editorial Committee was to limit the use of the code to nonfacility settings only. The commenter stated that correspondence from CMS indicated that CPT code 93351 would be billable only when provided in a physician's office or independent laboratory settings.

Response: As is our standard methodology, we review new CPT codes annually and assign status indicators to all new codes and provide APC assignments, if applicable, for codes that describe services that may be performed in the HOPD (which includes provider-based clinics located on and off campus). The CPT code descriptor for CPT code 93351 makes no mention that the code is restricted from use in the HOPD, or that its use is limited to nonfacility settings. Further, there are no additional CPT instructions that would limit the reporting of CPT code 93351 to nonfacility or nonhospital settings. We consider this CPT code to be part of the standard CPT code set hospitals use for reporting services under the OPSS, and the service described by the code is one that we believe could be furnished to a hospital outpatient and potentially covered and, therefore, paid by Medicare under the OPSS. CPT code 93351 describes a service that would previously have been reported with CPT codes 93350 and 93017 under the OPSS. While the commenter was correct that we do not recognize CPT code 93015 for payment under the OPSS, a code that describes a cardiovascular stress test with interpretation and report, we do recognize CPT code 93017, which describes the tracing only for the cardiovascular stress test. We incorporated CPT code 93351 in the CY 2009 OPSS/ASC final rule with comment period, assigning it a separately payable status indicator and APC, consistent with our belief that the service described by this code could be appropriately reported by hospitals when they furnish the service in the HOPD. Furthermore, we established professional component (PC) and technical component (TC) payments under the MPFS for CPT code 93351, also consistent with our belief that the

CPT code may be reported for services in facility settings, such as independent laboratory settings. We have communicated no information to the public that states that Medicare hospital outpatient payment would not be made if this CPT code were reported by a hospital for services furnished to hospital outpatients.

We proposed a methodology for identifying the hospital outpatient claims and isolating the hospital charges that would be associated with this procedure for CY 2010 in order to develop an appropriate hospital outpatient payment for the associated facility resources for the existing services that would be reported and paid under the new CPT code. Specifically, we proposed to use claims that contain both CPT codes 93350 and 93017 to simulate the median cost for CPT code 93351 and proposed to reassign CPT code 93351 from APC 0269 to revised APC 0270 for CY 2010 based on its simulated median cost. We continue to believe that this CPT code may be reported for OPSS services described by the code, and that our proposed CY 2010 cost estimation methodology accurately simulates a median cost for this new code that reflects the associated hospital resources for the component services that are newly described by this single CPT code.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to reassign CPT code 93351 to APC 0270 based on a simulated CPT-specific median cost identified from over 80,000 CY 2008 claims with both CPT code 93350 and CPT code 93017 on the same date of service and no other separately paid services appearing on the same date after applying our bypass processing logic, as discussed above. We calculated a final CPT-specific median cost of approximately \$605 for CPT code 93351 and a final CY 2010 APC median cost for APC 0270 of approximately \$591.

Comment: One commenter on the CY 2009 OPSS/ASC final rule with comment period requested that CMS delete HCPCS code C8930 (Transthoracic echocardiography, with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision) as the services

described by this code could be reported using CPT code 93017 (Cardiovascular stress test using maximal or submaximal treadmill or bicycle exercise, continuous electrocardiographic monitoring, and/or pharmacological stress; tracing only, without interpretation and report) and HCPCS code C8928 (Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report).

Response: As described above, we are continuing to recognize the service described by CPT code 93351, which is the noncontrast echocardiography procedure that is parallel to HCPCS code C8930 for the same procedures provided with contrast. As previously noted, we have developed parallel HCPCS C-codes to report each echocardiography procedure when furnished with contrast in order to provide payment through separate APCs for those echocardiography services furnished with and without contrast. While we understand that the service reported under HCPCS code C8930 may be reported using a combination of a CPT code and a HCPCS C-code, we do not believe that this would be appropriate because the noncontrast echocardiography service is reported with a single CPT code. Hospitals are generally instructed to use the HCPCS code that most appropriately and specifically describes the service that was provided, including not unbundling component services that could otherwise be separately reported. In this instance, HCPCS code C8930 would be the most specific code that describes the full service provided when the component services that would otherwise be reported by CPT code 93017 and HCPCS code C8928 are provided together. Our CY 2010 ratesetting methodology for HCPCS code C8928 is based on claims data and specifically excludes those cases when the service was furnished along with the procedure described by CPT code 93017. On the other hand, our CY 2010 ratesetting methodology for HCPCS code C8930 specifically includes cases where the services described by HCPCS code C8928 and CPT code 93017 were provided together. In that way, we are able to base CY 2010 payment for all of these services on their actual or simulated hospital costs in the context

of the CPT and HCPCS C-codes that will be reported in CY 2010.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to recognize HCPCS code C8930 for OPSS payment. For CY 2010, HCPCS code C8930 continues to be assigned to APC 0128, with a final CY 2010 APC median cost of approximately \$645.

Comment: One commenter on the CY 2009 OPSS/ASC final rule with comment period requested that CMS not recognize CPT code 93352 (Use of echocardiographic contrast agent during stress echocardiography), as the OPSS has already developed Level II HCPCS C-codes to identify echocardiography procedures performed with contrast.

Response: During our review of CPT code 93352 for the CY 2009 OPSS/ASC final rule with comment period, we assigned an interim final status indicator "M" (Not paid under the OPSS) to CPT code 93352 for CY 2009. In our CY 2010 OPSS/ASC proposed rule, we proposed to continue this status indicator assignment for CY 2010.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to assign status indicator "M" to CPT code 93352.

Comment: A few commenters requested that CPT code 93318 (Echocardiography, transesophageal (TEE) for monitoring purposes, including probe placement, real time 2-dimensional image acquisition and interpretation leading to ongoing (continuous) assessment of (dynamically changing) cardiac pumping function and to therapeutic measures on an immediate time basis) not be reassigned to APC 0269 as proposed for CY 2010. Instead, these commenters requested that CPT code 93318 continue to be assigned to APC 0270 for CY 2010. Commenters stated that CPT code 93318 is clinically similar to CPT code 93312 (Echocardiography, transesophageal, real time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report), and because CPT code 93312 is assigned to APC 0270, CPT code 93318 should be assigned to APC 0270 as well. While these commenters noted that the reassignment of CPT code 93318 to APC 0269 would be most consistent with its CPT-specific median cost presented in the proposed rule, they stated that the unexplained volatility in the cost of CPT code 93318 suggests that clinical homogeneity should be the deciding

factor when assigning this service to an APC.

Response: As is our standard process, for the CY 2010 proposed rule, we reviewed each APC for clinical cohesiveness and resource homogeneity. As the commenters noted, we proposed to reassign CPT code 93318 to APC 0269 as we believed that the proposed CPT-specific median cost more closely matched the median cost of APC 0269. While we continue to believe that the CPT-specific median cost of CPT 93318 (approximately \$472) closely resembles the median cost of APC 0269 (approximately \$447), upon further review, we agree with the commenter that the clinical characteristics of the procedure described by CPT code 93318 are similar to the procedure described by CPT code 93312. We also note that we have only 344 single and 593 total claims for CPT code 93318 from only 188 providers in comparison to 29,987 single and 52,342 total claims for CPT code 93312 from 2,093 providers. We believe the limited claims data from relatively few providers contribute to the variability in cost observed for CPT code 93318 and agree with the commenters that this procedure should remain assigned to APC 0270 for CY 2010.

After consideration of the public comments we received, we are not finalizing our proposal to reassign CPT code 93318 to APC 0269. Instead, for CY 2010, we are continuing to assign CPT code 93318 to APC 0270, with a final CY 2010 APC median cost of approximately \$591.

Comment: Several commenters supported the continuation of separate APCs for payment of echocardiography procedures with contrast and without contrast. While these commenters were generally supportive of the proposed ratesetting methodology, they were concerned that the proposed payment rate for APC 0128 of approximately \$683 was insufficient to cover the costs associated with providing the echocardiogram and the related contrast materials and services for HCPCS codes C8921 (Transthoracic echocardiography with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; complete); C8925 (Transthoracic echocardiography (TEE) with contrast, or without contrast followed by with contrast, real time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report); C8926 (Transthoracic echocardiography (TEE) with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; including probe

placement, image acquisition, interpretation and report); and C8930 (Transthoracic echocardiography, with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision). Specifically, the commenters noted that the noncontrast equivalent procedures (described by CPT codes 93303, 93312, 93315, and 93351) were all proposed for assignment to APC 0270, with a proposed payment rate of approximately \$600. The commenters believed that the difference between the proposed payment rate for these procedures with contrast and without contrast is too small to cover the cost of the contrast material used in these procedures. The commenters suggested that CMS reassign HCPCS codes C8921, C8925, C8926, and C8930 to a new APC for echocardiography procedures performed with contrast or that CMS provide separate payment for the contrast material used in these procedures.

Response: The final payment differential between APC 0270, where CPT codes 99303, 99312, 99315, and 99351 are assigned, and APC 0128, where the corresponding HCPCS codes for the same procedures with contrast (HCPCS codes C8921, C8925, C8926, and C8930) are assigned, is the difference between approximately \$645 and approximately \$591 of \$54. We believe this differential provides an appropriate higher payment to those

hospitals that furnish these procedures with contrast and appropriately accounts for the cost of the contrast material, which is required for all of the services assigned to APC 0128. HCPCS codes C8921, C8925, C8926, and C8930 have median costs that range from a low of approximately \$178 to a high of approximately \$712. Each of these HCPCS codes was reported by fewer than 170 providers in CY 2008. The median costs of these services span most of the range of median costs of HCPCS codes assigned to APC 0128, and they do not form a cluster of high cost procedures in the APC such that they would warrant assignment to a new clinical APC. In contrast, the median costs of CPT codes 99303, 99312, 99315, and 99351 span a much narrower range, from a low of approximately \$505 to a high of approximately \$605. Two of these CPT codes were reported by more than 1,500 providers in CY 2008. Clearly, fewer providers are reporting the echocardiogram procedures with contrast, and we expect that the hospital cost distribution for that subset of hospitals could be different than the cost distribution of the large number of providers reporting the procedures without contrast. Therefore, no conclusions can be drawn about the aggregate OPPS payment to that subset of hospitals for all of their echocardiogram services in comparison to the aggregate echocardiogram costs of the subset of hospitals specifically based on the payment rates for APCs 0128 and 0270. The OPPS is a prospective payment system that relies on hospital charge and cost report data from the hospitals that furnish the services in order to determine relative costs. Therefore, we believe that our

prospective payment rates calculated based on the costs of those providers furnishing the procedures in CY 2008 provide appropriate payment to the providers that will furnish the services in CY 2010.

In summary, after consideration of the public comments we received, we are finalizing our CY 2010 proposals for payment of echocardiography procedures with and without contrast, with modifications. We are finalizing our proposed methodologies for simulating the median costs of CPT codes 93306, 93307, 93351, and 93350 for which there are no CY 2008 hospital claims data for these specific CPT codes, as discussed above. In addition, we are finalizing our proposed methodologies for simulating the median costs of HCPCS codes C8929, C8923, C8930, and C8928 for which there are no CY 2008 hospital claims data for these specific HCPCS codes, as discussed above. We are not finalizing our proposal to reassign CPT code 93318 to APC 0369; instead, we are maintaining the assignment of CPT code 93318 to APC 0270 for CY 2010. Finally, we are finalizing our proposal to assign HCPCS codes C8921, C8925, C8926, and C8930 to APC 0128 for CY 2010.

Table 9 below shows CY 2010 CPT codes for billing echocardiography services without contrast, their final APC assignments for CY 2010, and the corresponding HCPCS codes for use when echocardiography services are performed with contrast (or without contrast followed by with contrast), along with their final APC assignments for CY 2010.

BILLING CODE 4120-01-P

**TABLE 9.—CY 2010 OPPS HCPCS CODES FOR BILLING
ECHOCARDIOGRAPHY SERVICES**

Echocardiography Without Contrast			Echocardiography With Contrast		
CY 2010 HCPCS Code	CY 2010 Descriptor	Final CY 2010 APC	CY 2010 HCPCS Code	CY 2010 Descriptor	Final CY 2010 APC
93303	Transthoracic echocardiography for congenital cardiac anomalies; complete	0270	C8921	Transthoracic echocardiography with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; complete	0128
93304	Transthoracic echocardiography for congenital cardiac anomalies; follow-up or limited study	0269	C8922	Transthoracic echocardiography with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; follow-up or limited study	0128
93306	Echocardiography, transthoracic real-time with image documentation (2D), includes M-mode recording, when performed, complete, with spectral Doppler echocardiography, and with color flow Doppler echocardiography	0269	C8929	Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, complete, with spectral Doppler echocardiography, and with color flow Doppler echocardiography	0128

Echocardiography Without Contrast			Echocardiography With Contrast		
93307	Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, complete, without spectral or color Doppler echocardiography	0697	C8923	Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, complete, without spectral or color Doppler echocardiography	0128
93308	Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, follow-up or limited study	0697	C8924	Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, follow-up or limited study	0128
93312	Echocardiography, transesophageal, real time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report	0270	C8925	Transesophageal echocardiography (TEE) with contrast, or without contrast followed by with contrast, real time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report	0128
93313	Echocardiography, transesophageal, real time with image documentation (2D) (with or without M-mode recording); placement of transesophageal probe only	0269	No corresponding C-code		

Echocardiography Without Contrast			Echocardiography With Contrast		
93315	Transesophageal echocardiography for congenital cardiac anomalies; including probe placement, image acquisition, interpretation and report	0270	C8926	Transesophageal echocardiography (TEE) with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; including probe placement, image acquisition, interpretation and report	0128
93316	Transesophageal echocardiography for congenital cardiac anomalies; placement of transesophageal probe only	0270	No corresponding C-code		
93318	Echocardiography, transesophageal (TEE) for monitoring purposes, including probe placement, real time 2-dimensional image acquisition and interpretation leading to ongoing (continuous) assessment of (dynamically changing) cardiac pumping function and to therapeutic measures on an immediate time basis	0270	C8927	Transesophageal echocardiography (TEE) with contrast, or without contrast followed by with contrast, for monitoring purposes, including probe placement, real time 2-dimensional image acquisition and interpretation leading to ongoing (continuous) assessment of (dynamically changing) cardiac pumping function and to therapeutic measures on an immediate time basis.	0128

Echocardiography Without Contrast			Echocardiography With Contrast		
93350	Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report	0269	C8928	Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report	0128
93351	Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision	0270	C8930	Transthoracic echocardiography, with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision	0128

BILLING CODE 4120-01-C

Finally, in the CY 2010 OPPS/ASC proposed rule (74 FR 35275), for CY 2010, based upon our proposed APC configurations, we also proposed to revise the titles of our existing series of echocardiography APCs to more accurately describe the groups of services identified by CPT codes 93303

through 93352 and HCPCS codes C8921 through C8930 that are assigned to these APCs. We proposed to rename APCs 0269, 0270, and 0697 as described in Table 7 of the proposed rule.

Comment: One commenter supported the proposed revisions to the echocardiography APC titles and configurations.

Response: We appreciate the support for our proposal.

We are finalizing our proposal to rename APCs 0269, 0270, and 0697 without modification. Therefore, we are adopting as final the titles of these APCs as reflected in Table 10 below:

TABLE 10—CY 2010 ECHOCARDIOGRAPHY APCs

Final CY 2010 APC	CY 2010 APC title	Final CY 2010 approximate APC median cost
0128	Echocardiogram With Contrast	\$645
0269	Level II Echocardiogram Without Contrast	447

TABLE 10—CY 2010 ECHOCARDIOGRAPHY APCs—Continued

Final CY 2010 APC	CY 2010 APC title	Final CY 2010 approximate APC median cost
0270	Level III Echocardiogram Without Contrast	591
0697	Level I Echocardiogram Without Contrast	262

(5) Nuclear Medicine Services

In CY 2008, we began packaging payment for diagnostic radiopharmaceuticals into the payment for the associated nuclear medicine procedure. (For a discussion regarding the distinction between diagnostic and therapeutic radiopharmaceuticals, we refer readers to the CY 2008 OPPTS/ASC final rule with comment period at 72 FR 66636.) Prior to the implementation of this policy, diagnostic radiopharmaceuticals were subject to the standard OPPTS drug packaging methodology whereby payments are packaged when the estimated mean per day product costs fall at or below the annual packaging threshold for drugs, biologicals (other than implantable biologicals), and radiopharmaceuticals.

Packaging costs into a single aggregate payment for a service, encounter, or episode-of-care is a fundamental principle that distinguishes a prospective payment system from a fee schedule. In general, packaging the costs of supportive items and services into the payment for the independent procedure or service with which they are associated encourages hospital efficiencies and also enables hospitals to manage their resources with maximum flexibility. All nuclear medicine procedures require the use of at least one radiopharmaceutical or other radiolabeled product, and there are only a small number of radiopharmaceuticals that may be appropriately billed with each diagnostic nuclear medicine procedure. For the OPPTS, we distinguish diagnostic radiopharmaceuticals from therapeutic radiopharmaceuticals for payment purposes, and this distinction is recognized in the Level II HCPCS codes for diagnostic radiopharmaceuticals that include the term “diagnostic” along with a radiopharmaceutical in their HCPCS code descriptors. As we stated in the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66635), we believe that our policy to package payment for diagnostic radiopharmaceuticals (other than those already packaged when their per day costs are below the packaging threshold for OPPTS drugs, biologicals, and radiopharmaceuticals) is consistent with

OPPTS packaging principles, provides greater administrative simplicity for hospitals, and encourages hospitals to use the most clinically appropriate and cost efficient diagnostic radiopharmaceutical for each study. For more background on this policy, we refer readers to discussions in the CY 2008 OPPTS/ASC proposed rule (72 FR 42667 through 42672) and the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66635 through 66641).

For CY 2008 ratesetting, we used only claims for nuclear medicine procedures that contained a diagnostic radiopharmaceutical in calculating the median costs for APCs that include nuclear medicine procedures (72 FR 66639). This is similar to the established methodology used for device-dependent APCs before claims reflecting the procedure-to-device edits were included in our claims data. For CY 2008, we also implemented claims processing edits (called procedure-to-radiolabeled product edits) requiring the presence of a radiopharmaceutical (or other radiolabeled product) HCPCS code when a separately payable nuclear medicine procedure is present on a claim. Similar to our practice regarding the procedure-to-device edits that have been in place for some time, we continually review comments and requests for changes related to these edits and, based on our review, may update the edit list during our quarterly update process if necessary. The radiolabeled product and procedure HCPCS codes that are included in these edits can be viewed on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/01_overview.asp.

The CY 2008 OPPTS claims that are subject to the procedure-to-radiolabeled product edits were not available for setting payment rates in CY 2009. Therefore, as described in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68545), we continued to use our established CY 2008 methodology for setting the payment rates for APCs that included nuclear medicine procedures for CY 2009. We used an updated list of radiolabeled products, including but not limited to diagnostic radiopharmaceuticals, from

the procedure-to-radiolabeled product edit file to identify single and “pseudo” single claims for nuclear medicine procedures that also included at least one eligible radiolabeled product. Using this subset of claims, we followed our standard OPPTS ratesetting methodology to calculate median costs for nuclear medicine procedures and their associated APCs. As in CY 2008, when we set APC median costs based on single and “pseudo” single claims that also included at least one radiolabeled product on our edit file, we observed an equivalent or higher median cost than that calculated from all single and “pseudo” single bills. We believe that this methodology appropriately ensured that the costs of diagnostic radiopharmaceuticals were included in the CY 2009 ratesetting process for these APCs.

As discussed in section II.A.4.b.(1) of the proposed rule (74 FR 35287) and this final rule with comment period, during the September 2007 APC Panel meeting, the APC Panel requested that CMS evaluate the impact of expanded packaging on beneficiaries. Also, during the March 2008 APC Panel meeting, the APC Panel requested that CMS report to the APC Panel at the first meeting in CY 2009 the impact of packaging on net payments for patient care. In response to these requests, we shared data with the APC Panel at the February 2009 APC Panel meeting that compared the frequency of the billing of diagnostic radiopharmaceuticals billed under the OPPTS in CY 2007, before the packaging of all diagnostic radiopharmaceuticals went into effect, to the frequency of the billing of those same products in CY 2008, their first year of packaged payment. We also reviewed information about the aggregate payment for diagnostic radiopharmaceuticals and nuclear medicine procedures during those same 2 years. A summary of these data analyses is provided in section II.A.4.b.(1) of this final rule with comment period.

In addition to these aggregate analyses of total frequency and payment, we also presented our analyses of the number of hospitals performing nuclear medicine scans and the specific diagnostic radiopharmaceuticals appearing with

cardiac and tumor imaging nuclear medicine procedures, excluding positron emission tomography (PET) scans, by classes of hospitals between the CY 2007 claims processed through September 30, 2007 and the CY 2008 claims processed through September 30, 2008. At the March 2008 APC Panel meeting, the APC Panel also recommended that we evaluate the usage and frequency, geographic distribution, and size and type of hospitals performing nuclear medicine studies using radioisotopes to assess beneficiaries' access and that we present these analyses at the first APC Panel meeting in CY 2009. The number of all hospitals reporting any nuclear medicine procedure declined by 2 percent between the CY 2007 claims data and the CY 2008 claims data. Across several classes of hospitals (urban and rural, teaching and nonteaching, and small and large OPPS service volume), the number of hospitals billing any nuclear medicine procedure declined by up to 4 percent over that same time period. With regard to the specific diagnostic radiopharmaceuticals reported with cardiac and tumor imaging nuclear medicine procedure, we generally observed comparable distributions of radiopharmaceuticals between the CY 2007 claims data and the CY 2008 claims data. However, the utility of this analysis was limited due to the introduction of the procedure-to-radiolabeled product claims processing edits discussed above. There are nuclear medicine procedures reported with a diagnostic radiopharmaceutical HCPCS code on the CY 2008 claims that would have not necessarily been billed with a diagnostic radiopharmaceutical HCPCS code on the CY 2007 claims. Specifically, we observed an increase in billing for many radiopharmaceuticals, some new and costly, between the CY 2007 claims data and the CY 2008 claims data. We do not know how much of this was attributable to changes in hospitals' use of radiopharmaceuticals or to the CY 2008 introduction of the procedure-to-radiolabeled product edits that require a radiolabeled product on the claim for payment of the nuclear medicine procedure. With the exception of the notable increases in the frequencies of certain radiopharmaceutical HCPCS codes that potentially resulted from the introduction of these edits, in general, hospital billing patterns for diagnostic radiopharmaceuticals associated with cardiac and tumor imaging nuclear medicine scans did not change dramatically between CY 2007 and CY

2008 for all hospitals and classes of hospitals. We concluded that very few hospitals stopped providing nuclear medicine procedures as a result of our CY 2008 policy to package payment for diagnostic radiopharmaceuticals and that, in general, hospitals did not decrease their use of expensive radiopharmaceuticals.

As a result of the discussions of the APC Panel following our presentation of the analyses of the impact of packaging payment for all diagnostic radiopharmaceuticals in the OPPS, the APC Panel further recommended that CMS continue to analyze the impact on beneficiaries of increased packaging of diagnostic radiopharmaceuticals and provide more detailed analyses at the next APC Panel meeting. Further, the APC Panel requested that, in the more detailed analyses of packaging of diagnostic radiopharmaceuticals by type of nuclear medicine scan, CMS analyze the data according to the specific CPT codes billed with the diagnostic radiopharmaceuticals. We stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35277) that we are accepting the APC Panel's recommendation and would provide additional data to the APC Panel at an upcoming meeting. We did not share additional data related to diagnostic radiopharmaceuticals with the APC Panel at the most recent August 2009 meeting because we believe the APC Panel's discussions would benefit from analyses of an additional year of claims data after CY 2008. Therefore, we plan to incorporate analysis of CY 2009 claims into the information we will present to the APC Panel for its review at the winter 2010 meeting.

At the February 2009 meeting of the APC Panel, the Panel commended CMS for its effort to date to tailor the resource-based APC system to facilitate appropriate payment for diagnostic and therapeutic radiopharmaceuticals. The APC Panel recommended that CMS continue its dialogue with professional societies, vendors, and other stakeholders to improve the accuracy of APC payments for these complex items and services, including consideration of developing composite APCs. We appreciate the support of the APC Panel, and we are accepting the APC Panel's recommendation to continue to communicate with interested stakeholders regarding payment for radiopharmaceuticals and the associated procedures. We regularly accept meetings from interested parties throughout the year, and we encourage stakeholders to continue a dialogue with us during the rulemaking cycle and throughout the year. Our response to the APC Panel's recommendation regarding

composite APCs is included in our response to the public comments summarized below.

For CY 2010 ratesetting, we are able to use CY 2008 OPPS claims that were subject to the procedure-to-radiolabeled product claims processing edits incorporated into the I/OCE prior to payment of claims in order to develop single and "pseudo" single claims for nuclear medicine procedures according to our standard methodology. We believe that using the CY 2008 claims data for these services without further editing for the presence of a radiolabeled product is now appropriate for CY 2010 because these claims reflect all possible relationships between the nuclear medicine procedures and their associated radiolabeled products that we have accommodated for payment of nuclear medicine procedures. Moreover, as we indicated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68548 through 68549), in the rare circumstance where a diagnostic radiopharmaceutical is not provided in association with a nuclear medicine procedure, for example, because a beneficiary receives a therapeutic radiopharmaceutical as part of a hospital inpatient stay and then returns to the HOPD for a nuclear medicine scan without needing a diagnostic radiopharmaceutical to be administered again for the study, we believe it is appropriate to use these claims for ratesetting purposes. We believe that just as these situations are representative of the performance of a nuclear medicine scan, it is also appropriate to include them for ratesetting purposes.

Comment: A number of commenters opposed CMS' proposed policy to package payment for all diagnostic radiopharmaceuticals into payment for their associated nuclear medicine procedures. They noted that the majority of diagnostic radiopharmaceuticals are not interchangeable and, for that reason, CMS' policy of packaging payment for all diagnostic radiopharmaceuticals into their associated nuclear medicine procedures does not foster hospital efficiencies. Some commenters expressed concern that packaging diagnostic radiopharmaceuticals into payment for associated nuclear medicine procedures results in overpayment of many procedures, especially those using existing low-cost radiopharmaceuticals, while the bundled payment would be insufficient for newer, and likely more expensive, radiopharmaceuticals.

In addition, the commenters requested that, if CMS continues to

package payment for diagnostic radiopharmaceuticals into payment for their associated nuclear medicine procedures. CMS revise the nuclear medicine APCs to provide differential payments for nuclear medicine procedures when used with different radiopharmaceuticals. Several commenters identified the series of tumor/infection imaging APCs, including APCs 0406 (Level I Tumor/Infection Imaging), 0408 (Level III Tumor/Infection Imaging), and 0414 (Level II Tumor/Infection Imaging), for CMS' attention to ensure appropriate payment for low volume, high cost radiopharmaceuticals. One commenter specifically suggested a composite APC for certain combinations of a tumor imaging scan and specific diagnostic radiopharmaceuticals.

Several commenters noted that there is wide variation in the costs of diagnostic radiopharmaceuticals, and that composite APCs for specific combinations of procedures and diagnostic radiopharmaceuticals would be necessary to ensure adequate payment to hospitals using expensive diagnostic radiopharmaceuticals. Other commenters suggested that the significant clinical and resource diversity of radiopharmaceuticals packaged into nuclear imaging procedures amounted to a violation of the 2 times rule. The commenters explained that, just as diagnostic radiopharmaceuticals are not interchangeable, certain radiopharmaceuticals are indicated for particular types of diseases, such as cancer, and are not clinically similar to other radiopharmaceuticals used for other purposes, such as tumor imaging.

Response: As we discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68547), we understand that the selection of a diagnostic radiopharmaceutical for a particular nuclear medicine procedure is a complex decision based on many factors, including patient-specific factors, and that not every diagnostic radiopharmaceutical is fully interchangeable with others. However, as stated in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66617) and in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68546), we believe that nonspecific packaging (as opposed to selected code packaging) based on combinations of items and services observed on hospital claims is fully appropriate because of the myriad combinations of items and services that can be appropriately provided together. Under the OPPS, we package payment for ancillary, supportive, and interrelated items and

services into payment for the independent services they accompany. As we discuss in section II.A.4. of this final rule with comment period, packaging promotes hospital efficiencies through numerous means, not only just through the choice of which radiopharmaceutical to use for a specific nuclear medicine scan. While all diagnostic radiopharmaceuticals may not be interchangeable, we believe that packaging the costs of diagnostic radiopharmaceuticals, however differential those costs may be, into the payment for nuclear medicine services that use these products is appropriate, whether there is one product or multiple products that could be used to furnish the particular service provided to an individual patient. The OPPS has a history of packaging items that are not necessarily interchangeable. It is our longstanding practice to package payment for nonpass-through implantable medical devices into payment for the procedure in which they are used, notwithstanding that there may be different devices or combinations of devices that could be used to furnish a service. (For a more complete discussion of the history of packaging items, we refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66639).) Therefore, in accordance with our understanding that a diagnostic radiopharmaceutical is never provided without an accompanying nuclear medicine scan, we believe that it is appropriate to package the payment for all diagnostic radiopharmaceuticals into the payment for the associated nuclear medicine procedure.

With regard to suggested composites or other revisions designed to isolate specific nuclear medicine scans with a subset of diagnostic radiopharmaceuticals, as we discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68546), we do not believe that the inability to substitute one diagnostic radiopharmaceutical for another is a compelling reason for creating composite APCs, as explained below. We developed composite APCs to provide a single payment for two or more services that are typically performed together during a single clinical encounter and that result in the provision of a complete service. Composite APCs differ from packaging. Composite APCs provide a single payment for specific combinations of independent services that would otherwise be separately payable if they were not provided together, while packaging entails associating the cost of

ancillary, supportive, and interrelated services and supplies with a distinct service or composite service. Composite APCs are intended to expand the OPPS payment bundles to encourage hospital efficiencies. Providing a single payment for a specific combination of a diagnostic radiopharmaceutical with a particular nuclear medicine procedure would not constitute a composite APC and would provide no incentives for hospital efficiency. Specifically, a diagnostic radiopharmaceutical would never be separately payable under the OPPS when furnished alone, so the combination of a diagnostic radiopharmaceutical and a nuclear medicine procedure would not meet the definition of a composite APC as described above. From the perspective of value-based purchasing, we see no benefit to paying for many individual diagnostic radiopharmaceutical and nuclear medicine procedure combinations over paying separately for both the item and service, beyond an appearance of bundling. Such an approach would add complexity to ratesetting and would create challenges and cost instability because payments would be based on data from small numbers of claims for certain HCPCS code pairs. As noted above, there are many items and services that we package under the OPPS that are similarly not interchangeable with other related items and services. Therefore, we are not accepting the APC Panel's recommendation to explore developing composite APCs for diagnostic radiopharmaceuticals and nuclear medicine procedures.

We understand that, by packaging payment for a range of products such as diagnostic radiopharmaceuticals, payment for the associated nuclear medicine procedure may be more or less than the hospital's cost for these services in a given case. As stated in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66639) and the CY 2009 OPPS/ASC final rule with comment period (73 FR 68546), we note that a fundamental characteristic of a prospective payment system is that payment is to be set at an average for the service which, by definition, means that some services are paid more or less than the average.

We discussed in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66640) and the CY 2009 OPPS/ASC final rule with comment period (73 FR 68546) the issue of variability in radiopharmaceutical costs or other packaged costs creating potential 2 times violations. We note that 2 times violations are specific to the total cost of the primary service, nuclear medicine

scans in this case, including packaged costs. We have performed our standard review of the APCs using updated CY 2008 claims data for this final rule with comment period and, as a result, have not identified any 2 times violations in the APCs containing nuclear medicine procedures, when calculated as described above. (For more information on the 2 times rule, we refer readers to sections III.B.2. and III.B.3. of this final rule with comment period.)

Comment: Several commenters expressed concern that CMS was relying on edits in the claims processing system in order to identify those claims that would be used for CY 2010 ratesetting purposes. These commenters suggested that CMS continue to require a diagnostic radiopharmaceutical in order to use a nuclear medicine claim for ratesetting purposes for at least another 2 years in order to ensure that the claims editing process is working properly and that all hospital costs are reflected in the median costs of nuclear medicine procedures.

One commenter noted that CMS' methodology for setting payment rates for nuclear medicine services may be flawed. This commenter contended that CMS should not solely rely on the claims processing edits in order to determine which claims are to be used for ratesetting purposes. The commenter suggested that, even though CMS is using claims that have passed the nuclear medicine-to-radiolabeled product edits, CMS' ratesetting methodology may exclude the cost of diagnostic radiopharmaceuticals when calculating median costs for associated nuclear medicine procedures. Specifically, the commenter stated that the program logic that creates "pseudo" single procedure claims may separate a nuclear medicine scan and the associated diagnostic radiopharmaceutical when the diagnostic radiopharmaceutical appears on a different day and, therefore, CMS would not package the cost of the diagnostic radiopharmaceutical when setting the median cost for the nuclear medicine procedure. The commenter added that CMS' ratesetting methodology for "pseudo" single procedure claims relies on the date of service to identify associated packaged costs. Therefore, the commenter requested that CMS use only single and "pseudo" single nuclear medicine procedure claims that also contain a diagnostic radiopharmaceutical in order to set payment rates for nuclear medicine procedures. More specifically, several commenters requested that CMS not reassign CPT code 78803 (Radiopharmaceutical localization of

tumor or distribution of tomographic agent(s); tomographic (SPECT)) to APC 0414 (Level II Tumor/Infection Imaging) as proposed, but instead assign CPT code 78803 to APC 0408 (Level III Tumor/Infection Imaging). One commenter believed that the use of "pseudo" single procedure claims to calculate payment rates may have neglected to include the cost of the radiopharmaceutical or other scans that may have been performed on other dates of service and reported on other claims.

Response: As we indicated in the CY 2008 OPPTS/ASC final rule with comment period (72 FR 42669), we are aware that several diagnostic radiopharmaceuticals may be used for multiple day studies; that is, a particular diagnostic radiopharmaceutical may be administered on one day and a related diagnostic nuclear medicine procedure may be performed on a subsequent day. While we understand that multiple-day episodes for diagnostic radiopharmaceuticals and the related diagnostic nuclear medicine procedures occur, we found the occurrence of nuclear medicine scans on a different date of service to be a small proportion of all diagnostic nuclear medicine imaging procedures appearing with the radiopharmaceutical. Specifically, our analysis at that time indicated that, roughly, 15 diagnostic radiopharmaceuticals have a half-life longer than one day such that they could support diagnostic nuclear medicine scans on different days. Excluding the 5 percent of diagnostic radiopharmaceutical claims that had no matching diagnostic nuclear medicine scan for the same beneficiary, we found that a diagnostic nuclear medicine scan was reported on the same day as a coded diagnostic radiopharmaceutical 90 percent or more of the time for 10 of these 15 diagnostic radiopharmaceuticals. Further, we found that between 80 and 90 percent of single bills for each of the remaining 5 diagnostic radiopharmaceuticals had a diagnostic nuclear medicine scan on the same day.

Moreover, as the commenter noted, the potential separation of a diagnostic radiopharmaceutical and the associated nuclear medicine procedure would only be relevant to the "pseudo" single procedure claims. In the "natural" single bills we use for ratesetting, we package costs across dates of service. Overall, in examining the CY 2008 claims data available for this final rule with comment period, we observed that "natural" single claims constituted a majority of all single procedure claims used to calculate median costs for APCs

with nuclear medicine procedures. Further, we acknowledge that we expect to lose packaged costs on a small proportion of claims when we create "pseudo" single procedure claims by splitting claims based on dates of service. This is an inevitable consequence of the "pseudo" single procedure claim creation process. We believe that the tradeoff is a minor one given the significant benefit of additional claims data, and the vast majority of commenters generally supported our "pseudo" single procedure claim methodology. Finally, we note that the nuclear medicine procedure-to-radiolabeled product I/OCE claims processing edits (http://www.cms.hhs.gov/HospitalOutpatientPPS/02_device_procedure.asp) to which the commenters referred include therapeutic radiopharmaceuticals and brachytherapy sources. Claims that pass these claims processing edits and enter into the ratesetting methodology without a diagnostic radiopharmaceutical reported on the claim are factored into ratesetting for nuclear medicine procedures as we do not expect that every nuclear medicine procedure would be billed with a diagnostic radiopharmaceutical, although we do expect each to be billed with a radiolabeled product. We note that the only time that we would not expect a nuclear medicine procedure to be billed with a radiolabeled product on an outpatient claim would be in the rare circumstance where a therapeutic radiopharmaceutical is provided in an inpatient setting and a nuclear medicine procedure associated with this radiopharmaceutical is subsequently furnished in the HOPD. In this specific circumstance, we would expect that hospitals would bill HCPCS code C9898 (Radiolabeled product provided during a hospital inpatient stay) in place of the radiolabeled product. Nuclear medicine scans are sometimes performed after the application of brachytherapy sources or the provision of a therapeutic radiopharmaceutical and in these cases the administration of an additional source of radioactivity (a diagnostic radiopharmaceutical) may not be required. While brachytherapy sources and therapeutic radiopharmaceuticals would be paid separately under the OPPTS, we believe it is appropriate for us to include the costs of the scans that include a brachytherapy source or therapeutic radiopharmaceutical (or where a therapeutic radiopharmaceutical used for the scan was furnished to an inpatient) but lack a diagnostic radiopharmaceutical in

calculating the median cost of the nuclear medicine procedure because these claims represent the hospital costs for the scans furnished under these circumstances. We previously discussed this issue in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68647 through 68648).

We believe that the single and “pseudo” single procedure claims resulting from our standard ratesetting methodology accurately capture the cost of providing nuclear medicine scans under a variety of clinical scenarios for several reasons discussed above and summarized again here. First, previous analyses demonstrated that a significant percentage of nuclear medicine procedures are reported on the same day as diagnostic radiopharmaceuticals with an extended half-life and, in these cases, our ratesetting methodology would capture these diagnostic radiopharmaceutical costs. We acknowledge that diagnostic radiopharmaceuticals with an extended half-life may be administered on a different day than the performance of the accompanying nuclear medicine scan. However, administration of the diagnostic radiopharmaceutical on a different day does not mean that these costs are not captured in our APC median costs for nuclear medicine procedures. The majority of the single procedure claims that we use to estimate APC median cost for APCs with nuclear medicine scans are “natural” single procedure claims that package all identified packaged costs (including diagnostic radiopharmaceuticals) into the nuclear medicine procedures, irrespective of the dates of service. While our standard ratesetting methodology also relies on “pseudo” single procedure claims that, by definition, represent only a single service date and potentially eliminate the cost of a packaged diagnostic radiopharmaceutical with an extended half-life billed on a different date of service than the nuclear medicine scan, the potential to ignore packaged costs on other dates of service is true for all procedures for which we use “pseudo” single procedure claims in ratesetting. This small loss of packaging is a tradeoff in adopting our methodology for breaking down multiple procedure claims through the bypass process, as discussed in section II.A.1.b. of this final rule with comment period. Finally, not all claims for nuclear medicine procedures should include a diagnostic radiopharmaceutical because they may include another type of radiolabeled product (such as a brachytherapy source or therapeutic radiopharmaceutical),

and these additional radiolabeled products are not packaged. In short, we believe that, overall, the single procedure claims for nuclear medicine scans, both “natural” and “pseudo” single procedure claims, together appropriately represent the full cost of providing various nuclear medicine procedures and result in accurate APC median costs. Therefore, our standard OPPTS ratesetting methodology of using median costs calculated from claims data according to our standard methodology from those claims that passed the I/OCE claims processing edits adequately captures the costs of diagnostic radiopharmaceuticals associated with diagnostic nuclear medicine procedures that are not provided on the same date of service.

Specifically with regard to our proposed reassignment of CPT code 78803, with a CPT code-specific median cost of approximately \$561, to APC 0414, with an APC median cost of approximately \$506, we note that we have almost 3,000 single claims upon which the median cost of CPT code 78803 is based. This CPT code-specific median cost is significantly lower than the median cost of APC 0408 of approximately \$954, the APC assignment requested by the commenters and the highest level APC in the tumor/infection imaging series. Therefore, we believe the most appropriate CY 2010 APC assignment for CPT code 78803 is APC 0414, as we proposed for CY 2010. As stated above, we believe that our standard ratesetting methodology adequately incorporates the packaged diagnostic radiopharmaceutical costs associated with nuclear medicine procedures, including the procedure described by CPT code 78803.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to package the costs of all diagnostic radiopharmaceuticals into payment for the associated nuclear medicine procedures utilizing our standard OPPTS ratesetting methodology that is applied to claims that passed the nuclear medicine procedure-to-radiolabeled product I/OCE claims processing edits in CY 2008. We also are finalizing our CY 2010 proposal, without modification, to reassign CPT code 78003 to APC 0414, with an APC median cost of approximately \$506.

Comment: A number of commenters cited concerns regarding the proposed APC assignments and proposed payment rates for a number of nuclear medicine procedures. These commenters believed that the proposed

APC assignments of certain nuclear medicine procedures led to clinically diverse procedures being grouped together for payment purposes.

Specifically, one commenter requested that: (1) CPT code 78645 (Cerebrospinal fluid flow, imaging (not including introduction of material); shunt evaluation) be reassigned from APC 0403 (Level I Nervous System Imaging) to APC 0402 (Level II Nervous System Imaging); (2) CPT code 78608 (Brain imaging, positron emission tomography (PET); metabolic evaluation) be reassigned from APC 0308 (Non-Myocardial Positron Emission Tomography (PET) Imaging) to a more appropriate APC; and (3) CPT codes 78000 (Thyroid uptake; single determination) and 78001 (Thyroid uptake; multiple determinations) be reassigned from APC 0389 (Level I Non-imaging Nuclear Medicine) to APC 0392 (Level II Non-Imaging Nuclear Medicine).

Response: We have performed our annual review of all the procedures and APC groupings for this final rule with comment period based on updated CY 2008 claims data. The CPT code-specific median cost of CPT code 78645 is approximately \$246 based on 434 single claims, which is reasonably close to the median cost of APC 0403 of approximately \$195, where we proposed to assign the service. The commenter recommended assignment of CPT code 78645 to APC 0402, in the same nervous system imaging series, with a significantly higher APC median cost of approximately \$573. Based on this review of the costs and clinical characteristics of other services assigned to these nervous system imaging APCs, we continue to believe CPT code 78645 is most appropriately assigned to APC 0403 as we proposed.

There is a single APC for nonmyocardial PET scans, APC 0308, with an APC median cost of approximately \$1,028. The median costs of all CPT codes assigned to that APC, including CPT codes for positron emission tomography (PET) scans and PET/computed tomography (CT) scans and CPT code 78608 for a metabolic evaluation of the brain using PET range from approximately \$849 to \$1,093, demonstrating very significant resource similarity across all of these procedures. Therefore, we do not agree with the commenter that the proposed configuration of APC 0308 should be modified because all of these nonmyocardial services that use PET technology demonstrate very similar costs and share clinical similarity as well.

With regard to the thyroid scans described by CPT codes 78000 and 78001, these procedures have CPT code-specific median costs of approximately \$91 and \$121 based on 1,167 and 982 single claims, respectively. The CPT code-specific median costs of these two procedures are very close to the median cost of APC 0389 of approximately \$112, where we proposed to assign them for CY 2010. CPT codes 78000 and 78001 are the only services assigned to this APC with significant volume, and the APC median cost is mostly a reflection of the costs of procedures reported with two codes. In contrast, the median cost of APC 0392, their recommended placement according to the commenter, is approximately \$179, substantially greater than the median costs of the two thyroid studies. Furthermore, if we were to reassign CPT codes 78000 and 78001 to APC 0392 as the commenter suggested, the median cost of APC 0392 would decrease to reflect the costs of these two procedures because, based on number of single claims for CPT codes 78000 and 78001, their costs would significantly affect the median cost of the APC. Therefore, we do not believe any changes to the proposed APC assignments of CPT codes 78000 or 78001 are justified.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to assign CPT code 78645 to APC 0403, CPT code 79608 to APC 0308, CPT code 78000 to APC 0389, and CPT code 78001 to APC 0389. The approximate APC median costs of these APCs are as follows: APC 0403 at \$195; APC 0308 at \$1,028; and APC 0389 at \$112.

Comment: A few commenters requested that CMS not reassign CPT code 78807 (Radiopharmaceutical localization of inflammatory process; tomographic (SPECT)) to APC 0406 (Level I Tumor/Infection Imaging) as proposed. These commenters noted that CPT code 78807 is more clinically similar to CPT codes 78805 (Radiopharmaceutical localization of inflammatory process; limited area) and 78806 (Radiopharmaceutical localization of inflammatory process; whole body) that are assigned to APC 0414. Therefore, the commenters requested that CMS continue to assign CPT code 78807 to APC 0414 for CY 2010.

Response: We proposed to assign CPT code 78807, with a CPT code-specific median cost of approximately \$371 based on 251 single claims, to APC 0406, with an APC median cost of approximately \$287. The significant individual services included in APC

0406 have a range of median costs, from approximately \$232 to approximately \$371. APC 0406 includes a number of tumor or infection imaging nuclear medicine procedures. Comparatively, APC 0414, where the commenters requested that we assign CPT code 78807, has an APC median cost of approximately \$506 and includes significant services with CPT code-specific median costs from approximately \$382 to approximately \$561. CPT codes 78805 and 78806 are both assigned to APC 0414 and have CPT code-specific median costs of approximately \$477 and \$538, respectively, significantly higher than the median cost of CPT code 78807. Therefore, we do not believe that there is a reason to assign CPT code 78807 to APC 0414, which principally includes services with significantly higher median costs than CPT code 78807. We note that CPT code 78807 is a SPECT scan to localize an inflammatory process, while the other two codes do not describe services that use SPECT technology. Therefore, we do not believe that CPT code 78807 is sufficiently similar to CPT codes 78805 and 78806 from clinical or resource perspectives to warrant assignment to the mid-level tumor/infection imaging APC along with the other two services.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to assign CPT code 78807 to APC 0406, with an APC median cost of approximately \$287.

Comment: Several commenters requested that CMS: (1) Not reassign CPT code 78610 (Brain imaging, vascular flow only) to APC 0403 as proposed but instead assign CPT code 78610 to APC 0402; (2) not reassign CPT code 78601 (Brain imaging, less than 4 static views; with vascular flow) to APC 0402 as proposed but instead assign CPT code 78601 to APC 0403; and (3) not reassign CPT code 78003 (Radiopharmaceutical localization of tumor or distribution of radiopharmaceutical agent(s); tomographic (SPECT)) to APC 0389 as proposed but instead assign CPT code 78003 to APC 0392.

Response: We proposed to assign CPT code 78610, with a CPT-specific median cost of approximately \$211, to APC 0403, with an APC median cost of approximately \$195. The significant services included in APC 0403 have a range of median costs, from approximately \$156 to approximately \$246. Comparatively, APC 0402, where the commenters requested that we assign CPT code 78610, has an APC median cost of approximately \$573 and

includes significant services with CPT code-specific median costs from approximately \$540 to approximately \$587. We do not believe that reassignment of CPT code 78610 to APC 0402 would be appropriate, given the procedure's relatively low median cost, although we recognize that we have few claims for the procedures. We continue to believe that payment for the resources required to provide CPT code 78610 is appropriately reflected through the procedure's assignment to APC 0403.

We proposed to assign CPT code 78601, with a CPT code-specific median cost of approximately \$436, to APC 0402 with an APC median cost of approximately \$573. The significant services included in APC 0402 have a range of median costs from approximately \$540 to approximately \$587. Comparatively, APC 0403, where the commenters requested that we assign CPT code 78601, has an APC median cost of approximately \$195 and includes significant services with CPT code-specific median costs ranging from approximately \$156 to approximately \$246. Although we have few claims for CPT code 78601, we continue to believe it is most appropriately assigned to APC 0402 for CY 2010.

We proposed to assign CPT code 78003, with a CPT code-specific median cost of approximately \$82, to APC 0389 with an APC median cost of approximately \$112. There are two services included in APC 0389 that have a significant volume, CPT codes 78000 and 78001. These two CPT codes both have higher CPT code-specific median costs than CPT code 78003, approximately \$91 and \$121, respectively. Comparatively, APC 0392, where the commenters requested that we assign CPT code 78003, has an APC median cost of approximately \$179. Based on its median cost, we continue to believe that the resources required for CPT code 78003 are appropriately reflected through its assignment to APC 0389.

Comment: A few commenters expressed their appreciation that the CY 2010 OPPS/ASC proposed rule included a proposed increase in payment for PET services compared to CY 2009 payment rates. These commenters also noted their concerns that hospital claims data for PET services are not predictable and that volatile data over the last several years may limit access to PET services. Some commenters urged CMS to use external data when setting payment rates for these services, while others suggested that CMS continue to monitor data to ensure that payment for these services is sufficient to cover the hospital costs for these resources.

Response: As we stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68547), while we utilized external data in the early years of the OPPS for ratesetting for a few services, we now rely on the cost data from claims as the system has matured and we have gained additional experience in ratesetting for HOPD services. The foundation of a system of relative weights like the OPPS is the relativity of the costs of all services to one another, as derived from a standardized system that uses standardized inputs and a consistent methodology. Further, the OPPS is a prospective payment system that relies on hospital charges and cost report data from the hospitals that furnish the services in order to determine relative costs. Therefore, we believe that our prospective payment rates, calculated based on the costs of those providers furnishing the procedures in CY 2008, provide appropriate payment to the providers who will furnish the services in CY 2010. We continue to believe that this standard ratesetting methodology accurately provides payment for PET services provided to hospital outpatients.

In summary, after consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, for the configuration of nuclear medicine APCs. The final CY 2010 median costs for these APCs, as proposed, are calculated according to the standard OPPS ratesetting methodology as applied to claims for nuclear medicine procedures that passed the CY 2008 nuclear medicine procedure-to-radiolabeled product I/OCE claims processing edits. These edits ensure that the claims that are taken through our standard ratesetting process, as described in section II.A.2.b. of this final rule with comment period, that incorporates the creation of "natural" single and "pseudo" single claims, include the radiolabeled product necessary for the performance of the associated nuclear medicine procedure.

(6) Hyperbaric Oxygen Therapy

Since the implementation of the OPPS in August 2000, the OPPS has recognized HCPCS code C1300 (Hyperbaric oxygen under pressure, full body chamber, per 30 minute interval) for hyperbaric oxygen therapy (HBOT) provided in the hospital outpatient setting. In the CY 2005 OPPS final rule with comment period (69 FR 65758 through 65759), we finalized a "per unit" median cost calculation for APC 0659 (Hyperbaric Oxygen) using only claims with multiple units or multiple

occurrences of HCPCS code C1300 because delivery of a typical HBOT service requires more than 30 minutes. We observed that claims with only a single occurrence of the code were anomalies, either because they reflected terminated sessions or because they were incorrectly coded with a single unit. In the same rule, we also established that HBOT would not generally be furnished with additional services that might be packaged under the standard OPPS APC median cost methodology. This enabled us to use claims with multiple units or multiple occurrences. Finally, we also used each hospital's overall CCR to estimate costs for HCPCS code C1300 from billed charges rather than the CCR for the respiratory therapy or other departmental cost centers. The public comments on the CY 2005 OPPS proposed rule effectively demonstrated that hospitals report the costs and charges for HBOT in a wide variety of cost centers. Since CY 2005, we have used this methodology to estimate the median cost for HBOT. The median costs of HBOT using this methodology have been relatively stable for the last 4 years. In the CY 2010 OPPS/ASC proposed rule (74 FR 35277), we proposed to continue using the same methodology to estimate a "per unit" median cost for HCPCS code C1300 for CY 2010 of approximately \$108, using 279,139 claims with multiple units or multiple occurrences.

We did not receive any public comments on our proposal to continue to use our established ratesetting methodology for calculating the median cost of APC 0659 for payment of HBOT. Therefore, we are finalizing, without modification, our CY 2010 proposal to continue to use our established ratesetting methodology for calculating the median cost of APC 0659 for payment of HBOT, with a final CY 2010 median cost of approximately \$106.

(7) Payment for Ancillary Outpatient Services When Patient Expires (-CA Modifier)

In the November 1, 2002 final rule with comment period (67 FR 66798), we discussed the creation of the new HCPCS CA modifier to address situations where a procedure on the OPPS inpatient list must be performed to resuscitate or stabilize a patient (whose status is that of an outpatient) with an emergent, life-threatening condition, and the patient dies before being admitted as an inpatient. In Transmittal A-02-129, issued on January 3, 2003, we instructed hospitals on the use of this modifier. For a complete description of the history of the policy and the development of the

payment methodology for these services, we refer readers to the CY 2007 OPPS/ASC final rule with comment period (71 FR 68157 through 68158).

In the CY 2010 OPPS/ASC proposed rule (74 FR 35277 through 35278), we proposed to continue to use our established ratesetting methodology for calculating the median cost of APC 0375 (Ancillary Outpatient Services When Patient Expires) and to continue to make one payment under APC 0375 for the services that meet the specific conditions for using modifier -CA. We proposed to calculate the relative payment weight for APC 0375 by using all claims reporting a status indicator "C" procedure appended with the -CA modifier, using estimated costs from claims data for line-items with a HCPCS code assigned status indicator "G," "H," "K," "N," "Q1," "Q2," "Q3," "R," "S," "T," "U," "V," and "X" and charges for packaged revenue codes without a HCPCS code. We continue to believe that this methodology results in the most appropriate aggregate median cost for the ancillary services provided in these unusual clinical situations.

We believe that hospitals are reporting the -CA modifier according to the policy initially established in CY 2003. We note that the claims frequency for APC 0375 has been relatively stable over the past few years. Although the median cost for APC 0375 has increased, the median in the CY 2008 data used for development of rates for CY 2010 was only slightly higher than that for CY 2009. Variation in the median cost for APC 0375 is expected because of the small number of claims and because the specific cases are grouped by the presence of the -CA modifier appended to an inpatient procedure and not according to the standard APC criteria of clinical and resource homogeneity. Cost variation for APC 0375 from year to year is anticipated and acceptable as long as hospitals continue judicious reporting of the -CA modifier. Table 8 of the proposed rule (74 FR 35278) showed the number of claims and the proposed median costs for APC 0375 for CYs 2007, 2008, and 2009. For CY 2010, we proposed a median cost for APC 0375 of approximately \$5,784.

We did not receive any public comments regarding this proposal. Therefore, we are finalizing our CY 2010 proposal, without modification, to continue to use our established ratesetting methodology for calculating the median cost of APC 0375, which has a final CY 2010 APC median cost of approximately \$5,911.

Table 11 below shows the number of claims and the final median cost for APC 0375 from CY 2007 to CY 2010.

TABLE 11—CLAIMS FOR ANCILLARY OUTPATIENT SERVICES WHEN PATIENT EXPIRES (-CA MODIFIER) FOR CYs 2007 TO 2010

Prospective payment year	Number of claims	APC median cost
CY 2007	260	\$3,549
CY 2008	183	4,945
CY 2009	168	5,545
CY 2010	182	5,911

e. Calculation of Composite APC Criteria-Based Median Costs

As discussed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66613), we believe it is important that the OPSS enhance incentives for hospitals to provide only necessary, high quality care and to provide that care as efficiently as possible. For CY 2008, we developed composite APCs to provide a single payment for groups of services that are typically performed together during a single clinical encounter and that result in the provision of a complete service. Combining payment for multiple independent services into a single OPSS payment in this way enables hospitals to manage their resources with maximum flexibility by monitoring and adjusting the volume and efficiency of services themselves. An additional advantage to the composite APC model is that we can use data from correctly coded multiple procedure claims to calculate payment rates for the specified combinations of services, rather than relying upon single procedure claims which may be low in volume and/or incorrectly coded. Under the OPSS, we currently have composite APC policies for extended assessment and management services, low dose rate (LDR) prostate brachytherapy, cardiac electrophysiologic evaluation and ablation services, mental health services, and multiple imaging services. We refer readers to the CY 2008 OPSS/ASC final rule with comment period for a full discussion of the development of the composite APC methodology (72 FR 66611 through 66614 and 66650 through 66652).

While we continue to consider the development and implementation of larger payment bundles, such as composite APCs (a long-term policy objective for the OPSS), and continue to explore other areas where this payment model may be utilized, in the CY 2010 OPSS/ASC proposed rule, we did not propose any new composite APCs for CY 2010 so that we may monitor the

effects of the existing composite APCs on utilization and payment. In response to our CY 2009 proposal to apply a composite payment methodology to multiple imaging procedures provided on the same date of service, several public commenters stated that we should proceed cautiously as we expand service bundling. They commented that we should not implement additional composite methodologies until adequate data are available to evaluate the composite policies' effectiveness and impact on beneficiary access to care (73 FR 68561 through 68562).

In response to the concerns of the public commenters and the APC Panel, in the CY 2010 OPSS proposed rule (74 FR 35278 through 35279) we reviewed the CY 2008 claims data for claims processed through September 30, 2008, for the services in the following composite APCs: APC 8000 (Cardiac Electrophysiologic Evaluation and Ablation Composite); APC 8001 (Low Dose Rate Prostate Brachytherapy Composite); APC 8002 (Level I Extended Assessment and Evaluation Composite); and APC 8003 (Level II Extended Assessment and Evaluation Composite). Our analyses did not consider inflation, changes in beneficiary population, or other comparable variables that can affect changes in aggregate payment from year to year. We found that the average payment for the package of services in both APC 8000 and APC 8001 increased from CY 2007, when payments were made for all individual services, to CY 2008 under the composite payment methodology. We also noted that the proposed median costs for these composite APCs for CY 2010 were higher than the median costs upon which the CY 2009 payments were based. We believe that, in part, this is because we used more claims data for common clinical scenarios to calculate the median costs of these APCs than we were able to use prior to the implementation of the composite payment methodology.

With regard to APCs 8002 and 8003, we compared payment for all visits appearing with observation services in CY 2007 with payments for all visits appearing with observation services in CY 2008 and found that total payment for visits and observation services increased from approximately \$197 million to \$270 million for claims processed through September 30 in each year. We attribute this increase in payments, in part, to the introduction of a composite payment for visits and observation through the extended assessment and management composite methodology that occurred for CY 2008 and that did not incorporate the International Classification of Diseases, Ninth Edition, Clinical Modification (ICD-9-CM) diagnosis criteria previously necessary for separate payment of observation.

At its February 2009 meeting, the APC Panel recommended that CMS evaluate the implications of creating composite APCs for cardiac resynchronization therapy (CRT) services with a defibrillator or pacemaker and report its findings to the APC Panel. The APC Panel also recommended at its August 2009 meeting that CMS reconsider creating a new composite APC or group of composite APCs for CRT procedures. While we did not propose any new composite APCs for CY 2010, we are accepting both of these APC Panel recommendations. We will reconsider creating composite APCs for CRT services and evaluate the implications of such a potential policy change, and report our findings to the APC Panel at a future meeting. We also will consider bringing other potential composite APCs to the APC Panel for further discussion.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35279), we proposed to continue for CY 2010 our established composite APC policies for extended assessment and management, LDR prostate brachytherapy, cardiac electrophysiologic evaluation and ablation, mental health services, and multiple imaging services, as discussed

in sections II.A.2.e.(1), II.A.2.e.(2), II.A.2.e.(3), II.A.2.e.(4), and II.A.2.e.(5), respectively, of this final rule with comment period.

Comment: Several commenters supported the development and implementation of the composite APC methodology, remarking that it is consistent with the principles of a prospective payment system and provides more appropriate payment rates through the use of multiple procedure claims for certain services. Many of these commenters also supported CMS' decision to monitor the existing composite APCs' effects on beneficiary access, utilization, and payment for at least another year before implementing additional composite APCs.

Other commenters, however, expressed disappointment that CMS did not propose additional composite APCs for CY 2010 in order to improve OPSS payment accuracy and include more correctly coded, multiple procedure claims in ratesetting. Some commenters recommended the development of composite APCs for nuclear medicine tumor or infection imaging services that encompass multiple days and multiple procedures, with separate payment for the associated diagnostic radiopharmaceuticals.

In addition, many commenters supported the development of composite APCs for CRT with defibrillator (CRT-D) or pacemaker (CRT-P) implantation. They indicated that the procedures involved in the implantation of CRT-D and CRT-P are separately payable services that, if coded correctly, are always represented by the submission of two CPT codes. According to the commenters, the number of single procedure CRT claims available for CY 2010 ratesetting is very low compared to the total number of claims submitted for CRT-D and CRT-P procedures. They argued that the establishment of a composite APC methodology for CRT-D and CRT-P would greatly increase the number of claims used in ratesetting, thereby lessening the year-to-year fluctuations in payment rates for CRT. The commenters also stated that the APC Panel advised CMS to use its discretion in forming one or a group of composite APCs for CRT without the need to report back to the APC Panel. They urged CMS to take this advice and move forward with the composite APC methodology for CRT-D and CRT-P for CY 2010.

Response: We appreciate the commenters' support of the composite APC methodology. As stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35279), we will continue to review the

claims data for the impact of all of the composite APCs on payments to hospitals and on services to beneficiaries and will take such data into consideration before proposing or implementing new composite APCs. We recognize the concerns expressed with respect to our CY 2009 proposal by the public commenters that moving ahead too quickly with any nonstandard OPSS payment methodology (even one such as composite APCs that may improve the accuracy of the OPSS payment rates by utilizing more complete claims for common clinical scenarios in ratesetting) could have unintended consequences and requires close monitoring. Because the multiple imaging composite APCs were implemented for the first time in CY 2009, we will not have data available for such monitoring until early CY 2010. Therefore, we continue to believe that it is in the best interest of hospitals and the continuing refinement of the OPSS that we not implement any new composite APC policies for at least one year.

As previously stated, we are accepting the recommendation made by the APC Panel at its August 2009 meeting that we reconsider creating a new composite APC or group of composite APCs for CRT-D and CRT-P procedures. We will evaluate the implications of such a potential policy change and report our findings to the APC Panel at a future meeting. We note that, while the APC Panel did recommend we reconsider creating a new composite APC or group of composite APCs for CRT-D and CRT-P, the Panel did not specify that we should move forward with the composite APC methodology for CRT-D and CRT-P for CY 2010 without first reporting back to the APC Panel, as some commenters indicated. We do not believe it would be appropriate to implement new composite APCs for CRT-D and CRT-P procedures for CY 2010 because neither we nor the public have had the opportunity to evaluate fully all of the implications of such a potential policy change, which may require complex claims processing logic or new claims processing edits and may have significant, unanticipated effects on the payment rates of other services. We also note that the total volume of claims that would qualify for a CRT-P composite APC in particular would be very low; in the past, we have explored composite APCs only for combinations of services that are commonly performed together (73 FR 68551). Because of the complex issues for these procedures with significant device costs, we believe that it is particularly

important that the APC Panel and the public, through the annual rulemaking cycle, have the opportunity to comment on the development of composite APCs for CRT-D and CRT-P procedures.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue our established composite APC policies for extended assessment and management, LDR prostate brachytherapy, cardiac electrophysiologic evaluation and ablation, mental health services, and multiple imaging services, as discussed in sections II.A.2.e.(1), II.A.2.e.(2), II.A.2.e.(3), II.A.2.e.(4), and II.A.2.e.(5), respectively, of this final rule with comment period.

(1) Extended Assessment and Management Composite APCs (APCs 8002 and 8003)

In the CY 2010 OPSS/SC proposed rule (74 FR 35279 through 35280), we proposed to continue to include composite APC 8002 (Level I Extended Assessment and Management Composite) and composite APC 8003 (Level II Extended Assessment and Management Composite) in the OPSS. For CY 2008, we created these two composite APCs to provide payment to hospitals in certain circumstances when extended assessment and management of a patient occur (an extended visit). In most circumstances, observation services are supportive and ancillary to the other services provided to a patient. In the circumstances when observation care is provided in conjunction with a high level visit or direct referral and is an integral part of a patient's extended encounter of care, payment is made for the entire care encounter through one of two composite APCs as appropriate.

As defined for the CY 2008 OPSS, composite APC 8002 describes an encounter for care provided to a patient that includes a high level (Level 5) clinic visit or direct referral for observation services in conjunction with observation services of substantial duration (72 FR 66648 through 66649). Composite APC 8003 describes an encounter for care provided to a patient that includes a high level (Level 4 or 5) Type A emergency department visit, a high level (Level 5) Type B emergency department visit, or critical care services in conjunction with observation services of substantial duration. HCPCS code G0378 (Observation services, per hour) is assigned status indicator "N," signifying that its payment is always packaged. As noted in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66648 through 66649), the Integrated Outpatient Code Editor

(I/OCE) evaluates every claim received to determine if payment through a composite APC is appropriate. If payment through a composite APC is inappropriate, the I/OCE, in conjunction with the OPSS Pricer, determines the appropriate status indicator, APC, and payment for every code on a claim. The specific criteria that must be met for the two extended assessment and management composite APCs to be paid are provided below in the description of the claims that were selected for the calculation of the proposed CY 2010 median costs for these composite APCs. We did not propose to change these criteria for the CY 2010 OPSS.

When we created composite APCs 8002 and 8003 for CY 2008, we retained as general reporting requirements for all observation services those criteria related to physician order and evaluation, documentation, and observation beginning and ending time as listed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66812). These are more general requirements that encourage hospitals to provide medically reasonable and necessary care and help to ensure the proper reporting of observation services on correctly coded hospital claims that reflect the full charges associated with all hospital resources utilized to provide the reported services. We did not propose to change these reporting requirements for the CY 2010 OPSS. However, as discussed below, the APC Panel at its February 2009 meeting requested that CMS issue guidance clarifying the correct method for reporting the starting time for observation services. The APC Panel noted that the descriptions of the start time for observation services located in the Medicare Claims Processing Manual (Pub. 100-4), Chapter 4, sections 290.2.2 through 290.5, cause confusion for hospitals. We accepted this recommendation and issued clarifying guidance in the Claims Processing Manual through Transmittal 1745, Change Request 6492, issued May 22, 2009 and implemented July 6, 2009.

As noted in detail in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66802 through 66805 and 66814), we saw a normal and stable distribution of clinic and emergency department visit levels in the OPSS claims data through CY 2006 available at that time. We stated that we did not expect to see an increase in the proportion of visit claims for high level visits as a result of the new composite APCs adopted for CY 2008. Similarly, we stated that we expected that hospitals would not purposely change their visit guidelines or otherwise

upcode clinic and emergency department visits reported with observation care solely for the purpose of composite payment. As stated in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66648), we expect to carefully monitor any changes in billing practices on a service-specific and hospital-specific level to determine whether there is reason to request that Quality Improvement Organizations (QIOs) review the quality of care furnished, or to request that Benefit Integrity contractors or other contractors review the claims against the medical record.

When we compared total payments for all visits appearing with observation services in CY 2007 to payments in CY 2008, using claims processed through September 30 in CY 2007 and CY 2008, we observed a 37 percent increase in total payments. We believe this increase is, in part, attributable to the expansion of payment under the extended assessment and management composites to all ICD-9-CM diagnoses. To confirm this, we calculated the percentage of visit HCPCS codes billed with HCPCS code G0378 (Observation services, per hour) between CY 2007 and CY 2008 and compared the percentage associated with visit codes included in the extended assessment and management composites in each year. If hospitals had inappropriately changed their visit reporting behavior to maximize payment through the new composite APCs, we would expect to see significant changes in the percentage of visit HCPCS codes included in the composite APCs billed with observation services relative to all other visit HCPCS codes billed with observation services between CY 2007 and CY 2008. We did not observe a sizable increase in the proportion of visit HCPCS codes included in the composite APCs relative to the proportion of all other visit HCPCS codes billed with observation services. For example, the percentage of claims billed with CPT code 99285 (Emergency department visit for the evaluation and management of a patient (Level 5)) and HCPCS code G0378 was 51 percent in the CY 2007 data and 54 percent in the CY 2008 data. Similarly, the percentage of claims billed with CPT code 99284 (Emergency department visit for the evaluation and management of a patient (Level 4)) and HCPCS code G0378 decreased only slightly from 28 percent in the CY 2007 data to 27 percent in the CY 2008 data. We concluded that, although the volume of visits billed with HCPCS code G0378 increased between CY 2007 and CY 2008, the overall pattern of billing visit

levels did not change significantly. We stated that we will continue to carefully monitor any changes in billing practices on a service-specific and hospital-specific level.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35280), we proposed to continue for CY 2010 the extended assessment and management composite APC payment methodology for APCs 8002 and 8003. As stated earlier, we also proposed to continue the general reporting requirements for observation services reported with HCPCS code G0378. We continue to believe that the composite APCs 8002 and 8003 and related policies provide the most appropriate means of paying for these services. We proposed to calculate the median costs for APCs 8002 and 8003 using all single and "pseudo" single procedure claims for CY 2008 that meet the criteria for payment of each composite APC.

Specifically, to calculate the proposed median costs for composite APCs 8002 and 8003, we selected single and "pseudo" single claims that met each of the following criteria:

1. Did not contain a HCPCS code to which we have assigned status indicator "T" that is reported with a date of service 1 day earlier than the date of service associated with HCPCS code G0378. (By selecting these claims from single and "pseudo" single claims, we had already assured that they would not contain a code for a service with status indicator "T" on the same date of service.);

2. Contained 8 or more units of HCPCS code G0378; and
3. Contained one of the following codes:

- In the case of composite APC 8002, HCPCS code G0379 (Direct referral of patient for hospital observation care) on the same date of service as G0378; or CPT code 99205 (Office or other outpatient visit for the evaluation and management of a new patient (Level 5)); or CPT code 99215 (Office or other outpatient visit for the evaluation and management of an established patient (Level 5)) provided on the same date of service or one day before the date of service for HCPCS code G0378. We refer readers to section XII.E. of the CY 2010 OPSS/ASC proposed rule (74 FR 35370 through 35371) and section XII.E. of this final rule with comment period for a full discussion of our proposed revision of the code descriptor for HCPCS code G0379 and the final policy for CY 2010.

- In the case of composite APC 8003, CPT code 99284 (Emergency department visit for the evaluation and management of a patient (Level 4)); CPT code 99285 (Emergency department visit for the

evaluation and management of a patient (Level 5)); CPT code 99291 (Critical care, evaluation and management of the critically ill or critically injured patient; first 30–74 minutes); or HCPCS code G0384 (Level 5 Hospital Emergency Department Visit Provided in a Type B Emergency Department) provided on the same date of service or one day before the date of service for HCPCS code G0378. (As discussed in detail in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68684), we finalized our proposal to add HCPCS code G0384 to the eligibility criteria for composite APC 8003 for CY 2009.)

We applied the standard packaging and trimming rules to the claims before calculating the proposed CY 2010 median costs. The proposed CY 2010 median cost resulting from this process for composite APC 8002 was approximately \$384, which was calculated from 14,981 single and “pseudo” single bills that met the required criteria. The proposed CY 2010 median cost for composite APC 8003 was approximately \$709, which was calculated from 154,843 single and “pseudo” single bills that met the required criteria. This is the same methodology we used to calculate the medians for composite APCs 8002 and 8003 for the CY 2008 OPPS (72 FR 66649).

As discussed further in section IX. of the CY 2010 OPPS/ASC proposed rule (74 FR 35350) and this final rule with comment period, and consistent with our CY 2008 and CY 2009 final policies, when calculating the median costs for the clinic, Type A emergency department visit, Type B emergency department visit, and critical care APCs (0604 through 0617 and 0626 through 0630), we utilize our methodology that excludes those claims for visits that are eligible for payment through the two extended assessment and management composite APCs, that is APC 8002 or APC 8003. We believe that this approach results in the most accurate cost estimates for APCs 0604 through 0617 and 0626 through 0630 for CY 2010.

At the August 2009 meeting of the APC Panel, the APC Panel recommended that CMS provide the Visits and Observation Subcommittee with an analysis of calendar year 2009 claims data for clinic, ED (Type A and B), and extended assessment and management composite APCs at the next meeting of the APC Panel. The APC Panel also recommended that CMS provide the Visits and Observation Subcommittee with continued analyses of observation services, as previously provided to the APC Panel, including

data on frequency, length of stay, and common diagnoses, as well as recovery audit contractor (RAC) data on these subjects if available. Furthermore, the APC Panel recommended that CMS provide the Visits and Observation Subcommittee with analyses of the most common diagnoses and services associated with Type A and Type B ED visits at the next meeting of the APC Panel, including analysis by hospital-specific characteristics. Finally, the APC Panel recommended that the work of the Visits and Observation Subcommittee continue. We accept all of these recommendations and will present the available requested data at the winter 2010 meeting of the APC Panel.

In summary, in the CY 2010 OPPS/ASC proposed rule (74 FR 35279 through 35280), we proposed to continue to include for CY 2010 composite APC 8002 (Level I Extended Assessment and Management Composite) and composite APC 8003 (Level II Extended Assessment and Management Composite) in the OPPS. We proposed to continue the extended assessment and management composite APC payment methodology and criteria that we finalized for CY 2009. We also proposed to calculate the median costs for APCs 8002 and 8003 using all single and “pseudo” single procedure claims from CY 2008 that meet the criteria for payment of each composite APC. We did not propose to change the reporting requirements for observation services for the CY 2010 OPPS. However, in CY 2009 we did issue further clarifying guidance in the Medicare Claims Processing Manual related to observation start time.

Comment: Several commenters expressed appreciation for CMS’ issuance of clarifying guidance for reporting the beginning and ending times of observation services.

Response: We appreciate these comments and note again that the guidance was issued in the Claims Processing Manual through Transmittal 1745, Change Request 6492, issued May 22, 2009, and implemented July 6, 2009.

Comment: Several commenters requested clarification of the reporting of observation services in relation to maternity care paid under another payer’s policies and in relation to changes in patient status from inpatient to outpatient using Condition Code 44. One commenter pointed out that references to “observation status” versus “inpatient admission” are potentially confusing for beneficiaries and physicians.

Response: Each of these comments/questions is outside of the scope of the proposals in the CY 2010 OPPS/ASC

proposed rule. However, we will consider the possibility of addressing these concerns through other available mechanisms, as appropriate. We note that we have continued to emphasize that observation care is a hospital outpatient service, ordered by a physician and reported with a HCPCS code, like any other outpatient service. It is not a patient status for Medicare purposes.

After consideration of the public comments we received, we are finalizing, without modification, our CY 2010 proposal to continue to include composite APC 8002 and composite APC 8003 in the OPPS and to continue the extended assessment and management composite APC payment methodology and criteria that we finalized for CY 2009. We also are calculating the median costs for APCs 8002 and 8003 using all single and “pseudo” single procedure claims from CY 2008 that meet the criteria for payment of each composite APC. The final CY 2010 median cost resulting from this methodology for composite APC 8002 is approximately \$378, which was calculated from 17,074 single and “pseudo” single bills that met the required criteria. The final CY 2010 median cost for composite APC 8003 is approximately \$699, which was calculated from 176,226 single and “pseudo” single bills that met the required criteria.

(2) Low Dose Rate (LDR) Prostate Brachytherapy Composite APC (APC 8001)

LDR prostate brachytherapy is a treatment for prostate cancer in which hollow needles or catheters are inserted into the prostate, followed by permanent implantation of radioactive sources into the prostate through the needles/catheters. At least two CPT codes are used to report the composite treatment service because there are separate codes that describe placement of the needles/catheters and the application of the brachytherapy sources: CPT code 55875 (Transperineal placement of needles or catheters into prostate for interstitial radioelement application, with or without cystoscopy) and CPT code 77778 (Interstitial radiation source application; complex). Generally, the component services represented by both codes are provided in the same operative session in the same hospital on the same date of service to the Medicare beneficiary being treated with LDR brachytherapy for prostate cancer. As discussed in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66653), OPPS payment rates for CPT code 77778, in

particular, had fluctuated over the years. We were frequently informed by the public that reliance on single procedure claims to set the median costs for these services resulted in use of only incorrectly coded claims for LDR prostate brachytherapy because a correctly coded claim should include, for the same date of service, CPT codes for both needle/catheter placement and application of radiation sources, as well as separately coded imaging and radiation therapy planning services (that is, a multiple procedure claim).

In order to base payment on claims for the most common clinical scenario, and to further our goal of providing payment under the OPPTS for a larger bundle of component services provided in a single hospital encounter, beginning in CY 2008, we provide a single payment for LDR prostate brachytherapy when the composite service, reported as CPT codes 55875 and 77778, is furnished in a single hospital encounter. We base the payment for composite APC 8001 (LDR Prostate Brachytherapy Composite) on the median cost derived from claims for the same date of service that contain both CPT codes 55875 and 77778 and that do not contain other separately paid codes that are not on the bypass list. In uncommon occurrences in which the services are billed individually, hospitals continue to receive separate payments for the individual services. We refer readers to the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66652 through 66655) for a full history of OPPTS payment for LDR prostate brachytherapy and a detailed description of how we developed the LDR prostate brachytherapy composite APC.

In the CY 2010 OPPTS/ASC proposed rule (74 FR 35281), we proposed for CY 2010 to continue paying for LDR prostate brachytherapy services using the composite APC methodology proposed and implemented for CY 2008 and CY 2009. That is, we proposed to use CY 2008 claims on which both CPT codes 55875 and 77778 were billed on the same date of service with no other separately paid procedure codes (other than those on the bypass list) to calculate the payment rate for composite APC 8001. Consistent with our CY 2008 and CY 2009 practice, we proposed not to use the claims that meet these criteria in the calculation of the median costs for APCs 0163 (Level IV Cystourethroscopy and Other Genitourinary Procedures) and 0651 (Complex Interstitial Radiation Source Application), the APCs to which CPT codes 55875 and 77778 are assigned, respectively. The median costs for APCs 0163 and 0651 would continue to be

calculated using single and “pseudo” single procedure claims. We continue to believe that this composite APC contributes to our goal of creating hospital incentives for efficiency and cost containment, while providing hospitals with the most flexibility to manage their resources. We also continue to believe that data from claims reporting both services required for LDR prostate brachytherapy provide the most accurate median cost upon which to base the composite APC payment rate.

Using partial year CY 2008 claims data available for the CY 2010 OPPTS/ASC proposed rule, we were able to use 669 claims that contained both CPT codes 77778 and 55875 to calculate the median cost upon which the proposed CY 2010 payment for composite APC 8001 was based. The proposed median cost for composite APC 8001 for CY 2010 was approximately \$3,106. This was an increase compared to the CY 2009 OPPTS/ASC final rule with comment period in which we calculated a final median cost for this composite APC of approximately \$2,967 based on a full year of CY 2007 claims data. The CY 2010 proposed median cost for this composite APC was slightly less than \$3,268, the sum of the proposed median costs for APCs 0163 and 0651 (\$2,453+\$815), the APCs to which CPT codes 55875 and 77778 map if one service is billed on a claim without the other. We stated in the CY 2010 OPPTS/ASC proposed rule (74 FR 35281) that we believe the proposed CY 2010 median cost for composite APC 8001 of approximately \$3,106, calculated from claims we believe to be correctly coded, would result in a reasonable and appropriate payment rate for this service in CY 2010.

Comment: Several commenters requested changes to the bypass list that could potentially affect the number of claims used to calculate the median costs upon which payments for several APCs involving radiation oncology services, including APC 8001, are based. In particular, some commenters requested CMS add CPT code 77470 (Special treatment procedure (eg, total body irradiation, hemibody radiation, per oral, endocavitary or intraoperative cone irradiation)), CPT code 77328 (Brachytherapy isodose plan; complex (multiplane isodose plan, volume implant calculations, over 10 sources/ribbons used, special spatial reconstruction, remote afterloading brachytherapy, over 12 sources), and CPT code 77295 (Therapeutic radiology simulation-aided field setting; 3-dimensional) to the bypass list in order to utilize more single claims in

calculating the median costs of APC 8001 and other APCs for radiation oncology services. According to one commenter's analysis, the addition of these CPT codes to the bypass list would result in a 17 percent increase in the median cost for APC 8001.

Response: As discussed in detail in section II.A.1.b. of this final rule with comment period, we are not adding CPT codes 77470, 77328, and 77295 to the list of bypass codes for CY 2010 ratesetting, but we are adding several other CPT codes for radiation oncology services. The addition of these codes to the bypass list results in a modest increase in the number of single claims used to calculate the median cost upon which the final payment rate for CY 2010 for APC 8001 is based, but does not result in a significant increase or decrease in the median cost itself.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue paying for LDR prostate brachytherapy services using the composite APC methodology implemented in CY 2008. We were able to use 906 claims that contained both CPT codes 77778 and 55875 to calculate the median cost upon which the CY 2010 final payment for composite APC 8001 is based. The final median cost for composite APC 8001 for CY 2010 is approximately \$3,084. We note that this is slightly less than \$3,303, the approximate sum of the median costs for APC 0163 and APC 0651 (\$2,418 + \$885), the APCs to which CPT codes 55875 and 77778 map if one service is billed on a claim without the other. These CPT codes are assigned status indicator “Q3” in Addendum B to this final rule with comment period to identify their status as potentially payable through a composite APC. Their composite APC assignment is identified in Addendum M to this final rule with comment period.

(3) Cardiac Electrophysiologic Evaluation and Ablation Composite APC (APC 8000)

Cardiac electrophysiologic evaluation and ablation services frequently are performed in varying combinations with one another during a single episode-of-care in the hospital outpatient setting. Therefore, correctly coded claims for these services often include multiple codes for component services that are reported with different CPT codes and that, prior to CY 2008, were always paid separately through different APCs (specifically, APC 0085 (Level II Electrophysiologic Evaluation), APC 0086 (Ablate Heart Dysrhythm Focus),

and APC 0087 (Cardiac Electrophysiologic Recording/Mapping). As a result, there would never be many single bills for cardiac electrophysiologic evaluation and ablation services, and those that are reported as single bills would often represent atypical cases or incorrectly coded claims. As described in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66655 through 66659), the APC Panel and the public expressed persistent concerns regarding the limited and reportedly unrepresentative single bills available for use in calculating the median costs for these services according to our standard OPPS methodology.

Effective January 1, 2008, we established APC 8000 (Cardiac Electrophysiologic Evaluation and Ablation Composite) to pay for a composite service made up of at least one specified electrophysiologic evaluation service and one specified electrophysiologic ablation service. Calculating a composite APC for these services allowed us to utilize many more claims than were available to establish the individual APC median costs for these services, and we also saw this composite APC as an opportunity to advance our stated goal of promoting hospital efficiency through larger payment bundles. In order to calculate the median cost upon which the payment rate for composite APC 8000 is based, we used multiple procedure claims that contained at least one CPT code from group A for evaluation services and at least one CPT code from group B for ablation services reported on the same date of service on an individual claim. Table 9 in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66656) identified the CPT codes that are assigned to groups A and B. For a full discussion of how we identified the group A and group B procedures and established the payment rate for the cardiac electrophysiologic evaluation and ablation composite APC, we refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66655 through 66659). Where a service in group A is furnished on a date of service that is different from the date of service for a code in group B for the same beneficiary, payments are made under

the appropriate single procedure APCs and the composite APC does not apply.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35282), we proposed to continue for CY 2010 to pay for cardiac electrophysiologic evaluation and ablation services using the composite APC methodology proposed and implemented for CY 2008 and CY 2009. Consistent with our CY 2008 and CY 2009 practice, we proposed not to use the claims that meet the composite payment criteria in the calculation of the median costs for APC 0085 and APC 0086, to which the CPT codes in both groups A and B for composite APC 8000 are otherwise assigned. Median costs for APCs 0085 and 0086 would continue to be calculated using single procedure claims. We continue to believe that the composite APC methodology for cardiac electrophysiologic evaluation and ablation services is the most efficient and effective way to use the claims data for the majority of these services and best represents the hospital resources associated with performing the common combinations of these services that are clinically typical. Furthermore, this approach creates incentives for efficiency by providing a single payment for a larger bundle of major procedures when they are performed together, in contrast to continued separate payment for each of the individual procedures.

Using partial year CY 2008 claims data available for the proposed rule, we were able to use 6,975 claims containing a combination of group A and group B codes and calculated a proposed median cost of approximately \$10,105 for composite APC 8000. This was an increase compared to the CY 2009 OPPS/ASC final rule with comment period in which we calculated a final median cost for this composite APC of approximately \$9,206 based on a full year of CY 2007 claims data. We stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35282) that we believe the proposed median cost of \$10,105 calculated from a high volume of correctly coded multiple procedure claims would result in an accurate and appropriate proposed payment for cardiac electrophysiologic evaluation and ablation services when at least one evaluation service is furnished during the same clinical encounter as at least one ablation service. Table 9 of the CY

2010 OPPS/ASC proposed rule (74 FR 35282) listed the groups of procedures upon which we proposed to base composite APC 8000 for CY 2010.

Comment: Several commenters supported CMS' proposal to continue using the composite APCs created in CY 2008, in particular the composite APC for cardiac electrophysiologic evaluation and ablation services. One commenter also supported the modest increase in payment for this APC, stating that it is reflective of the increased costs of providing these important services to patients.

Response: We appreciate commenters' support for the composite payment methodology in general and the composite APC for cardiac electrophysiologic evaluation and ablation in particular.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue paying for cardiac electrophysiologic evaluation and ablation services using the composite APC methodology implemented for CY 2008. For this final rule with comment period, we were able to use 7,599 claims from CY 2008 containing a combination of group A and group B codes and calculated a final median cost of approximately \$10,026 for composite APC 8000. This is an increase compared to the CY 2009 OPPS/ASC final rule with comment period in which we calculated a final median cost of approximately \$9,206 based on a full year of CY 2007 claims data. We believe that the final median cost of \$10,026 calculated from a high volume of correctly coded multiple procedure claims results in an accurate and appropriate final payment for cardiac electrophysiologic evaluation and ablation services when at least one evaluation service is furnished during the same clinical encounter as at least one ablation service. Table 12 below lists the groups of procedures upon which we are basing composite APC 8000 for CY 2010. These CPT codes are assigned status indicated "Q3" in Addendum B to this final rule with comment period to identify their status as potentially payable through a composite APC. Their composite APC assignment is identified in Addendum M to this final rule with comment period.

TABLE 12—GROUPS OF CARDIAC ELECTROPHYSIOLOGIC EVALUATION AND ABLATION PROCEDURES UPON WHICH COMPOSITE APC 8000 IS BASED

Codes used in combinations: At least one in Group A and one in Group B	CY 2010 CPT code	Final single code CY 2010 APC	Final CY 2010 SI (composite)
Group A: Comprehensive electrophysiologic evaluation with right atrial pacing and recording, right ventricular pacing and recording, His bundle recording, including insertion and repositioning of multiple electrode catheters, without induction or attempted induction of arrhythmia	93619	0085	Q3.
Comprehensive electrophysiologic evaluation including insertion and repositioning of multiple electrode catheters with induction or attempted induction of arrhythmia; with right atrial pacing and recording, right ventricular pacing and recording, His bundle recording	93620	0085	Q3.
Group B: Intracardiac catheter ablation of atrioventricular node function, atrioventricular conduction for creation of complete heart block, with or without temporary pacemaker placement	93650	0085	Q3.
Intracardiac catheter ablation of arrhythmogenic focus; for treatment of supraventricular tachycardia by ablation of fast or slow atrioventricular pathways, accessory atrioventricular connections or other atrial foci, singly or in combination	93651	0086	Q3.
Intracardiac catheter ablation of arrhythmogenic focus; for treatment of ventricular tachycardia	93652	0086	Q3.

(4) Mental Health Services Composite APC (APC 0034)

In the CY 2010 OPSS/ASC proposed rule (74 FR 35282 through 35283), we proposed to continue our longstanding policy of limiting the aggregate payment for specified less resource-intensive mental health services furnished on the same date to the payment for a day of partial hospitalization, which we consider to be the most resource-intensive of all outpatient mental health treatment for CY 2010. We refer readers to the April 7, 2000 OPSS final rule with comment period (65 FR 18455) for the initial discussion of this longstanding policy. We stated in the CY 2010 OPSS/ASC proposed rule that we continue to believe that the costs associated with administering a partial hospitalization program represent the most resource-intensive of all outpatient mental health treatment. Therefore, we do not believe that we should pay more for a day of individual mental health services under the OPSS than the partial hospitalization per diem payment.

As discussed in the CY 2010 OPSS/ASC proposed rule (74 FR 35356 through 35357), for CY 2010 we proposed to continue using the two tiered payment approach for partial hospitalization services that we implemented in CY 2009: one APC for days with three services (APC 0172) (Level I Partial Hospitalization (3 services)) and one APC for days with four or more services (APC 0173) (Level II Partial Hospitalization (4 or more services)) (74 FR 35282 through 35283). When a CMHC or hospital provides three units of partial hospitalization services and meets all other partial hospitalization payment criteria, we

proposed that the CMHC or hospital be paid through APC 0172. When the CMHC or hospital provides 4 or more units of partial hospitalization services and meets all other partial hospitalization payment criteria, we proposed that the CMHC or hospital be paid through APC 0173. We proposed to set the CY 2010 payment rate for mental health services composite APC 0034 (Mental Health Services Composite) at the same rate as we proposed for APC 0173, which is the maximum partial hospitalization per diem payment. We stated in the CY 2010 OPSS/ASC proposed rule that we believe this APC payment rate would provide the most appropriate payment for composite APC 0034, taking into consideration the intensity of the mental health services and the differences in the HCPCS codes for mental health services that could be paid through this composite APC compared with the HCPCS codes that could be paid through partial hospitalization APC 0173. When the aggregate payment for specified mental health services provided by one hospital to a single beneficiary on one date of service based on the payment rates associated with the APCs for the individual services exceeds the maximum per diem partial hospitalization payment, we proposed that those specified mental health services would be assigned to APC 0034. We proposed that APC 0034 would continue to have the same payment rate as APC 0173 and that the hospital would continue to be paid one unit of APC 0034. The I/OCE currently determines, and we proposed for CY 2010 that it would continue to determine, whether to pay these

specified mental health services individually or to make a single payment at the same rate as the APC 0173 per diem rate for partial hospitalization for all of the specified mental health services furnished by the hospital on that single date of service.

We also proposed to continue assigning status indicator “Q3” (Codes that May be Paid Through a Composite APC) to the HCPCS codes that are assigned to composite APC 0034 in Addendum M, and to continue assigning status indicator “S” (Significant Procedure, Not Discounted when Multiple), as adopted for CY 2009, to APC 0034 for CY 2010 (74 FR 35283).

Comment: One commenter expressed concern that using claims data from CMHCs and hospitals to calculate the payment rate for APC 0173 would result in reduced access not only for hospital-based partial hospitalization services but also for other less intensive mental health services provided in hospital outpatient departments. The commenter stated that CMS should use hospital data to calculate the payment rates for hospital services.

Response: As discussed in section X. of this final rule with comment period, the final CY 2010 payment rates for APCs 0172 and 0173 are calculated using hospital-only cost data for CY 2010, rather than using both hospital and CMHC cost data. This final policy results in an increase in the median cost for APC 0173 from approximately \$200 in CY 2009 to approximately \$209. As noted in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66739), we continue to believe that the costs associated with administering a partial hospitalization program

represent the most resource intensive of all outpatient mental health treatment, and we do not believe that we should pay more for a day of individual mental health services under the OPPS. The mental health payment limitation will rise and fall in the same manner as payment for partial hospitalization services.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to limit the aggregate payment for specified less intensive outpatient mental health services furnished on the same date by a hospital to the payment for a day of partial hospitalization, specifically APC 0173. For CY 2010, we also are finalizing our proposal, without modification, to assign status indicator "Q3" to those HCPCS codes that describe the specified mental health services to which APC 0034 applies in Addendum B to this final rule with comment period. Lastly, we are finalizing our proposal to continue assigning status indicator "S" (Significant Procedure, Not Discounted When Multiple) to APC 0034.

(5) Multiple Imaging Composite APCs (APCs 8004, 8005, 8006, 8007, and 8008)

Prior to CY 2009, hospitals received a full APC payment for each imaging service on a claim, regardless of how many procedures were performed during a single session using the same imaging modality. Based on extensive data analysis, we determined that this practice neither reflected nor promoted the efficiencies hospitals can achieve when performing multiple imaging procedures during a single session (73 FR 41448 through 41450). As a result of our data analysis, and in response to ongoing recommendations from MedPAC to improve payment accuracy for imaging services under the OPPS, we expanded the composite APC model developed in CY 2008 to multiple imaging services. Effective January 1, 2009, we provide a single payment each time a hospital bills more than one imaging procedure within an imaging family on the same date of service. We utilize three imaging families based on imaging modality for purposes of this methodology: ultrasound, computed tomography (CT) and computed tomographic angiography (CTA), and magnetic resonance imaging (MRI) and magnetic resonance angiography (MRA). The HCPCS codes subject to the multiple imaging composite policy, and their respective families, are listed in Table 8 of the CY 2009 OPPS/ASC final rule with comment period (73 FR 68567 through 68569).

While there are three imaging families, there are five multiple imaging composite APCs due to the statutory requirement at section 1833(t)(2)(G) of the Act that we differentiate payment for OPPS imaging services provided with and without contrast. While the ultrasound procedures included in the policy do not involve contrast, both CT/CTA and MRI/MRA scans can be provided either with or without contrast. The five multiple imaging composite APCs established in CY 2009 are: APC 8004 (Ultrasound Composite); APC 8005 (CT and CTA without Contrast Composite); APC 8006 (CT and CTA with Contrast Composite); APC 8007 (MRI and MRA without Contrast Composite); and APC 8008 (MRI and MRA with Contrast Composite). We define the single imaging session for the "with contrast" composite APCs as having at least one or more imaging procedures from the same family performed with contrast on the same date of service. For example, if the hospital performs an MRI without contrast during the same session as at least one other MRI with contrast, the hospital will receive payment for APC 8008, the "with contrast" composite APC.

Hospitals continue to use the same HCPCS codes to report imaging procedures, and the I/OCE determines when combinations of imaging procedures qualify for composite APC payment or map to standard (sole service) APCs for payment. We make a single payment for those imaging procedures that qualify for composite APC payment, as well as any packaged services furnished on the same date of service. The standard (noncomposite) APC assignments continue to apply for single imaging procedures and multiple imaging procedures performed across families. For a full discussion of the development of the multiple imaging composite APC methodology, we refer readers to the CY 2009 OPPS/ASC final rule with comment period (73 FR 68559 through 68569).

As we discussed in the CY 2010 OPPS/ASC proposed rule (74 FR 35283), during the February 2009 meeting of the APC Panel, the APC Panel heard from stakeholders who claimed that a composite payment is not appropriate when multiple imaging procedures are provided on the same date of service but at different times. Some APC Panel members expressed concern that the same efficiencies that may be gained when multiple imaging procedures are performed during the same sitting may not be gained if a significant amount of time passes between the second and subsequent imaging procedures, when

the patient may leave not only the scanner, but also the radiology department or hospital. The APC Panel recommended that CMS continue to work with stakeholders to examine different options for APCs for multiple imaging sessions and multiple imaging procedures.

We accepted the APC Panel recommendation that CMS continue to work with stakeholders to examine different options for APCs for multiple imaging sessions and multiple imaging procedures. However, as we stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35283 through 35284), we do not believe it is appropriate to make modifications to the multiple imaging composite policy for CY 2010. We indicated that we continue to believe that composite payment is appropriate even when procedures are provided on the same date of service but at different times, because hospitals do not expend the same facility resources each and every time a patient is seen for a distinct imaging service in a separate imaging session. In most cases, we expect that patients in those circumstances would receive imaging procedures at different times during a single prolonged hospital outpatient encounter, and that the efficiencies that may be gained from providing multiple imaging procedures during a single session are achieved in such ways as not having to register the patient again, or not having to re-establish new intravenous access for an additional study when contrast is required. Furthermore, we stated that even if the same level of efficiencies could not be gained for multiple imaging procedures performed on the same date of service but at different times, we expect that any higher costs associated with these cases would be reflected in the claims data and cost reports we use to calculate the median costs for the multiple imaging composite APCs and, therefore, in their payment rates.

In summary, in the CY 2010 OPPS/ASC proposed rule (74 FR 35284), for CY 2010 we proposed to continue paying for all multiple imaging procedures within an imaging family performed on the same date of service using the multiple imaging composite payment methodology, and we proposed no changes from the final CY 2009 policy. The proposed CY 2010 payment rates for the five multiple imaging composite APCs (APC 8004, APC 8005, APC 8006, APC 8007, and APC 8008) were based on median costs calculated from the partial year CY 2008 claims available for the proposed rule that would have qualified for composite payment under the current policy (that

is, those claims with more than one procedure within the same family on a single date of service). To calculate the proposed median costs, we used the same methodology that we used to calculate the final CY 2009 median costs for these composite APCs. That is, we removed any HCPCS codes in the OPPI imaging families that overlapped with codes on our bypass list ("overlap bypass codes") to avoid splitting claims with multiple units or multiple occurrences of codes in an OPPI imaging family into new "pseudo" single claims. The imaging HCPCS codes that we removed from the bypass list for purposes of calculating the proposed multiple imaging composite APC median costs appeared in Table 11 of the CY 2010 OPPI/ASC proposed rule (74 FR 35286). We integrated the identification of imaging composite "single session" claims, that is, claims with multiple imaging procedures within the same family on the same date of service, into the creation of "pseudo" single claims to ensure that claims were split in the "pseudo" single process into accurate reflections of either a composite "single session" imaging service or a standard sole imaging service resource cost. Like all single bills, the new composite "single session" claims were for the same date of service and contained no other separately paid services in order to isolate the session imaging costs. Our last step after processing all claims through the "pseudo" single process was to reassess the remaining multiple procedure claims using the full bypass list and bypass process in order to determine if we could make other "pseudo" single bills. That is, we assessed whether a single separately paid service remained on the claim after removing line-items for the "overlap bypass codes."

We were able to identify 1.7 million "single session" claims out of an estimated 2.5 million potential composite cases from our ratesetting claims data, or well over half of all eligible claims, to calculate the proposed CY 2010 median costs for the multiple imaging composite APCs. The HCPCS codes subject to the proposed multiple imaging composite policy and their respective families were listed in Table 10 of the proposed rule (74 FR 35284 through 35285).

Comment: Many commenters asserted that a single composite APC payment is not appropriate when multiple imaging services of the same modality are provided on the same date of service but at different times. They argued that the same efficiencies that may be gained when multiple imaging procedures are

performed during the same sitting may not be realized if a significant amount of time passes between the first and subsequent imaging procedures, when the patient may have to be repositioned or may have to leave not only the scanner, but also the radiology department or hospital. The commenters stated that, in such cases, facilities must expend equivalent facility resources in each sitting as if the sittings occurred on different dates of service. They noted that not all of these costs are reflected in claims data and, therefore, would not be reflected in the multiple imaging composite APC payment rates. The commenters requested that CMS allow hospitals to report modifier -59 when multiple imaging services of the same modality are provided at different times on the same date of service, and that such cases be excluded from the multiple imaging composite payment methodology. They stated that such an approach is necessary to recognize the provider costs when imaging services must be provided at different sittings due to clinical need or safety requirements. One commenter also asked CMS to work with the AMA to create new CPT codes that describe combined procedures so that providers could use those codes when they provide multiple imaging services in a single session. The commenter argued that utilization of such codes would be easier for providers and would facilitate the capturing of charge data that could be used to create new APCs or payment policies that reflect economies of scale for combined procedures reported through claims data.

Response: We do not agree with the commenters that multiple imaging procedures of the same modality provided on the same date of service but at different times should be exempt from the multiple imaging composite payment methodology. As we indicated in the CY 2010 OPPI/ASC proposed rule (74 FR 35283 through 35284) and the CY 2009 OPPI/ASC final rule with comment period (73 FR 68565), we believe that composite payment is appropriate even when procedures are provided on the same date of service but at different times because hospitals do not expend the same facility resources each and every time a patient is seen for a distinct imaging service in a separate imaging session. In most cases, we expect that patients in these circumstances would receive imaging procedures at different times during a single prolonged hospital outpatient encounter. The efficiencies that may be gained from providing multiple imaging

procedures during a single session are achieved in ways other than merely not having to reposition the patient. For example, a patient who has two MRI procedures 3 hours apart during a single hospital outpatient encounter would not have to be registered again, and hospital staff might not have to explain the procedure in detail prior to the second scan. In the case of multiple procedures involving contrast that are provided at different times during a single hospital outpatient encounter, establishment of new intravenous access for the second study would not be necessary. Even if the same level of efficiencies could not be gained for multiple imaging procedures performed on the same date of service but at different times, we expect that any higher costs associated with these cases would be reflected in the claims data and cost reports we use to calculate the median costs for the multiple imaging composite APCs and, therefore, in the payment rates for the multiple imaging composite APCs. We do not believe it is necessary or appropriate for hospitals to report imaging procedures provided on the same date of service but during different sittings any differently than they would report imaging procedures performed consecutively in one sitting with no time in between the imaging services.

We also do not agree with the commenter that it is necessary to create new CPT codes that describe combined services to ease the burden of hospital billing and improve claims data for ratesetting. As we stated in the CY 2009 OPPI/ASC final rule with comment period (73 FR 68565), certain combination CPT codes, specifically those single codes that describe imaging procedures without contrast and then followed by contrast, already allow for hospitals to report commonly performed combinations of imaging procedures in one anatomic area using a single CPT code. Hospitals can continue to use existing codes to report multiple imaging services by reporting multiple HCPCS codes, and for ratesetting, we use the charges reported to us by hospitals on claims for those multiple imaging services to calculate composite APC payment rates. The I/OCE determines whether composite APC payment applies to a claim, so the composite payment policy creates no additional administrative burden for hospitals.

Comment: Several commenters asserted that the multiple imaging composite payment methodology could have a disproportionately negative effect on cancer centers and trauma units, where patients frequently require more than two imaging services during a

hospital encounter. They argued that the use of a single composite APC payment for an imaging modality regardless of the number of services provided is only appropriate if the underlying claims data used to set the "average" payment rate reflect an average number of services furnished by all providers. According to the commenters, certain providers, such as cancer centers and trauma hospitals, face systematic underpayment of multiple imaging services due to their unique patient population because they routinely provide a greater than average number of imaging services in one sitting or multiple sittings on the same date of service. The commenters stated that, at the same time, all other hospitals experience systematic overpayment.

Response: We do not agree with the commenters that the underlying claims data used to calculate the median costs upon which the payment rates for the multiple imaging composite APCs are inappropriate for payment of all hospitals, or that the multiple imaging composite payment methodology is likely to have a disproportionately negative effect on cancer centers and trauma units. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68562 through 68563), we explored data from CY 2007 claims in response to similar concerns from commenters and a recommendation by the APC Panel at its August 2008 meeting. An analysis of diagnosis codes present on the CY 2007 multiple imaging "single session" claims did show more variability in the number of scans for cancer patients compared to patients with noncancer diagnoses, consistent with commenters' concerns. We observed that, for several of the more commonly reported cancer diagnoses, more than half of the patients received more than two imaging procedures on the same day, while generally lower proportions of patients with noncancer diagnoses received more than two imaging procedures on a single date of service. We did not observe the same pattern for trauma diagnoses. As we stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68562), we do not believe that the higher rate of variability that we observed in the number of scans cancer patients receive was so extreme, however, that the mix of services hospitals provide to patients with diagnoses other than cancer would not balance out higher numbers of scans for cancer patients.

We continue to believe that OPPS hospitals demonstrate sufficient variability in the number of imaging procedures they provide to a single patient on the same day that it is

unlikely any particular class of hospital would experience disproportionate financial effects from the multiple imaging composite payment methodology. For CY 2009, the first year of implementation of the multiple imaging composite APC methodology, the modeled impacts of payment changes by class of hospital due to APC recalibration (where the effects of the multiple imaging composite payment methodology and other APC recalibration would be observed), were very modest across classes of hospital, ranging from -2.5 percent to +1.9 percent (73 FR 68799 through 68800).

The goal of the multiple imaging composite payment methodology is to establish incentives for efficiency through larger payment bundles based on the practice patterns of OPPS hospitals as a whole. We acknowledge that there may be a small number of dedicated cancer centers that, relative to other hospitals paid under the OPPS, may provide a higher proportion of imaging services to cancer patients that involve three or more scans. However, as discussed above, our prior analyses do not lead us to believe that any class of hospitals would experience significantly negative effects from the multiple imaging composite payment methodology. We note that we establish national payment policies for the OPPS and, while certain policies may have greater or lesser impact on individual hospitals, on average we believe that the total OPPS payment to a hospital for all of its services is appropriate. Our modeled estimates of changes in total payment for classes of hospitals between CY 2008 and CY 2009 support this conclusion. We do not believe it would be appropriate to establish national policy based on considerations of the service mix of individual hospitals, or to exclude individual hospitals from national policy because of the impact a specific policy may have on one component of a hospital's operations as a result of a particular hospital's service mix. Furthermore, we note that several cancer centers are held permanently harmless under section 1833(t)(7)(D)(ii) of the Act in order to account for the fact that they may be more costly and have different practice patterns than other hospitals paid under the OPPS.

Comment: Some commenters questioned the adequacy of the proposed multiple imaging composite APC payment rates for sessions involving three or more imaging procedures, and expressed general concern that multiple imaging composite payment methodology would limit beneficiary access to imaging

services. For example, these commenters asserted that the multiple imaging composite payment methodology could create incentives for hospitals to require patients who need more than two imaging procedures to return for additional sittings on other days if the costs for sessions in which more than two procedures are performed far exceed the multiple imaging composite APC payment rates.

Response: As we stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68562), we do not believe that, in aggregate, OPPS payment for multiple imaging services will be inadequate under the multiple imaging composite payment methodology so as to limit beneficiary access, even considering the minority of cases in which hospitals provide more than two imaging procedures on a single date of service. The median costs upon which the payment rates for the multiple imaging composite APCs are based are calculated using CY 2008 claims that would have qualified for composite payment, including those with only two imaging procedures and those with substantially higher numbers of imaging procedures. Payment based on a measure of central tendency is a principle of any prospective payment system. In some individual cases payment exceeds the average cost and in other cases payment is less than the average cost. On balance, however, payment should approximate the relative cost of the average case, recognizing that, as a prospective payment system, the OPPS is a system of averages.

We also do not agree with the commenters that the multiple imaging composite payment methodology will result in hospitals requiring patients who need more than two imaging procedures to return for additional sittings on other days. As we stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68562), we do not believe that, in general, hospitals would routinely and for purposes of financial gain put patients at unnecessary risk of harm from radiation or contrast exposure, or inconvenience them or risk lack of timely followup to the point of making them return to the hospital on separate days to receive medically necessary diagnostic studies. However, we again note that we do have the capacity to examine our claims data for patterns of fragmented care. If we were to find a pattern in which a hospital appears to be fragmenting imaging services across multiple days for individual beneficiaries, we could refer it for review by the Quality Improvement Organizations (QIOs) with

respect to the quality of care furnished, or for review by the Program Safeguard Contractors of claims against the medical record, as appropriate to the circumstances we found.

Comment: Several commenters urged CMS to standardize cost reporting for both advanced imaging procedures and other problematic cost centers before it makes any methodological changes to OPSS payment methodologies, including a composite policy for multiple imaging procedures. One commenter was concerned that observed efficiencies in the multiple imaging composite APC median costs are the result of inaccurate cost report data only and do not reflect true efficiencies from multiple imaging services provided during a single session. According to the commenter, CMS should implement separate cost centers for CT and MRI procedures and the revised revenue code-to-cost center crosswalk, as recommended in the July 2008 report by RTI International (RTI) entitled, "Refining Cost to Charge Ratios for Calculating APC and DRG Relative Payment Weights." The commenter stated that the creation of the new standard cost centers and the adoption of the revised revenue code-to-cost center crosswalk would provide much more accurate charge and cost data for these imaging modalities, and that the true efficiencies associated with providing multiple imaging procedures in a single session may only be discernable once these data are available. The commenter also remarked that the adoption of these changes would result in significant shifts in the underlying CCRs for all APCs, thereby impacting all relative weights and payment rates across all services over time.

Response: We published information regarding the proposed draft hospital cost report CMS-2552-10 in the **Federal Register** on July 2, 2009 and the proposed agency information collection activities were open for a 60-day review and comment period (74 FR 31738). The comment period ended August 31, 2009. The proposed cost report can be viewed at: <http://www.cms.hhs.gov/PaperworkReductionActof1995/PRAL/itemdetail.asp?filterType=none&filterByDID=-99&sortByDID=2&sortOrder=descending&itemID=CMS1224069&intNumPerPage=10>. We will consider all comments received during the comment period in our determination of whether to create new modality-specific standard diagnostic radiology cost centers.

As noted in our response to a comment regarding the recommendations included in RTI's July 2008 report entitled, "Refining Cost to

Charge Ratios for Calculating APC and DRG Relative Payment Weights" (73 FR 68526), the current cost report form already includes nonstandard cost centers for CT Scanning and MRI. We also explained that under the principle of departmental apportionment of costs at \$413.53 hospitals are required to report separately the costs and charges for each ancillary department for which charges are customarily billed if the corresponding cost and charge information is accumulated separately in the provider's accounting system. We believe the nonstandard cost center information for CT Scanning and MRI that we currently collect reflects costs and charges for CT Scanning and MRI and we use these data to estimate median costs for ratesetting.

In the meantime, we believe it is fully appropriate to continue the multiple imaging composite payment methodology, which we believe improves the accuracy of OPSS payment rates and promotes efficiency among hospitals. As we stated in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68563), the most recent hospital cost report data are the best and most consistent estimate of relative costs that we have available to us for all hospitals for all hospital services. We will continue to use these data to estimate APC median costs. Should revised cost report data become available for CT and MRI procedures, our composite methodology would automatically incorporate that additional precision into the multiple imaging composite APC median cost estimates.

Comment: One commenter stated that the differential in the CY 2010 proposed payment rates for APC 8007 and APC 8008 appears adequate to account for the substantial differences in costs between magnetic resonance procedures when performed with and without contrast. The commenter asked CMS to evaluate the claims available for the CY 2010 OPSS/ASC final rule with comment period to ensure that payment rates for the two APCs reflect the incremental costs for the contrast agent and contrast administration included in APC 8008.

Response: We agree with the commenter regarding the appropriateness of the proposed differential in payment rates for APC 8007 and APC 8008 for CY 2010. The median costs upon which the CY 2010 final payment rates for APC 8007 and APC 8008 are based (\$706 and \$986, respectively) also appropriately reflect differences in costs for MRI and MRA imaging sessions with and without the administration of contrast.

Comment: One commenter stated that there was a discrepancy in CMS' estimated volume of APC 8005 single claims for the CY 2010 OPSS/ASC proposed rule. The commenter indicated that CMS' estimated volume of APC 8005 single claims increased by approximately one-third from the CY 2007 claims used in CY 2009 ratesetting to the CY 2008 claims available for the CY 2010 OPSS/ASC proposed rule. The commenter noted that this increase was inconsistent with the commenter's data analysis, which indicated that the total volume of single claims for APC 8005 did not increase significantly over this same time period.

Response: We reviewed the CY 2007 "single session" claims data used in ratesetting for APC 8005 for the CY 2009 OPSS/ASC final rule with comment period, and the CY 2008 "single session" claims data used in ratesetting for APC 8005 for the CY 2010 OPSS/ASC proposed rule. For the CY 2009 final rule, we identified 429,525 "single session" claims out of 809,483 potential composite cases to calculate the median cost for APC 8005. For the CY 2010 OPSS/ASC proposed rule, we identified 423,890 "single session" claims out of 810,469 potential composite cases to calculate the median cost for APC 8005. These published data do not demonstrate an increase of approximately one-third in the volume of "single session" claims from the CY 2007 claims used to calculate the median costs upon which the CY 2009 final payment rates are based compared to the CY 2008 claims used to calculate the median costs upon which the CY 2010 proposed payment rates are based, as the commenter indicated. For this final rule with comment period, we identified 455,191 "single session" claims (an increase of approximately 6 percent compared to CY 2009) out of 882,581 potential composite cases (an increase of approximately 9 percent compared to CY 2009) to calculate the median cost of APC 8005.

Comment: Many commenters requested that CMS thoroughly evaluate the impact of the multiple imaging composite payment methodology and commended CMS for not proposing to expand the multiple imaging composite payment methodology for CY 2010. Commenters asked CMS to review claims data to ensure that hospitals are being adequately paid for providing multiple imaging services, that patients are not being required by hospitals to return to the hospital on multiple days for imaging services, and that certain types or classes of hospitals are not being negatively affected before moving forward with any additional imaging

composite policies. One commenter noted that while CMS will have data available from CY 2009 to analyze for the winter 2010 APC Panel meeting, the commenter believed that such analyses would be more meaningful if claims data through CY 2012 are used to show impacts and a change in hospital behavior under the composite payment policy. Commenters also stated that any expansion of the multiple imaging composite payment methodology should be subject to full public comment.

Response: We appreciate the commenters' support of our proposal not to implement any significant changes to the composite APC methodology for CY 2010 so that we may monitor the effects of the existing composite APCs on utilization and payment. We also appreciate the commenters' thoughtful suggestions for data analysis that can be performed toward that end once CY 2009 claims data become available and in the longer term. We will take commenters' suggestions into consideration as we review the CY 2009 claims data for the impact of the multiple imaging composite APCs on payments to hospitals and on services to beneficiaries.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue paying for all multiple imaging procedures within an imaging family performed on the same date of service using the multiple imaging composite payment methodology. The CY 2010 payment rates for the five multiple imaging composite APCs (APC 8004, APC 8005, APC 8006, APC 8007, and APC 8008) are based on median costs calculated from the CY 2008 claims that would have qualified for composite payment under the current policy (that is, those claims with more than one procedure within the same family on a single date of service). Using the same ratesetting methodology described in the CY 2010 OPPS/ASC proposed rule (74 FR 35284), we were able to identify 1.8 million "single session" claims out of an estimated 2.7 million potential composite cases from our ratesetting claims data, or well over half of all eligible claims, to calculate the final CY 2010 median costs for the multiple imaging composite APCs.

Table 13 below lists the HCPCS codes subject to the final multiple imaging composite policy and their respective families for CY 2010. We note that we

have updated Table 13 to reflect HCPCS coding changes for CY 2010. Specifically, we added CPT code 74261 (Computed tomographic (CT) colonography, diagnostic, including image postprocessing; without contrast material) and CPT code 74262 (Computed tomographic (CT) colonography, diagnostic, including image postprocessing, with contrast material(s) including non-contrast images, if performed) to the CT and CTA family, and removed CPT code 0067T (Computed tomographic (CT) colonography (ie, virtual colonoscopy); diagnostic), which was replaced by these CPT codes. The HCPCS codes listed in Table 13 are assigned status indicated "Q3" in Addendum B to this final rule with comment period to identify their status as potentially payable through a composite APC. Their composite APC assignment is identified in Addendum M to this final rule with comment period. Table 14 below lists the imaging services subject to the composite methodology that overlap with HCPCS codes on the CY 2010 bypass list.

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TABLE 13.--OPPS IMAGING FAMILIES AND MULTIPLE IMAGING PROCEDURE COMPOSITE APCs

Family 1 - Ultrasound	
Final CY 2010 APC 8004 (Ultrasound Composite)	Final CY 2010 Approximate APC Median Cost = \$190
76604	Us exam, chest
76700	Us exam, abdom, complete
76705	Echo exam of abdomen
76770	Us exam abdo back wall, comp
76775	Us exam abdo back wall, lim
76776	Us exam k transpl w/Doppler
76831	Echo exam, uterus
76856	Us exam, pelvic, complete
76870	Us exam, scrotum
76857	Us exam, pelvic, limited
Family 2 - CT and CTA with and without Contrast	
Final CY 2010 APC 8005 (CT and CTA without Contrast Composite)*	Final CY 2010 Approximate APC Median Cost = \$416
70450	Ct head/brain w/o dye
70480	Ct orbit/ear/fossa w/o dye
70486	Ct maxillofacial w/o dye
70490	Ct soft tissue neck w/o dye
71250	Ct thorax w/o dye
72125	Ct neck spine w/o dye
72128	Ct chest spine w/o dye
72131	Ct lumbar spine w/o dye
72192	Ct pelvis w/o dye
73200	Ct upper extremity w/o dye
73700	Ct lower extremity w/o dye
74150	Ct abdomen w/o dye
74261	Ct colonography, w/o dye
Final CY 2010 APC 8006 (CT and CTA with Contrast Composite)	Final CY 2010 Approximate APC Median Cost = \$623
70487	Ct maxillofacial w/dye
70460	Ct head/brain w/dye

70470	Ct head/brain w/o & w/dye
70481	Ct orbit/ear/fossa w/dye
70482	Ct orbit/ear/fossa w/o&w/dye
70488	Ct maxillofacial w/o & w/dye
70491	Ct soft tissue neck w/dye
70492	Ct sft tsue nck w/o & w/dye
70496	Ct angiography, head
70498	Ct angiography, neck
71260	Ct thorax w/dye
71270	Ct thorax w/o & w/dye
71275	Ct angiography, chest
72126	Ct neck spine w/dye
72127	Ct neck spine w/o & w/dye
72129	Ct chest spine w/dye
72130	Ct chest spine w/o & w/dye
72132	Ct lumbar spine w/dye
72133	Ct lumbar spine w/o & w/dye
72191	Ct angiograph pelv w/o&w/dye
72193	Ct pelvis w/dye
72194	Ct pelvis w/o & w/dye
73201	Ct upper extremity w/dye
73202	Ct uppr extremity w/o&w/dye
73206	Ct angio upr extrm w/o&w/dye
73701	Ct lower extremity w/dye
73702	Ct lwr extremity w/o&w/dye
73706	Ct angio lwr extr w/o&w/dye
74160	Ct abdomen w/dye
74170	Ct abdomen w/o & w/dye
74175	Ct angio abdom w/o & w/dye
74262	Ct colonography, w/dye
75635	Ct angio abdominal arteries
* If a "without contrast" CT or CTA procedure is performed during the same session as a "with contrast" CT or CTA procedure, the I/OCE will assign APC 8006 rather than APC 8005.	
Family 3 - MRI and MRA with and without Contrast	
Final CY 2010 APC 8007 (MRI and MRA without Contrast Composite)*	Final CY 2010 Approximate APC Median Cost = \$706

70336	Magnetic image, jaw joint
70540	Mri orbit/face/neck w/o dye
70544	Mr angiography head w/o dye
70547	Mr angiography neck w/o dye
70551	Mri brain w/o dye
70554	Fmri brain by tech
71550	Mri chest w/o dye
72141	Mri neck spine w/o dye
72146	Mri chest spine w/o dye
72148	Mri lumbar spine w/o dye
72195	Mri pelvis w/o dye
73218	Mri upper extremity w/o dye
73221	Mri joint upr extrem w/o dye
73718	Mri lower extremity w/o dye
73721	Mri jnt of lwr extre w/o dye
74181	Mri abdomen w/o dye
75557	Cardiac mri for morph
75559	Cardiac mri w/stress img
C8901	MRA w/o cont, abd
C8904	MRI w/o cont, breast, uni
C8907	MRI w/o cont, breast, bi
C8910	MRA w/o cont, chest
C8913	MRA w/o cont, lwr ext
C8919	MRA w/o cont, pelvis
Final CY 2010 APC 8008 (MRI and MRA with Contrast Composite)	Final CY 2010 Approximate APC Median Cost = \$986
70549	Mr angiograph neck w/o&w/dye
70542	Mri orbit/face/neck w/dye
70543	Mri orbt/fac/nck w/o & w/dye
70545	Mr angiography head w/dye
70546	Mr angiograph head w/o&w/dye
70548	Mr angiography neck w/dye
70552	Mri brain w/dye
70553	Mri brain w/o & w/dye
71551	Mri chest w/dye
71552	Mri chest w/o & w/dye
72142	Mri neck spine w/dye
72147	Mri chest spine w/dye

72149	Mri lumbar spine w/dye
72156	Mri neck spine w/o & w/dye
72157	Mri chest spine w/o & w/dye
72158	Mri lumbar spine w/o & w/dye
72196	Mri pelvis w/dye
72197	Mri pelvis w/o & w/dye
73219	Mri upper extremity w/dye
73220	Mri uppr extremity w/o&w/dye
73222	Mri joint upr extrem w/dye
73223	Mri joint upr extr w/o&w/dye
73719	Mri lower extremity w/dye
73720	Mri lwr extremity w/o&w/dye
73722	Mri joint of lwr extr w/dye
73723	Mri joint lwr extr w/o&w/dye
74182	Mri abdomen w/dye
74183	Mri abdomen w/o & w/dye
75561	Cardiac mri for morph w/dye
75563	Card mri w/stress img & dye
C8900	MRA w/cont, abd
C8902	MRA w/o fol w/cont, abd
C8903	MRI w/cont, breast, uni
C8905	MRI w/o fol w/cont, brst, un
C8906	MRI w/cont, breast, bi
C8908	MRI w/o fol w/cont, breast,
C8909	MRA w/cont, chest
C8911	MRA w/o fol w/cont, chest
C8912	MRA w/cont, lwr ext
C8914	MRA w/o fol w/cont, lwr ext
C8918	MRA w/cont, pelvis
C8920	MRA w/o fol w/cont, pelvis

* If a "without contrast" MRI or MRA procedure is performed during the same session as a "with contrast" MRI or MRA procedure, the I/OCE will assign APC 8008 rather than 8007.

TABLE 14.—OPPS IMAGING FAMILY SERVICES OVERLAPPING WITH HCPCS CODES ON THE CY 2010 BYPASS LIST

Family 1 – Ultrasound	
76700	Us exam, abdom, complete
76705	Echo exam of abdomen
76770	Us exam abdo back wall, comp
76775	Us exam abdo back wall, lim
76776	Us exam k transpl w/doppler
76856	Us exam, pelvic, complete
76870	Us exam, scrotum
76857	Us exam, pelvic, limited
Family 2 - CT and CTA with and without Contrast	
70450	Ct head/brain w/o dye
70480	Ct orbit/ear/fossa w/o dye
70486	Ct maxillofacial w/o dye
70490	Ct soft tissue neck w/o dye
71250	Ct thorax w/o dye
72125	Ct neck spine w/o dye
72128	Ct chest spine w/o dye
72131	Ct lumbar spine w/o dye
72192	Ct pelvis w/o dye
73200	Ct upper extremity w/o dye
73700	Ct lower extremity w/o dye
74150	Ct abdomen w/o dye
Family 3 - MRI and MRA with and without Contrast	
70336	Magnetic image, jaw joint
70544	Mr angiography head w/o dye
70551	Mri brain w/o dye
72141	Mri neck spine w/o dye
72146	Mri chest spine w/o dye
72148	Mri lumbar spine w/o dye
73218	Mri upper extremity w/o dye
73221	Mri joint upr extrem w/o dye
73718	Mri lower extremity w/o dye
73721	Mri jnt of lwr extre w/o dye

BILLING CODE 4120-01-C**3. Calculation of OPPS Scaled Payment Weights**

Using the APC median costs discussed in sections II.A.1. and II.A.2. of this final rule with comment period, we calculated the final relative payment weights for each APC for CY 2010 shown in Addenda A and B to this final rule with comment period. In years

prior to CY 2007, we standardized all the relative payment weights to APC 0601 (Mid Level Clinic Visit) because mid-level clinic visits were among the most frequently performed services in the hospital outpatient setting. We assigned APC 0601 a relative payment weight of 1.00 and divided the median cost for each APC by the median cost for APC 0601 to derive the relative payment weight for each APC.

Beginning with the CY 2007 OPPS (71 FR 67990), we standardized all of the relative payment weights to APC 0606 (Level 3 Clinic Visits) because we deleted APC 0601 as part of the reconfiguration of the clinic visit APCs. We selected APC 0606 as the base because APC 0606 was the mid-level clinic visit APC (that is, Level 3 of five levels). Therefore, for CY 2010, to maintain consistency in using a median

for calculating unscaled weights representing the median cost of some of the most frequently provided services, we proposed to continue to use the median cost of the mid-level clinic visit APC, APC 0606, to calculate unscaled weights. Following our standard methodology, but using the proposed CY 2010 median cost for APC 0606, for CY 2010 we assigned APC 0606 a relative payment weight of 1.00 and divided the median cost of each APC by the proposed median cost for APC 0606 to derive the proposed unscaled relative payment weight for each APC. The choice of the APC on which to base the proposed relative weights for all other APCs did not affect the payments made under the OPSS because we scale the weights for budget neutrality.

Section 1833(t)(9)(B) of the Act requires that APC reclassification and recalibration changes, wage index changes, and other adjustments be made in a budget neutral manner. Budget neutrality ensures that the estimated aggregate weight under the OPSS for CY 2010 is neither greater than nor less than the estimated aggregate weight that would have been made without the changes. To comply with this requirement concerning the APC changes, we proposed to compare the estimated aggregate weight using the CY 2009 scaled relative weights to the estimated aggregate weight using the CY 2010 unscaled relative weights. For CY 2009, we multiply the CY 2009 scaled APC relative weight applicable to a service paid under the OPSS by the volume of that service from CY 2008 claims to calculate the total weight for each service. We then add together the total weight for each of these services in order to calculate an estimated aggregate weight for the year. For CY 2010, we perform the same process using the CY 2010 unscaled weights rather than scaled weights. We then calculate the weight scaler by dividing the CY 2009 estimated aggregate weight by the CY 2010 estimated aggregate weight. The service mix is the same in the current and prospective years because we use the same set of claims for service volume in calculating the aggregate weight for each year. For a detailed discussion of the weight scaler calculation, we refer readers to the OPSS claims accounting document available on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/>. We included payments to CMHCs in our comparison of estimated unscaled weight in CY 2010 to estimated total weight in CY 2009 using CY 2008 claims data and holding all other things constant. Based

on this comparison, we adjusted the unscaled relative weights for purposes of budget neutrality. In our proposal for CY 2010, the proposed CY 2010 unscaled relative payment weights were adjusted by multiplying them by a proposed weight scaler of 1.2863 to ensure budget neutrality of the proposed CY 2010 relative weights.

Section 1833(t)(14)(H) of the Act, as added by section 621(a)(1) of Public Law 108–173, states that, “Additional expenditures resulting from this paragraph shall not be taken into account in establishing the conversion factor, weighting and other adjustment factors for 2004 and 2005 under paragraph (9) but shall be taken into account for subsequent years.” Section 1833(t)(14) of the Act provides the payment rates for certain “specified covered outpatient drugs.” Therefore, the cost of those specified covered outpatient drugs (as discussed in section V.B.3. of the proposed rule (74 FR 35324 through 35333) and this final rule with comment period) was included in the proposed budget neutrality calculations for the CY 2010 OPSS.

We did not receive any public comments on the proposed methodology for calculating scaled weights from the median costs for the CY 2010 OPSS. Therefore, we are finalizing our proposed methodology, without modification, including updating of the budget neutrality scaler for this final rule with comment period. Under this methodology, the final unscaled payment weights were adjusted by a weight scaler of 1.3222 for this final rule with comment period. The final scaled relative payment weights listed in Addenda A and B to this final rule with comment period incorporate the recalibration adjustments discussed in sections II.A.1. and II.A.2. of this final rule with comment period.

4. Changes to Packaged Services

a. Background

The OPSS, like other prospective payment systems, relies on the concept of averaging, where the payment may be more or less than the estimated cost of providing a service or bundle of services for a particular patient, but with the exception of outlier cases, the payment is adequate to ensure access to appropriate care. Packaging and bundling payment for multiple interrelated services into a single payment create incentives for providers to furnish services in the most efficient way by enabling hospitals to manage their resources with maximum flexibility, thereby encouraging long-

term cost containment. For example, where there are a variety of supplies that could be used to furnish a service, some of which are more expensive than others, packaging encourages hospitals to use the least expensive item that meets the patient’s needs, rather than to routinely use a more expensive item. Packaging also encourages hospitals to negotiate carefully with manufacturers and suppliers to reduce the purchase price of items and services or to explore alternative group purchasing arrangements, thereby encouraging the most economical health care. Similarly, packaging encourages hospitals to establish protocols that ensure that necessary services are furnished, while carefully scrutinizing the services ordered by practitioners to maximize the efficient use of hospital resources. Finally, packaging payments into larger payment bundles promotes the stability of payment for services over time. Packaging and bundling also may reduce the importance of refining service-specific payment because there is more opportunity for hospitals to average payment across higher cost cases requiring many ancillary services and lower cost cases requiring fewer ancillary services.

Decisions about packaging and bundling payment involve a balance between ensuring that payment is adequate to enable the hospital to provide quality care and establishing incentives for efficiency through larger units of payment. In the CY 2008 OPSS/ASC final rule with comment period (72 FR 66610 through 66659), we adopted the packaging of payment for items and services in the seven categories listed below into the payment for the primary diagnostic or therapeutic modality to which we believe these items and services are typically ancillary and supportive. The seven categories are guidance services, image processing services, intraoperative services, imaging supervision and interpretation services, diagnostic radiopharmaceuticals, contrast media, and observation services. We specifically chose these categories of HCPCS codes for packaging because we believe that the items and services described by the codes in these categories are the HCPCS codes that are typically ancillary and supportive to a primary diagnostic or therapeutic modality and, in those cases, are an integral part of the primary service they support.

We assign status indicator “N” to those HCPCS codes that we believe are always integral to the performance of the primary modality; therefore, we always package their costs into the costs

of the separately paid primary services with which they are billed. Services assigned status indicator "N" are unconditionally packaged.

We assign status indicator "Q1" ("STVX-Packaged Codes"), "Q2" ("T-Packaged Codes"), or "Q3" (Codes that may be paid through a composite APC) to each conditionally packaged HCPCS code. An "STVX-packaged code" describes a HCPCS code whose payment is packaged when one or more separately paid primary services with the status indicator of "S," "T," "V," or "X" are furnished in the hospital outpatient encounter. A "T-packaged code" describes a code whose payment is packaged when one or more separately paid surgical procedures with the status indicator of "T" are provided during the hospital encounter. "STVX-packaged codes" and "T-packaged codes" are paid separately in those uncommon cases when they do not meet their respective criteria for packaged payment. "STVX-packaged codes" and "T-packaged HCPCS codes" are conditionally packaged. We refer readers to section XIII.A.1. of this final rule with comment period for a complete listing of status indicators.

We use the term "dependent service" to refer to the HCPCS codes that represent services that are typically ancillary and supportive to a primary diagnostic or therapeutic modality. We use the term "independent service" to refer to the HCPCS codes that represent the primary therapeutic or diagnostic modality into which we package payment for the dependent service. We note that, in future years as we consider the development of larger payment groups that more broadly reflect services provided in an encounter or episode-of-care, it is possible that we might propose to bundle payment for a service that we now refer to as "independent."

In addition, in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66650 through 66659), we finalized additional packaging for the CY 2008 OPPS, which included the establishment of new composite APCs for CY 2008, specifically APC 8000 (Cardiac Electrophysiologic Evaluation and Ablation Composite), APC 8001 (LDR Prostate Brachytherapy Composite), APC 8002 (Level I Extended Assessment & Management Composite), and APC 8003 (Level II Extended Assessment & Management Composite). In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68559 through 68569), we expanded the composite APC model to one new clinical area, multiple imaging services. We created five multiple imaging composite APCs for payment in CY

2009 that incorporate statutory requirements to differentiate between imaging services provided with contrast and without contrast as required by section 1833(t)(2)(G) of the Act. The multiple imaging composite APCs are: APC 8004 (Ultrasound Composite); APC 8005 (CT and CTA without Contrast Composite); APC 8006 (CT and CTA with Contrast Composite); APC 8007 (MRI and MRA without Contrast Composite); and APC 8008 (MRI and MRA with Contrast Composite). We discuss composite APCs in more detail in section II.A.2.e. of this final rule with comment period.

Hospitals include charges for packaged services on their claims, and the estimated costs associated with those packaged services are then added to the costs of separately payable procedures on the same claims in establishing payment rates for the separately payable services. We encourage hospitals to report all HCPCS codes that describe packaged services that were provided, unless the CPT Editorial Panel or CMS provide other guidance. If a HCPCS code is not reported when a packaged service is provided, it can be challenging to track utilization patterns and resource costs.

b. Packaging Issues

(1) Packaged Services Addressed by the February 2009 APC Panel Recommendations

The Packaging Subcommittee of the APC Panel was established to review packaged HCPCS codes. In deciding whether to package a service or pay for a code separately, we have historically considered a variety of factors, including whether the service is normally provided separately or in conjunction with other services; how likely it is for the costs of the packaged code to be appropriately mapped to the separately payable codes with which it was performed; and whether the expected cost of the service is relatively low. As discussed in section II.A.4.a. of this final rule with comment period regarding our packaging approach for CY 2008, we established packaging criteria that apply to seven categories of codes whose payments are packaged.

During the September 2007 APC Panel meeting, the APC Panel requested that CMS evaluate the impact of expanded packaging on beneficiaries. During the March 2008 APC Panel meeting, the APC Panel requested that CMS report to the Panel at the first Panel meeting in CY 2009 regarding the impact of packaging on net payments for patient care. In response to these requests, we shared data with the APC

Panel at the February 2009 APC Panel meeting that compared the frequency of specific categories of services billed under the OPPS in CY 2007, before the expanded packaging went into effect, to the frequency of those same categories of services in CY 2008, their first year of packaged payment. In each category, the HCPCS codes that we compared are the ones that we identified in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66659 through 66664) as fitting into one of the seven packaging categories listed in section II.A.4.a. of this final rule with comment period. The data shared with the APC Panel at the February 2009 APC Panel meeting compared CY 2007 claims processed through September 30, 2007 to CY 2008 claims processed through September 30, 2008. We did not make any adjustments for inflation, changes in Medicare population, or other variables that potentially influenced billing between CY 2007 and CY 2008. These data represent about 60 percent of the full year data. A summary of these data analyses is provided below.

Analysis of the diagnostic radiopharmaceuticals category showed that the frequency of the reporting of diagnostic radiopharmaceuticals increased by 1 percent between the first 9 months of CY 2007 and the first 9 months of CY 2008. In CY 2007, some diagnostic radiopharmaceuticals were packaged and others were separately payable, depending on whether their per day mean costs fell above or below the \$55 drug packaging threshold for CY 2007. All diagnostic radiopharmaceuticals were uniformly packaged in CY 2008. Two percent more hospitals reported one or more diagnostic radiopharmaceuticals during CY 2008 as compared to CY 2007. Effective for CY 2008, we first required reporting of a radiolabeled product (including diagnostic radiopharmaceuticals) when billing a nuclear medicine procedure, and we believe that the increases in frequency and the number of reporting hospitals reflect hospitals meeting this reporting requirement.

We also found that nuclear medicine procedures (into which diagnostic radiopharmaceuticals were packaged) and associated diagnostic radiopharmaceuticals were billed approximately 3 million times during the first 9 months of both CY 2007 and CY 2008. Further analysis revealed that we paid hospitals over \$637 million for nuclear medicine procedures and diagnostic radiopharmaceuticals during the first 9 months of CY 2007, when diagnostic radiopharmaceuticals were separately payable, and over \$619

million for nuclear medicine procedures and diagnostic radiopharmaceuticals during the first 9 months of CY 2008, when payment for diagnostic radiopharmaceuticals was packaged. This represented a 3 percent decrease in aggregate payment between the first 9 months of CY 2007 and the first 9 months of CY 2008.

Using the same data, we calculated an average payment per service or item billed (including nuclear medicine procedures and packaged or separately payable diagnostic radiopharmaceuticals) of \$203 in CY 2007 and \$198 in CY 2008 for nuclear medicine procedures. This represented a decrease of 2 percent in average payment per item or service billed between CY 2007 and CY 2008. It is unclear how much of the decrease in estimated aggregate or average per service or item billed payment may be due to packaging payment for diagnostic radiopharmaceuticals (and other services that were newly packaged for CY 2008) and how much may be due to the usual annual APC recalibration and typical fluctuations in service frequency. However, we believe that all of these factors likely contributed to the slight decrease in aggregate payment in CY 2008, as compared to CY 2007. Overall, the observed changes between CY 2007 and CY 2008 were very small and indicated that there has been very little change in frequency or aggregate payment in this clinical area between CY 2007 and CY 2008.

We similarly analyzed 9 months of CY 2007 and CY 2008 data related to all services that were packaged during CY 2008 because they were categorized as guidance services. Analysis of the guidance category (which includes image-guided radiation therapy services) showed that the frequency of guidance services increased by 2 percent between the first 9 months of CY 2007 and the first 9 months of CY 2008. One percent fewer hospitals reported one or more guidance services during CY 2007 as compared to CY 2008.

We further analyzed 9 months of CY 2007 and CY 2008 claims data for radiation oncology services that would be accompanied by radiation oncology guidance. We found that radiation oncology services (including radiation oncology guidance services) were billed approximately 4 million times in CY 2007 and 3.9 million times in CY 2008, representing a decrease in frequency of approximately 5 percent between CY 2007 and CY 2008. These numbers represented each instance where a radiation oncology service or a radiation oncology guidance service was billed.

Our analysis indicated that hospitals were paid over \$818 million for radiation oncology services and radiation oncology guidance services under the OPSS during the first 9 months of CY 2007, when radiation oncology guidance services were separately payable. During the first 9 months of CY 2008, when payments for radiation oncology guidance were packaged, hospitals were paid over \$740 million for radiation oncology services under the OPSS. This \$740 million included packaged payment for radiation oncology guidance services and represented a 10 percent decrease in aggregate payment from CY 2007 to CY 2008. Using the first 9 months of data for both CY 2007 and CY 2008, we calculated an average payment per radiation oncology service or item billed of \$201 in CY 2007 and \$190 in CY 2008, representing a decrease of 5 percent from CY 2007 to CY 2008. It is unclear how much of the decrease in aggregate payment and the decrease in average payment per service provided may be due to packaging payment for radiation oncology guidance services (and other services that were newly packaged for CY 2008) and how much may be due to the usual annual APC recalibration and typical fluctuations in service frequency. This analysis is discussed in further detail below, under "Recommendation 1" in this section of this final rule with comment period. In that analysis, we demonstrated that the volume of some packaged radiation oncology guidance services increased during the period, leading us to conclude that, irrespective of the decline in the frequency of radiation oncology services in general, hospitals did not appear to be changing their practice patterns specifically in response to packaged payment for radiation oncology guidance services.

We similarly analyzed 9 months of CY 2007 and CY 2008 data related to all services that were packaged during CY 2008 because they were categorized as intraoperative services. Analysis of the intraoperative category (which includes intravascular ultrasound (IVUS), intracardiac echocardiography (ICE), and coronary fractional flow reserve (FFR)) showed minimal changes in the frequency and the number of reporting hospitals between CY 2007 and CY 2008.

We found that cardiac catheterization and other percutaneous vascular procedures that would typically be accompanied by IVUS, ICE and FFR (including IVUS, ICE, and FFR) were billed approximately 375,000 times in CY 2007 and approximately 400,000 times in CY 2008, representing an

increase of 8 percent in the number of services and items billed between CY 2007 and CY 2008. Further analysis revealed that the OPSS paid hospitals over \$912 million for cardiac catheterizations, other related services, and IVUS, ICE, and FFR in CY 2007, when IVUS, ICE, and FFR were separately payable. In the first 9 months of CY 2008, the OPSS paid hospitals approximately \$1.1 billion for cardiac catheterization and other percutaneous vascular procedures and IVUS, ICE, and FFR, when payments for IVUS, ICE, and FFR were packaged. This represented a 25 percent increase in payment from CY 2007 to CY 2008. Using the 9 months of data for both CY 2007 and CY 2008, we calculated an average payment per service or item provided of \$2,430 in CY 2007 and \$2,800 in CY 2008 for cardiac catheterization and other related services. This represented an increase of 15 percent in average payment per item or service from CY 2007 to CY 2008.

We could not determine how much of the 25 percent increase in aggregate payment for these services may be due to the packaging of payment for IVUS, ICE, and FFR (and other services that were newly packaged for CY 2008) and how much may be due to the usual annual APC recalibration and typical fluctuations in service frequency. However, we believe that all of these factors contributed to the increase in payment between these 2 years.

The three remaining packaging categories (excluding observation services, which are further discussed in section II.A.2.e.(1) of this final rule with comment period), contrast agents, image processing services, and imaging supervision and interpretation services, showed minimal changes in frequency between CY 2007 and CY 2008, ranging from a 2 percent increase to a 1 percent decrease in frequency. Similarly, when examining the number of hospitals reporting these services, the data showed similar numbers of hospitals reporting these services in CY 2007, when these services were separately payable, and CY 2008, when they were packaged. Specifically, the percentage change in the number of reporting hospitals for these categories between CY 2007 and CY 2008 ranged from 0 percent to a decrease of 1 percent.

In summary, these preliminary data indicated that hospitals in aggregate did not appear to have significantly changed their service reporting patterns as a result of the expanded packaging adopted for the OPSS beginning in CY 2008.

The APC Panel's Packaging Subcommittee reviewed the packaging status of several CPT codes and reported

its findings to the APC Panel at its February 2009 meeting. The full report of the February 18 and 19, 2009 APC Panel meeting can be found on the CMS Web site at: http://www.cms.hhs.gov/FACA/05_AdvisoryPanelonAmbulatoryPaymentClassificationGroups.asp. The APC Panel accepted the report of the Packaging Subcommittee, heard several presentations related to packaged services, discussed the deliberations of the Packaging Subcommittee, and recommended that—

1. CMS pay separately for radiation therapy guidance services performed in the treatment room for 2 years and then reevaluate packaging on the basis of claims data. (Recommendation 1)

2. CMS continue to analyze the impact of increased packaging on beneficiaries and provide more detailed versions of the analyses presented at the February 2009 meeting of services initially packaged in CY 2008 at the next Panel meeting. In addition, the Panel requested that, in the more detailed analyses of radiation oncology services that would be accompanied by radiation oncology guidance, CMS stratify the data according to the type of radiation oncology service, specifically, intensity modulated radiation therapy, stereotactic radiosurgery, brachytherapy, and conventional radiation therapy. (Recommendation 2)

3. CMS continue to analyze the impact on beneficiaries of increased packaging of diagnostic radiopharmaceuticals and provide more detailed analyses at the next Panel meeting. In addition, the Panel requested that, in the more detailed analyses of packaging of diagnostic radiopharmaceuticals by type of nuclear medicine scan, CMS break down the data according to the specific CPT codes billed with the diagnostic radiopharmaceuticals. (Recommendation 3)

4. CPT code 36592 (Collection of blood specimen using established central or peripheral catheter, venous, not otherwise specified) remain assigned to APC 0624 (Phlebotomy and Minor Vascular Access Device Procedures) for CY 2010. (Recommendation 4)

5. The Packaging Subcommittee continue its work until the next APC Panel meeting. (Recommendation 5)

In the proposed rule, we addressed each of these recommendations in turn in the discussion that follows.

Recommendation 1

In the CY 2010 OPPS/ASC proposed rule (74 FR 35289), we did not propose to pay separately for radiation therapy guidance services provided in the

treatment room for CY 2010, which would have been consistent with the APC Panel's recommendation. Instead, we proposed to maintain the packaged status of radiation therapy guidance services performed in the treatment room for CY 2010.

As discussed below in this section, during the February 2009 APC Panel meeting, we presented data that estimated that aggregate payment for radiation oncology services, including the payment for radiation oncology guidance services, decreased by approximately 10 percent between the first 9 months of CY 2007 (before the expanded packaging went into effect) and the first 9 months of CY 2008 (after the expanded packaging went into effect). This decline may be attributable to many factors, including lower payment rates for common radiation oncology services in CY 2008 specifically and generally reduced volume for separately paid radiation oncology services. The APC Panel expressed concern that this aggregate payment decrease could inhibit patient access to technologically advanced and clinically valuable radiation oncology guidance services whose payment became packaged effective January 1, 2008.

While we presented data to the APC Panel comparing payment between CY 2007 and CY 2008 in response to past APC Panel recommendations, we note that we made changes to the bypass list for CY 2009 to ensure that we more fully captured all packaged costs on each claim, which resulted in significantly increased payment rates for many of these radiation oncology services for CY 2009, as compared to the CY 2008 payment rates for these services.

Specifically, as discussed in detail in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68575), in response to public comments received, several radiation oncology CPT codes had been included on the bypass list for the CY 2008 OPPS, although they failed to meet the empirical criteria for inclusion on the bypass list. For CY 2009, we removed from the bypass list those radiation oncology codes that did not meet the empirical criteria. As a result of these changes to the bypass list, the CY 2009 median costs for several common radiation oncology APCs increased by more than 9 percent as compared to the CY 2008 median costs, while the median costs for some of the other lower volume radiation oncology APCs, most notably the brachytherapy source application APCs, declined. For example, as noted in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68575), these changes to the bypass

list resulted in payment for the common combination of intensity modulated radiation therapy (IMRT) and image-guided radiation therapy (IGRT) increasing from \$348 in CY 2008 to \$411 in CY 2009. Notably, the CY 2007 total payment rate for this combination of services, before the expanded packaging went into effect, was \$403.

We do not yet have CY 2009 claims data reflecting utilization based on the payment rates in effect for CY 2009. However, we do not expect that an overall per-service payment comparison between CY 2007 and CY 2009 would likely demonstrate a significant decrease in payment for radiation oncology services because we have adopted a significant increase in the CY 2009 payment rates for the most common radiation oncology services. In addition, we note that CY 2010 proposed rule data indicated that the CY 2010 APC median costs applicable to most radiation oncology services experienced increases of approximately 2 to 15 percent when compared to their CY 2009 final rule median costs. Although a small number of other lower volume radiation oncology APCs, most notably the brachytherapy and stereotactic radiosurgery APCs, experienced declines in median costs, we do not expect that an overall per-service payment comparison between CY 2007 and CY 2010 would likely demonstrate a significant decrease in payment for radiation oncology services over this time period.

While we understand that the CY 2007 to CY 2008 aggregate payment comparison provided to the APC Panel during the February 2009 meeting may have contributed to the APC Panel's particular concern about payment for radiation oncology services for CY 2010, we do not believe that packaging payment for radiation oncology guidance services has primarily caused this decline. In addition, we do not believe that beneficiaries' access to these services has been limited as a result of packaging payment for radiation oncology guidance services. In the data presented to the APC Panel at the February 2009 meeting, the number of all packaged guidance services provided during the first 9 months of CY 2008 represented a 2 percent increase from the number of guidance services provided during the first 9 months of CY 2007. Further, although the CY 2008 volume of the radiation oncology guidance codes that we newly packaged for CY 2008 varied, with some of the services experiencing increases in volume and others experiencing decreases in volume, in aggregate, the reporting of radiation oncology

guidance services increased by 4 percent in the first 9 months of claims for CY 2008, as compared to the first 9 months of CY 2007, and the number of hospitals reporting these services also increased. This further supports our belief that, irrespective of the decline in the frequency of radiation oncology services in general, hospitals do not appear to be changing their practice patterns specifically in response to packaged payment for radiation oncology guidance services.

Therefore, in the CY 2010 OPPS/ASC proposed rule (74 FR 35289), we did not propose to pay separately for radiation therapy guidance services performed in the treatment room for 2 years as the APC Panel recommended. Instead, for CY 2010, we proposed to maintain the packaged status of all radiation therapy guidance services, including those radiation therapy guidance services performed in the treatment room.

A summary of the public comments and our response on the CY 2010 proposal to package payment for radiation therapy guidance services are included in section II.A.4.b.(2) of this final rule with comment period.

Recommendation 2

In the CY 2010 OPPS/ASC proposed rule (74 FR 35290), we stated that we are accepting the APC Panel recommendation to continue to analyze the impact of increased packaging on beneficiaries and to share more data with the APC Panel. We noted that we would carefully consider which additional data would be most informative for the APC Panel and would discuss these data with the APC Panel at the next CY 2009 APC Panel meeting and/or the first CY 2010 APC Panel meeting. We did not share additional packaging data with the APC Panel at the most recent August 2009 meeting because we believe the APC Panel's discussions would benefit from analyses of an additional year of claims data after CY 2008. Therefore, we plan to incorporate analysis of CY 2009 claims into the information we will bring to the APC Panel for its review at the winter 2010 meeting. Similarly, in the proposed rule, we noted that we would determine what additional detailed data related to radiation oncology services would be helpful to the APC Panel and would share these data at the next CY 2009 APC Panel meeting and/or the first CY 2010 APC Panel meeting. We did not share additional data related to radiation oncology services with the APC Panel at the most recent August 2009 meeting because we believe the APC Panel's discussions would benefit from analyses

of an additional year of claims data after CY 2008. Therefore, we plan to incorporate analysis of CY 2009 claims into the information we will bring to the APC Panel for its review at the winter 2010 meeting.

A summary of the public comments and our response regarding the impact of the CY 2010 packaging proposal are included in section II.A.4.b.(2) of this final rule with comment period.

Recommendation 3

In the CY 2010 OPPS/ASC proposed rule (74 FR 35290), we stated that we are accepting the APC Panel's recommendation that CMS continue to analyze the impact on beneficiaries of increased packaging of diagnostic radiopharmaceuticals and provide more detailed analyses at the next APC Panel meeting. In these analyses of diagnostic radiopharmaceuticals by type of nuclear medicine scan, the APC Panel further recommended that CMS analyze the data according to the specific CPT codes billed with the diagnostic radiopharmaceuticals. This APC Panel recommendation is discussed in detail in section II.A.2.d.(5) of this final rule with comment period. In the proposed rule, we noted that we are accepting the APC Panel's recommendation and would provide additional data to the APC Panel at an upcoming meeting. We did not share additional data related to diagnostic radiopharmaceuticals and nuclear medicine scans with the APC Panel at the most recent August 2009 meeting because we believe the APC Panel's discussions would benefit from analyses of an additional year of claims data after CY 2008. Therefore, we plan to incorporate analysis of CY 2009 claims into the information we will bring to the APC Panel for its review at the winter 2010 meeting.

A summary of the public comments and our response on the CY 2010 proposal to package payment for diagnostic radiopharmaceuticals into payment for the associated nuclear medicine procedures are included in sections II.A.2.d.(5) and V.B.2.d. of this final rule with comment period.

Recommendation 4

In the CY 2010 OPPS/ASC proposed rule (74 FR 35290), we proposed to continue for CY 2010 to treat CPT code 36592 (Collection of blood specimen using established central or peripheral catheter, venous, not otherwise specified) as an "STVX packaged code" and to assign it to APC 0624 (Phlebotomy and Minor Vascular Access Device Procedures), the same APC to which CPT code 36591 (Collection of blood specimen from a completely

implantable venous access device) is currently assigned as the APC Panel recommended. CPT code 36592 became effective January 1, 2008 and was assigned interim status indicator "N" in the CY 2008 OPPS/ASC final rule with comment period. For CY 2009, in response to public comments, we proposed to treat CPT code 36592 as a conditionally packaged code, with assignment to APC 0624. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68576), we discussed the public comments we received regarding our proposed treatment of CPT code 36592. Several of these commenters supported our proposal to treat CPT code 36592 as a conditionally packaged code with assignment to APC 0624. We stated in the CY 2009 OPPS/ASC final rule with comment period that when cost data for CPT code 36592 became available for the CY 2010 OPPS annual update, we would reevaluate whether assignment to APC 0624 continued to be appropriate.

Based on our analysis of claims data, our clinical understanding of the service, and our discussion with the APC Panel Packaging Subcommittee, in the CY 2010 OPPS/ASC proposed rule (74 FR 35290), we proposed to maintain the assignment of CPT code 36592 to APC 0624 for CY 2010, consistent with the APC Panel recommendation, and we proposed to continue to treat CPT code 36592 as an "STVX packaged code" and assign it to APC 0624. We noted that we expect hospitals to follow the CPT guidance related to CPT codes 36591 and 36592 regarding when these services should be appropriately reported.

We received no public comments on the CY 2010 proposal to maintain the assignment of CPT code 36592 to APC 0624 and treat it as an "STVX packaged code," so we are finalizing our proposal, without modification.

Recommendation 5

In response to the APC Panel's recommendation for the Packaging Subcommittee to remain active until the next APC Panel meeting, in the CY 2010 OPPS/ASC proposed rule (74 FR 35290) we noted that we have accepted this recommendation and the APC Panel Packaging Subcommittee remains active. We stated that additional issues and new data concerning the packaging status of codes would be shared for its consideration as information becomes available. We continue to encourage submission of common clinical scenarios involving currently packaged HCPCS codes to the Packaging Subcommittee for its ongoing review. We also encourage recommendations of

specific services or procedures whose payment would be most appropriately packaged under the OPPS. Additional detailed suggestions for the Packaging Subcommittee should be submitted by e-mail to APCPanel@cms.hhs.gov with Packaging Subcommittee in the subject line.

The Packaging Subcommittee has remained active; the Subcommittee's last meeting to discuss packaging issues was the August 2009 meeting.

(2) Packaged Services Addressed by the August 2009 APC Panel Recommendations

The APC Panel met again on August 5 and 6, 2009 to hear public presentations on the proposals set forth in the CY 2010 OPPS/ASC proposed rule. The APC Panel's Packaging Subcommittee reviewed the packaging status of several CPT codes and reported its findings to the APC Panel. The full report of the August 5 and 6, 2009 APC Panel meeting can be found on the CMS Web site at: http://www.cms.hhs.gov/FACA/05_Advisory_PanelonAmbulatoryPaymentClassificationGroups.asp. The APC Panel accepted the report of the Packaging Subcommittee, heard several presentations related to packaged services, discussed the deliberations of the Packaging Subcommittee, and recommended that—

1. CMS submit to the Packaging Subcommittee, for its ongoing review, common clinical scenarios involving currently packaged HCPCS codes and recommendations of specific services or procedures for which payment would be most appropriately packaged under the OPPS. (Recommendation 6)

2. When CMS changes the dollar amount of the drug packaging threshold and determines that some drugs within a single therapeutic class fall on either side of the packaging threshold, CMS consider packaging all of the drugs within that class on the basis of feedback from providers, the APC Panel, and stakeholders. (Recommendation 7)

3. CMS continue to study the impact of increased packaging on beneficiaries. (Recommendation 8)

4. The work of the APC Packaging Subcommittee continue. (Recommendation 9)

With respect to these August 2009 APC Panel recommendations, we are accepting recommendations 6, 8, and 9. We are continuing the work of the APC Panel Packaging Subcommittee, and we appreciate the Packaging Subcommittee's expertise and experience regarding packaging under the OPPS and the valuable advice the Subcommittee continues to provide to us. We will

continue to bring to the Subcommittee's attention clinical scenarios identified by us or the public regarding services that are currently packaged or are candidates for future packaging under the OPPS. As discussed above, we also will continue to study the impact of increased packaging on Medicare beneficiaries, as the APC Panel has previously recommended to us. We did not share additional packaging data with the APC Panel at the most recent August 2009 meeting because we believe the APC Panel's discussions would benefit from analyses of an additional year of claims data after CY 2008. Therefore, we plan to incorporate analysis of CY 2009 claims into the information we will bring to the APC Panel for its review at the winter 2010 meeting. Finally, our response to recommendation 7 regarding the packaging of payment for all drugs in the same therapeutic class is discussed in section V.B.2.c. of this final rule with public comment.

Comment: Many commenters expressed a wide range of views on the existing policies for packaging payment for categories of services that CMS proposed to continue for CY 2010. One commenter claimed that while packaging provides an incentive for providers to deliver services in the most efficient, cost-effective manner possible, payment bundles that are too small do not enhance efficiencies, while payment bundles that are too large may carry excessive copayments for patients who need only a small proportion of services in the bundle. Another commenter suggested that CMS' packaging policy is likely to lead to less efficient use of resources, limited access to innovative treatment options, and greater instability in payments because, unlike the incentives from packaging under the IPPS, under the OPPS, the hospital would receive greater payment by bringing the outpatient back for a second visit or admitting the patient for inpatient care than by utilizing a more costly approach to providing an outpatient service that would be paid the same, regardless of the approach. The commenter also stated that when an APC's payment rate is significantly less than the cost of a technology, hospitals have a strong disincentive to use that technology, even if it could reduce the costs of care at a later date and provide better care to the patient.

Several commenters asserted that the implications of OPPS packaging policies are unknown due to a lack of transparency in the OPPS ratesetting process and methodology used to determine payment for packaged services, potentially leading to inappropriate payment and

underutilization of image-guidance services. The commenters believed that packaging payment for image-guidance leads to hospitals discouraging physicians from using guidance services and that, therefore, CMS should not package payment for image-guidance services. Several commenters urged CMS to consider establishing a 2 to 3 year data collection period during which separate payment would be made for new technology or new applications of existing technology. The commenters further suggested that the data could then be used to evaluate the impact of packaging on clinical utilization and payment and could also be used to determine whether to package or maintain separate payment for the services in the future. Another commenter recommended that CMS adopt a threshold policy that would be similar to the existing policy used to identify packaged drugs, under which separate payment would be made for all services with a median cost in excess of a nominal threshold amount.

Response: We continue to believe that packaging creates incentives for hospitals and their physician partners to work together to establish appropriate protocols that eliminate unnecessary services where they exist and institutionalize approaches to providing necessary services more efficiently. With respect to new services or new applications of existing technology, we believe that packaging payment for ancillary and dependent services creates appropriate incentives for hospitals to seriously consider whether a new service or a new technology offers a benefit that is sufficient to justify the cost of the new service or technology. Where this review results in reductions in services that are only marginally beneficial or hospitals' choices not to utilize certain technologies, we believe that this could improve, rather than harm, the quality of care for Medicare beneficiaries because every service furnished in a hospital carries some level of risk to the patient. Moreover, we believe that hospitals strive to provide the best care they can to the patients they serve so that when new technologies are proven to improve the quality of care, their utilization will increase appropriately, whether the payment for them is packaged or not.

However, we are aware that there are financial pressures on hospitals that might motivate some providers to split services among different hospital encounters in such a way as to maximize payments. While we do not expect that hospitals would routinely change the way they furnish services or the way they bill for services in order

to maximize payment, we recognize that it would be possible and we consider that possibility as we annually review hospital claims data. We will to continue examine claims data for patterns of fragmented care, and if we find a pattern in which a hospital appears to be dividing care across multiple days, we will refer it for investigation to the QIO or to the program safeguard contractor, as appropriate to the circumstances we find.

In section II.A.1. of this final rule with comment period, we discuss the established methodology we use to incorporate the costs of packaged services into payment for the associated independent procedures. In response to those commenters with concerns about transparency of the ratesetting process that incorporates packaged costs, in general, we package the costs of services into the payment for the major separately paid procedure on the same claim on which the packaged service appears. Hence, it is the practice of hospitals with regard to reporting and charging for packaged services that determines the separately paid service into which the cost of a packaged service is incorporated and the amount of packaged cost included the payment for that separately paid procedure.

Regarding the recommendation that we establish a cost threshold that would guide the packaging of services, we do not agree that this approach would result in appropriate packaging of costs for dependent ancillary services. A threshold policy could create incentives for hospitals to increase charges to ensure that payment for certain services was made separately, and the result would be contrary to the creation of incentives for prudent assessment of the costs and benefits of these services. Furthermore, as we stated in the CY 2009 OPPS/ASC final with comment period (73 FR 68572), it is not clear whether one set of packaging principles or one threshold could apply to the wide variety of services under the OPPS. Finally, to adopt a policy that would only package services that are low cost ancillary and supportive services would essentially negate the concept of averaging that is an underlying premise of a prospective payment system because hospitals would not have a particular incentive to provide care more efficiently.

We believe it is important to continue to advance value-based purchasing by Medicare in the hospital outpatient setting by furthering the focus on value of care rather than volume. While we acknowledge the concerns of the commenters and, as discussed below,

are committed to considering the impact of packaging payment on Medicare beneficiaries further in the future, we must balance the concerns of the commenters with our goal of continuing to encourage efficient use of hospital resources. As we noted in the CY 2009 OPPS/ASC final rule with comment period in our response to comments on the CY 2009 OPPS/ASC proposed rule (73 FR 68572) and as we note in our responses to public comments on the CY 2010 OPPS/ASC proposed rule, the suggestions and packaging criteria recommended by most commenters are focused almost exclusively on preventing packaging, rather than on determining when packaging would be appropriate. We also welcome suggestions from the public on approaches to packaging that would encourage efficient use of hospital resources.

Comment: Several commenters commended CMS for reviewing and accepting the APC Panel's February 2009 recommendation that CMS continue to analyze the impact of increased packaging on Medicare beneficiaries. The commenters expressed concern about CMS' current packaging policy and urged CMS to conduct a more detailed review of the hospital claims data in order to verify that current OPPS packaging policies and methodologies are accomplishing CMS' goals. A few commenters offered recommendations for additional data analyses for CMS to consider in the ongoing efforts to study the impact of increased packaging under the OPPS. The commenters recommended that CMS compare utilization of currently packaged services billed and paid separately under the OPPS in CY 2007, before the packaging of additional categories of services went into effect, to the frequency of those same services that were packaged in CY 2008 and later, after the packaging of additional categories of services went into effect. The commenters requested that CMS conduct these studies at the CPT code level. The commenters also recommended that CMS conduct a hospital-level review of the data, in addition to an overall review, and compare overall utilization by packaged HCPCS code for CYs 2005 and 2006 to CYs 2007, 2008, and 2009. Another commenter, in support of a provider-level review of the data, asserted that reviewing the data for packaged services at a national aggregate level can easily mask the behavioral changes of classes of hospitals and, therefore, concluded that more detailed analysis is needed to determine the impact of the policy.

Several commenters requested that CMS present its analyses in the final rule with comment period and at upcoming APC Panel meetings and consult with relevant stakeholders before proposing any additional packaging changes. The commenters also recommended that CMS make the data underlying payments for packaged services, including utilization rates and median costs, publicly available to enhance the transparency of its decision making so that stakeholders could assess whether the payment rates truly reflect the costs of providing the bundle of services.

Response: We agree that it is important to examine our claims data to assess the impact of packaging to the extent we can do so. During the September 2007 APC Panel meeting, the APC Panel requested that CMS evaluate the impact of expanded packaging on Medicare beneficiaries. At the March 2008 APC Panel meeting, the APC Panel requested that CMS report to the APC Panel at the first meeting in CY 2009 regarding the impact of packaging on net payments for patient care. In response to these requests, we shared the first available CY 2008 claims data with the APC Panel at the February 2009 APC Panel meeting. In that analysis, we compared the frequency of specific categories of services we newly packaged for CY 2008 as they were billed under the OPPS in CY 2007, before expanded packaging went into effect, to the frequency of those same categories of services in CY 2008, their first year of packaged payment. In each category, the HCPCS codes that we compared are the ones that we identified in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66659 through 66664) as fitting into one of the seven packaging categories listed in section II.A.4.a. of this final rule with comment period. The data shared with the APC Panel at the February 2009 APC Panel meeting compared CY 2007 claims processed through September 30, 2007, to CY 2008 claims processed through September 30, 2008, and represented about 60 percent of the full year data. We did not make any adjustments for inflation, changes in Medicare population, or other variables that potentially influenced billing between CY 2007 and CY 2008. A summary of these data analyses was included in the CY 2010 OPPS/ASC proposed rule (74 FR 35287 through 35289) and is reiterated above.

We note that we plan to present subsequent analyses that compare CY 2007 claims processed through September 30, 2007, to CY 2008 claims processed through September 30, 2008, and to CY 2009 claims processed

through September 30, 2009, to the APC panel at the APC Panel's winter 2010 meeting. We do not anticipate providing analyses using claims for services furnished during CY 2005 or CY 2006 because the packaging of the seven categories of services was effective for services furnished on and after January 1, 2008, and, therefore, we view CY 2007, the year immediately preceding the year that the packaging expansion went into effect, to be the base year for our comparisons. In addition, we do not anticipate providing the analyses at a provider-specific level or at a HCPCS code level. It is not clear to us how we would be able to use an analysis at the provider-specific level or the HCPCS code level or what value such an analysis would have in the context of national packaging policies for the OPSS.

We note that we make available a considerable amount of data for public analysis each year through the supporting data files that are posted on the CMS Web site in association with the display of the proposed and final rules. In addition, we make available the public use files of claims and a detailed narrative description of our data process for the annual OPSS/ASC proposed and final rules that the public can use to perform any desired analyses. Therefore, commenters are able to examine and analyze these data to develop specific information to support their requests for changes to payments under the OPSS, whether with regard to separate payment for a packaged service or other issues. We understand that the OPSS is a complex payment system and that it may be difficult to determine the quantitative amount of packaged cost included in the median cost for every independent service. However, based on the complex and detailed public comments on prior proposed rules that we have received, some commenters have performed meaningful analyses at a detailed and service-specific level based on the public claims data available.

With regard to the commenters' request that we not expand OPSS packaging until after we have produced data on the impact of packaging policy changes and consulted with stakeholders, we note that we establish all significant OPSS payment policies, including the packaging status of each HCPCS code, through the annual rulemaking process. Integral to this process is a detailed explanation of the claims data on which we base our proposals and the availability of the claims from which we develop those data for the use of the public to perform any level of analysis they choose.

Moreover, the OPSS/ASC annual rulemaking process provides a 60-day public comment period, as well as public presentations and discussion of the proposals at the summer APC Panel meeting. We also reply to all public comments that are within the scope of the OPSS/ASC proposed rule when we issue the OPSS/ASC final rule with comment period. In addition, we regularly meet with parties throughout the year who want to share their views on topics of interest to them. All of these activities and discussions provide significant information and opportunities for the public to influence and inform policy changes that we may be considering.

Comment: Some commenters expressed concern about the impact that packaging payment for services described by separate HCPCS codes could have on the submission of claims data by hospitals for those packaged codes and, therefore, with the validity of conclusions that could be drawn from impact analyses performed by CMS. One commenter questioned CMS' assumption that the OPSS packaging policies would allow continued collection of the data necessary to set appropriate, stable payment rates in the future. The commenter believed that greater packaging may eliminate hospitals' incentive to code for items and services for which separate payment is not made. The commenter further argued that CMS' past experiences with packaging payment for ancillary items and services indicate that hospitals do not report HCPCS codes for items and services that do not directly affect hospital payment. Similarly, the commenter explained that, under the IPPS, hospitals report only the data required to assign a case to the highest paying appropriate diagnosis-related group (DRG), even though other data might affect payment in the long term. The commenter saw no reason to believe that the current OPSS packaging approach would have a different outcome unless CMS gives clear instructions that hospitals should continue coding for all items and services used in the care of patients and provides an incentive to report packaged items and services.

Several commenters argued that the costs of new services are not reflected in the historical claims data CMS uses to set payment rates. The commenters believed that if CMS were to package payment for a new imaging service under the same criteria proposed for many existing imaging services, not only would CMS have no basis for determining how much the new service costs in its first 2 years of availability,

but also CMS would provide no incentive to hospitals to report codes and charges for the new service for use in future OPSS ratesetting. The commenters further asserted that the resulting incomplete data would lead to inappropriate payments for independent services that, in turn, would limit access to care and would discourage continued innovation to improve patient care.

Response: We do not believe that there will be a significant change in what hospitals report and charge for the outpatient services they furnish to Medicare beneficiaries and other patients as a result of our current packaging methodology. Medicare cost reporting standards specify that hospitals must impose the same charges for Medicare patients as for other patients. We are often told by hospitals that many private payers pay based on a percentage of charges and that, in accordance with Medicare cost reporting rules and generally accepted accounting principles, hospital chargemasters do not differentiate between the charges to Medicare patients and other patients. Therefore, we have no reason to believe that hospitals will stop reporting HCPCS codes and charges for packaged services they provide to Medicare beneficiaries. As we stated in the CY 2009 OPSS/ASC final rule with comment period (74 FR 68575), we strongly encourage hospitals to report a charge for each packaged service they furnish, either by billing the packaged HCPCS code and a charge for that service if separate reporting is consistent with CPT and CMS instructions, by increasing the charge for the separately paid associated service to include the charge for the packaged service, or by reporting the charge for the packaged service with an appropriate revenue code but without a HCPCS code. Any of these means of charging for the packaged service will result in the cost of the packaged service being incorporated into the cost we estimate for the separately paid service. If a HCPCS code is not reported when a packaged service is provided, we acknowledge that it can be challenging to specifically track the utilization patterns and resource cost of the packaged service itself. However, we have no reason to believe that hospitals have not considered the cost of the packaged service in reporting charges for the independent, separately paid service.

We expect that hospitals, as other prudent businesses, have a quality review process that ensures that they accurately and completely report the services they furnish, with appropriate charges for those services to Medicare

and all other payers. We encourage hospitals to report all HCPCS codes that describe packaged services that were furnished, unless the CPT Editorial Panel or CMS provides other guidance. To the extent that hospitals include separate charges for packaged services on their claims, the estimated costs of those packaged services are then added to the costs of separately paid procedures on the same claims and used

in establishing payment rates for the separately paid services.

Comment: One commenter argued that CMS' packaging methodology for guidance services used in radiation oncology procedures is not transparent. Specifically, the commenter claimed that CMS packaged payment for the radiation oncology image-guidance services (shown in Table 15) into the payment for independent radiation

therapy services (shown in Table 16) without publishing its packaging methodology. The commenter further stated that the lack of transparency regarding CMS' packaging methodology is of concern to the radiation oncology community, and that it would be helpful if CMS published the information used in the APC Panel's determination of packaging and payment rates.

TABLE 15—PACKAGED RADIATION ONCOLOGY GUIDANCE SERVICES

CY 2010 CPT code	CY 2010 Long descriptor
77421	Stereoscopic X ray guidance for localization of target volume for the delivery of radiation therapy.
77014	Computed tomography guidance for placement of radiation fields.
77417	Therapeutic radiology port film(s).
76950	Ultrasonic guidance for aspiration of ova, imaging supervision and interpretation.

TABLE 16—SEPARATELY PAID RADIATION THERAPY SERVICES

CY 2010 CPT code	CY 2010 Long descriptor
77402	Radiation treatment delivery, single treatment area, single port or parallel opposed ports, simple blocks or no blocks; up to 5MeV.
77403	Radiation treatment delivery, single treatment area, single port or parallel opposed ports, simple blocks or no blocks; 6–10MeV.
77404	Radiation treatment delivery, single treatment area, single port or parallel opposed ports, simple blocks or no blocks; 11–19 MeV.
77406	Radiation treatment delivery, single treatment area, single port or parallel opposed ports, simple blocks or no blocks; 20 MeV or greater.
77407	Radiation treatment delivery, 2 separate treatment areas, 3 or more ports on a single treatment area, use of multiple blocks; up to 5 MeV.
77408	Radiation treatment delivery, 2 separate treatment areas, 3 or more ports on a single treatment area, use of multiple blocks; 6–10 MeV.
77409	Radiation treatment delivery, 2 separate treatment areas, 3 or more ports on a single treatment area, use of multiple blocks; 11–19 MeV.
77411	Radiation treatment delivery, 2 separate treatment areas, 3 or more ports on a single treatment area, use of multiple blocks; 20 MeV or greater.
77412	Radiation treatment delivery, 3 or more separate treatment areas, custom blocking, tangential ports, wedges, rotational beam, compensators, electron beam; up to 5 MeV.
77413	Radiation treatment delivery, 3 or more separate treatment areas, custom blocking, tangential ports, wedges, rotational beam, compensators, electron beam; 6–10 MeV.
77414	Radiation treatment delivery, 3 or more separate treatment areas, custom blocking, tangential ports, wedges, rotational beam, compensators, electron beam; 11–19 MeV.
77416	Radiation treatment delivery, 3 or more separate treatment areas, custom blocking, tangential ports, wedges, rotational beam, compensators, electron beam; 20 MeV or greater.
77417	Therapeutic radiology port film(s).
77418	Intensity modulated treatment delivery, single or multiple fields/arcs, via narrow spatially and temporally modulated beams, binary, dynamic MLC, per treatment session.

Response: Although the APC Panel provides valuable advice with regard to the establishment of OPPS payment policies and payment rates, the APC Panel does not, as the commenter suggested, determine what services are packaged under the OPPS or establish OPPS payment rates. We adopt the OPPS payment policies regarding packaging and other issues and establish payment rates through the annual rulemaking cycle.

In general, payment for a packaged HCPCS code is included in the payment for the independent service with which it is associated, to the extent that the cost of the packaged service is reflected

on the single procedure claims that are used to calculate the median cost for the independent, separately paid service. We intend to further examine the packaging of image-guidance for radiation therapy in the analyses of the impact of packaging that we plan to discuss with the APC Panel at the winter 2010 meeting. However, as we describe earlier in this section, we make available a considerable amount of data for public analysis each year, provide the claims we use to calculate median costs, and provide a detailed narrative description of our data process that the public can use to analyze any topic of interest to them.

Comment: One commenter supported CMS' goal of increased efficiency in hospital outpatient care. However, the commenter was concerned that packaging payment for services too soon could create access problems for technologies that would otherwise improve patient outcomes and reduce costs. The commenter urged CMS to reinstate separate payment in CY 2010 for ICE, FFR, and IVUS until a thorough analysis has been performed on the impact of packaging payment for these services, including the rate of change in their utilization over time and market penetration.

Response: As discussed earlier in this section, in response to the request from the APC Panel that CMS evaluate the impact of expanded packaging on Medicare beneficiaries, we analyzed 9 months of CY 2007 and CY 2008 data related to all services that were packaged during CY 2008. Analysis of the intraoperative category (which includes IVUS, ICE, and FFR) showed minimal changes in the frequency and the number of hospitals reporting these packaged services between CY 2007 and CY 2008. The IVUS, ICE, and FFR services studied specifically included CPT codes 37250 (Intravascular ultrasound (non-coronary vessel) during diagnostic evaluation and/or therapeutic intervention; initial vessel); 37251 (Intravascular ultrasound (non-coronary vessel) during diagnostic evaluation and/or therapeutic intervention; each additional vessel); 92978 (Intravascular ultrasound (coronary vessel or graft) during diagnostic evaluation and/or therapeutic intervention including imaging supervision, interpretation and report; initial vessel); 92979 (Intravascular ultrasound (coronary vessel or graft) during diagnostic evaluation and/or therapeutic intervention including imaging supervision, interpretation and report; each additional vessel); 93662 (Intracardiac echocardiography during therapeutic/diagnostic intervention, including imaging supervision and interpretation); 93571 (Intravascular Doppler velocity and/or pressure derived coronary flow reserve measurement (coronary vessel or graft) during coronary angiography including pharmacologically induced stress, initial vessel); and 93572 (Intravascular Doppler velocity and/or pressure derived coronary flow reserve measurement (coronary vessel or graft) during coronary angiography including pharmacologically induced stress, each additional vessel).

As discussed previously, in February 2009 we presented an analysis to the APC Panel that showed an increase of 8 percent in the number of services billed and an increase in aggregate payment of 25 percent in CY 2008, when IVUS, ICE and FFR were packaged, in comparison to CY 2007 when IVUS, ICE and FFR were paid separately. Additionally, we intend to continue our analysis of the impact of greater packaging on Medicare beneficiaries and to present additional data to the APC Panel at the winter 2010 meeting.

We note that IVUS, ICE, and FFR services are existing, established technologies and that hospitals have provided some of these services in the

HOPD since the implementation of the OPPI in CY 2000. IVUS, FFR, and ICE are all dependent services that are always provided in association with independent services. Given the increase in the number of services furnished and the associated payment between CY 2007 and CY 2008, we have seen no evidence from our claims data that beneficiary access to care is being harmed by packaging payment for IVUS, ICE, and FFR services. We believe that packaging creates appropriate incentives for hospitals and their physician partners to carefully consider the technologies that are used in the care of patients, in order to ensure that technologies are selected for use in each case based on their expected benefit to a particular Medicare beneficiary.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to packaged payment for the seven categories of services, including guidance services, image processing services, intraoperative services, imaging supervision and interpretation services, diagnostic radiopharmaceuticals, contrast media, and observation services. We refer readers to section V.B.2.d. of this final rule with comment period for further discussion of our final policy to package payment for contrast agents and diagnostic radiopharmaceuticals. We refer readers to section II.A.2.e.(1) for further discussion of our final policy to pay for observation services through extended assessment and management composite APCs under certain circumstances. We plan to discuss with the APC Panel additional analyses of the impact of packaging these categories of services at the winter 2010 APC Panel meeting.

(3) Other Service-Specific Packaging Issues

The APC Panel also recommended that CMS reassign CPT code 76098 (Radiological examination, surgical specimen) from APC 0317 (Level II Miscellaneous Radiology Procedures) to APC 0260 (Level I Plain Film), and to place CPT code 76098 on the bypass list. Based on our analysis of the CY 2010 claims containing CPT 76098 and clinical review of the services being furnished, in the CY 2010 OPPI/ASC proposed rule (74 FR 35241), we proposed to treat CPT code 76098 as a "T-packaged" code for CY 2010 with continued assignment to APC 0317. As discussed above, a "T-packaged code," identified with status indicator "Q2," describes a code whose payment is packaged when one or more separately paid surgical procedures with a status

indicator of "T" are provided during the hospital encounter. The assignment of status indicator "Q2" to CPT code 76098 would result in more claims data being available to set the median costs for the surgical procedures with which CPT code 76098 is most commonly billed (for example, CPT code 19101 (Biopsy of breast, percutaneous, needle core, not using image guidance; open incisional)), while continuing to provide appropriate separate payment that reflects the costs of the service, including its packaged costs, when it is not billed with a surgical procedure. Further discussion related to the proposal is included in section II.A.1.b. of this final rule with comment period.

Comment: One commenter requested that CPT code 76098 remain separately payable instead of conditionally packaged. The commenter acknowledged that radiological examination of a surgical specimen is performed in conjunction with a surgical procedure most of the time but asserted that when the service is conditionally packaged, surgical procedure payment would not cover the cost of the radiological examination of a surgical specimen.

Response: We continue to believe that when CPT code 76098 is furnished on the same date of service as a major separately payable procedure, CPT code 76098 is a dependent service that is ancillary and supportive to the independent service with which it is performed and that, therefore, it is most appropriate to package the cost of CPT code 76098 into the payment for the independent, separately paid procedure. The full cost of CPT code 76098 is packaged into the cost of the independent, separately paid procedure to the extent that the hospital's charge for the packaged service, when reduced to cost by the hospital's applicable CCR, results in an accurate reflection of the cost of the packaged service. As we stated in the CY 2009 OPPI/ASC final rule with comment period (74 FR 68575), we strongly encourage hospitals to report a charge for each packaged service they furnish, either by billing the packaged HCPCS code and a charge for that service if separate reporting is consistent with CPT and CMS instructions, by increasing the charge for the separately paid associated service to include the charge for the packaged service, or by reporting the charge for the packaged service with an appropriate revenue code but without a HCPCS code. Any of these means of charging for the packaged service will result in the costs of the packaged service being incorporated into the cost we estimate for the separately paid

service. We note that further discussion of CPT code 76098 as it relates to the commenters' requests to add this code to the bypass list is included in section II.A.1.b. of this final rule with comment period.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to assign CPT code 76098 status indicator "Q2" to signify that the service is packaged when it is reported with a separately paid procedure that has a status indicator of "T" on the same date of service and separately paid under APC 0317 when it is not reported on the same date of service with a separately paid surgical procedure that has a status indicator of "T." The final CY 2010 APC median cost of APC 0317 is approximately \$374.

Comment: A number of commenters disagreed with CMS' proposal to package electrodiagnostic guidance for chemodenervation procedures. The commenters asserted that paying chemodenervation procedures at the same rate, regardless of the use of electrodiagnostic guidance, may discourage providers from using guidance to place a needle filled with a potentially fatal substance like botulinum toxin. They urged CMS to consider providing a separate payment for electrodiagnostic needle guidance to ensure that quality of care is not compromised.

Response: While the commenters did not identify specific chemodenervation guidance CPT codes, we note that the costs of chemodenervation guidance services, specifically CPT codes 95873 (Electrical stimulation for guidance in conjunction with chemodenervation (List separately in addition to code for primary procedure)) and 95874 (Needle electromyography for guidance in conjunction with chemodenervation (list separately in addition to code for primary procedure)) are reflected in the median costs of the independent, separately paid chemodenervation procedures as a function of the frequency that chemodenervation services are reported with a particular guidance CPT code. We recognize that, in some cases, supportive and ancillary dependent services are furnished at a high frequency with independent services, and in other cases, they are furnished with independent services at a low frequency. Nonetheless, we believe that packaging should reflect the reality of how these services are furnished. Moreover, we believe that hospitals make prudent and appropriate patient care decisions with regard to when they furnish packaged services.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to unconditionally package payment for chemodenervation guidance services described by CPT codes 95873 and 95874. These CPT codes are, therefore, assigned status indicator "N" in Addendum B to this final rule with comment period.

Comment: One commenter objected to the assignment of status indicator "N" to a number of guidance procedures and requested that CMS conditionally package these services so that they could be paid separately if they are the only services furnished in a hospital encounter. The commenter believed that it is not appropriate that the hospital receives no payment when these services are furnished in preparation for a surgical procedure that is canceled after the services have been furnished but before the patient is taken to the operating room. The procedures of concern to the commenter include CPT codes 19290 (Preoperative placement of needle localization wire, breast;); 19291 (Preoperative placement of needle localization wire, breast; each additional lesion (List separately in addition to code for primary procedure)); 19295 (Image guided placement, metallic localization clip, percutaneous, during breast biopsy (List separately in addition to code for primary procedure)); 77031 (Stereotactic localization guidance for breast biopsy or needle placement (e.g., for wire localization or for injection), each lesion, radiological supervision and interpretation); 77032 (Mammographic guidance for needle placement, breast (e.g., for wire localization or for injection), each lesion, radiological supervision and interpretation); and 76942 (Ultrasonic guidance for needle placement (e.g., biopsy, aspiration, injection, localization device), imaging supervision and interpretation).

Response: We appreciate the commenter's submission of this clinical scenario for our review. The APC Panel Packaging Subcommittee provides substantive advice to us on the packaging of services, either conditionally or unconditionally under the OPPS, and the Subcommittee members bring broad and deep expertise and experience to their review of clinical scenarios. Therefore, we will review these services and the scenario described by the commenter with the APC Panel's Packaging Subcommittee at the winter 2010 APC Panel meeting.

After review of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to

unconditionally package payment for CPT codes 19290, 19291, 19295, 77031, 77032, and 76942. These CPT codes are assigned status indicator "N" in Addendum B to this final rule with comment period. We will review the OPPS treatment of these CPT codes with the APC Panel Packaging Subcommittee at the winter 2010 APC Panel meeting.

Comment: One commenter suggested that it is very challenging for hospitals to determine they were paid correctly for services furnished because of CMS' "Q" status indicators and the complexity of determining which HCPCS codes should be separately paid. The commenter asked that CMS make the claims processing system more transparent.

Response: We acknowledge that the OPPS is a complex payment system and that it is difficult to determine the correct payment for a service that is subject to conditional packaging. Addendum D1 to this final rule with comment period describes how services that appear in Addendum B with status indicators "Q1," "Q2," and "Q3" are treated in claims processing. In the case of conditionally packaged codes with status indicators "Q1" or "Q2," where the criteria for separate payment are not met, these codes are treated as packaged services. We assign status indicators "Q1" and "Q2" to conditionally packaged services to indicate that they are usually packaged, except for special circumstances when they are separately payable. Through the I/OCE claims processing logic, the status indicator of a conditionally packaged service reported on a claim is changed either to "N" or the status indicator of the APC to which the code is assigned for separate payment, depending upon the presence or absence of other OPPS services also reported on the claim with the same date of service. Status indicator "Q3" indicates that the code is a member of a composite APC. Addendum M includes the HCPCS codes for all services that are paid either through single code APCs or composite APCs when the criteria for composite APC payment are met. A full discussion of the composite criteria for each composite APC (to which status indicator "Q3" applies) is included in section II.A.2.e. of this final rule with comment period.

In addition to the availability of these resources that describe whether a service is separately payable or packaged (in the case of services with status indicators "Q1" or "Q2") or a member of a composite APC (in the case of services with status indicator "Q3"), the quarterly I/OCE and the OPPS Pricer that are used by the Fiscal Intermediary

Standard System (FISS) to process claims paid under the OPSS are both available to the public each calendar quarter. The I/OCE instructions and specifications that are utilized for OPSS and non-OPSS payment for hospital outpatient services are available quarterly for download on the CMS Web site at: http://www.cms.hhs.gov/OutpatientCodeEdit/02_OCEQtrReleaseSpecs.asp#TopOfPage. Providers interested in purchasing the I/OCE software should visit the authorized distributor's Web site at <http://www.ntis.gov/products/oceapc.aspx> for more information on how to obtain the software. There is no OPSS Pricer application for personal computers at this time. However, providers can download the files that contain the logic, rates, wage indices, and off-set amounts used by the OPSS Pricer program to calculate APC rates, copayments, and deductibles from the CMS Web site at: http://www.cms.hhs.gov/PCPricer/08_OPSS.asp.

B. Conversion Factor Update

Section 1833(t)(3)(C)(ii) of the Act requires us to update the conversion factor used to determine payment rates under the OPSS on an annual basis. Under the authority in section 1833(t)(3)(C)(iv) of the Act, for CY 2010, the update is equal to the hospital inpatient market basket percentage increase applicable to hospital discharges under section 1886(b)(3)(B)(iii) of the Act. The final hospital market basket increase for FY 2010 published in the FY 2010 IPPS/LTCH PPS final rule (74 FR 44002) is 2.1 percent. To set the OPSS conversion factor for CY 2010, we increased the CY 2009 conversion factor of \$66.059, as specified in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68584 through 68585), by 2.1 percent. Hospitals that fail to meet the reporting requirements of the Hospital Outpatient Quality Data Reporting Program (HOP QDRP) are subject to a reduction of 2.0 percentage points from the market basket update to the conversion factor. For a complete discussion of the HOP QDRP requirements and the payment reduction for hospitals that fail to meet those requirements, we refer readers to section XVI. of this final rule with comment period.

In accordance with section 1833(t)(9)(B) of the Act, we further adjusted the conversion factor for CY 2010 to ensure that any revisions we made to our updates for a revised wage index and rural adjustment are made on a budget neutral basis. We calculated an overall budget neutrality factor of

0.9997 for wage index changes by comparing total payments from our simulation model using the FY 2010 IPPS final wage index values to those payments using the current (FY 2009) IPPS wage index values. For CY 2010, we did not propose a change to our rural adjustment policy. Therefore, the budget neutrality factor for the rural adjustment is 1.0000.

For this final rule with comment period, we estimated that pass-through spending for both drugs and biologicals and devices for CY 2010 will equal approximately \$45.5 million, which represents 0.14 percent of total projected CY 2010 OPSS spending. Therefore, the conversion factor was also adjusted by the difference between the 0.11 percent estimate of pass-through spending set aside for CY 2009 and the 0.14 percent estimate of CY 2010 pass-through spending. Finally, estimated payments for outliers remain at 1.0 percent of total OPSS payments for CY 2010.

The market basket increase update factor of 2.1 percent for CY 2010, the required wage index budget neutrality adjustment of approximately 0.9997, and the adjustment of 0.03 percent of projected OPSS spending for the difference in the pass-through spending set aside resulted in a full market basket conversion factor for CY 2010 of \$67.406. To calculate the CY 2010 reduced market basket conversion factor for those hospitals that fail to meet the requirements of the HOP QDRP for the full CY 2010 payment update, we made all other adjustments discussed above, but used a reduced market basket increase update factor of 0.1 percent. This resulted in a reduced market basket conversion factor for CY 2010 of \$66.086 for those hospitals that fail to meet the HOP QDRP requirements.

We did not receive any public comments on the calculation of the conversion factor. Therefore, we are finalizing our CY 2010 proposal, without modification, to update the OPSS conversion factor by the FY 2010 IPPS market basket increase update factor of 2.1 percent, resulting in a final full conversion factor of \$67.406 and in a reduced conversion factor of \$66.086 for those hospitals that fail to meet the HOP QDRP reporting requirements for the full CY 2010 payment update.

C. Wage Index Changes

Section 1833(t)(2)(D) of the Act requires the Secretary to determine a wage adjustment factor to adjust, for geographic wage differences, the portion of the OPSS payment rate, which includes the copayment standardized amount, that is attributable to labor and

labor-related cost. This adjustment must be made in a budget neutral manner and budget neutrality is discussed in section II.B. of this final rule with comment period.

The OPSS labor-related share is 60 percent of the national OPSS payment. This labor-related share is based on a regression analysis that determined that approximately 60 percent of the costs of services paid under the OPSS were attributable to wage costs. We confirmed that this labor-related share for outpatient services is still appropriate during our regression analysis for the payment adjustment for rural hospitals in the CY 2006 OPSS final rule with comment period (70 FR 68553). Therefore, in the CY 2010 OPSS/ASC proposed rule (74 FR 35291), we did not propose to revise this policy for the CY 2010 OPSS. We refer readers to section II.G. of this final rule with comment period for a description and example of how the wage index for a particular hospital is used to determine the payment for the hospital.

As discussed in section II.A.2.c. of this final rule with comment period, for estimating national median APC costs, we standardize 60 percent of estimated claims costs for geographic area wage variation using the same FY 2010 pre-reclassified wage indices that the IPPS uses to standardize costs. This standardization process removes the effects of differences in area wage levels from the determination of a national unadjusted OPSS payment rate and the copayment amount.

As published in the original OPSS April 7, 2000 final rule with comment period (65 FR 18545), the OPSS has consistently adopted the final IPPS wage indices as the wage indices for adjusting the OPSS standard payment amounts for labor market differences. Thus, the wage index that applies to a particular acute care short-stay hospital under the IPPS also applies to that hospital under the OPSS. As initially explained in the September 8, 1998 OPSS proposed rule, we believed and continue to believe that using the IPPS wage index as the source of an adjustment factor for the OPSS is reasonable and logical, given the inseparable, subordinate status of the HOPD within the hospital overall. In accordance with section 1886(d)(3)(E) of the Act, the IPPS wage index is updated annually. Therefore, in accordance with our established policy, we proposed to use the final FY 2010 version of the IPPS wage indices used to pay IPPS hospitals to adjust the CY 2010 OPSS payment rates and copayment amounts for geographic differences in labor cost for all providers that participate in the

OPPS, including providers that are not paid under the IPPS (referred to in this section as “non-IPPS” providers).

We note that the final FY 2010 IPPS wage indices continue to reflect a number of adjustments implemented over the past few years, including revised Office of Management and Budget (OMB) standards for defining geographic statistical areas (Core-Based Statistical Areas or CBSAs), reclassification to different geographic areas, rural floor provisions and the accompanying budget neutrality adjustment, an adjustment for out-migration labor patterns, an adjustment for occupational mix, and a policy for allocating hourly wage data among campuses of multicampus hospital systems that cross CBSAs. For the FY 2010 wage indices, these changes include a continuing transition to the new reclassification threshold criteria that were finalized in the FY 2009 IPPS final rule (73 FR 48568 through 48570), updated 2007–2008 occupational mix survey data, and a continuing transition to state-level budget neutrality for the rural and imputed floors. We refer readers to the FY 2010 IPPS/LTCH PPS final rule (74 FR 43823) for a detailed discussion of all final changes to the FY 2010 IPPS wage indices. In addition, we refer readers to the CY 2005 OPPS final rule with comment period (69 FR 65842 through 65844) and subsequent OPPS rules for a detailed discussion of the history of these wage index adjustments as applied under the OPPS.

The IPPS wage indices that we proposed to adopt in the CY 2010 OPPS/ASC proposed rule (74 FR 35291) include all reclassifications that are approved by the Medicare Geographic Classification Review Board (MGCRB) for FY 2010.

As noted in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68585), after issuance of the CY 2009 OPPS/ASC proposed rule, section 124 of Public Law 110–275 further extended geographic reclassifications under section 508 and certain special exception reclassifications until September 30, 2009. We did not make any proposals related to these provisions for the CY 2009 OPPS wage indices in our CY 2009 proposed rule because Public Law 110–275 was enacted after issuance of the CY 2009 OPPS/ASC proposed rule. In accordance with section 124 of Public Law 110–275, for CY 2009, we adopted all section 508 geographic reclassifications through September 30, 2009. Similar to our treatment of section 508 reclassifications extended under Public Law 110–173 as described in the CY 2009 OPPS/ASC final rule with

comment period (73 FR 68586), hospitals with section 508 reclassifications revert to their home area wage index, with out-migration adjustment if applicable, or a current MGCRB reclassification, from October 1, 2009 to December 31, 2009. As we did for CY 2008, we also have extended the special exception wage indices for certain hospitals through December 31, 2009, under the OPPS, in order to give these hospitals the special exception wage indices under the OPPS for the same time period as under the IPPS. We refer readers to the **Federal Register** notice published subsequent to the FY 2009 IPPS final rule for a detailed discussion of the changes to the wage indices as required by section 124 of Public Law 110–275 (73 FR 57888). Because the provisions of section 124 of Public Law 110–275 expire in 2009 and are not applicable to FY 2010, we did not make any proposals related to those provisions for the OPPS wage indices for CY 2010.

For purposes of the OPPS, we proposed to continue our policy in CY 2010 to allow non-IPPS hospitals paid under the OPPS to qualify for the out-migration adjustment if they are located in a section 505 out-migration county. We note that because non-IPPS hospitals cannot reclassify, they are eligible for the out-migration wage adjustment. Table 4J in the FY 2010 IPPS final rule (74 FR 44118 through 44125), as subsequently corrected at 74 FR 51506, identifies counties eligible for the out-migration adjustment and providers receiving the adjustment. As we have done in prior years, we are reprinting Table 4J, as corrected, as Addendum L to this final rule with comment period, with the addition of non-IPPS hospitals that will receive the section 505 out-migration adjustment under the CY 2010 OPPS.

As stated earlier in this section, we continue to believe that using the IPPS wage indices as the source of an adjustment factor for the OPPS is reasonable and logical, given the inseparable, subordinate status of the HOPD within the hospital overall. Therefore, we proposed to use the final FY 2010 IPPS wage indices for calculating the OPPS payments in CY 2010. With the exception of the out-migration wage adjustment table (Addendum L to this final rule with comment period), which includes non-IPPS hospitals paid under the OPPS, we are not reprinting the FY 2010 IPPS final wage indices referenced in this discussion of the wage index. We refer readers to the CMS Web site for the OPPS at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/>. At this link,

readers will find a link to the FY 2010 IPPS final wage index tables.

Comment: Several commenters expressed support for the CMS proposal to extend the IPPS wage indices to the OPPS in CY 2010, consistent with prior year policies under the OPPS.

Response: We appreciate the support expressed by commenters for our proposed CY 2010 wage index policies.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to use the final FY 2010 IPPS wage indices to adjust the OPPS standard payment amounts for labor market differences.

D. Statewide Average Default CCRs

In addition to using CCRs to estimate costs from charges on claims for ratesetting, CMS uses overall hospital-specific CCRs calculated from the hospital's most recent cost report to determine outlier payments, payments for pass-through devices, and monthly interim transitional corridor payments under the OPPS during the PPS year. Medicare contractors cannot calculate a CCR for some hospitals because there is no cost report available. For these hospitals, CMS uses the statewide average default CCRs to determine the payments mentioned above until a hospital's Medicare contractor is able to calculate the hospital's actual CCR from its most recently submitted Medicare cost report. These hospitals include, but are not limited to, hospitals that are new, have not accepted assignment of an existing hospital's provider agreement, and have not yet submitted a cost report. CMS also uses the statewide average default CCRs to determine payments for hospitals that appear to have a biased CCR (that is, the CCR falls outside the predetermined ceiling threshold for a valid CCR) or for hospitals whose most recent cost report reflects an all-inclusive rate status (Medicare Claims Processing Manual, Pub. 100–04, Chapter 4, Section 10.11). In the CY 2010 OPPS/ASC proposed rule (74 FR 35292), we proposed to update the default ratios for CY 2010 using the most recent cost report data. We discuss our policy for using default CCRs, including setting the ceiling threshold for a valid CCR, in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68594 through 68599) in the context of our adoption of an outlier reconciliation policy for cost reports beginning on or after January 1, 2009.

For CY 2010, we used our standard methodology of calculating the statewide average default CCRs using the same hospital overall CCRs that we

use to adjust charges to costs on claims data for setting the CY 2010 proposed OPPS relative weights. Table 12 that was published in the CY 2010 OPPS/ASC proposed rule (74 FR 35293 through 35294) listed the proposed CY 2010 default urban and rural CCRs by State and compared them to last year's default CCRs. These CCRs are the ratio of total costs to total charges from each hospital's most recently submitted cost report, for those cost centers relevant to outpatient services weighted by Medicare Part B charges. We also adjusted ratios from submitted cost reports to reflect final settled status by applying the differential between settled to submitted overall CCR for the cost centers relevant to outpatient services from the most recent pair of final settled and submitted cost reports. We then weighted each hospital's CCR by the volume of separately paid line-items on

hospital claims corresponding to the year of the majority of cost reports used to calculate the overall CCRs. We refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66680 through 66682) and prior OPPS rules for a more detailed discussion of our established methodology for calculating the statewide average default CCRs, including the hospitals used in our calculations and our trimming criteria.

For this CY 2010 OPPS/ASC final rule with comment period, approximately 44 percent of the submitted cost reports utilized in the default ratio calculations represented data for cost reporting periods ending in CY 2008 and 55 percent were for cost reporting periods ending in CY 2007. For Maryland, we used an overall weighted average CCR for all hospitals in the nation as a substitute for Maryland CCRs. Few hospitals in Maryland are eligible to

receive payment under the OPPS, which limits the data available to calculate an accurate and representative CCR. In general, observed changes in the statewide average default CCRs between CY 2009 and CY 2010 are modest and the few significant changes are associated with areas that have a small number of hospitals.

We did not receive any public comments concerning our CY 2010 proposal to apply our standard methodology of calculating the statewide average default CCRs using the same hospital overall CCRs that we use to adjust charges to costs on claims data for setting the CY 2010 proposed OPPS relative weights. Therefore, we are finalizing the statewide average default CCRs as shown in Table 17 below for OPPS services furnished on or after January 1, 2010.

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TABLE 17.—CY 2010 STATEWIDE AVERAGE CCRs

State	Urban/Rural	Final CY 2010 Default CCR	Previous Default CCR (CY 2009 OPPS Final Rule)
ALASKA	RURAL	0.499	0.562
ALASKA	URBAN	0.328	0.345
ALABAMA	RURAL	0.220	0.221
ALABAMA	URBAN	0.193	0.202
ARKANSAS	RURAL	0.251	0.256
ARKANSAS	URBAN	0.263	0.268
ARIZONA	RURAL	0.251	0.267
ARIZONA	URBAN	0.217	0.226
CALIFORNIA	RURAL	0.208	0.219
CALIFORNIA	URBAN	0.210	0.218
COLORADO	RURAL	0.345	0.346
COLORADO	URBAN	0.255	0.248
CONNECTICUT	RURAL	0.375	0.372
CONNECTICUT	URBAN	0.319	0.322
DISTRICT OF COLUMBIA	URBAN	0.324	0.329
DELAWARE	RURAL	0.320	0.302
DELAWARE	URBAN	0.363	0.349
FLORIDA	RURAL	0.198	0.204
FLORIDA	URBAN	0.184	0.189
GEORGIA	RURAL	0.265	0.267
GEORGIA	URBAN	0.246	0.251
HAWAII	RURAL	0.359	0.367
HAWAII	URBAN	0.307	0.344
IOWA	RURAL	0.332	0.439
IOWA	URBAN	0.302	0.294
IDAHO	RURAL	0.507	0.449
IDAHO	URBAN	0.409	0.419
ILLINOIS	RURAL	0.273	0.280
ILLINOIS	URBAN	0.253	0.266
INDIANA	RURAL	0.299	0.298
INDIANA	URBAN	0.296	0.295

State	Urban/Rural	Final CY 2010 Default CCR	Previous Default CCR (CY 2009 OPPS Final Rule)
KANSAS	RURAL	0.291	0.300
KANSAS	URBAN	0.226	0.238
KENTUCKY	RURAL	0.223	0.236
KENTUCKY	URBAN	0.254	0.255
LOUISIANA	RURAL	0.271	0.283
LOUISIANA	URBAN	0.259	0.258
MARYLAND	RURAL	0.294	0.303
MARYLAND	URBAN	0.267	0.276
MASSACHUSETTS	URBAN	0.323	0.328
MAINE	RURAL	0.433	0.452
MAINE	URBAN	0.452	0.428
MICHIGAN	RURAL	0.318	0.317
MICHIGAN	URBAN	0.320	0.321
MINNESOTA	RURAL	0.502	0.488
MINNESOTA	URBAN	0.330	0.348
MISSOURI	RURAL	0.266	0.269
MISSOURI	URBAN	0.270	0.282
MISSISSIPPI	RURAL	0.244	0.261
MISSISSIPPI	URBAN	0.192	0.209
MONTANA	RURAL	0.438	0.455
MONTANA	URBAN	0.462	0.439
NORTH CAROLINA	RURAL	0.270	0.272
NORTH CAROLINA	URBAN	0.285	0.292
NORTH DAKOTA	RURAL	0.333	0.369
NORTH DAKOTA	URBAN	0.361	0.354
NEBRASKA	RURAL	0.340	0.345
NEBRASKA	URBAN	0.260	0.283
NEW HAMPSHIRE	RURAL	0.329	0.350
NEW HAMPSHIRE	URBAN	0.285	0.296
NEW JERSEY	URBAN	0.235	0.257
NEW MEXICO	RURAL	0.259	0.263
NEW MEXICO	URBAN	0.329	0.328
NEVADA	RURAL	0.296	0.312
NEVADA	URBAN	0.187	0.192
NEW YORK	RURAL	0.423	0.412

State	Urban/Rural	Final CY 2010 Default CCR	Previous Default CCR (CY 2009 OPPS Final Rule)
NEW YORK	URBAN	0.383	0.388
OHIO	RURAL	0.350	0.353
OHIO	URBAN	0.250	0.258
OKLAHOMA	RURAL	0.267	0.278
OKLAHOMA	URBAN	0.225	0.238
OREGON	RURAL	0.303	0.318
OREGON	URBAN	0.344	0.374
PENNSYLVANIA	RURAL	0.280	0.284
PENNSYLVANIA	URBAN	0.223	0.232
PUERTO RICO	URBAN	0.514	0.519
RHODE ISLAND	URBAN	0.299	0.294
SOUTH CAROLINA	RURAL	0.232	0.242
SOUTH CAROLINA	URBAN	0.242	0.240
SOUTH DAKOTA	RURAL	0.320	0.336
SOUTH DAKOTA	URBAN	0.261	0.267
TENNESSEE	RURAL	0.233	0.244
TENNESSEE	URBAN	0.214	0.221
TEXAS	RURAL	0.251	0.257
TEXAS	URBAN	0.222	0.238
UTAH	RURAL	0.397	0.413
UTAH	URBAN	0.400	0.430
VIRGINIA	RURAL	0.242	0.257
VIRGINIA	URBAN	0.255	0.266
VERMONT	RURAL	0.413	0.406
VERMONT	URBAN	0.397	0.422
WASHINGTON	RURAL	0.365	0.349
WASHINGTON	URBAN	0.340	0.342
WISCONSIN	RURAL	0.384	0.399
WISCONSIN	URBAN	0.329	0.346
WEST VIRGINIA	RURAL	0.283	0.293
WEST VIRGINIA	URBAN	0.339	0.349
WYOMING	RURAL	0.407	0.418
WYOMING	URBAN	0.315	0.331

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E. OPSS Payment to Certain Rural and Other Hospitals

1. Hold Harmless Transitional Payment Changes Made by Public Law 110-275 (MIPPA)

When the OPSS was implemented, every provider was eligible to receive an

additional payment adjustment (called either transitional corridor payments or transitional outpatient payment (TOPs)) if the payments it received for covered OPD services under the OPSS were less than the payments it would have received for the same services under the

prior reasonable cost-based system (referred to as the pre-BBA amount). Section 1833(t)(7) of the Act provides that the transitional corridor payments are temporary payments for most providers and were intended to ease their transition from the prior reasonable cost-based payment system to the OPSS system. There are two exceptions to this provision, cancer hospitals and children's hospitals, and those hospitals receive the transitional corridor payments on a permanent basis. Section 1833(t)(7)(D)(i) of the Act originally provided for transitional corridor payments to rural hospitals with 100 or fewer beds for covered OPD services furnished before January 1, 2004. However, section 411 of Public Law 108-173 amended section 1833(t)(7)(D)(i) of the Act to extend these payments through December 31, 2005, for rural hospitals with 100 or fewer beds. Section 411 also extended the transitional corridor payments to SCHs located in rural areas for services furnished during the period that began with the provider's first cost reporting period beginning on or after January 1, 2004, and ended on December 31, 2005. Accordingly, the authority for making transitional corridor payments under section 1833(t)(7)(D)(i) of the Act, as amended by section 411 of Public Law 108-173, for rural hospitals having 100 or fewer beds and SCHs located in rural areas expired on December 31, 2005.

Section 5105 of Public Law 109-171 reinstated the TOPs for covered OPD services furnished on or after January 1, 2006, and before January 1, 2009, for rural hospitals having 100 or fewer beds that are not SCHs. When the OPSS payment was less than the provider's pre-BBA amount, the amount of payment was increased by 95 percent of the amount of the difference between the two payment systems for CY 2006, by 90 percent of the amount of that difference for CY 2007, and by 85 percent of the amount of that difference for CY 2008.

For CY 2006, we implemented section 5105 of Public Law 109-171 through Transmittal 877, issued on February 24, 2006. In the Transmittal, we did not specifically address whether TOPs apply to essential access community hospitals (EACHs), which are considered to be SCHs under section 1886(d)(5)(D)(iii)(III) of the Act. Accordingly, under the statute, EACHs are treated as SCHs. In the CY 2007 OPSS/ASC final rule with comment period (71 FR 68010), we stated that EACHs were not eligible for TOPs under Public Law 109-171. However, we stated they were eligible for the adjustment for rural SCHs. In the CY

2007 OPSS/ASC final rule with comment period (71 FR 68010 and 68228), we updated § 419.70(d) of our regulations to reflect the requirements of Public Law 109-171.

In the CY 2009 OPSS/ASC proposed rule (73 FR 41461), we stated that, effective for services provided on or after January 1, 2009, rural hospitals having 100 or fewer beds that are not SCHs would no longer be eligible for TOPs, in accordance with section 5105 of Public Law 109-171. However, subsequent to issuance of the CY 2009 OPSS/ASC proposed rule, section 147 of Public Law 110-275 amended section 1833(t)(7)(D)(i) of the Act by extending the period of TOPs to rural hospitals with 100 beds or fewer for 1 year, for services provided before January 1, 2010. Section 147 of Public Law 110-275 also extended TOPs to SCHs (including EACHs) with 100 or fewer beds for covered OPD services provided on or after January 1, 2009, and before January 1, 2010. In accordance with section 147 of Public Law 110-275, when the OPSS payment is less than the provider's pre-BBA amount, the amount of payment is increased by 85 percent of the amount of the difference between the two payment systems for CY 2009.

For CY 2009, we revised our regulations at §§ 419.70(d)(2) and (d)(4) and added a new paragraph (d)(5) to incorporate the provisions of section 147 of Public Law 110-275. In addition, we made other technical changes to § 419.70(d)(2) to more precisely capture our existing policy and to correct an inaccurate cross-reference. We also made technical corrections to the cross-references in paragraphs (e), (g), and (i) of § 419.70. In the CY 2010 OPSS/ASC proposed rule (74 FR 35295), for CY 2010, we proposed to make a technical correction to the heading of § 419.70(d)(5) to correctly identify the policy as described in the subsequent regulation text. The paragraph heading should indicate that the adjustment applies to small SCHs, rather than to rural SCHs.

Effective for services provided on or after January 1, 2010, rural hospitals and SCHs (including EACHs) having 100 or fewer beds will no longer be eligible for hold harmless TOPs, in accordance with section 147 of Public Law 110-275.

2. Adjustment for Rural SCHs Implemented in CY 2006 Related to Public Law 108-173 (MMA)

In the CY 2006 OPSS final rule with comment period (70 FR 68556), we finalized a payment increase for rural SCHs of 7.1 percent for all services and procedures paid under the OPSS, excluding drugs, biologicals,

brachytherapy sources, and devices paid under the pass-through payment policy in accordance with section 1833(t)(13)(B) of the Act, as added by section 411 of Public Law 108-173. Section 411 gave the Secretary the authority to make an adjustment to OPSS payments for rural hospitals, effective January 1, 2006, if justified by a study of the difference in costs by APC between hospitals in rural areas and hospitals in urban areas. Our analysis showed a difference in costs for rural SCHs. Therefore, for the CY 2006 OPSS, we finalized a payment adjustment for rural SCHs of 7.1 percent for all services and procedures paid under the OPSS, excluding separately payable drugs and biologicals, brachytherapy sources, and devices paid under the pass-through payment policy, in accordance with section 1833(t)(13)(B) of the Act.

In CY 2007, we became aware that we did not specifically address whether the adjustment applies to EACHs, which are considered to be SCHs under section 1886(d)(5)(D)(iii)(III) of the Act. Thus, under the statute, EACHs are treated as SCHs. Therefore, in the CY 2007 OPSS/ASC final rule with comment period (71 FR 68010 and 68227), for purposes of receiving this rural adjustment, we revised § 419.43(g) to clarify that EACHs are also eligible to receive the rural SCH adjustment, assuming these entities otherwise meet the rural adjustment criteria. Currently, fewer than 10 hospitals are classified as EACHs and as of CY 1998, under section 4201(c) of Public Law 105-33, a hospital can no longer become newly classified as an EACH.

This adjustment for rural SCHs is budget neutral and applied before calculating outliers and copayment. As stated in the CY 2006 OPSS final rule with comment period (70 FR 68560), we would not reestablish the adjustment amount on an annual basis, but we may review the adjustment in the future and, if appropriate, would revise the adjustment. We provided the same 7.1 percent adjustment to rural SCHs, including EACHs, again in CY 2008 and CY 2009. Further, in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68590), we updated the regulations at § 419.43(g)(4) to specify, in general terms, that items paid at charges adjusted to costs by application of a hospital-specific CCR are excluded from the 7.1 percent payment adjustment.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35295), for the CY 2010 OPSS, we proposed to continue our policy of a budget neutral 7.1 percent payment adjustment for rural SCHs, including EACHs, for all services and procedures paid under the OPSS,

excluding separately payable drugs and biologicals, devices paid under the pass-through payment policy, and items paid at charges reduced to costs. We intend to reassess the 7.1 percent adjustment in the near future by examining differences between urban and rural hospitals' costs using updated claims, cost reports, and provider information.

Comment: A number of commenters generally supported the proposal to continue the rural SCH (including EACHs) adjustment for CY 2010 OPPS. Several commenters also asked that CMS extend for CY 2010 the TOPs payment policies that were in effect for CY 2009. The commenters recommended that CMS evaluate the differences in cost between urban and rural hospitals over an extended 3-year period using updated claims, cost reports, and provider information. They further suggested that, during the 3-year period in which CMS would be gathering data, CMS pay SCHs and rural hospitals with less than 100 beds that are not SCHs the greater of the TOPs payment in effect for CY 2009 or the OPPS payment for the applicable calendar year plus the 7.1 percent rural adjustment, whichever is greater. The commenters claimed that CMS' reversal of the TOPs allowance after only 1 year of reimplementation for certain rural hospitals was unreasonable and could irreparably harm those rural hospitals absent a safety net mechanism in place.

Response: We agree that it is appropriate to continue the 7.1 percent adjustment for rural SCHs (including EACHs) as we proposed for CY 2010. However, we are not extending the CY 2009 TOPs payment policies for rural hospitals with 100 beds or less and for SCHs (including EACHs) with 100 or fewer beds for CY 2010. Section 1833(t)(7)(D)(i)(II) of the Act provides that, in the case of a hospital located in a rural area with 100 beds or fewer and that is not a sole community hospital, for covered OPD services furnished on or after January 1, 2006 and before January 1, 2010, for which the PPS amount is less than the pre-BBA amount, the amount of payment should be increased by the applicable percentage of the amount of such difference. Section 1833(t)(7)(D)(i)(III) of the Act also extends TOPs to SCHs (including EACHs) with 100 or fewer beds for covered OPD services provided on or after January 1, 2009 and before January 1, 2010, under the specific circumstances outlined in the statute. Therefore, sections 1833(t)(D)(i)(II) and (III) of the Act specifically expire TOPs payment to these categories of hospitals for services furnished on and after January 1, 2010. Accordingly, in CY

2010, neither rural SCHs nor rural hospitals with less than 100 beds will receive payment at whichever is greater, the TOPs payment in place for CY 2009 or payment for CY 2010, which includes the rural adjustment for rural SCHs, because sections 1833(t)(7)(D)(i)(II) and (III) of the Act expire TOPs payments as explained above. As we indicate above, we intend to reassess the 7.1 percent rural adjustment in the near future by examining differences between urban and rural hospitals' costs using updated claims, cost reports, and provider information.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to apply the 7.1 percent payment adjustment to rural SCHs for most services paid under the CY 2010 OPPS, excluding drugs, biologicals, and devices paid under the pass-through payment policy, and items paid at charges adjusted to cost. We also are making a technical correction to the heading of § 419.70(d)(5) to correctly identify the policy described in the regulation text of § 419.70(d)(5). The paragraph heading indicates that the adjustment applies to small SCHs, rather than to rural SCHs.

F. Hospital Outpatient Outlier Payments

1. Background

Currently, the OPPS pays outlier payments on a service-by-service basis. For CY 2009, the outlier threshold is met when the cost of furnishing a service or procedure by a hospital exceeds 1.75 times the APC payment amount and exceeds the APC payment rate plus a \$1,800 fixed-dollar threshold. We introduced a fixed-dollar threshold in CY 2005 in addition to the traditional multiple threshold in order to better target outliers to those high cost and complex procedures where a very costly service could present a hospital with significant financial loss. If the cost of a service meets both of these conditions, the multiple threshold and the fixed-dollar threshold, the outlier payment is calculated as 50 percent of the amount by which the cost of furnishing the service exceeds 1.75 times the APC payment rate. Before CY 2009, this outlier payment had historically been considered a final payment by longstanding OPPS policy. We implemented a reconciliation process similar to the IPPS outlier reconciliation process for cost reports with cost reporting periods beginning on or after January 1, 2009 (73 FR 68594 through 68599).

It has been our policy for the past several years to report the actual amount

of outlier payments as a percent of total spending in the claims being used to model the proposed OPPS. We previously estimated that CY 2008 outlier payments were approximately 0.73 percent of the total CY 2008 OPPS payments (73 FR 68592). Our current estimate of total outlier payments as a percent of total CY 2008 OPPS payment, using CY 2008 claims processed through June 30, 2009, and the revised OPPS expenditure estimate for the 2009 Trustees Report, is approximately 1.2 percent of the total aggregated OPPS payments. Therefore, for CY 2008, we estimate that we paid approximately 0.2 percent more than the CY 2008 outlier target of 1.0 percent of total aggregated OPPS payments.

As explained in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68594), we set our projected target for aggregate outlier payments at 1.0 percent of the aggregate total payments under the OPPS for CY 2009. The outlier thresholds were set so that estimated CY 2009 aggregate outlier payments would equal 1.0 percent of the total aggregated payments under the OPPS. Using our final rule CY 2008 claims data and CY 2009 payment rates, we currently estimate that the aggregate outlier payments for CY 2009 would be approximately 1.03 percent of the total CY 2009 OPPS payments. The difference between 1.0 percent and 1.03 percent is reflected in the regulatory impact analysis in section XXI.B. of this final rule with comment period. We note that we provide estimated CY 2010 outlier payments for hospitals and CMHCs with claims included in the claims data that we used to model impacts in the Hospital-Specific Impacts—Provider-Specific Data file on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/>.

2. Outlier Calculation

In the CY 2010 OPPS/ASC proposed rule (74 FR 35296), we proposed to continue our policy of estimating outlier payments to be 1.0 percent of the estimated aggregate total payments under the OPPS in CY 2010. We proposed that a portion of that 1.0 percent, specifically 0.02 percent, would be allocated to CMHCs for PHP outlier payments. This is the amount of estimated outlier payments that would result from the proposed CMHC outlier threshold as a proportion of total estimated outlier payments. As discussed in section X.C. of this final rule with comment period, for CMHCs, we proposed that if a CMHC's cost for partial hospitalization services, paid under either APC 0172 (Level I Partial

Hospitalization (3 services)) or APC 0173 (Level II Partial Hospitalization (4 or more services)), exceeds 3.40 times the payment for APC 0173, the outlier payment would be calculated as 50 percent of the amount by which the cost exceeds 3.40 times the APC 0173 payment rate. For further discussion of CMHC outlier payments, we refer readers to section X.C. of this final rule with comment period.

To ensure that the estimated CY 2010 aggregate outlier payments would equal 1.0 percent of estimated aggregate total payments under the OPSS, we proposed that the hospital outlier threshold be set so that outlier payments would be triggered when the cost of furnishing a service or procedure by a hospital exceeds 1.75 times the APC payment amount and exceeds the APC payment rate plus a \$2,225 fixed-dollar threshold. This proposed threshold reflected the methodology discussed below in this section, as well as the proposed APC recalibration for CY 2010.

We calculated the fixed-dollar threshold for the CY 2010 OPSS/ASC proposed rule using largely the same methodology as we did in CY 2009 (73 FR 41462). For purposes of estimating outlier payments for the CY 2010 OPSS/ASC proposed rule, we used the hospital-specific overall ancillary CCRs available in the April 2009 update to the Outpatient Provider-Specific File (OPSF). The OPSF contains provider-specific data, such as the most current CCR, which are maintained by the Medicare contractors and used by the OPSS Pricer to pay claims. The claims that we use to model each OPSS update lag by 2 years. For the CY 2010 OPSS/ASC proposed rule, we used CY 2008 claims to model the CY 2010 OPSS. In order to estimate the CY 2010 hospital outlier payments for the CY 2010 OPSS/ASC proposed rule, we inflated the charges on the CY 2008 claims using the same inflation factor of 1.1511 that we used to estimate the IPPS fixed-dollar outlier threshold for the FY 2010 IPPS/LTCH PPS proposed rule (74 FR 24245). For 1 year, the inflation factor we used was 1.0729. The methodology for determining this charge inflation factor was discussed in the FY 2010 IPPS/LTCH PPS proposed rule (74 FR 24245). As we stated in the CY 2005 OPSS final rule with comment period (69 FR 65845), we believe that the use of this charge inflation factor is appropriate for the OPSS because, with the exception of the routine service cost centers, hospitals use the same cost centers to capture costs and charges across inpatient and outpatient services.

As noted in the CY 2007 OPSS/ASC final rule with comment period (71 FR 68011), we are concerned that we could systematically overestimate the OPSS hospital outlier threshold if we did not apply a CCR inflation adjustment factor. Therefore, we proposed to apply the same CCR inflation adjustment factor that we proposed to apply for the FY 2010 IPPS outlier calculation to the CCRs used to simulate the CY 2010 OPSS outlier payments that determine the fixed-dollar threshold. Specifically, for CY 2010, we proposed to apply an adjustment of 0.9840 to the CCRs that were in the April 2009 OPSF to trend them forward from CY 2009 to CY 2010. The methodology for calculating this adjustment is discussed in the FY 2010 IPPS/LTCH PPS proposed rule (74 FR 24245 through 24247) and the FY 2010 IPPS/LTCH PPS final rule (74 FR 44007 through 44011).

Therefore, to model hospital outlier payments for the CY 2010 OPSS/ASC proposed rule, we applied the overall CCRs from the April 2009 OPSF file after adjustment (using the proposed CCR inflation adjustment factor of 0.9840 to approximate CY 2010 CCRs) to charges on CY 2008 claims that were adjusted (using the proposed charge inflation factor of 1.1511 to approximate CY 2010 charges). We simulated aggregated CY 2010 hospital outlier payments using these costs for several different fixed-dollar thresholds, holding the 1.75 multiple threshold constant and assuming that outlier payment would continue to be made at 50 percent of the amount by which the cost of furnishing the service would exceed 1.75 times the APC payment amount, until the total outlier payments equaled 1.0 percent of aggregated estimated total CY 2010 OPSS payments. We estimated that a proposed fixed-dollar threshold of \$2,225, combined with the proposed multiple threshold of 1.75 times the APC payment rate, would allocate 1.0 percent of aggregated total OPSS payments to outlier payments. We proposed to continue to make an outlier payment that equals 50 percent of the amount by which the cost of furnishing the service exceeds 1.75 times the APC payment amount when both the 1.75 multiple threshold and the proposed fixed-dollar \$2,225 threshold are met. For CMHCs, if a CMHC's cost for partial hospitalization services, paid under either APC 0172 or APC 0173, exceeds 3.40 times the payment for APC 0173, the outlier payment would be calculated as 50 percent of the amount by which the cost exceeds 3.40 times the APC 0173 payment rate.

Section 1833(t)(17)(A) of the Act, which applies to hospitals as defined under section 1886(d)(1)(B) of the Act, requires that hospitals that fail to report data required for the quality measures selected by the Secretary, in the form and manner required by the Secretary under 1833(t)(17)(B) of the Act, incur a 2.0 percentage point reduction to their OPD fee schedule increase factor, that is, the annual payment update factor. The application of a reduced OPD fee schedule increase factor results in reduced national unadjusted payment rates that will apply to certain outpatient items and services furnished by hospitals that are required to report outpatient quality data and that fail to meet the HOP QDRP requirements. For hospitals that fail to meet the HOP QDRP requirements, we proposed to continue our policy that we implemented in CY 2009 that the hospitals' costs would be compared to the reduced payments for purposes of outlier eligibility and payment calculation. For more information on the HOP QDRP, we refer readers to section XVI. of this final rule with comment period.

Comment: Several commenters supported the proposal to increase the outlier fixed-dollar threshold to maintain a target outlier spending percentage of 1.0 percent. One commenter requested that CMS not overestimate the fixed-dollar outlier threshold by decreasing the CY 2010 proposed threshold proportionally to only account for the amount Medicare paid in excess of the 1 percent target outlier percentage in CY 2009. A few commenters suggested that the target outlier spending percentage be raised. One commenter recommended that the target outlier spending percentage be raised to maintain the \$1,800 fixed-dollar threshold that is in effect for CY 2009. Another commenter requested that CMS increase the amount of outlier payment from 50 percent to 80 percent of the difference between the OPSS payment and the estimated provider cost for the service to make OPSS outlier policy more consistent with IPPS outlier policy. One commenter expressed concern that changes in outlier payments disproportionately affected the safety net hospitals. One commenter supported the proposal to use the same assumptions regarding charge inflation and CCR inflation as under the IPPS.

Response: We appreciate the commenters' support regarding the development of the OPSS outlier policy. We are not raising the threshold to recover the 0.03 percent of OPSS payment that we estimate was paid in

addition to the target outlier percent of 1 percent for CY 2009 because we do not adjust the fixed-dollar threshold in future years for either paying too much or too little in outlier payments in past years. We are not increasing the percent of total OPPS payment that we attribute to outlier payments, either for general purposes or to maintain the \$1,800 threshold for CY 2010, because we continue to believe that it is appropriate to maintain the target outlier percentage of 1 percent of total payment under the OPPS and to have a fixed-dollar threshold so that OPPS outlier payments are made only where the hospital would experience a significant loss for supplying a particular service. Similarly, we are not increasing the outlier payment percentage from 50 percent to 80 percent of the difference between the amount by which the cost of furnishing the service exceeds 1.75 times the APC payment rate because we do not believe that hospitals carry the same level of risk when they furnish outpatient hospital services as when they furnish inpatient hospital services. OPPS outlier payments are intended to protect hospitals from excessive losses when providing an extraordinarily costly service, and we believe that the potential for loss when furnishing OPPS services is limited. Payment bundles under the OPPS are small relative to those under the IPPS, and the OPPS pays separately for many services. The OPPS would pay hospitals for many individual services provided to a very costly patient reducing their financial risk. Patients for whom a hospital may incur extraordinary costs for providing individual OPPS services would usually require hospital admission. As described above, outlier payments are designed to protect hospitals from financial risk in providing services to costly patients, and are not designed to affect any specific hospital classes, such as safety net hospitals. With regard to the application of charge inflation factors, we agree that the charge inflation factors that apply to inpatient hospital services are equally applicable to services provided under the OPPS. Therefore, as specified below, we are applying the charge inflation factors that were used to calculate the outlier fixed-dollar threshold for the IPPS in the calculation of the fixed-dollar threshold for the CY 2010 OPPS.

Comment: Several commenters asked that CMS eliminate outlier payments for CMHCs and use the funds allocated to outlier payments for CMHCs to increase payments for services provided by CMHCs.

Response: Outlier payments to CMHCs are discussed in section X.C. of

this final rule with public comment. We respond to this comment as part of that discussion.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal for the outlier calculation, without modification, as outlined below.

3. Final Outlier Calculation

For CY 2010, we are applying the overall CCRs from the July 2009 OPSF file with a CCR adjustment factor of 0.988 to approximate CY 2010 CCRs to charges on the final CY 2008 claims that were adjusted to approximate CY 2010 charges (using the final 2-year charge inflation factor of 1.1418). We simulated aggregated CY 2010 hospital outlier payments using these costs for several different fixed-dollar thresholds, holding the 1.75 multiple threshold constant and assuming that outlier payment would continue to be made at 50 percent of the amount by which the cost of furnishing the service would exceed 1.75 times the APC payment amount, until the total outlier payments equaled 1.0 percent of aggregated estimated total CY 2010 OPPS payments. We estimate that a fixed-dollar threshold of \$2,175, combined with the multiple threshold of 1.75 times the APC payment rate, will allocate 1.0 percent of aggregated total OPPS payments to outlier payments.

In summary, for CY 2010, we will continue to make an outlier payment that equals 50 percent of the amount by which the cost of furnishing the service exceeds 1.75 times the APC payment amount when both the 1.75 multiple threshold and the final fixed-dollar \$2,175 threshold are met. For CMHCs, if a CMHC's cost for PHP services, paid under either APC 0172 or APC 0173, exceeds 3.40 times the payment for APC 0173, the outlier payment is calculated as 50 percent of the amount by which the cost exceeds 3.40 times the APC 0173 payment rate. We estimate that this threshold will allocate 0.03 percent of outlier payments to CMHCs for PHP outlier payments.

4. Outlier Reconciliation

In the CY 2009 OPPS/ASC final rule with comment period (73 CFR 68599), we adopted as final policy a process to reconcile hospital or CMHC outlier payments at cost report settlement for services furnished during cost reporting periods beginning in CY 2009. OPPS outlier reconciliation ensures accurate outlier payments for those facilities whose CCRs fluctuate significantly relative to the CCRs of other facilities, and who receive a significant amount of outlier payments. OPPS outlier

reconciliation thresholds are provided in the Medicare Claims Processing Manual (Pub. 100-4), Chapter 4, Section 10.7.2.1, reevaluated annually, and modified if necessary. When the cost report is settled, reconciliation of outlier payments will be based on the hospital-specific overall ancillary CCR, calculated as the ratio of costs and charges computed from the cost report at the time the cost report coinciding with the service dates is settled. Reconciling outlier payments ensures that the outlier payments made are appropriate and that final outlier payments reflect the most accurate cost data. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68599), we also finalized a proposal to adjust the amount of final outlier payments determined during reconciliation for the time value of money. The OPPS outlier reconciliation process will require recalculating outlier payments for individual claims in order to accurately determine the net effect of a change in a hospital's or CMHC's overall CCR on the facility's total outlier payments. For cost reporting periods beginning in CY 2009, Medicare contractors will begin to identify cost reports that require outlier reconciliation as a component of cost report settlement. At this time, CMS continues to develop a method for reexamining claims to calculate the change in total outlier payments in order to reconcile outlier payments for these cost reports.

As under the IPPS, we do not adjust the fixed-dollar threshold or amount of total OPPS payment set aside for outlier payments for reconciliation activity. The predictability of the fixed-dollar threshold is an important component of a prospective payment system. We do not adjust the prospectively set outlier threshold for the amount of outlier payment reconciled at cost report settlement because such action would be contrary to the prospective nature of the system. Our outlier threshold calculation assumes that overall ancillary CCRs accurately estimate hospital costs based on the information available to us at the time we set the prospective fixed-dollar outlier threshold. For these reasons, we are not incorporating any assumptions about the effects of reconciliation into our calculation of the OPPS fixed-dollar outlier threshold.

Comment: A number of commenters asked that CMS report the amount of outlier reconciliation activity, including aggregate amounts recovered by provider type and region. They suggested that, if the reconciled amounts are significant, these amounts

should be factored into the annual fixed-dollar outlier threshold. Several commenters supported the current reconciliation thresholds identified in the CMS manual (Medicare Claims Processing Manual (Pub. 100-04), Chapter 4, Section 10.7.2.1). One commenter asked that CMS apply the outlier reconciliation thresholds established in manual instructions to the claims used for estimating outlier payment and the fixed-dollar threshold to achieve the most accurate estimates possible.

Response: We revised Worksheet E, Part B, of the Medicare hospital cost report form CMS 2552-10 to collect OPSS outlier reconciliation information for cost reports beginning on or after January 1, 2009. This information will be available to the public through the Hospital Cost Report Information System (HCRIS). We do not expect to take outlier reconciliation amounts into account in our projections of future outlier payments. We believe that the reconciliation CCR and outlier payment thresholds implemented in the final rule (73 CFR 68599) are generous and that most hospitals will not be subject to outlier reconciliation upon cost report settlement. Further, it is difficult to predict the specific hospitals that will have CCRs and outlier payments reconciled in any given year. We also note that reconciliation occurs because hospitals' actual CCRs for the cost reporting period are different than the interim CCRs used to calculate outlier payment when a bill is processed. Our fixed-dollar threshold calculation assumes that CCRs accurately estimate hospital costs based on information available to us at the time we set the prospective fixed-dollar outlier threshold. We do not believe that estimating the fixed-dollar threshold to estimate the amount of payment that may be recovered as a result of outlier reconciliation in any given year would necessarily result in a more accurate estimate of outlier payments or a more accurate calculation of the fixed-dollar threshold for outlier payment for the prospective payment year. For these reasons, we will not make any assumptions about the amount of anticipated reconciliation of outlier payments on the outlier threshold calculation.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, for an OPSS outlier reconciliation policy. We are implementing the outlier reconciliation policy for each hospital and CMHC for services furnished during cost reporting periods beginning in CY 2010, and we

are including an adjustment for the time value of money.

G. Calculation of an Adjusted Medicare Payment From the National Unadjusted Medicare Payment

The basic methodology for determining prospective payment rates for HOPD services under the OPSS is set forth in existing regulations at 42 CFR part 419, subparts C and D. The payment rate for most services and procedures for which payment is made under the OPSS is the product of the conversion factor calculated in accordance with section II.B. of this final rule with comment period and the relative weight determined under section II.A. of this final rule with comment period. Therefore, the final national unadjusted payment rate for most APCs contained in Addendum A to this final rule with comment period and for most HCPCS codes to which separate payment under the OPSS has been assigned in Addendum B to this final rule with comment period was calculated by multiplying the final CY 2010 scaled weight for the APC by the final CY 2010 conversion factor.

We note that section 1833(t)(17) of the Act, which applies to hospitals as defined under section 1886(d)(1)(B) of the Act, requires that hospitals that fail to submit data required to be submitted on quality measures selected by the Secretary, in the form and manner and at a time specified by the Secretary, receive a 2.0 percentage point reduction to their OPD fee schedule increase factor, that is, the annual payment update factor. The application of a reduced OPD fee schedule increase factor results in reduced national unadjusted payment rates that apply to certain outpatient items and services provided by hospitals that are required to report outpatient quality data and that fail to meet the Hospital Outpatient Quality Data Reporting Program (HOP QDRP) requirements. For further discussion of the payment reduction for hospitals that fail to meet the requirements of the HOP QDRP, we refer readers to section XVI.D. of this final rule with comment period.

We demonstrate in the steps below how to determine the APC payments that would be made in a calendar year under the OPSS to a hospital that fulfills the HOP QDRP requirements and to a hospital that fails to meet the HOP QDRP requirements for a status indicator has any of the following status indicator assignments: "P," "Q1," "Q2," "Q3," "R," "S," "T," "U," "V," or "X" (as defined in Addendum D1 to this final rule with comment period), in a circumstance in which the multiple

procedure discount does not apply, the procedure is not bilateral, and conditionally packaged services (status indicator of "Q1" and "Q2") qualify for separate payment. We note that blood and blood products with status indicator "R" are not subject to wage adjustment but are subject to reduced payments when a hospital fails to meet the HOP QDRP requirements, as outlined in the steps and examples below.

Individual providers interested in calculating the payment amount that they would receive for a specific service from the national unadjusted payment rates presented in Addenda A and B to this final rule with comment period should follow the formulas presented in the following steps. For purposes of the payment calculations below, we refer to the national unadjusted payment rate for hospitals that meet the requirements of the HOP QDRP as the "full" national unadjusted payment rate. We refer to the national unadjusted payment rate for hospitals that fail to meet the requirements of the HOP QDRP as the "reduced" national unadjusted payment rate. The reduced national unadjusted payment rate is calculated by multiplying the reporting ratio of 0.98 times the "full" national unadjusted payment rate. The national unadjusted payment rate used in the calculations below is either the full national unadjusted payment rate or the reduced national unadjusted payment rate, depending on whether the hospital met its HOP QDRP requirements in order to receive the full CY 2010 OPSS increase factor.

Step 1. Calculate 60 percent (the labor-related portion) of the national unadjusted payment rate. Since the initial implementation of the OPSS, we have used 60 percent to represent our estimate of that portion of costs attributable, on average, to labor. We refer readers to the April 7, 2000 OPSS final rule with comment period (65 FR 18496 through 18497) for a detailed discussion of how we derived this percentage. We confirmed that this labor-related share for hospital outpatient services is still appropriate during our regression analysis for the payment adjustment for rural hospitals in the CY 2006 OPSS final rule with comment period (70 FR 68553).

The formula below is a mathematical representation of Step 1 and identifies the labor-related portion of a specific payment rate for a specific service.

X is the labor-related portion of the national unadjusted payment rate.
 $X = .60 * (\text{national unadjusted payment rate})$

Step 2. Determine the wage index area in which the hospital is located and identify the wage index level that applies to the specific hospital. The wage index values assigned to each area reflect the geographic statistical areas (which are based upon OMB standards) to which hospitals are assigned for FY 2010 under the IPPS, reclassifications through the MGCRB, section 1886(d)(8)(B) "Lugar" hospitals, reclassifications under section 1886(d)(8)(E) of the Act, as defined in § 412.103 of the regulations and hospitals designated as urban under section 601(g) of Public Law 98–21. We note that the reclassifications of hospitals under section 508 of Public Law 108–173, as extended by section 124 of Public Law 110–275, expired on September 30, 2009, and will not be applicable under the IPPS for FY 2010. Therefore, these reclassifications will not apply to the CY 2010 OPSS. For further discussion of the changes to the FY 2010 IPPS wage indices, as applied to the CY 2010 OPSS, we refer readers to section II.C. of this final rule with comment period. The wage index values include the occupational mix adjustment described in section II.C. of this final rule with comment period that was developed for the FY 2010 IPPS final payment rates published in the **Federal Register** on August 27, 2009 (74 FR 43827).

Step 3. Adjust the wage index of hospitals located in certain qualifying counties that have a relatively high percentage of hospital employees who reside in the county, but who work in a different county with a higher wage index, in accordance with section 505 of Public Law 108–173. Addendum L to this final rule with comment period contains the qualifying counties and the final wage index increase developed for the FY 2010 IPPS and published as Table 4J in the FY 2010 IPPS final rule (74 FR 44118 through 44125), as corrected in the **Federal Register** on October 2, 2009 (74 FR 51506) This step is to be followed only if the hospital is not reclassified or redesignated under section 1886(d)(8) or section 1886(d)(10) of the Act.

Step 4. Multiply the applicable wage index determined under Steps 2 and 3 by the amount determined under Step 1 that represents the labor-related portion of the national unadjusted payment rate.

The formula below is a mathematical representation of Step 4 and adjusts the labor-related portion of the national payment rate for the specific service by the wage index.

X_a is the labor-related portion of the national unadjusted payment rate (wage adjusted).

$X_a = .60 * (\text{national unadjusted payment rate}) * \text{applicable wage index}.$

Step 5. Calculate 40 percent (the nonlabor-related portion) of the national unadjusted payment rate and add that amount to the resulting product of Step 4. The result is the wage index adjusted payment rate for the relevant wage index area.

The formula below is a mathematical representation of Step 5 and calculates the remaining portion of the national payment rate, the amount not attributable to labor, and the adjusted payment for the specific service.

Y is the nonlabor-related portion of the national unadjusted payment rate.

$Y = .40 * (\text{national unadjusted payment rate})$

Adjusted Medicare Payment = $Y + X_a$

Step 6. If a provider is a SCH, set forth in the regulations at § 412.92, or an EACH, which is considered to be a SCH under section 1886(d)(5)(D)(iii)(III) of the Act, and located in a rural area, as defined in § 412.64(b), or is treated as being located in a rural area under § 412.103, multiply the wage index adjusted payment rate by 1.071 to calculate the total payment.

The formula below is a mathematical representation of Step 6 and applies the rural adjustment for rural SCHs.

Adjusted Medicare Payment (SCH or EACH) = Adjusted Medicare Payment * 1.071

We have provided examples below of the calculation of both the full and reduced national unadjusted payment rates that would apply to certain outpatient items and services performed by hospitals that meet and that fail to meet the HOP QDRP requirements, using the steps outlined above. For purposes of this example, we use a provider that is located in Wayne, New Jersey that is assigned to CBSA 35644. This provider bills one service that is assigned to APC 0019 (Level I Excision/Biopsy). The CY 2010 full national unadjusted payment rate for APC 0019 is \$294.06. The reduced national unadjusted payment rate for a hospital that fails to meet the HOP QDRP requirements is \$288.17. This reduced rate is calculated by multiplying the reporting ratio of 0.98 by the full unadjusted payment rate for APC 0019.

The FY 2010 wage index for a provider located in CBSA 35644 in New Jersey is 1.3005. The labor-related portion of the full national unadjusted payment is \$229.45 (.60 * \$294.06 * 1.3005). The labor-related portion of the reduced national unadjusted payment is

\$224.85 (.60 * \$288.17 * 1.3005). The nonlabor-related portion of the full national unadjusted payment is \$117.62 (.40 * \$294.06). The nonlabor-related portion of the reduced national unadjusted payment is \$115.26 (.40 * \$288.17). The sum of the labor-related and nonlabor-related portions of the full national adjusted payment is \$347.07 (\$229.45 + \$117.62). The sum of the reduced national adjusted payment is \$340.11 (\$224.85 + \$115.26).

We did not receive any public comments concerning our proposed methodology for calculating an adjusted payment from the national unadjusted Medicare payment amount for CY 2010. Therefore, we are finalizing our proposed CY 2010 methodology, without modification.

H. Beneficiary Copayments

1. Background

Section 1833(t)(3)(B) of the Act requires the Secretary to set rules for determining the unadjusted copayment amounts to be paid by beneficiaries for covered OPD services. Section 1833(t)(8)(C)(ii) of the Act specifies that the Secretary must reduce the national unadjusted copayment amount for a covered OPD service (or group of such services) furnished in a year in a manner so that the effective copayment rate (determined on a national unadjusted basis) for that service in the year does not exceed a specified percentage. As specified in section 1833(t)(8)(C)(ii)(V) of the Act, for all services paid under the OPSS in CY 2010, and in calendar years thereafter, the percentage is 40 percent of the APC payment rate.

Section 1833(t)(3)(B)(ii) of the Act provides that, for a covered OPD service (or group of such services) furnished in a year, the national unadjusted copayment amount cannot be less than 20 percent of the OPD fee schedule amount. Sections 1834(d)(2)(C)(ii) and (d)(3)(C)(ii) of the Act further require that the copayment for screening flexible sigmoidoscopies and screening colonoscopies be equal to 25 percent of the payment amount. Since the beginning of the OPSS, we have applied the 25-percent copayment to screening flexible sigmoidoscopies and screening colonoscopies.

2. Copayment Policy

In the CY 2010 OPSS/ASC proposed rule (74 FR 35298), for CY 2010, we proposed to determine copayment amounts for new and revised APCs using the same methodology that we implemented beginning in CY 2004. (We refer readers to the November 7,

2003 OPPS final rule with comment period (68 FR 63458)). In addition, we proposed to use the same standard rounding principles that we have historically used in instances where the application of our standard copayment methodology would result in a copayment amount that is less than 20 percent and cannot be rounded, under standard rounding principles, to 20 percent. (We refer readers to the CY 2008 OPPS/ASC final rule with comment period (72 FR 66687) in which we discuss our rationale for applying these rounding principles.) The national unadjusted copayment amounts for services payable under the OPPS that will be effective January 1, 2010, are shown in Addenda A and B to this final rule with comment period. As discussed in section XVI.D. of this final rule with comment period, as we proposed, we are providing that, for CY 2010, the Medicare beneficiary's minimum unadjusted copayment and national unadjusted copayment for a service to which a reduced national unadjusted payment rate applies will equal the product of the reporting ratio and the national unadjusted copayment, or the product of the reporting ratio and the minimum unadjusted copayment, respectively, for the service.

Comment: One commenter recommended that CMS continue its educational outreach and keep Medicare beneficiaries informed about the benefits of supplemental/secondary insurance in reducing their out-of-pocket costs for orthopedic procedures.

Response: We appreciate the commenter's support for our educational efforts on the availability of supplemental/secondary insurance and refer beneficiaries seeking information about their Medicare benefits and supplemental/secondary insurance coverage to the Web site at: <http://www.medicare.gov>.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, for determining APC copayment amounts.

3. Calculation of an Adjusted Copayment Amount for an APC Group

Individuals interested in calculating the national copayment liability for a Medicare beneficiary for a given service provided by a hospital that met or failed to meet its HOP QDRP requirements should follow the formulas presented in the following steps.

Step 1. Calculate the beneficiary payment percentage for the APC by dividing the APC's national unadjusted copayment by its payment rate. For example, using APC 0019, \$64.51 is 22

percent of the full national unadjusted payment rate of \$294.06. For APCs with only a minimum unadjusted copayment in Addendum A and B of this final rule with comment period, identify a beneficiary payment percentage of 20 percent.

The formula below is a mathematical representation of Step 1 and calculates national copayment as a percentage of national payment for a given service.

B is the beneficiary payment percentage.
 $B = \text{National unadjusted copayment for APC} / \text{national unadjusted payment rate for APC}$

Step 2. Calculate the appropriate wage-adjusted payment rate for the APC for the provider in question, as indicated in Steps 2 through 4 under section II.G. of this final rule with comment period. Calculate the rural adjustment for eligible providers as indicated in Step 6 under section II.G. of this final rule with comment period.

Step 3. Multiply the percentage calculated in Step 1 by the payment rate calculated in Step 2. The result is the wage-adjusted copayment amount for the APC.

The formula below is a mathematical representation of Step 3 and applies the beneficiary percentage to the adjusted payment rate for a service calculated under section II.G. of this final rule with comment period, with and without the rural adjustment, to calculate the adjusted beneficiary copayment for a given service.

Wage-adjusted copayment amount for the APC = Adjusted Medicare Payment * B

Wage-adjusted copayment amount for the APC (SCH or EACH) = (Adjusted Medicare Payment * 1.071) * B

Step 4. For a hospital that failed to meet its HOP QDRP requirements, multiply the copayment calculated in Step 3 by the reporting ratio of 0.98.

The unadjusted copayments for services payable under the OPPS that will be effective January 1, 2010, are shown in Addenda A and B to this final rule with comment period. We note that the national unadjusted payment rates and copayment rates shown in Addenda A and B to this final rule with comment period reflect the full market basket conversion factor increase, as discussed in section XVI.D. of this final rule with comment period.

III. OPPS Ambulatory Payment Classification (APC) Group Policies

A. OPPS Treatment of New CPT and Level II HCPCS Codes

CPT and Level II HCPCS codes are used to report procedures, services,

items, and supplies under the hospital OPPS. Specifically, CMS recognizes the following codes on OPPS claims: (1) Category I CPT codes, which describe medical services and procedures; (2) Category III CPT codes, which describe new and emerging technologies, services, and procedures; and (3) Level II HCPCS codes, which are used primarily to identify products, supplies, temporary procedures, and services not described by CPT codes. CPT codes are established by the AMA and the Level II HCPCS codes are established by the CMS HCPCS Workgroup. These codes are updated and changed throughout the year. CPT and HCPCS code changes that affect the OPPS are published both through the annual rulemaking cycle and through the OPPS quarterly update Change Requests (CRs). CMS releases new Level II HCPCS codes to the public or recognizes the release of new CPT codes by the AMA and makes these codes effective (that is, the codes can be reported on Medicare claims) outside of the formal rulemaking process via OPPS quarterly update CRs. This quarterly process offers hospitals access to codes that may more accurately describe items or services furnished and/or provides payment or more accurate payment for these items or services in a timelier manner than if CMS waited for the annual rulemaking process. We solicit comments on these new codes and finalize our proposals related to these codes through our annual rulemaking process.

We note that we sought public comments in the CY 2009 OPPS/ASC final rule with comment period on the new CPT and Level II HCPCS codes that were effective January 1, 2009. We also sought public comments in the CY 2009 OPPS/ASC final rule with comment period on the new Level II HCPCS codes effective October 1, 2008. These new codes with an effective date of October 1, 2008 or January 1, 2009 were flagged with comment indicator "NI" (New code, interim APC assignment; comments will be accepted on the interim APC assignment for the new code) in Addendum B to the CY 2009 OPPS/ASC final rule with comment period to indicate that we were assigning them an interim payment status and an APC and payment rate, if applicable, which were subject to public comment following publication of the CY 2009 OPPS/ASC final rule with comment period. Summaries of public comments on the codes flagged with comment indicator "NI" in the CY 2009 OPPS/ASC final rule with comment period and our responses are included in the sections of this final rule with

comment period that are relevant to the services described by those codes.
 In Table 13 of the CY 2010 OPPS/ASC proposed rule (74 FR 35299), which is

reproduced as Table 18 in this final rule with comment period, we summarized our process for updating codes through

our OPPS quarterly update CRs, seeking public comment, and finalizing their treatment under the OPPS.

TABLE 18—COMMENT TIMEFRAME FOR NEW OR REVISED HCPCS CODES

OPPS quarterly update CR	Type of code	Effective date	Comments sought	When finalized
April 1, 2009	Level II HCPCS Codes	April 1, 2009	CY 2010 OPPS/ASC proposed rule.	CY 2010 OPPS/ASC final rule with comment period.
July 1, 2009	Level II HCPCS Codes	July 1, 2009	CY 2010 OPPS/ASC proposed rule.	CY 2010 OPPS/ASC final rule with comment period.
	Category I (certain vaccine codes) and Category III CPT Codes.	July 1, 2009	CY 2010 OPPS/ASC proposed rule.	CY 2010 OPPS/ASC final rule with comment period.
October 1, 2009	Level II HCPCS Codes	October 1, 2009	CY 2010 OPPS/ASC final rule with comment period.	CY 2011 OPPS/ASC final rule with comment period.
January 1, 2010	Level II HCPCS Codes	January 1, 2010	CY 2010 OPPS/ASC final rule with Comment Period.	CY 2011 OPPS/ASC final rule with comment period.
	Category I and Category III CPT Codes.	January 1, 2010	CY 2010 OPPS/ASC final rule with comment period.	CY 2011 OPPS/ASC final rule with comment period.

1. Treatment of New Level II HCPCS Codes and Category I CPT Vaccine Codes and Category III CPT Codes

In the April 1 and July 1 CRs for CY 2009, we made effective a total of 13 new Level II HCPCS codes that were not addressed in the CY 2009 OPPS/ASC final rule with comment period that updated the OPPS and we allowed separate payment for 12 of these new codes. Through the April 1, 2009 CR, we also changed the OPPS status indicator for one existing Level II HCPCS code from the interim status indicator designated in the CY 2009 OPPS/ASC final rule with comment period to a status indicator that allowed separate pass-through payment for this code. In addition to the changes for Level II HCPCS codes, we made effective 5 new Category I vaccine and Category III CPT codes that were not addressed in the CY 2009 OPPS/ASC final rule with comment period that updated the OPPS and we allowed separate payment for 3 of these new codes.

Through the April 2009 OPPS quarterly update CR (Transmittal 1702, Change Request 6416, dated March 13, 2009), we allowed separate payment for a total of 2 additional Level II HCPCS codes, specifically existing HCPCS code C9247 (Iobenguane, I-123, diagnostic, per study dose, up to 10 millicuries) and new HCPCS code C9249 (Injection, certolizumab pegol, 1 mg). HCPCS code C9249, which received separate payment as a result of its pass-through status under the OPPS, was made effective on April 1, 2009. HCPCS code C9247 was released January 1, 2009

through the January 2009 OPPS quarterly update CR (Transmittal 1657, Change Request 6320, dated December 31, 2008). From January 1, 2009 through March 31, 2009, HCPCS code C9247 was packaged under the OPPS and assigned status indicator “N” (Items and Services Packaged into APC Rates). We note that between January 1, 2009 through March 31, 2009, HCPCS code C9247 was recognized as a nonpass-through diagnostic radiopharmaceutical. Because nonpass-through diagnostic radiopharmaceuticals are packaged under the OPPS, there was no separate APC payment for HCPCS code C9247 from January 1, 2009 through March 31, 2009. However, effective April 1, 2009, HCPCS code C9247 was allowed separate pass-through payment and its status indicator was revised from “N” to “G” (Pass-Through Drugs and Biologicals).

In the CY 2010 OPPS/ASC proposed rule, we solicited public comments on the status indicators and APC assignments of HCPCS codes C9247 and C9249, which were listed in Table 14 of that proposed rule (74 FR 35301) and now appear in Table 19 of this final rule with comment period.

We did not receive any public comments on the proposed APC assignments and status indicators for HCPCS codes C9247 and C9249. However, for CY 2010, the HCPCS Workgroup replaced both HCPCS C-codes with permanent HCPCS codes. Specifically, C9247 was replaced with A9582 (Iodine I-123 iobenguane, diagnostic, per study dose, up to 15

millicuries) and C9249 was replaced with J0718 (Injection, certolizumab pegol, 1 mg). Consistent with our general policy of using permanent HCPCS codes if appropriate rather than HCPCS C-codes for the reporting of drugs under the OPPS in order to streamline coding, we are showing the replacement HCPCS codes in Table 19 that will replace the HCPCS C-codes effective January 1, 2010. Both HCPCS C-codes will be deleted December 31, 2009. Because HCPCS code J0718 describes the same drug and the same dosage currently designated by HCPCS code C9249 and this drug will continue on pass-through status in CY 2010, we are assigning HCPCS code J0718 the same status indicator and APC as its predecessor C-code, as shown in Table 19. Although the dosage descriptor of HCPCS code A9582 indicates “per study dose, up to 15 millicuries” and the descriptor of its predecessor C-code designates “per study dose, up to 10 millicuries,” because we believe that the reporting of one unit for a study dose would be the same in almost all cases under either HCPCS code, we are assigning HCPCS code A9582 to the same APC as its predecessor C-code, as shown in Table 19. The recommended dose of I-123 iobenguane is 10 millicuries for adult patients, so we expect that hospitals would report 1 unit of new HCPCS code A9582 for the typical dose in CY 2010, just as they would have reported one unit of HCPCS code C9247 previously for the typical dose. We also note this diagnostic radiopharmaceutical will continue on

pass-through status in CY 2010; therefore, its CY 2010 status indicator remains as “G.” Because we did not receive any public comments on the new Level II HCPCS codes that were

implemented in April 2009, we are adopting as final, without modification, our proposal to assign the Level II HCPCS codes listed in Table 19 to the

APCs and status indicators as proposed for CY 2010.

Table 19 below shows the final APC and status indicator assignments for both HCPCS codes A9582 and J0718.

TABLE 19—LEVEL II HCPCS CODES WITH A CHANGE IN OPSS STATUS INDICATOR OR NEWLY IMPLEMENTED IN APRIL 2009

CY 2010 HCPCS code	CY 2009 HCPCS code	CY 2010 long descriptor	Final CY 2010 status indicator	Final CY 2010 APC
A9582	C9247	Iodine I-123 iobenguane, diagnostic, per study dose, up to 15 millicuries	G	9247
J0718	C9249	Injection, certolizumab pegol, 1 mg	G	9249

Through the July 2009 OPSS quarterly update CR (Transmittal 107, Change Request 6492, dated May 22, 2009), which included HCPCS codes that were made effective July 1, 2009, we allowed separate payment for a total of 11 new Level II HCPCS codes for pass-through drugs and biologicals and nonpass-through drugs and nonimplantable biologicals. Specifically, we provided separate payment for HCPCS codes C9250 (Human plasma fibrin sealant, vapor-heated, solvent-detergent (Artiss), 2ml); C9251 (Injection, C1 esterase inhibitor (human), 10 units); C9252 (Injection, plerixafor, 1 mg); C9253 (Injection, temozolomide, 1 mg); C9360 (Dermal substitute, native, non-denatured collagen, neonatal bovine origin (SurgiMend Collagen Matrix), per 0.5 square centimeters); C9361 (Collagen matrix nerve wrap (NeuroMend Collagen Nerve Wrap), per 0.5 centimeter length); C9362 (Porous purified collagen matrix bone void filler (Integra Mozaik Osteoconductive Scaffold Strip), per 0.5 cc); C9363 (Skin substitute, Integra Meshed Bilayer Wound Matrix, per square centimeter); C9364 (Porcine implant, Permacol, per square centimeter); Q2023 (Injection, factor viii (antihemophilic factor, recombinant) (Xyntha), per i.u.); and Q4116 (Skin substitute, Alloderm, per square centimeter).

Although HCPCS code Q4115 (Skin substitute, Alloskin, per square centimeter) was made effective July 1, 2009, because ASP pricing information was not available at the time the code was made effective, the HCPCS code was not paid separately and it was assigned status indicator “M” (Items and Services Not Billable to the Fiscal Intermediary/MAC) in the CY 2010 OPSS/ASC proposed rule (74 FR 35300 through 35301). For the October 2009 OPSS quarterly update, the status indicator for HCPCS code Q4115 was revised from “M” to “K” (Nonpass-Through Drugs and Biologicals)

effective October 1, 2009 because pricing information was available, and this product was paid separately as a new biological HCPCS code based on the ASP methodology, consistent with the final CY 2009 policy and the final CY 2010 policy for payment of new drug and biological HCPCS codes without pass-through status. The change in status indicator assignment was announced through the October 2009 OPSS quarterly update CR (Transmittal 1803, Change Request 6626, dated August 28, 2009).

In the CY 2010 OPSS/ASC proposed rule, we solicited public comments on the status indicators, APC assignments, and payment rates of these codes, which were listed in Table 15 of that proposed rule (74 FR 35301) and now appear in Table 20 of this final rule with comment period. Because of the timing of the proposed rule, the codes implemented in the July 2009 OPSS update were not included in Addendum B of that proposed rule, while those codes based upon the April 2009 OPSS update were included in Addendum B. In the CY 2009 OPSS/ASC proposed rule, we proposed to assign the new HCPCS codes for CY 2010 to the designated APCs listed in Table 15 for each HCPCS code and incorporate them into our final rule with comment period for CY 2010, which is consistent with our annual APC updating policy.

We did not receive any public comments on the proposed APC assignments, payment rates, and status indicators designated for the codes listed in Table 15 of the proposed rule. However, for CY 2010, the HCPCS Workgroup created permanent HCPCS J-codes for 4 of the 11 separately payable drug codes. Consistent with our general policy of using permanent HCPCS codes if appropriate rather than HCPCS C-codes for the reporting of drugs under the OPSS in order to streamline coding, we are showing the HCPCS J-codes in Table 20 of this final rule with comment

period that will replace the HCPCS C-codes effective January 1, 2010. HCPCS code C9251 is replaced with J0598 (Injection, C1 esterase inhibitor (human), 10 units); C9252 with J2562 (Injection, plerixafor, 1 mg); C9253 is replaced with J9328 (Injection, temozolomide, 1 mg); and Q2023 is replaced with J7185 (Injection, factor viii (antihemophilic factor, recombinant) (Xyntha), per i.u.). The HCPCS J-codes describe the same drugs and the same dosages as the HCPCS C-codes that will be deleted December 31, 2009. We note that HCPCS C-codes are temporary national HCPCS codes. To avoid duplication, temporary national HCPCS codes, such as “C,” “G,” “K,” and “Q” codes, are generally deleted once permanent national HCPCS codes are created that describe the same item, service, or procedure. Because three of the four new HCPCS J-codes describe the same drugs and the same dosages that are currently designated by HCPCS codes C9251, C9252, and C9253 and all three of these drugs will continue on pass-through status in CY 2010, we are assigning the HCPCS J-codes to the same APCs and status indicators as their predecessor HCPCS C-codes, as shown in Table 20. That is, HCPCS code J0598 is assigned to the same APC and status indicator as HCPCS code C9251 (APC 9251); HCPCS code J2562 is assigned to APC 9252; and HCPCS J9328 is assigned to APC 9253. Also, we note that, effective January 1, 2010, HCPCS code Q2023 will be replaced with HCPCS code J7185, which has the same descriptor and is assigned to the same APC and status indicator as HCPCS code Q2023.

Because we did not receive any public comments on the new Level II HCPCS codes that were implemented in July 2009, we are adopting as final, without modification, our proposal to assign the Level II HCPCS codes listed in Table 20 to the APCs and status indicators as proposed for CY 2010.

TABLE 20—NEW LEVEL II HCPCS CODES IMPLEMENTED IN JULY 2009

CY 2010 HCPCS code	CY 2009 HCPCS code	CY 2010 Long descriptor	Final CY 2010 status indicator	Final CY 2010 APC
C9250	C9250	Human plasma fibrin sealant, vapor-heated, solvent-detergent (Artiss), 2ml.	G	9250
J0598	C9251	Injection, C1 esterase inhibitor (human), 10 units	G	9251
J2562	C9252	Injection, plerixafor, 1 mg	G	9252
J9328	C9253	Injection, temozolomide, 1 mg	G	9253
C9360	C9360	Dermal substitute, native, non-denatured collagen, neonatal bovine origin (SurgiMend Collagen Matrix), per 0.5 square centimeters.	G	9360
C9361	C9361	Collagen matrix nerve wrap (NeuroMend Collagen Nerve Wrap), per 0.5 centimeter length.	G	9361
C9362	C9362	Porous purified collagen matrix bone void filler (Integra Mozaik Osteoconductive Scaffold Strip), per 0.5 cc.	G	9362
C9363	C9363	Skin substitute, Integra Meshed Bilayer Wound Matrix, per square centimeter.	G	9363
C9364	C9364	Porcine implant, Permacol, per square centimeter	G	9364
J7185	Q2023	Injection, factor viii (antihemophilic factor, recombinant) (Xyntha), per i.u..	K	1268
Q4115	Q4115	Skin substitute, Alloskin, per square centimeter	K	1287
Q4116	Q4116	Skin substitute, Alloderm, per square centimeter	K	1270

In the CY 2010 OPPS/ASC proposed rule (74 FR 35300), we proposed to continue our established policy of recognizing Category I CPT vaccine codes for which FDA approval is imminent and Category III CPT codes that the AMA releases in January of each year for implementation in July through the OPPS quarterly update process. Under the OPPS, Category I vaccine codes and Category III CPT codes that are released on the AMA Web site in January are made effective in July of the same year through the July OPPS quarterly update CR, consistent with the AMA's implementation date for the codes. Through the July 2009 OPPS quarterly update CR, we allowed separate payment for 3 of the 5 new Category I vaccine and Category III CPT Codes effective July 1, 2009. Specifically, as displayed in Table 16 of the CY 2010 OPPS/ASC proposed rule (74 FR 35301) and reproduced in this final rule with comment period as Table 21, we allowed payment for CPT codes 0199T (Physiologic recording of tremor using accelerometer(s) and gyroscope(s), (including frequency and amplitude) including interpretation and report); 0200T (Percutaneous sacral augmentation (sacroplasty), unilateral

injection(s), including the use of a balloon or mechanical device (if utilized), one or more needles); and 0201T (Percutaneous sacral augmentation (sacroplasty), bilateral injections, including the use of a balloon or mechanical device (if utilized), two or more needles). We note that CPT code 0202T (Posterior vertebral joint(s) arthroplasty (e.g., facet joint[s] replacement) including facetectomy, laminectomy, foraminotomy and vertebral column fixation, with or without injection of bone cement, including fluoroscopy, single level, lumbar spine) was assigned status indicator "C" (Inpatient Procedures) because we believe that this procedure may only be safely performed on Medicare beneficiaries in the hospital inpatient setting. In addition, CPT code 90670 (Pneumococcal conjugate vaccine, 13 valent, for intramuscular use), a Category I CPT vaccine code, was assigned status indicator "E" (Items, Codes, and Services not paid by Medicare when submitted on outpatient claims (any outpatient bill type)) because the drug has not yet been approved by the FDA for marketing. Because the July 2009 OPPS quarterly update CR was issued close to the

publication of the CY 2010 OPPS/ASC proposed rule, the Category I vaccine and Category III CPT codes implemented through the July 2009 OPPS quarterly update CR were not be included in Addendum B to the proposed rule, but these codes were listed in Table 16 of the proposed rule. Additionally, we proposed to incorporate them into Addendum B to this CY 2010 OPPS/ASC final rule with comment period, which is consistent with our annual OPPS update policy.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35301), we solicited public comments on the proposed status indicators, APC assignments, and payment rates for the new Category I and III CPT codes. We did not receive any public comments on our proposals for CPT codes 0199T, 0200T, 0201T, 0202T, and 90670. Therefore, we are finalizing our CY 2010 proposals for these codes, without modification. The final CY 2010 status indicators and APC assignments for CPT codes 0199T, 0200T, 0201T, 0202T, and 90670 are listed in Table 21 below, as well as in Addendum B to this final rule with comment period.

TABLE 21—CATEGORY I VACCINE AND CATEGORY III CPT CODES IMPLEMENTED IN JULY 2009

CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 status indicator	Final CY 2010 APC
0199T	Physiologic recording of tremor using accelerometer(s) and gyroscope(s), (including frequency and amplitude) including interpretation and report.	S	0215
0200T	Percutaneous sacral augmentation (sacroplasty), unilateral injection(s), including the use of a balloon or mechanical device (if utilized), one or more needles.	T	0049

TABLE 21—CATEGORY I VACCINE AND CATEGORY III CPT CODES IMPLEMENTED IN JULY 2009—Continued

CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 status indicator	Final CY 2010 APC
0201T	Percutaneous sacral augmentation (sacroplasty), bilateral injections, including the use of a balloon or mechanical device (if utilized), two or more needles.	T	0050
0202T	Posterior vertebral joint(s) arthroplasty (e.g., facet joint[s] replacement) including facetectomy, laminectomy, foraminotomy and vertebral column fixation, with or without injection of bone cement, including fluoroscopy, single level, lumbar spine.	C	Not applicable.
90670	Pneumococcal conjugate vaccine, 13 valent, for intramuscular use	E	Not applicable.

2. Process for New Level II HCPCS Codes and Category I and Category III CPT Codes for Which We Are Soliciting Public Comments on the CY 2010 OPPS/ASC Final Rule With Comment Period

As has been our practice in the past, we incorporate those new Category I and III CPT codes and new Level II HCPCS codes that are effective January 1 in the final rule with comment period updating the OPPS for the following calendar year. These codes are released to the public via the CMS HCPCS (for Level II HCPCS codes) and AMA Web sites (for CPT codes), and also through the January OPPS quarterly update CRs. In the past, we also have released new Level II HCPCS codes that are effective October 1 through the October OPPS quarterly update CRs and incorporated these new codes in the final rule with comment period updating the OPPS for the following calendar year. All of these codes are flagged with comment indicator “NI” in Addendum B to the OPPS/ASC final rule with comment period to indicate that we are assigning them an interim payment status which is subject to public comment. Specifically, the status indicator and the APC assignment, and payment rate, if applicable, for all such codes flagged with comment indicator “NI” are open to public comment in this final rule with comment period, and we respond to these comments in the final rule with comment period for the next calendar year’s OPPS/ASC update. In the CY 2010 OPPS/ASC proposed rule (74 FR 35302), we proposed to continue this process for CY 2010. Specifically, for CY 2010, we proposed to include in Addendum B to the CY 2010 OPPS/ASC final rule with comment period the new Category I and III CPT codes effective January 1, 2010 (including those Category I vaccine and Category III CPT codes that were released by the AMA in July 2009) that would be incorporated in the January 2010 OPPS quarterly update CR and the new Level II HCPCS codes, effective October 1, 2009 or January 1,

2010, that would be released by CMS in its October 2009 and January 2010 OPPS quarterly update CRs. Excluding those Category I vaccine and Category III CPT codes that were released by the AMA in July 2009 which were subject to comment in the CY 2010 OPPS/ASC proposed rule as described above, these codes would be flagged with comment indicator “NI” in Addendum B to this CY 2010 OPPS/ASC final rule with comment period to indicate that we have assigned them an interim OPPS payment status. We proposed that their status indicators and their APC assignments and payment rates, if applicable, would be open to public comment in the CY 2010 OPPS/ASC final rule with comment period and would be finalized in the CY 2011 OPPS/ASC final rule with comment period.

Comment: One commenter requested that CMS solicit public comments on APC assignments for the newly implemented CPT codes that go into effect January 1 and, when necessary, revise their APC assignments and implement the changes in the next quarterly OPPS update to promote payment accuracy.

Response: For new HCPCS codes with an interim final APC and/or status indicator designation in a final rule, we are only able to finalize their assignments in another OPPS final rule in order to allow for the necessary public notice and comment period and to allow for CMS to respond to such comments. Therefore, we only assign HCPCS codes permanently for the year through the annual regulatory process. Because we are not able to revise APC and/or status indicator assignments for the newly implemented HCPCS codes in CY 2010 that are assigned an interim final status in this CY 2010 OPPS/ASC final rule with comment period outside of the rulemaking process, the next available opportunity to update an APC or status indicator for these codes is in the CY 2011 OPPS update. These HCPCS codes retain their interim final APC and status indicator assignments

for all of CY 2010. Therefore, only in the CY 2011 OPPS/ASC final rule with comment period will we be able to finalize the APC and/or status indicator assignments of the new CY 2010 HCPCS codes and respond to all public comments received on their interim designations.

After consideration of the public comment we received, we are finalizing our proposal, without modification, to provide interim final status indicators and APC assignments and payment rates, if applicable, for all CPT codes newly implemented in January 2010 and all HCPCS codes newly implemented in October 2009 or January 2010 in Addendum B to this final rule with comment period. The interim final OPPS treatment of these codes is open to public comment in the CY 2010 OPPS/ASC final rule with comment period and will be finalized in the CY 2011 OPPS/ASC final rule with comment period.

B. OPPS Changes—Variations Within APCs

1. Background

Section 1833(t)(2)(A) of the Act requires the Secretary to develop a classification system for covered outpatient department services. Section 1833(t)(2)(B) of the Act provides that the Secretary may establish groups of covered outpatient department services within this classification system, so that services classified within each group are comparable clinically and with respect to the use of resources (and so that an implantable item is classified to the group that includes the service to which the item relates). In accordance with these provisions, we developed a grouping classification system, referred to as APCs, as set forth in § 419.31 of the regulations. We use Level I and Level II HCPCS codes and descriptors to identify and group the services within each APC. The APCs are organized such that each group is homogeneous both clinically and in terms of resource use. Using this classification system, we have

established distinct groups of similar services, as well as medical visits. We also have developed separate APC groups for certain medical devices, drugs, biologicals, therapeutic radiopharmaceuticals, and brachytherapy devices.

We have packaged into payment for each procedure or service within an APC group the costs associated with those items or services that are directly related to and supportive of performing the main independent procedures or furnishing the services. Therefore, we do not make separate payment for these packaged items or services. For example, packaged items and services include: (1) Use of an operating, treatment, or procedure room; (2) use of a recovery room; (3) observation services; (4) anesthesia; (5) medical/surgical supplies; (6) pharmaceuticals (other than those for which separate payment may be allowed under the provisions discussed in section V.3. of this final rule with comment period); (7) incidental services such as venipuncture; and (8) guidance services, image processing services, intraoperative services, imaging supervision and interpretation services, diagnostic radiopharmaceuticals, and contrast media. Further discussion of packaged services is included in section II.A.4. of this final rule with comment period.

In CY 2008 (72 FR 66650), we implemented composite APCs to provide a single payment for groups of services that are typically performed together during a single clinical encounter and that result in the provision of a complete service. Under our CY 2010 OPPS policy, we provide composite APC payment for certain extended assessment and management services, low dose rate (LDR) prostate brachytherapy, cardiac electrophysiologic evaluation and ablation, mental health services, and multiple imaging services. Further discussion of composite APCs is included in section II.A.2.e. of this final rule with comment period.

Under the OPPS, we generally pay for hospital outpatient services on a rate-per-service basis, where the service may be reported with one or more HCPCS codes. Payment varies according to the APC group to which the independent service or combination of services is assigned. Each APC weight represents the hospital median cost of the services included in that APC relative to the hospital median cost of the services included in APC 0606 (Level 3 Hospital Clinic Visits). The APC weights are scaled to APC 0606 because it is the middle level clinic visit APC (that is,

where the Level 3 clinic visit CPT code of five levels of clinic visits is assigned), and because middle level clinic visits are among the most frequently furnished services in the hospital outpatient setting.

Section 1833(t)(9)(A) of the Act requires the Secretary to review not less often than annually and revise the groups, relative payment weights, and the wage and other adjustments under the OPSS to take into account changes in medical practice, changes in technology, the addition of new services, new cost data, and other relevant information and factors. Section 1833(t)(9)(A) of the Act, as amended by section 201(h) of the BBRA, also requires the Secretary to consult with an outside panel of experts to review (and advise the Secretary concerning) the clinical integrity of the APC groups and the relative payment weights (the APC Panel recommendations for specific services for the CY 2010 OPPS and our responses to them are discussed in the relevant specific sections throughout this final rule with comment period).

Finally, section 1833(t)(2) of the Act provides that, subject to certain exceptions, the items and services within an APC group cannot be considered comparable with respect to the use of resources if the highest median cost (or mean cost as elected by the Secretary) for an item or service in the group is more than 2 times greater than the lowest median cost (or mean cost, if so elected) for an item or service within the same group (referred to as the "2 times rule"). We use the median cost of the item or service in implementing this provision. In performing this analysis, we examine data from the significant services assigned to an APC, specifically those HCPCS codes with a single claim frequency of greater than 1,000 or a frequency of greater than 99 and a percentage of all single claims that is equal to or greater than 2 percent. Because, as a matter of policy, HCPCS codes that are unlisted procedures, not otherwise classified, or not otherwise specified codes are assigned to the lowest level APC that is appropriate to the clinical nature of the service (69 FR 65724 through 65725), we do not consider the costs of these services in assessing APCs for 2 times violations. Section 1833(t)(2) of the Act authorizes the Secretary to make exceptions to the 2 times rule in unusual cases, such as low-volume items and services (but the Secretary may not make such an exception in the case of a drug or biological that has been designated as an orphan drug under section 526 of the Federal Food, Drug, and Cosmetic Act).

2. Application of the 2 Times Rule

In accordance with section 1833(t)(2) of the Act and § 419.31 of the regulations, we annually review the items and services within an APC group to determine, with respect to comparability of the use of resources, if the median cost of the highest cost item or service within an APC group is more than 2 times greater than the median of the lowest cost item or service within that same group. In the CY 2010 OPPS/ASC proposed rule (74 FR 35303), we proposed to make exceptions to this limit on the variation of costs within each APC group in unusual cases, such as low-volume items and services for CY 2010.

During the APC Panel's February 2009 meeting, we presented median cost and utilization data for services furnished during the period of January 1, 2008 through September 30, 2008, because we had concerns or the public had raised concerns regarding their APC assignments, status indicator assignments, or payment rates. In addition to the assignment of specific services to APCs that we discussed with the APC Panel, we also identified APCs with 2 times violations that were not specifically discussed with the APC Panel but for which we proposed changes to their HCPCS codes' APC assignments in Addendum B to the CY 2010 OPPS/ASC proposed rule. In these cases, to eliminate a 2 times violation or to improve clinical and resource homogeneity, we proposed to reassign the codes to APCs that contain services that are similar with regard to both their clinical and resource characteristics. We also proposed to rename existing APCs or create new clinical APCs to complement proposed HCPCS code reassignments. In many cases, the proposed HCPCS code reassignments and associated APC reconfigurations for CY 2010 included in the proposed rule were related to changes in median costs of services that were observed in the CY 2008 claims data newly available for CY 2010 ratesetting. In addition, we proposed changes to the status indicators for some codes that were not specifically and separately discussed in the proposed rule. In these cases, we proposed to change the status indicators for some codes because we believed that another status indicator would more accurately describe their payment status from an OPPS perspective based on the policies that we proposed for CY 2010.

Addendum B to the CY 2010 OPPS/ASC proposed rule identified, with comment indicator "CH," those HCPCS codes for which we proposed a change to the APC assignment or status

indicator that were initially assigned in the April 2009 Addendum B update (Transmittal 1702, Change Request 6416, dated March 13, 2009).

Comment: One commenter generally supported CMS' adherence to the 2 times rule to ensure appropriate payment for OPPS services and to provide incentives for patients to be treated in the most suitable clinical setting. In particular, the commenter supported the proposed reassignments of the following CPT codes: CPT code 20103 (Exploration of penetrating wound (separate procedure); extremity) from APC 0136 (Level IV Skin Repair) to APC 0007 (Level II Incision & Drainage); and CPT code 29888 (Arthroscopically aided anterior cruciate ligament repair/augmentation or reconstruction) and CPT code 29889 (Arthroscopically aided posterior cruciate ligament repair/augmentation or reconstruction) from APC 0042 (Level II Arthroscopy) to APC 0052 (Level IV Musculoskeletal Procedures Except Hand and Foot).

Response: We appreciate the commenter's support for the 2 times rule in general and for the proposed APC reassignments of CPT codes 20103, 29888, and 29889 in particular. We agree with the commenter that the 2 times rule is an important mechanism for ensuring appropriate payment for OPPS services. Based on the CY 2008 claims and cost report data available for this final rule with comment period, we also agree with the commenter that we should adopt the proposed reassignments of CPT codes 20103, 29888, and 29889 in order to improve clinical and resource homogeneity.

Therefore, after consideration of the public comment we received, we are finalizing, without modification, our CY 2010 proposals to reassign CPT code 20103 to APC 0007, with a final CY 2010 APC median cost of approximately \$843 and CPT codes 29888 and 29889 to APC 0052, with a final CY 2010 APC median cost of approximately \$5,921.

3. Exceptions to the 2 Times Rule

As discussed earlier, we may make exceptions to the 2 times limit on the variation of costs within each APC group in unusual cases such as low-volume items and services. We stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35303) that we took into account the APC changes that we proposed for CY 2010 based on the APC Panel recommendations, the other proposed changes to status indicators and APC assignments as identified in Addendum B to the proposed rule, and the use of CY 2008 claims data available for the proposed rule to calculate the median

costs of procedures classified in the APCs. We then reviewed all of the APCs to determine which APCs would not satisfy the 2 times rule and to determine which APCs should be proposed as exceptions to the 2 times rule for CY 2010. We used the following criteria to decide whether to propose exceptions to the 2 times rule for affected APCs:

- Resource homogeneity;
- Clinical homogeneity;
- Hospital outpatient setting;
- Frequency of service (volume); and
- Opportunity for upcoding and code fragments.

For a detailed discussion of these criteria, we refer readers to the April 7, 2000 OPPS final rule with comment period (65 FR 18457).

Table 17 of the CY 2010 OPPS/ASC proposed rule (74 FR 35303) listed 14 APCs that we proposed to exempt from the 2 times rule for CY 2010 based on the criteria cited above. For cases in which a recommendation by the APC Panel appeared to result in or allow a violation of the 2 times rule, we generally accepted the APC Panel's recommendation because those recommendations were based on explicit consideration of resource use, clinical homogeneity, hospital specialization, and the quality of the CY 2008 claims data used to determine the APC payment rates that we proposed for CY 2010. The median costs for hospital outpatient services for these and all other APCs that were used in the development of the CY 2010 OPPS/ASC proposed rule and this final rule with comment period can be found on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/01_overview.asp.

For the CY 2010 OPPS/ASC proposed rule, we based the listed exceptions to the 2 times rule on claims data for dates of service between January 1, 2008, and December 31, 2008, that were processed before January 1, 2009. For this final rule with comment period, we used claims data for dates of service between January 1, 2008, and December 31, 2008, that were processed on or before June 30, 2008, and updated CCRs, if available. Thus, after responding to all of the public comments on the CY 2010 OPPS/ASC proposed rule and making changes to APC assignments based on those comments, we analyzed the CY 2008 claims data used for this final rule with comment period to identify the APCs with 2 times violations. Based on the final CY 2008 claims data, we found that there are 15 APCs with 2 times rule violations, a cumulative increase of 1 APC from the proposed rule. We applied the criteria as described earlier

to identify the APCs that are exceptions to the 2 times rule for CY 2010, and identified 4 additional APCs that meet the criteria for exception to the 2 times rule for this final rule with comment period: APC 0057 (Bunion Procedures); APC 0060 (Manipulation Therapy); APC 0341 (Skin Tests); and APC 0409 (Red Blood Cell Tests). These APC exceptions are listed in Table 22 below. We also determined that there are 3 APCs that no longer violate the 2 times rule: APC 0237 (Level II Posterior Segment Eye Procedures); APC 0325 (Group Psychotherapy); and APC 0436 (Level I Drug Administration). We have not included in this count those APCs where a 2 times violation is not a relevant concept, such as APC 0375 (Ancillary Outpatient Services when Patient Expires), with an APC median cost set based on multiple procedure claims, so that we have identified only final APCs, including those with criteria-based median costs, such as device-dependent APCs, with 2 times violations.

Comment: One commenter requested that CMS not exempt any imaging and radiation therapy APCs from the 2 times rule. According to the commenter, violations of the 2 times rule should demonstrate to CMS that a particular APC is incorrectly constructed. The commenter recommended that, rather than exempting these APCs from the 2 times rule, CMS review the configurations of the APCs and make any necessary revisions.

Response: We do not agree with the commenter that we should not exempt any imaging and radiation therapy APCs from the 2 times rule. As stated earlier in this section, we may make exceptions to the 2 times limit on the variation of costs within each APC group in unusual cases such as low-volume items and services. In the CY 2010 OPPS/ASC proposed rule (74 FR 35303), we proposed to exempt one imaging APC and one radiation therapy APC from the 2 times rule, specifically APC 0128 (Echocardiogram with Contrast) and APC 0664 (Level I Proton Beam Radiation Therapy), respectively. As discussed in greater detail in section II.A.2.d.(4) of this final rule with comment period, we believe that the median costs of the echocardiography procedures assigned to APC 0128 do not warrant assignment of any of those procedures to a new clinical APC. APC 0664 includes CPT code 77520 (Proton treatment delivery; simple, without compensation), with a median cost of approximately \$396 (based on 243 single claims of 251 total claims), and CPT code 77522 (Proton treatment delivery; simple, with compensation),

with a median cost of approximately \$934 (based on 11,012 single claims of 12,252 total claims). We continue to believe that the resources and clinical characteristics of the low-volume procedure described by CPT code 77520 are sufficiently similar to the procedure described by CPT code 77522 to warrant continued assignment of both CPT codes to APC 0664. Therefore, we are finalizing our proposal to exempt APC 0128 and APC 0664 from the 2 times rule for CY 2010. Consistent with our

standard policy, we will continue to review, on an annual basis, the APC assignments for all OPSS services to ensure appropriate placement.

We also received a number of specific public comments regarding some of the procedures assigned to APCs that we proposed to exempt from the 2 times rule for CY 2010. Discussions of those public comments are included elsewhere in this final rule with comment period in the specific sections

related to the types of procedures that were the subjects of the comments.

After consideration of the public comments we received and our review of the CY 2008 costs from claims available for this final rule with comment period, we are exempting 15 APCs from the 2 times rule for CY 2010, as described previously in this section. Our final list of 15 APCs exempted from the 2 times rule is displayed in Table 22 below.

TABLE 22—FINAL APC EXCEPTIONS TO THE 2 TIMES RULE FOR CY 2010

Final CY 2010 APC	Final CY 2010 APC Title
0057	Bunion Procedures.
0060	Manipulation Therapy.
0080	Diagnostic Cardiac Catheterization.
0105	Repair/Revision/Removal of Pacemakers, AICDs, or Vascular Devices.
0128	Echocardiogram with Contrast.
0141	Level I Upper GI Procedures.
0142	Small Intestine Endoscopy.
0245	Level I Cataract Procedures without IOL Insert.
0303	Treatment Device Construction.
0341	Skin Tests.
0381	Single Allergy Tests.
0409	Red Blood Cell Tests.
0432	Health and Behavior Services.
0604	Level 1 Hospital Clinic Visits.
0664	Level I Proton Beam Radiation Therapy.

C. New Technology APCs

1. Background

In the November 30, 2001 final rule (66 FR 59903), we finalized changes to the time period a service was eligible for payment under a New Technology APC. Beginning in CY 2002, we retain services within New Technology APC groups until we gather sufficient claims data to enable us to assign the service to a clinically appropriate APC. This policy allows us to move a service from a New Technology APC in less than 2 years if sufficient data are available. It also allows us to retain a service in a New Technology APC for more than 2 years if sufficient data upon which to base a decision for reassignment have not been collected.

We note that the cost bands for New Technology APCs range from \$0 to \$50 in increments of \$10, from \$50 to \$100 in increments of \$50, from \$100 through \$2,000 in increments of \$100, and from \$2,000 through \$10,000 in increments of \$500. These cost bands identify the APCs to which new technology procedures and services with estimated service costs that fall within those cost bands are assigned under the OPSS. Payment for each APC is made at the mid-point of the APC's assigned cost

band. For example, payment for New Technology APC 1507 (New Technology—Level VII (\$500–\$600)) is made at \$550. Currently, there are 82 New Technology APCs, ranging from the lowest cost band assigned to APC 1491 (New Technology—Level IA (\$0–\$10)) through the highest cost band assigned to APC 1574 (New Technology—Level XXXVII (\$9,500–\$10,000)). In CY 2004 (68 FR 63416), we last restructured the New Technology APCs to make the cost intervals more consistent across payment levels and refined the cost bands for these APCs to retain two parallel sets of New Technology APCs, one set with a status indicator of “S” (Significant Procedures, Not Discounted when Multiple. Paid under OPSS; separate APC payment) and the other set with a status indicator of “T” (Significant Procedure, Multiple Reduction Applies. Paid under OPSS; separate APC payment). These current New Technology APC configurations allow us to price new technology services more appropriately and consistently.

2. Movement of Procedures From New Technology APCs to Clinical APCs

As we explained in the November 30, 2001 final rule (66 FR 59902), we

generally keep a procedure in the New Technology APC to which it is initially assigned until we have collected sufficient data to enable us to move the procedure to a clinically appropriate APC. However, in cases where we find that our original New Technology APC assignment was based on inaccurate or inadequate information (although it was the best information available at the time), or where the New Technology APCs are restructured, we may, based on more recent resource utilization information (including claims data) or the availability of refined New Technology APC cost bands, reassign the procedure or service to a different New Technology APC that most appropriately reflects its cost.

Consistent with our current policy, in the CY 2010 OPSS/ASC proposed rule (74 FR 35304), for CY 2010, we proposed to retain services within New Technology APC groups until we gather sufficient claims data to enable us to assign the service to a clinically appropriate APC. The flexibility associated with this policy allows us to move a service from a New Technology APC in less than 2 years if sufficient data are available. It also allows us to retain a service in a New Technology APC for more than 2 years if sufficient

hospital claims data upon which to base a decision for reassignment have not been collected.

Table 18 of the proposed rule (74 FR 35304) listed the HCPCS code and its associated status indicator that we proposed to reassign from a New Technology APC to a clinically appropriate APC for CY 2010. Based on the CY 2008 OPPS claims data available for the proposed rule, we believe we had sufficient claims data to propose reassignment of CPT code 0182T to a clinically appropriate APC. Specifically, we proposed to reassign this electronic brachytherapy service from APC 1519 (New Technology—Level IXX (\$1,700–\$1,800)) to APC 0313 (Brachytherapy), where other brachytherapy services also reside. Based on hospital claims data for CPT code 0182T, its hospital resource costs are similar to those of other services assigned to APC 0313.

The proposed CY 2010 APC reassignment of CPT code 0182T was discussed with the APC Panel at its August 2009 meeting. One public presenter indicated that CPT code 0182T describes both single-fraction and multiple-fraction electronic brachytherapy, and that most of the claims on which CMS based its CY 2010 proposal were from hospitals reporting multi-fraction electronic brachytherapy. The presenter believed that the hospital resources required for these two types of brachytherapy were significantly different from one another. The presenter also stated that, unlike the conventional brachytherapy procedures that are assigned to APC 0313 and for which the associated brachytherapy sources are all paid separately under the OPPS, payment for the brachytherapy source associated with CPT code 0182T is packaged into the procedure payment. The APC Panel noted that the problem of distinguishing single-fraction from multiple-fraction electronic brachytherapy is a coding issue that would not be resolved by additional claims data because the two types of procedures are reported with the same CPT code. After discussion of the median cost of CPT code 0182T observed in claims data and the potential contribution of the brachytherapy source cost to the overall procedure cost, the APC Panel made no recommendation on the CY 2010 APC assignment for CPT code 0182T.

Comment: Several commenters disagreed with the CMS proposal to reassign CPT code 0182T to APC 0313 for CY 2010. Commenters responding regarding both single-fraction and multiple-fraction electronic brachytherapy procedures reported with CPT code 0182T asserted that the

proposed payment of approximately \$747 does not cover the full costs of providing these services. They indicated that the current payment rate of \$1,750 that is associated with APC 1519, where CPT code 0182T is assigned, includes the cost of the electronic brachytherapy source, whereas the payment rate for APC 0313 does not. Several commenters noted that the claims data used in determining the reassignment for CPT code 0182T are very limited and pointed out that the claims data for this service came from only a few hospitals that were early adopters of the multiple-source electronic brachytherapy technology. They argued that data from these hospitals represented pre-commercial clinical site data and that, therefore, these hospitals had received equipment and sources at reduced cost. One commenter suggested that CMS should develop coding that would distinguish between single-fraction and multiple-fraction electronic brachytherapy procedures in order to pay appropriately for each type of service because the current single payment for both technologies provides a financial incentive for the use of multiple-fraction electronic brachytherapy. Most commenters urged CMS to continue to assign CPT code 0182T to APC 1519 for at least another year to enable CMS to gather sufficient claims data from more hospitals in order to appropriately reassign CPT code 0182T to a clinical APC based on the clinical and resource costs of the procedure.

Response: CPT code 0182T was initially assigned to New Technology APC 1519 with a payment rate of \$1,750 when the code was implemented in July 2007, and it has been assigned to that same APC through CY 2009. For CY 2010, we proposed to reassign CPT code 0182T from New Technology APC 1519 to APC 0313, which had a proposed payment rate of approximately \$747 and a proposed median cost of approximately \$753 (and now has a final rule median cost of \$770). Analysis of hospital claims data from CY 2007 and CY 2008 revealed that the procedure described by CPT code 0182T is not commonly performed on Medicare patients. For CY 2008, claims data show 223 total claims with a median cost of about \$506 for this procedure. For CY 2007, claims data (6 months due to implementation of the code in July 2007) show only 21 total claims with a median cost of about \$495. Therefore, we believe that the hospital resources required for CPT code 0182T are consistent with the costs of other services assigned to APC 0313

and payment for CPT code 0182T would be appropriately made through that clinical APC.

We are not creating a new Level II HCPCS code for single-fraction electronic brachytherapy at this time because we believe that the two forms of electronic brachytherapy, whether provided in a single-fraction or multiple-fraction regimen, depending on the technology, are both described by CPT code 0182T, which is appropriately assigned to a single APC. We note that the payment is per-fraction according to the code descriptor for CPT code 0182T, and that would include a single-fraction treatment as well. While we recognize that CPT code 0182T describes both single- and multiple-fraction electronic brachytherapy, we commonly pay for different technologies under the OPPS that are reported in a single CPT code and we expect the hospital claims data to reflect the resources required for all of the different technologies reported under the one code. To the extent that one technology is predominantly used by hospitals, then the costs of that technology will have a greater effect on the procedure's median cost, but we do not believe payment through such groupings inappropriately encourages the use of certain technologies, such as the provision of multiple-fraction electronic brachytherapy in the case of CPT code 0182T. Our standard OPPS ratesetting methodology provides a single payment based on historical hospital costs that reflect utilization patterns of the various technologies, consistent with prospective payment for groups of similar services in order to encourage hospital efficiencies. The hospital cost information for services always reflects the discounts available to hospitals in the claims year, such as the commenters indicate was the case for multiple-fraction electronic brachytherapy in CY 2008, that may not be available in the payment year for those services, and those discounts may vary from year to year for different HOPD services. Nevertheless, we rely on the relativity of median costs as reflected in claims data to be appropriate. Payment based on a measure of central tendency is a principle of any prospective payment system like the OPPS. In some individual cases payment exceeds the average cost and in other cases payment is less than the average cost. On balance, however, payment should approximate the relative cost of the average case, recognizing that, as a prospective payment system, the OPPS is a system of averages. In the case of CPT code 0182T, we believe that its assignment to

APC 0313 for CY 2010 is fully consistent with our standard ratesetting methodology that provides appropriate incentives for efficiency.

As of January 2010, CPT code 0182T will have been assigned to New Technology APC 1519 for 2½ years. While we have relatively few claims data from CY 2007 and CY 2008 for electronic brachytherapy, a commenter on a prior rule has indicated that this service may only be used to treat a small number of patients (72 FR 66691). To the extent that more hospitals furnish electronic brachytherapy in future years and that hospital costs from commercialization of the technology change, we expect to see those costs reflected in our claims data for those future years, which we will annually review for electronic brachytherapy, just as we do for all OPPS services. Moreover, while we acknowledge that, in the case of conventional brachytherapy procedures where distinct radioactive sources are implanted, the statute requires separate payment of the associated radioactive brachytherapy source so that APC 0313 only pays for the application of those sources, we have no reason to believe that reported hospital costs for CPT code 0182T do not include the cost of the source. As we stated in the CY 2007 OPPS/ASC final rule with comment period (71 FR 68113), we do not consider specific devices, beams of radiation, or equipment that do not constitute separate sources that utilize radioactive isotopes to deliver radiation to be brachytherapy sources for separate

payment, as such items do not meet the statutory requirements provided in section 1833(t)(2)(H) of the Act. Electronic brachytherapy, described by CPT code 0182T, utilizes such devices, beams of radiation, or equipment to generate the radiation for the treatment, rather than distinct radioactive sources. Therefore, in CY 2008, hospitals would have included the costs of the devices or equipment that are necessary to generate the radiation in their charges for the electronic brachytherapy procedures, in contrast to the conventional brachytherapy procedures also assigned to APC 0313 where the sources would have been separately reported and paid. Therefore, just as in the case of other OPPS services, our ratesetting methodology relies upon hospitals' consideration of all costs associated with furnishing services, including the costs of those items and services for which payment is packaged, in setting hospital charges for separately payable procedures such as electronic brachytherapy. Although the hospital median costs for other HCPCS codes assigned to APC 0313 do not include the cost of radioactive brachytherapy sources that are separately paid, we believe the hospital cost of CPT code 0182T includes the cost of the devices or equipment used to generate the radiation for the treatment. This difference in packaging source payment between conventional and electronic brachytherapy procedures alone does not lead us to conclude that these procedures do not share sufficient cost and clinical similarity to be assigned to

the same clinical APC. The overall hospital costs for conventional and electronic brachytherapy procedures, including the associated packaged costs, that are paid through the procedure codes for electronic and conventional brachytherapy are comparable and the procedures are clinically similar so we believe that their assignment to the same APC is appropriate, regardless of the differences in their packaged costs.

Therefore, we continue to believe that APC 0313 is an appropriate APC for assignment of CPT code 0182T based on our consideration of the clinical characteristics of electronic brachytherapy and hospital costs from claims data. Maintaining CPT code 0182T in APC 0313 for another year would pay at a rate that is three times the cost of this service as reflected in the hospital outpatient claims data, and we do not believe continued payment at \$1,750 is appropriate. To the extent that hospitals' costs change over time if the procedure is more broadly furnished, consistent with our current policy to annually assess the appropriateness of the APC assignments for all services under the hospital OPPS, we will continue to monitor our claims data for CPT code 0182T in the future.

After consideration of the public comments we received, we are finalizing our proposal, without modification, to assign CPT code 0182T to APC 0313, which has a final CY 2010 APC median cost of approximately \$770. Table 23 below lists the HCPCS code and its associated status indicator for CY 2010.

TABLE 23.—CY 2010 REASSIGNMENT OF A NEW TECHNOLOGY PROCEDURE TO A CLINICAL APC

CY 2010 HCPCS code	Short descriptor	CY 2009 SI	CY 2009 APC	Final CY 2010 SI	Final CY 2010 APC
0182T	Hdr elect brachytherapy	S	1519	S	0313

D. OPPS APC-Specific Policies

In this section, we discuss HCPCS codes and their proposed status indicators and APC reassignments for which we provided explicit discussion in section III.D. of the CY 2010 OPPS/ASC proposed rule or for which we received public comments on their proposed CY 2010 OPPS treatment. Certain HCPCS codes are discussed in other sections of this final rule with comment period, as appropriate to the items or services they describe. The final CY 2010 OPPS/ASC treatment of all other HCPCS not explicitly discussed in this final rule with comment period

is displayed in Addendum B to this final rule with comment period.

1. Cardiovascular Services
a. Cardiovascular Telemetry (APC 0209)

For CY 2010, we proposed to continue to assign CPT code 93229 (Wearable mobile cardiovascular telemetry with electrocardiographic recording, concurrent computerized real time data analysis and greater than 24 hours of accessible ECG data storage (retrievable with query) with ECG-triggered and patient-selected events transmitted to a remote attended surveillance center for up to 30 days; technical support for connection and patient instructions for

use, attended surveillance, analysis and physician prescribed transmission of daily and emergent data reports) to APC 0209 (Level II Extended EEG, Sleep, and Cardiovascular Studies), with a proposed payment rate of approximately \$774. Because CPT code 93229 was a new code for CY 2009, in the CY 2009 OPPS/ASC final rule with comment period, we finalized an interim final APC assignment for this code of APC 0209, with a payment rate of approximately \$754.

Comment: Some commenters recommended that CMS assign status indicator "A" (Services furnished to a hospital outpatient that are paid under a fee schedule or payment system other

than OPPS) to CPT code 93229 in order to make this service nonpayable under the OPPS for CY 2010. The commenters argued that there are currently no hospitals that can provide the type of constant monitoring the service described by CPT code 93229 requires. For this reason, according to the commenters, any claims submitted for CPT code 93229 by hospitals are incorrectly coded. The commenters stated that, if CMS chose not to adopt their recommendation and instead chose to continue recognizing CPT code 93229 as payable under the OPPS, CMS should reconsider the proposed assignment of the service to APC 0209. According to the commenters, the service described by CPT code 93229 is not similar clinically or in terms of resource utilization to the other procedures assigned to APC 0209, in particular, the polysomnography procedures described by CPT codes 95810 (Polysomnography; sleep staging with 4 or more additional parameters of sleep, attended by a technologist) and 95811 (Polysomnography; sleep staging with 4 or more additional parameters of sleep, with initiation of continuous positive airway pressure therapy or bilevel ventilation, attended by a technologist), which are the most commonly reported procedures in APC 0209 with the highest number of single claims contributing to the APC's median cost. The commenters urged CMS to assign CPT code 93229 to New Technology APC 1513 (New Technology—Level XIII (\$1,100–\$1,200)) with a payment rate of \$1,150, or New Technology APC 1514 (New Technology—Level XIV (\$1,200–\$1,300)) with a payment rate of \$1,250. The commenters argued that, if any hospitals were to provide the remote cardiac monitoring service described by CPT code 93229, the proposed payment rate for APC 0209 would be less than hospitals' costs for providing this service.

Response: We do not agree with the commenters that we should assign status indicator "A" to CPT code 93229 in order to make the service nonpayable under the OPPS for CY 2010. For each new calendar year, we typically recognize for OPPS payment purposes new HCPCS codes describing services that could be covered by Medicare when provided to hospital outpatients, regardless of whether those services are actually being provided by hospitals at the time the OPPS/ASC final rule with comment period for the upcoming year is issued. We believe that CPT code 93229 describes a diagnostic study that could be provided to Medicare

beneficiaries in the hospital outpatient setting and, therefore, could be covered by Medicare. We also do not agree that the service described by CPT code 93229 is not similar clinically and in terms of resource utilization to the other procedures assigned to APC 0209 for CY 2010. For example, similar to the remote cardiac monitoring service described by CPT code 93229, the polysomnography procedures described by CPT codes 95810 and 95811 involve continuous and simultaneous monitoring and recording of various physiological and pathophysiological parameters, with attendance by a technologist.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to assign CPT code 93229 to APC 0209, with a final CY 2010 APC median cost of approximately \$764.

b. Implantable Loop Recorder Monitoring (APC 0689)

For CY 2010, we proposed to reassign CPT code 93299 (Interrogation device evaluation(s), (remote) up to 30 days; implantable cardiovascular monitor system or implantable loop recorder system, remote data acquisition(s), receipt of transmissions and technician review, technical support and distribution of results) to APC 0689 (Level II Electronic Analysis of Devices), with a proposed payment rate of approximately \$40. In CY 2009, this CPT code was assigned to APC 0209 (Level II Extended EEG, Sleep, and Cardiovascular Studies), with a payment rate of approximately \$754.

Comment: One commenter supported the proposed reassignment of CPT code 93299 to APC 0689 for CY 2010. According to the commenter, the procedure described by CPT code 93299 is similar clinically and in terms of resource utilization to other procedures assigned to APC 0689.

Response: We appreciate the commenter's support of our proposal to reassign CPT code 93299 to APC 0689 for CY 2010. We agree that the procedure described by CPT code 93299 is similar clinically and in terms of resource utilization to other procedures assigned to APC 0689.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to reassign CPT code 93299 to APC 0689, which has a final CY 2010 APC median cost of approximately \$38.

c. Transluminal Balloon Angioplasty (APC 0279)

For CY 2010, we proposed to reassign CPT code 75978 (Transluminal balloon angioplasty, venous (eg, subclavian stenosis), radiological supervision and interpretation) from APC 0083 (Coronary or Non-Coronary Angioplasty and Percutaneous Valvuloplasty) to APC 0279 (Level II Angiography and Venography), with a proposed payment rate of approximately \$2,000.

Comment: Some commenters disagreed with the proposed APC reassignment of CPT code 75978. The commenters noted that CPT code 75978 is a therapeutic interventional procedure that is not similar to the other diagnostic procedures assigned to APC 0279. The commenters requested that CMS continue to assign CPT code 75978 to APC 0083 where they believe the service is most appropriately placed based on considerations of clinical coherence and resource costs.

Response: The proposed CY 2010 median cost for APC 0083 of approximately \$3,380 is significantly higher than the proposed CY 2010 median cost of CPT code 75978 of approximately \$2,597. Given the difference in median costs, we do not believe that we should continue to assign this procedure to APC 0083. After further analysis and review by CMS medical advisors, we believe that CPT code 75978 would be most appropriately assigned to APC 0093 (Vascular Reconstruction/Fistula Repair without Device) for CY 2010 because it is a therapeutic procedure performed on veins, similar to other therapeutic blood vessel procedures that are currently assigned to APC 0093. Further, the CY 2010 final median cost of CPT code 75978 of approximately \$2,597 is very similar to the CY 2010 final median cost of APC 0093 of approximately \$2,378.

After consideration of the public comments we received, we are modifying our proposed CY 2010 reassignment of CPT code 75978 from APC 0083 to APC 0279. In this final rule with comment period, for CY 2010, we are reassigning CPT code 75978 from APC 0083 to APC 0093, which has a final CY 2010 APC median cost of approximately \$2,378.

2. Gastrointestinal Services

a. Change of Gastrostomy Tube (APC 0676)

For CY 2010, we proposed to reassign CPT code 43760 (Change of gastrostomy tube, percutaneous, without imaging or endoscopic guidance) from APC 0121 (Level I Tube or Catheter Changes or Repositioning) to APC 0676

(Thrombolysis and Other Device Revisions), with a proposed CY 2010 payment rate of approximately \$160.

Comment: Some commenters disagreed with the proposed APC reassignment and requested that CMS continue to assign CPT code 43760 to APC 0121 because they believe the gastrostomy tube change procedure shares significant clinical and resource characteristics with other procedures assigned to APC 0121.

Response: Prior to CY 2008, OPSS payment for CPT code 43760 captured both the procedure and the imaging or endoscopic guidance, if used, that was associated with percutaneously changing a gastrostomy tube because its descriptor read, "Change of gastrostomy tube," and the OPSS packages payment for all guidance into payment for the associated procedures. However, effective January 1, 2008, the CPT Editorial Panel revised the code descriptor by adding the words "without imaging or endoscopic guidance" to further clarify that the code should be reported for tube change procedures that do not require imaging or endoscopic guidance. The CPT Editorial Panel further determined that gastrostomy tube placement requiring fluoroscopic or endoscopic guidance should be reported with either CPT code 49450 (Replacement of gastrostomy or cecostomy (or other colonic) tube, percutaneous, under fluoroscopic guidance including contrast injection(s), image documentation and report) or CPT code 43246 (Upper gastrointestinal endoscopy including esophagus, stomach, and either the duodenum and/or jejunum as appropriate; with directed placement of percutaneous gastrostomy tube). Based on the median cost from CY 2008 claims data that reflects the new service reported under CPT code 43760, we believe a reassignment of the CPT code for CY 2010 is necessary.

We disagree with the commenters that CPT code 43760 would be appropriately assigned to APC 0121, which has a median cost of approximately \$426 for CY 2010. We note that the median cost for CPT code 43760 from CY 2007, when the code represented services provided with and without guidance, was higher at approximately \$216, compared with the CY 2008 median cost of the revised procedure code. Claims data from CY 2008 reveal that we have 21,178 single claims (out of 38,246 total claims) for CPT code 43760, with a lower median cost of approximately \$160. The median cost for CPT code 43760 closely aligns with the median cost of approximately \$160 for APC 0676. We note that the procedure for gastrostomy tube placement using

fluoroscopic guidance, specifically CPT code 49450, is assigned to APC 0121, and the procedure for gastrostomy tube placement using endoscopic guidance, specifically CPT code 43246, is assigned to APC 0141 (Level I Upper GI Procedures). We believe that both of these other procedures are appropriately assigned to APCs 0121 and 0141, respectively, based on considerations of clinical and resource homogeneity. As expected, their CPT code-specific median costs that include the cost of fluoroscopy or endoscopic guidance are significantly higher than the median cost of CPT code 43760, which is provided without guidance.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to reassign CPT code 43760 from APC 0121 to APC 0676, which has a final CY 2010 APC median cost of approximately \$160.

b. Laparoscopic Liver Cryoablation (APC 0131)

For CY 2010, we proposed to continue to assign CPT code 47371 (Laparoscopy, surgical, ablation of one or more liver tumor(s); cryosurgical) to APC 0131 (Level II Laparoscopy), with a proposed payment rate of approximately \$3,181.

Comment: One commenter requested that CMS reassign CPT code 47371 from APC 0131 to APC 0174 (Level IV Laparoscopy), which had a proposed payment rate of approximately \$7,766, to better reflect the actual costs of the procedure. The commenter stated that CPT code 47371 is neither similar in resource costs nor clinical characteristics to the other procedures assigned to APC 0131, and that the four single claims for procedure available for ratesetting are not reflective of the costs of the procedure. In addition, the commenter indicated that most of the procedures assigned to APC 0131 describe abdominal biopsy or repair procedures, in contrast to CPT code 47371, which describes cryosurgical ablation of a liver tumor. The commenter noted that there are other similar laparoscopic liver tumor ablation procedures already assigned to APC 0174.

Response: We agree with the commenter's argument that CPT code 47371 would be more appropriately assigned to APC 0174, where other laparoscopic liver and renal ablation procedures are assigned. Although we have few CY 2008 claims for CPT code 47371, our claims data show a higher median cost of approximately \$4,229 for CPT code 47371 based on four single claims out of seven total claims,

compared to the APC median cost of approximately \$3,128 for APC 0131.

We also note that since CPT code 47371 was made effective in CY 2002, the procedure for the code is rarely performed on Medicare beneficiaries in the HOPD based on analysis of our hospital outpatient claims data. Based upon OPSS claims submitted from CY 2002 through CY 2008, the median cost for this code has varied widely, perhaps due to the small volume of claims annually. Specifically, the historical median cost for CPT code 47371 has ranged from \$1,850 based on two single claims to \$6,839 based on one single claim. Although this procedure is not commonly performed on Medicare beneficiaries in the HOPD, because we believe CPT code 47371 is similar in clinical characteristics and resource costs to the other procedures currently assigned to APC 0174, we agree with the commenter's recommendation.

After consideration of the public comment we received, we are modifying our CY 2010 proposal and assigning CPT code 47371 to APC 0174, which has a final CY 2010 APC median cost of approximately \$7,342.

c. Cholangioscopy (APC 0151)

The CPT Editorial Panel created a new add-on code for cholangioscopy, CPT code 43273 (Endoscopic cannulation of papilla with direct visualization of common bile duct(s) and/or pancreatic ducts (List separately in addition to code(s) for primary procedure), effective January 1, 2009. We assigned CPT code 43273 to APC 0151 (Endoscopic Retrograde Cholangio-Pancreatography (ERCP)) on an interim final basis in the CY 2009 OPSS/ASC final rule with comment period (73 FR 69030), and the CPT code was flagged with comment indicator "NI" to indicate that its OPSS treatment was open to comment on that final rule with comment period. For CY 2010, we proposed to continue the assignment of CPT code 43273 to APC 0151, with a proposed payment rate of approximately \$1,527.

At the August 2009 APC Panel meeting, the APC Panel heard a public presentation recommending that CPT code 43273 be reassigned to APC 0152 (Level I Percutaneous Abdominal and Biliary Procedures). However, the APC Panel recommended that CPT code 43273 continue to be assigned to APC 0151.

Comment: One commenter on the CY 2009 OPSS/ASC final rule with comment period and again on the CY 2010 OPSS/ASC proposed rule recommended that CPT code 43273 be reassigned to APC 0152 and that CMS

rename APCs 0151 and 0152 to accommodate the reassignment. The commenter provided cost information for CPT code 43273 in a New Technology APC application previously filed with CMS, estimating that the cost of the cholangioscopy procedure is approximately \$2,958. Because the procedure is always performed with an ERCP procedure, the commenter estimated that the combined cost of ERCP and cholangioscopy is approximately \$4,484 by adding the CY 2008 median cost of APC 0151 to the estimated cost of the cholangioscopy procedure. The commenter pointed out that, because all ERCP CPT codes, which are assigned to APC 0151, have a status indicator of "T" (Significant Procedure, Multiple Reduction Applies), under CMS' proposal, CPT code 43273 would be paid at 50 percent of the CY 2010 proposed payment rate for APC 0151, or \$757, a rate that is approximately \$2,200 less than the reported cost of \$2,958 for the cholangioscopy procedure alone. The commenter further estimated the cost of disposable devices alone for cholangioscopy in combination with ERCP to be approximately \$2,064. The commenter argued that the proposed CY 2010 payment hospitals would receive for the two procedures of approximately \$2,271 would barely cover the device costs of \$2,064, and not the additional procedure costs. The commenter maintained that if CMS reassigned CPT code 43273 to APC 0152, payment for the combination of the two procedures would be approximately \$2,821, partially closing the gap between OPPS payment and the commenter's estimated combined cost of the two procedures of \$4,484.

The commenter explained that cholangioscopy is a complex, resource-intensive procedure requiring additional physician training, which adds 45 minutes to the ERCP procedure, for a total of approximately 112 minutes for the two procedures. The commenter also asserted that the clinical intensity of cholangioscopy is closer to the non-draining percutaneous procedures assigned to APC 0152 than to the ERCP procedures assigned to APC 0151. Further, the commenter explained that the primary clinical difference between non-draining percutaneous procedures and CPT code 43273 is the method of access to the biliary and pancreatic area, noting that a small incision is required for the percutaneous procedures and the use of an additional endoscope for cholangioscopy.

The commenter also recommended that APC 0151 be renamed "Level I Hepatobiliary Procedures" and APC

0152 be renamed "Level II Hepatobiliary Procedures," and that lower complexity hepatobiliary procedures be assigned to APC 0151 and higher complexity hepatobiliary procedures, including percutaneous procedures and cholangioscopy, be assigned to APC 0152. The commenter believed that renaming and reconfiguring APCs 0151 and 0152 would improve the clinical homogeneity of the APCs and appropriately account for differences in the resource costs of biliary procedures. The commenter argued that CMS has previously renamed existing APCs, created multilevel APCs for specific clinical areas, and configured APCs to incorporate surgical procedures that include a variety of access types.

Response: Because CPT code 43273 was new for CY 2009, we do not yet have cost information for the procedure based upon hospital claims for CY 2010 ratesetting. According to our established policy, a New Technology APC applicant's cost estimate is only one source of information we consider in determining the cost of a new service for purposes of its initial APC assignment under the OPPS (66 FR 59900; 73 FR 68614). We generally assign new CPT codes to an APC based on input from a variety of sources, including, but not limited to, review of the resource costs and clinical similarity of the service to existing procedures; input from CMS medical advisors; information from interested specialty societies; and review of all other information available to us. We note that, while CPT code 43273 is new for CY 2009, cholangioscopy in association with ERCP procedures has been performed, using a variety of technologies, for many years. We expect that its costs have already been incorporated into the OPPS, either packaged into payment for the associated ERCP procedures or under an unlisted CPT procedure code.

We continue to believe that APC 0151 is an appropriate APC assignment for CPT code 43273 for CY 2010, based on consideration of the procedure's clinical and resource characteristics. CPT code 43273, which is an add-on code to ERCP procedures, clinically resembles the ERCP procedures that also are assigned to APC 0151 because they all use an endoscope to examine various components of the hepatobiliary system. While cholangioscopy extends ERCP procedures to visualize the common bile duct(s) and/or pancreatic duct(s) through use of an additional endoscope, many of the other ERCP procedures assigned to APC 0151 also have additional procedures associated with them that require the use of other devices or equipment as well. We do not

agree with the commenter that percutaneous biliary procedures that require incisions and, in some cases drainage tubes, are more clinically similar to cholangioscopy than ERCP procedures that also examine the hepatobiliary system with an endoscope.

Furthermore, we understand that there are a variety of technologies that can be used to perform cholangioscopy with variable resource costs. Therefore, we do not believe that the commenter's estimate based on one type of new cholangioscopy technology is necessarily an appropriate representation of the cost of the procedure described by CPT code 43273 to hospitals. We believe that the cost of cholangioscopy is similar to the cost of ERCP procedures that require similar procedure time and devices, and that CPT code 43273 is appropriately assigned to APC 0151 along with these ERCP procedures. Moreover, we believe that applying the multiple procedure discount to payment for cholangioscopy as a result of its status indicator "T" and its routine performance with ERCP procedures that also are assigned status indicator "T" is appropriate because cholangioscopy is performed directly after ERCP and much of the preparatory procedure work is performed during the ERCP.

Finally, because we are not reassigning CPT code 43273 to APC 0152, no renaming of APCs 0151 and 0152 is warranted because we are maintaining the endoscopic and percutaneous biliary procedures in separate APCs.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to maintain the assignment of CPT code 43273 to APC 0151, which has a final CY 2010 APC median cost of approximately \$1,510.

d. Laparoscopic Hernia Repair (APC 0131)

For CY 2010, we proposed to reassign the following six laparoscopic hernia repair CPT codes that were new for CY 2009 from APC 0130 (Level I Laparoscopy), with a proposed payment rate of approximately \$2,538, to APC 0131 (Level II Laparoscopy), with a proposed payment rate of approximately \$3,181: CPT code 49652 (Laparoscopy, surgical, repair, ventral, umbilical, spigelian or epigastric hernia (includes mesh insertion, when performed); reducible); CPT code 49653 (Laparoscopy, surgical, repair, ventral, umbilical, spigelian or epigastric hernia (includes mesh insertion, when performed); incarcerated or

strangulated); CPT code 49654 (Laparoscopy, surgical, repair, incisional hernia (includes mesh insertion, when performed); reducible); CPT code 49655 (Laparoscopy, surgical, repair, incisional hernia (includes mesh insertion, when performed); incarcerated or strangulated); CPT code 49656 (Laparoscopy, surgical, repair, recurrent incisional hernia (includes mesh insertion, when performed); reducible); and CPT code 49657 (Laparoscopy, surgical, repair, recurrent incisional hernia (includes mesh insertion, when performed); incarcerated or strangulated).

Comment: One commenter indicated that the resource costs associated with the six laparoscopic hernia repair CPT codes are significantly greater than the proposed payment rate of approximately \$3,181 for APC 0131 and suggested that the procedures in APC 0132 (Level III Laparoscopy) are more similar to the six laparoscopic hernia repair codes because they have similar resource costs. The commenter requested that CMS review the clinical characteristics and resource costs associated with the six CPT codes and consider reassigning these codes to APC 0132, which had a CY 2010 proposed payment rate of approximately \$4,903. In addition, the commenter provided an analysis of CY 2008 claims data for cases reported under the unlisted CPT code that would previously have been reported for these procedures (CPT code 49659 (Unlisted laparoscopy procedure, hernioplasty, herniorrhaphy, herniotomy)), combined with the ICD-9 diagnoses codes specific to hernia of the abdominal cavity. Because these codes were new for CY 2009, they were assigned comment indicator "NI" in the CY 2009 OPSS/ASC final rule with comment period to indicate that they were subject to comment. For the CY 2009 OPSS/ASC final rule with comment period, the same commenter submitted a similar comment and data analysis of CY 2007 OPSS claims. The commenter found 3,456 claims that met the same case criteria in the analysis, with a median cost of approximately \$4,261. The commenter believed that this cost from historical hospital claims data represented the hospital cost of procedures that would be reported with one of the laparoscopic hernia repair CPT codes in CY 2010.

Response: We have no hospital claims data for CPT codes 49652 through 49657 because these CPT codes were new for CY 2009. However, we agree with the commenter that procedures described by these CPT codes were likely commonly furnished in the HOPD in CY 2008 and reported under CPT code

49659. Taking into consideration the commenter's analyses of CY 2007 and CY 2008 claims and performing a detailed clinical review, we agree with the commenter that the resource costs for the six laparoscopic hernia repair CPT codes, specifically CPT codes 49652 through 49657, more closely align with other services assigned to APC 0132. In addition, from a clinical perspective, we also believe that APC 0132 is an appropriate APC assignment for these codes because APC 0132 most accurately recognizes the complexity of the laparoscopic hernia repair codes.

After consideration of the public comments we received, we are modifying our CY 2010 proposals and reassigning CPT codes 49652, 49653, 49654, 49655, 49656, and 49657 from APC 0130 to APC 0132, which has a final CY 2010 APC median cost of approximately \$4,873.

3. Genitourinary Services

a. Percutaneous Renal Cryoablation (APC 0423)

For CY 2010, we proposed to continue to assign CPT code 50593 (Ablation, renal tumor(s), unilateral, percutaneous, cryotherapy) to APC 0423 (Level II Percutaneous Abdominal and Biliary Procedures), with a proposed payment rate of approximately \$3,329. This CPT code was new in CY 2008. However, the same service was previously described by CPT code 0135T (Ablation renal tumor(s), unilateral, percutaneous, cryotherapy). We note that, for CY 2007, based upon the APC Panel's recommendation made at its March 2006 meeting, we reassigned CPT code 50593 (then CPT code 0135T) from APC 0163 (Level IV Cystourethroscopy and other Genitourinary Procedures) to APC 0423, effective January 1, 2007.

Comment: One commenter expressed concern that the proposed payment rate of approximately \$3,329 for CPT code 50593 is inadequate because the payment does not accurately account for the costs incurred by hospitals in performing this procedure. The commenter argued that the low proposed payment rate for CPT code 50593 is attributable to claims data that do not accurately capture the full costs of CPT code 50593 because almost half of the single claims do not contain the HCPCS code and associated charge for the required device, specifically HCPCS code C2618 (Probe, cryoablation). The commenter requested that CMS designate CPT code 50593 as a device-dependent procedure, which would require hospitals to submit claims with the appropriate device HCPCS code, assign the procedure to its own APC,

and set the payment rate for that APC based on claims for CPT code 50593 reported with HCPCS code C2618. The commenter's analysis concluded that the median cost on which payment for CPT code 50593 would be based if these recommendations were adopted would be approximately \$5,469, resulting in more accurate payment for the procedure and continued Medicare beneficiary access to percutaneous renal cryoablation in the HOPD.

Response: We believe that CPT code 50593 is appropriately assigned to APC 0423 based on clinical and resource considerations when compared to other procedures also proposed for assignment to APC 0423 for CY 2010. As we stated in the CY 2007 OPSS final rule with comment period (71 FR 68049 through 68050), the CY 2008 OPSS/ASC final rule with comment period (72 FR 66709), and the CY 2009 OPSS/ASC final rule with comment period (73 FR 68611), we initially revised the APC assignment for the percutaneous renal cryoablation procedure from APC 0163 to APC 0423 in CY 2007 based on the APC Panel's recommendation to reassign the procedure to APC 0423. The median costs of the four HCPCS codes assigned to APC 0423 for CY 2010 range from approximately \$3,159 to \$4,670, well within the 2 times rule for the OPSS payment groups. Even if we were to calculate the median cost for CPT code 50593 using only claims that also contain HCPCS code C2618, estimated by the commenter to be approximately \$5,469 using proposed rule data, the grouping of these procedures in the same APC would not violate the 2 times rule. Further, we note that all four of these procedures are relatively low volume, with fewer than 1,100 total claims each for CY 2008 and fewer than 700 single claims each for ratesetting. We believe that grouping these clinically similar, low volume procedures for the percutaneous ablation of renal, liver, or pulmonary tumors in the same payment group helps to promote payment stability for these low volume services.

We also do not agree that CPT code 50593 should be designated as a device-dependent procedure and assigned to its own separate APC. We have only 226 single claims (out of 513 total claims) for CPT code 50593 from CY 2008 and, as such, the procedure has the second lowest frequency of the four procedures assigned to APC 0423. We believe this relatively low volume procedure should be assigned to a payment group with similar services, as we have proposed, in order to promote payment stability and encourage hospital efficiency. In addition, we do not identify individual

HCPCS codes as device-dependent HCPCS codes under the OPPS. Rather, we first consider the clinical and resource characteristics of a procedure and determine the most appropriate APC assignment. When we determine that we should assign a procedure to an APC that is device-dependent, based on whether that APC has been historically identified under the OPPS as having very high device costs, we then consider the implementation of device edits, as appropriate. We note that the identification of device-dependent APCs was particularly important in the early years of the OPPS when separate pass-through payment for many implantable devices expired. At that time, a variety of methodologies to package the costs of those devices into procedural APCs was utilized over several years to ensure appropriate incorporation of the device costs into the procedure payments. At this point in time, hospitals have significantly more experience reporting HCPCS codes for packaged and separately payable items and services under the OPPS and the payment groups are more mature. We believe our standard ratesetting methodology typically results in appropriate payment rates for new procedures that utilize devices, as well as those that do not use high cost devices. In recent years, we have not encountered circumstances whereby we have had to establish new device-dependent APCs because we were not able to accommodate the clinical and resource characteristics of a procedure by assigning it to an existing APC (whether device-dependent or non-device-dependent), and the procedure described by CPT code 50593 is no exception.

While all of the procedures assigned to APC 0423 require the use of implantable devices, for many of the procedures, there are no Level II HCPCS codes that describe all of the technologies that may be used in the procedures. Therefore, it would not be possible for us to develop procedure-to-device edits for all of the CPT codes assigned to APC 0423. Under the OPPS, there are many other procedures that require the use of implantable devices that, because they are assigned to OPPS APCs that are not device-dependent, do not have procedure-to-device edits applied, even if those claims processing edits would be feasible. We believe that our payments for procedures that utilize high cost devices are appropriate for those services, even when those services are grouped with other procedures that either do not require the use of implantable devices or that utilize

devices that are not described by specific Level II HCPCS codes.

When reporting CPT code 50593, we expect hospitals to also report the device HCPCS code C2618, which is associated with this procedure. We also remind hospitals that they must report all of the HCPCS codes that appropriately describe the items used to provide services, regardless of whether the HCPCS codes are packaged or paid separately. If hospitals use more than one probe in performing CPT code 50593, we expect hospitals to report this information on the claim and adjust their charges accordingly. Hospitals should report the number of cryoablation probes used to perform CPT code 50593 as the units of HCPCS code C2618 which describes these devices, with their charges for the probes. Since CY 2005, we have required hospitals to report device HCPCS codes for all devices used in procedures if there are appropriate HCPCS codes available. In this way, we can be confident that hospitals have included charges on their claims for costly devices used in procedures when they submit claims for those procedures.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign CPT code 50593 to APC 0423, which has a final CY 2010 APC median cost of approximately \$3,430.

b. Hemodialysis (APC 0170)

Currently, APC 0170 (Dialysis) contains two HCPCS codes: CPT code 90935 (Hemodialysis procedure with single physician evaluation) and HCPCS code G0257 (Unscheduled or emergency dialysis treatment for an ESRD patient in an HOPD that is not certified as an ESRD facility). Hospital outpatient and emergency departments sometimes must furnish hemodialysis to patients who do not have ESRD and, in these cases, they would report CPT code 90935 for the service. Under the Medicare ESRD benefit, to be covered by Medicare, routine dialysis required by a Medicare ESRD beneficiary must be furnished in a certified ESRD facility. Most HOPDs and emergency departments are not certified by Medicare to furnish routine dialysis to Medicare ESRD patients and, therefore, are not paid under the ESRD benefit. However, there are a limited number of specific cases in which Medicare pays under the OPPS for an ESRD patient to receive unscheduled dialysis in an outpatient department of a hospital that does not have an ESRD-certified facility. These provisions were established in the CY 2003 OPPS final rule with comment period (67 FR 66803

through 66805). Specifically, Medicare pays hospitals under the OPPS for dialysis for ESRD patients under the following limited circumstances as specified in the Medicare Claims Processing Manual, Pub. 100-04, Chapter 4, Section 200.2:

- Dialysis is performed following or in connection with a dialysis-related procedure such as a vascular access procedure or a blood transfusion;
- Dialysis is performed following treatment for an unrelated medical emergency; or
- Emergency dialysis is performed for ESRD patients who would otherwise have to be admitted as inpatients in order for the hospital to receive payment.

When these criteria are met, the hospital reports HCPCS code G0257, which we proposed to assign to APC 0170 for CY 2010, with a proposed payment of approximately \$442.

Comment: One commenter, who recognized that Medicare pays under the OPPS for hemodialysis for ESRD patients under very specific, limited circumstances, asked whether hospitals are permitted to bill and be paid under the OPPS for routine hemodialysis for ESRD patients who are unable to arrange for routine hemodialysis at a Medicare-certified ESRD facility.

Response: As the commenter noted, Medicare pays under OPPS for dialysis for a beneficiary with ESRD only under the exceptional circumstances specified in the Medicare Claims Processing Manual, Pub. 100-04, Chapter 4, Section 200.2 that are listed above. Routine treatments in hospitals that do not have an ESRD facility are not payable under the OPPS. A hospital that would like to provide routine hemodialysis to ESRD patients should contact the State survey and certification agency to pursue ESRD certification of an outpatient dialysis unit.

After consideration of the public comment we received, we continue to believe that our policy governing the payment of dialysis services in HOPDs and emergency departments is appropriate. We are not making any change to this policy for CY 2010. The final CY 2010 median cost of APC 0170 for hemodialysis is approximately \$456.

c. Radiofrequency Remodeling of Bladder Neck (APC 0165)

For CY 2010, we proposed to continue to assign Category III CPT code 0193T (Transurethral, radiofrequency micro-remodeling of the female bladder neck and proximal urethra for stress urinary incontinence) to APC 0165 (Level IV Urinary and Anal Procedures) with a

proposed payment rate of approximately \$1,353. This CPT code was new for CY 2009 and was assigned to APC 0165 on an interim final basis in the CY 2009 OPPI/ASC final rule with comment period.

At the August 2009 APC Panel meeting, a presenter requested that the APC Panel recommend that CMS reassign CPT code 0193T to either APC 0202 (Level VII Female Reproductive Procedures) or APC 0168 (Level II Urethral Procedures) for CY 2010 based on resource intensity and therapeutic benefit. The presenter claimed that the device cost associated with CPT code 0193T is comparable to the single-use devices that are used with certain procedures assigned to APC 0202, specifically those procedures described by CPT codes 58356 (Endometrial cryoablation with ultrasonic guidance, including endometrial curettage, when performed); 58565 (Hysteroscopy, surgical; with bilateral fallopian tube cannulation to induce occlusion by placement of permanent implants); and 57288 (Sling operation for stress incontinence (e.g., fascia or synthetic)). The presenter indicated that, unlike procedures assigned to APC 0202 that require costly medical devices, the costs of single-use medical devices for procedures assigned to APC 0165 are very minimal. After a discussion, the APC Panel recommended that CMS maintain the APC assignment of CPT code 0193T to APC 0165, as proposed, for CY 2010.

Comment: Some commenters disagreed with CMS' proposal to continue to assign CPT code 0193T to APC 0165. The commenters believed that the proposed payment for the procedure would not pay appropriately for the costs incurred by hospitals to perform the procedure, especially because the procedure utilizes a costly, single-use, disposable medical device. The commenters argued that APC 0202, which had a proposed CY 2010 proposed payment rate of approximately \$2,991, contains procedures that are very similar to CPT code 0193T. Specifically, the commenters indicated that CPT code 0193T is similar in clinical characteristics and resource costs to CPT codes 58356, 58565, and 57288. The commenters added that the probe used in the procedure reported with CPT code 0193T costs \$1,095 and, overall, the total procedure cost that includes the cost of the probe is approximately \$2,473, which is comparable to the proposed CY 2010 payment rate for APC 0202.

Another commenter was concerned that, at the August 2009 APC Panel meeting, the APC Panel members may

have been confused about the surgical nature of CPT code 0193T. Specifically, the commenter believed that the APC Panel concluded that all of the procedures assigned to APC 0202 are surgical in nature, whereas the procedure described by CPT code 0193T is not, which resulted in the APC Panel's recommendation to continue to assign this code to APC 0165. The commenter clarified that CPT code 0193T is similar to surgical CPT codes 58565 and 58356, which are both assigned to APC 0202 based on their resource use and clinical characteristics. The commenter further noted that, although CPT code 0193T may be performed in the physician's office, in the Medicare population, this procedure is more likely to be performed in the hospital outpatient setting because of medical conditions and comorbidities experienced by Medicare patients.

Response: As a new Category III CPT code for CY 2009, we do not yet have hospital claims data for the procedure. Category III CPT codes are temporary codes that describe emerging technology, procedures, and services, and they are created by the AMA to allow for data collection for new services or procedures. Under the OPPI, we generally assign a payment rate to a new Category III CPT code based on input from a variety of sources, including but not limited to, review of resource costs and clinical homogeneity of the service to existing procedures, information from specialty societies, input from CMS medical advisors, and other information available to us. Based on our review of the clinical characteristics of CPT code 0193T, as well as the other procedures assigned to APC 0165 and APC 0202 that was recommended by the commenters, and the APC Panel discussion and recommendation regarding the procedure, we continue to believe that APC 0165 is the most appropriate APC assignment for CPT code 0193T for CY 2010. We understand that CPT code 0193T is a minimally invasive procedure for female stress urinary incontinence that requires a relative brief time in the procedure room. We do not agree with the commenters that the procedures assigned to APC 0202 that involve fallopian tube cannulation, endometrial ablation, or implantation of a sling for stress urinary incontinence are sufficiently similar to the procedure described by CPT code 0193T based on procedure duration, device utilization, use of guidance, or other characteristics to warrant reassignment of CPT code 0193T to APC 0202 based on considerations of clinical homogeneity.

Rather, we believe that assignment to APC 0165 will appropriately account for the device and procedure costs of CPT code 0193T.

After consideration of the public comments we received and the APC Panel recommendation from the August 2009 meeting, we are finalizing our CY 2010 proposal, without modification, to continue to assign CPT code 0193T to APC 0165, which has a final CY 2010 APC median cost of approximately \$1,337.

d. Change of Bladder Tube (APC 0121)

For CY 2010, we proposed to reassign CPT code 51710 (Change of cystostomy tube; complicated) from APC 0427 (Level II Tube or Catheter Changes or Repositioning) to APC 0121 (Level I Tube or Catheter Changes or Repositioning), with a proposed CY 2010 payment rate of approximately \$428.

Comment: One commenter supported the proposed APC reassignment of CPT code 51710 from APC 0427 to APC 0121.

Response: We appreciate the commenter's support. Hospital outpatient claims data revealed that we have approximately 267 single claims (out of 431 total claims) for CPT code 51710, with a final CY 2010 median cost of approximately \$446. The final CY 2010 median cost for CPT code 51710 closely aligns with the final CY 2010 median cost of approximately \$426 for APC 0121. We believe that CPT code 51710 is appropriately reassigned to APC 0121 based on clinical and resource considerations.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to reassign CPT code 51710 from APC 0427 to APC 0121, which has a final CY 2010 APC median cost of approximately \$426.

4. Nervous System Services

a. Pain-Related Procedures (APCs 0203, 0204, 0206, 0207, 0221, 0224, and 0388)

We proposed to set the CY 2010 payment rates for APCs 0203 (Level IV Nerve Injections), 0204 (Level I Nerve Injections), 0206 (Level II Nerve Injections), 0207 (Level III Nerve Injections), 0221 (Level II Nerve Procedures), 0224 (Implantation of Catheter/Reservoir/Shunt) and 0388 (Discography) based on the median costs determined under the OPPI standard ratesetting. Among the CPT codes included in these APCs are: 62350 (Implantation, revision, or repositioning of tunneled intrathecal or epidural catheter for long-term medication

administration via an external pump or implantable reservoir/infusion pump; with laminectomy); 62355 (Removal of previously implanted intrathecal or epidural catheter); 62365 (Removal of subcutaneous reservoir or pump, previously implanted for intrathecal or epidural infusion); 64472 (Injection, anesthetic agent and/or steroid, paravertebral facet joint or facet joint nerve; cervical or thoracic, each additional level (list separately in addition to code for primary procedure)); 64476 (Injection, anesthetic agent and/or steroid, paravertebral facet joint or facet joint nerve; lumbar or sacral, each additional level (list separately in addition to code for primary procedure)); 64480 (Injection, anesthetic agent and/or steroid, transforminal epidural; cervical or thoracic, each additional level (list separately in addition to code for primary procedure)); 64623, (Destruction by neurolytic agent, paravertebral facet joint nerve; lumbar or sacral, each additional level, (list separately in addition to code for primary procedure)); 64627 (Destruction by neurolytic agent, paravertebral facet joint nerve; cervical or thoracic, each additional level, (list separately in addition to code for primary procedure)); 72285 (Discography, cervical or thoracic, radiological supervision and interpretation); and 72295 (Discography, lumbar, radiological supervision and interpretation).

Comment: One commenter objected to the proposed CY 2010 payment rates for CPT codes 64472, 64476, 64480, 64623, and 64627, which the commenter believed have declined 28 percent to 48 percent since CY 2007. The commenter also objected to the proposed CY 2010 increase in payments for CPT codes 72285 and 72295 on the basis that their proposed payment rates are unreasonable because they are not procedures. The commenter added that CPT codes 62290 (Injection procedure for discography, each level, lumbar) and 62291 (Injection procedure for discography, each level, cervical or thoracic) are the related procedures,

which are paid at an unreasonably low rate.

Response: OPPS payment rates fluctuate based on a variety of factors, including, but not limited to, changes in the mix of hospitals billing the services, differential changes in hospital charges and costs for the services, and changes in the volumes of services reported. Therefore, the median costs on which the OPPS payment rates are based vary from one year to another. We note that the median costs of all of the APCs to which CPT codes 64472, 64476, 64480, 64623, and 64627 are assigned increased between CY 2009 and CY 2010. For CPT codes 64472 and 64480, the median cost of APC 0206 to which they are assigned increased from approximately \$236 in CY 2009 to approximately \$249 in CY 2010. In the case of CPT codes 64476 and 64627, the median cost of APC 0204 to which they are assigned increased from approximately \$161 in CY 2009 to approximately \$171 in CY 2010. Lastly, for CPT code 64623, the median cost of APC 0207 to which the code is assigned increased from approximately \$463 in CY 2009 to approximately \$481 in CY 2010.

CPT codes 72285 and 72295, both of which are assigned to APC 0388, are “T” packaged codes and, as such, are paid separately only if there is no separately paid surgical procedure with a status indicator of “T” on the same claim. When there is a separate payment made for these codes, the payment is not only payment for the code itself but also includes payment for all services reported on the claim that are always packaged (that is, those with a status indicator of “N”). The median cost of APC 0388 to which CPT codes 72285 and 72295 are assigned for payment when separate payment can be made increased from approximately \$1,470 in CY 2009 to approximately \$1,727 in CY 2010, reflecting the cost of all conditionally and unconditionally packaged services on the claim. Payment for CPT codes 62290 and 62291 is always packaged into payment for the independent, separately paid procedures with which these codes are reported because we believe that these codes are ancillary and supportive to other major separately paid procedures

and that they are furnished only as an ancillary and dependent part of an independent separately paid procedure.

Comment: One commenter disagreed with the proposed CY 2010 payment rates for CPT codes 62355, 62350, 62363 (we note that this code did not exist in CY 2008 and does not exist in CY 2009; the commenter did not provide a description of the procedure that would enable us to identify the code and respond to the comment), and 62365. The commenter believed that access to these services is very limited as a result of payment reductions for these procedures.

Response: The final median costs for the APCs to which CPT codes 62350 and 62365 are assigned for CY 2010 increased from CY 2009 to CY 2010, while the final median cost for APC 0203 to which CPT code 62355 is assigned declined over that same time period. Specifically, CPT code 62350 is assigned to APC 0224, which has a CY 2009 median cost of approximately \$2,715 and a final CY 2010 median cost of approximately \$2,740. Similarly, CPT code 62365 is assigned to APC 0221, which has a CY 2009 median cost of approximately \$2,322 and a final CY 2010 median cost of approximately \$2,490. In contrast, CPT code 62355 is assigned to APC 0203, which has a CY 2009 median cost of approximately \$928 that declined to approximately \$885 in CY 2010. The increased median costs of APCs 0221 and 0224 do not create barriers to care for these procedures. Moreover, we do not believe that the modest reduction in median cost for APC 0203 would cause hospitals to cease to furnish the service.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to pay for CPT codes 62350, 62355, 62365, 64472, 64476, 64480, 64623, 64627, 72285, and 72295 through APCs 0203, 0204, 0206, 0207, 0221, 0224, and 0388. The final CY 2010 median costs of the relevant APCs are displayed in Table 24 below. For comparative purposes, we also are showing in the table the median costs on which the CY 2009 OPPS payments are based.

TABLE 24—MEDIAN COSTS FOR SELECTED APCs FOR PAIN-RELATED PROCEDURES MENTIONED BY COMMENTERS

APC	APC title	CY 2009 approximate median cost	Proposed CY 2010 approximate median cost	Final CY 2010 approximate median cost
0203	Level IV Nerve Injections	\$929	\$1,066	\$885
0204	Level I Nerve Injections	161	181	171
0206	Level IV Nerve Injections	236	254	249
0207	Level III Nerve Injections	463	504	481

TABLE 24—MEDIAN COSTS FOR SELECTED APCs FOR PAIN-RELATED PROCEDURES MENTIONED BY COMMENTERS—
Continued

APC	APC title	CY 2009 approximate median cost	Proposed CY 2010 approximate median cost	Final CY 2010 approximate median cost
0221	Level II Nerve Procedures	2,322	2,521	2,490
0224	Implantation of Catheter/Reservoir/Shunt	2,715	2,769	2,740
0388	Discography	1,470	1,769	1,727

b. Magnetoencephalography (APCs 0065 and 0067)

Three CPT codes describe magnetoencephalography services: 95965 (Magnetoencephalography (MEG), recording and analysis; for spontaneous brain magnetic activity (e.g. epileptic cerebral cortex localization)); 95966 (Magnetoencephalography (MEG), recording and analysis; for spontaneous brain magnetic activity (e.g. epileptic cerebral cortex localization) for evoked magnetic fields, single modality (e.g. sensory, motor, language or visual cortex localization)); and 95967 (Magnetoencephalography (MEG), recording and analysis; for spontaneous brain magnetic activity (e.g. epileptic cerebral cortex localization), for evoked magnetic fields, each additional modality (e.g. sensory, motor language, or visual cortex localization (List separately in addition to code for primary procedure)). These CPT codes were originally assigned to New Technology APCs but, beginning in CY 2006 and for every year thereafter, these codes have been assigned to clinical APCs on the basis of the clinical and resource characteristics of the services. For CY 2010, we proposed to continue to assign CPT code 95965 to APC 0067 (Level III Stereotactic Radiosurgery, MRgFUS and MEG) with a proposed payment rate of approximately \$3,507, and we proposed to continue to assign CPT codes 95966 and 96967 to APC 0065 (Level II Stereotactic Radiosurgery, MRgFUS and MEG), with a proposed payment rate of approximately \$894.

Comment: Several commenters requested that CMS restore the payment rates for CPT codes 95965, 95966, and 95967 to the levels at which they were paid under New Technology APCs in CY 2005 of \$5,250, \$1,450, and \$950, respectively. They believed the payment rates for CYs 2006, 2007, 2008, and 2009 and the proposed rate for CY 2010 were based on median cost calculations that understated the full costs of the services. The commenters asked CMS to create a cost center on the Medicare cost report that would be used solely to house hospitals' costs of MEG and

indicated that the NUBC had approved a request for a dedicated revenue code for the reporting of charges for MEG. The commenters argued that if CMS would create a cost center for the costs of MEG from which a specific CCR could be developed for application to MEG charges, the resulting median cost would be a more accurate reflection of the cost of MEG and would, therefore, result in more appropriate payment. One commenter submitted eight claims for MEG, stating that its Medicare contractor had approved use of a subscript for a specific cost center on the cost report to house the costs of MEG. The commenter asked if these claims were used in ratesetting, provided the CCR the commenter calculated using the costs and charges for MEG that would be reported on the cost report line that contained only MEG costs, and asked if CMS calculated the costs of these claims using the specific CCR for MEG services.

Response: We assign new services to New Technology APCs only until we believe that we have sufficient historical hospital claims data reflecting hospital costs to reassign them to appropriate clinical APCs. We initially assigned MEG services to New Technology APCs based on the information available to us at the time about the expected hospital costs. For CY 2006, we believed that we had sufficient claims data to enable us to make informed decisions regarding the proper clinical APCs for assignment of MEG services. We note that the volumes of claims for MEG services have remained stable since we moved them to clinical APCs in CY 2006. We have no reason to believe that the costs that we have derived from our standard cost estimation process for the CY 2010 OPPS fail to appropriately reflect the relative costs of MEG services in relation to the costs of other services paid under the OPPS, nor do we have reason to believe that payment at the rates under which these services were paid under the New Technology APCs in CY 2005 are justified.

With regard to whether individual claims that were submitted by one commenter were used to set the median

costs on which the CY 2010 MEG payment rates are based, we note that the claims we use to set the payment rates under the OPPS are available for purchase and a provider that wishes to see if particular claims were used can attain the claims file and perform any analysis they choose. We are not able to create provider-specific revenue code-to-cost center crosswalks that would use unique cost report subscripts that hospitals choose to create for particular services. In the case of a hospital reporting MEG costs on a subscripted line 54.01, the costs would be included as costs in cost center 5400 (the cost center to which 54.01 is a subscripted line), the standard cost center for electroencephalography. In accordance with our standard revenue code-to-cost center crosswalk, we would apply the CCR for this cost center to the charges reported under revenue code 0740 (EEG (Electroencephalogram); General Classification)) if there is no CCR available for nonstandard cost center 3280 (EKG and EEG).

We recognize that the NUBC created a new revenue code for MEG on August 11, 2009, to be effective for services reported on or after April 1, 2010, if a hospital chooses to use it. We anticipate that we will propose to use claims for services furnished in CY 2010 to calculate OPPS payment rates for CY 2012. Therefore, for the CY 2012 OPPS, we expect that we will propose to determine the primary, secondary and tertiary (if any) CCRs to be applied to the new revenue code as part of our standard ratesetting process for the CY 2012 OPPS. With regard to requests for a dedicated cost center for MEG services, the revised draft hospital cost report Form CMS-2552-10 went on public display through the **Federal Register** (74 FR 31738), with a comment period that ended on August 31, 2009. We will consider whether creation of such a cost center is appropriate in our review of all public comments on the proposed revisions to the cost report.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to continue to

assign CPT code 95965 to APC 0067, with a final CY 2010 median cost of approximately \$3,539, and to continue to assign CPT codes 95966 and 96967 to APC 0065, with a final CY 2010 median cost of approximately \$954.

5. Ocular Services

a. Insertion of Anterior Segment Aqueous Drainage Device (APC 0234)

The CPT Editorial Panel created Category III CPT code 0191T (Insertion of anterior segment aqueous drainage device, without extraocular reservoir; internal approach), effective on July 1, 2008. We assigned CPT code 0191T to APC 0234 (Level III Anterior Segment Eye Procedures), effective July 1, 2008, and maintained this APC assignment for CY 2009. For CY 2010, we proposed to continue the assignment of CPT code 0191T to APC 0234, with a proposed payment rate of approximately \$1,639.

Comment: One commenter asserted that the assignment of CPT code 0191T to APC 0234 for CY 2010 would not provide sufficient payment to hospitals and ASCs to cover the cost of the procedure and, therefore, is inappropriate. The commenter indicated that the manufacturer of the device system inserted in the procedure reported by CPT code 0191T currently has an Investigational Device Exemption (IDE) from the FDA and has filed a premarket approval (PMA) application with the FDA with the expectation that the device will be available for use in the United States as early as the first quarter of CY 2010. The commenter noted that the CY 2010 proposed median cost of CPT code 0191T of approximately \$2,380, based on only three single claims, was much higher than the CY 2010 proposed median cost of APC 0234 of approximately \$1,639 and the CY 2010 proposed ASC payment of approximately \$962. The commenter explained that the relatively low number of Medicare hospital outpatient claims for CPT code 0191T resulted from the limited use of the procedure in IDE studies and its predominant performance in ASCs in association with cataract surgery. The commenter also noted that none of the other procedures assigned to APC 0234 involve the placement of an implantable device, while CPT code 0191T requires the insertion of a device that costs about \$2,500.

Response: CPT code 0191T is a new CPT code with very few Medicare claims from CY 2008, possibly because this procedure has been limited to IDE studies, as noted by the commenter. Furthermore, because this CPT code was effective on July 1, 2008, CY 2008

claims reflect only 6 months of hospital data, rather than a full year. We note that there are a number of other surgical eye procedures to treat glaucoma that are also assigned to APC 0234 for CY 2010. Moreover, the final CY 2010 median cost of CPT code 0191T based on a small number of CY 2008 claims is approximately \$1,962, close to the final CY 2010 median cost of APC 0234 of approximately \$1,630. Therefore, based on considerations of clinical and resource homogeneity, we continue to believe that APC 0234 is the most appropriate APC assignment for CPT code 0191T for CY 2010.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign CPT code 0191T to APC 0234, with a final CY 2010 APC median cost of approximately \$1,630.

b. Backbench Preparation of Corneal Allograft

For CY 2010, we proposed to continue to assign CPT code 65757 (Backbench preparation of corneal endothelial allograft prior to transplantation) status indicator "N" (Items and Services Packaged into APC Rates). In the CY 2009 OPPTS/ASC final rule with comment period (73 FR 69076), we assigned CPT code 65757 status indicator "N" and flagged the code with comment indicator "NI" to indicate that, as a new CPT code for CY 2009, its interim final CY 2009 OPPTS treatment was subject to comment on that final rule with comment period.

Comment: One commenter requested that CMS pay separately for CPT code 65757 under the OPPTS through an APC. According to the commenter, this service represents the preparation process for corneal transplants. The commenter argued that because this service is time-consuming and requires specialized skills and equipment, CPT code 65757 should not be packaged under the OPPTS but, instead, should be paid separately.

Response: We packaged CPT code 65757 because we consider it to be an intraoperative service that is ancillary and supportive to another service that is paid separately under the OPPTS, specifically the corneal transplant. Our general packaging policies for certain categories of services are discussed in section II.A.4. of this final rule with comment period. Although OPPTS payment for CPT code 65757 is packaged, we will consider its costs in setting the payment rates for the associated surgical procedures under the OPPTS, according to the standard OPPTS cost estimation methodology that

is discussed in section II.A.2. of this final rule with comment period.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign CPT code 65757 status indicator "N."

6. Orthopedic and Musculoskeletal Services

a. Arthroscopic Procedures (APCs 0041 and 0042)

For CY 2010, we proposed to continue the assignment of various arthroscopy procedures to APCs 0041 (Level I Arthroscopy) and APC 0042 (Level II Arthroscopy), with proposed payment rates of approximately \$2,014 and \$3,279, respectively.

Comment: One commenter expressed concern about the variety of procedures assigned to APC 0041, whose HCPCS code-specific median costs ranged from \$50 to \$22,000, and to APC 0042, whose HCPCS code-specific median costs ranged from \$143 to \$20,000. In particular, the commenter indicated that the current designation of only two APCs for the more than 60 distinct arthroscopic procedures assigned to these APCs does not appropriately reflect the unique clinical and resource characteristics associated with arthroscopic procedures that are provided to Medicare beneficiaries. The commenter urged CMS to create several new APCs to ensure clinical homogeneity and similar resource utilization for the arthroscopy procedures assigned to them and provided recommended APC configurations.

To pay appropriately for arthroscopic procedures under the OPPTS, the commenter recommended that CMS restructure the arthroscopy procedures into 11 new APCs based on the following three clinical categories: (1) Diagnostic arthroscopies; (2) lower extremity versus upper extremity arthroscopies; and (3) arthroscopies with implants. The commenter further recommended specific payment rates associated with each of the 11 recommended APCs, ranging from \$1,400 to \$5,400. According to the commenter, the recommended clinical distinctions parallel the distinctions CMS has created for other classes of procedures, including other orthopedic procedures, and would more accurately reflect the clinical characteristics and resource utilization of the services provided.

Alternatively, the commenter provided, in the event a reconfiguration of APCs 0041 and 0042 is not possible at this time, two more limited

suggestions: Finalize the proposal to reassign CPT codes 29888 (Arthroscopically aided anterior cruciate ligament repair/augmentation or reconstruction) and 29889 (Arthroscopically aided posterior cruciate ligament repair/augmentation or reconstruction) from APC 0042 to APC 0052 (Level IV Musculoskeletal Procedures Except Hand and Foot) and reassign CPT code 29892 (Arthroscopically aided repair of large osteochondritis dissecans lesion, talar dome fracture, or tibial plafond fracture, with or without internal fixation (includes arthroscopy)) from APC 0042 to APC 0052.

Response: We believe the existing clinical APCs 0041 and 0042 sufficiently account for the different clinical and resource characteristics of the procedures assigned to them. To reduce the size of the APC payment groups and establish new APC payment groups to pay more precisely would be inconsistent with our overall strategy to encourage hospitals to use resources more efficiently by increasing the size of the payment bundles. Moreover, many of the services that are assigned to APCs 0041 and 0042 are low volume services, with even fewer single claims available for ratesetting. Including low volume services in APCs with clinically similar higher volume services and similar median costs generates more stability in the payment rates that are set for these low volume services.

For APC 0041, based on significant services with a total claim frequency of greater than 1,000 or a frequency of greater than 99 and percentage of single claims equal to or greater than 2 percent, CY 2008 hospital outpatient claims data showed that the median cost of the lowest cost service is approximately \$1,463 and the median cost of the highest cost service is approximately \$2,086. Likewise, for APC 0042, claims data showed that the median cost of the lowest cost significant procedure is approximately \$2,730 and the median cost of the highest cost significant procedure is approximately \$4,592. Based on the CY 2008 claims data, there is no 2 times violation in either APC 0041 or APC 0042. Therefore, we see no reason for a reconfiguration into many more APCs in light of our interest in promoting hospital efficiency, as discussed earlier.

With respect to the reassignment of CPT code 29892 from APC 0042 to APC 0052 as recommended by the commenter, we agree that this reassignment would be appropriate for CY 2010. While we have very few claims for this procedure upon which to accurately estimate its cost, we

reviewed the clinical characteristics associated with CPT code 29892 and agree that, based on the complexity of this procedure, it would be more appropriately assigned to APC 0052 based on its clinical characteristics and expected resource utilization. Furthermore, we appreciate the commenter's support for our proposed reassignment of CPT codes 29888 and 29889 from APC 0042 to APC 0052 for CY 2010.

After consideration of the public comment we received, we are finalizing our CY 2010 proposals to reassign CPT codes 29888 and 29889 from APC 0042 to APC 0052, with a final CY 2010 APC median cost of approximately \$5,921. In addition, we are also finalizing the reassignment of CPT code 29892 from APC 0042 to APC 0052 for CY 2010. We are making no other changes to the proposed configurations of APC 0041 and 0042 for CY 2010. The final CY 2010 APC median cost for APC 0041 is approximately \$1,998 and approximately \$3,261 for APC 0042.

b. Knee Arthroscopy (APCs 0041 and 0042)

For CY 2010, we proposed to continue to assign CPT codes 29882 (Arthroscopy, knee, surgical; with meniscus repair (medial or lateral)) and 29883 (Arthroscopy, knee, surgical; with meniscus repair (medial and lateral)) to APC 0041 (Level I Arthroscopy), with a proposed payment rate of approximately \$2,014. In addition, we proposed to continue to assign CPT code 29867 (Arthroscopy, knee, surgical; osteochondral allograft (eg, mosaicplasty)) to APC 0042 (Level II Arthroscopy), with a proposed payment rate of approximately \$3,279.

Comment: One commenter recommended that CMS reassign CPT code 29882 and 29883 from APC 0041 to APC 0042 because of their similarity to procedures assigned to APC 0042. The commenter also requested that CMS reassign CPT code 29867 from APC 0042 to APC 0052 (Level IV Musculoskeletal Procedures Except Hand and Foot), with a proposed payment rate of approximately \$5,889. The commenter believed that CPT code 29867 is clinically comparable to the other procedures assigned to APC 0052.

Response: We reviewed the clinical and resource characteristics of CPT codes 29882 and 29883 and continue to believe these CPT codes are appropriately assigned to APC 0041 for CY 2010. Analysis of CY 2008 claims data showed that the median cost for CPT code 29882, based on 165 single claims (out of 334 total claims), is approximately \$2,224 and for CPT code

29883, based on 116 claims (out of 182 total claims), is approximately \$2,075. These median costs are consistent with the final CY 2010 median cost of APC 0041, which is approximately \$1,998. Furthermore, these procedures are clinically similar to the majority of other knee arthroscopy procedures that are also assigned to APC 0041.

In addition, we do not agree with the commenter's assertion that CPT code 29867 is similar to the other procedures in APC 0052. Our claims data show that CPT code 29867 has a median cost of approximately \$3,652, which is significantly lower than the median cost of approximately \$5,921 for APC 0052, but close to the median cost of approximately \$3,261 for APC 0042, where we proposed to assign the code for CY 2010. Furthermore, the knee arthroscopy procedure described by CPT code 29867 is not clinically similar to other procedures assigned to APC 0052, which are generally not performed arthroscopically.

After consideration of the public comment we received, we are finalizing our CY 2010 proposals, without modification, to continue to assign CPT codes 29882 and 29883 to APC 0041, which has a final CY 2010 APC median cost of approximately \$1,998, and to continue to assign CPT code 29867 to APC 0042, which has a final CY 2010 APC median cost of approximately \$3,261.

c. Shoulder Arthroscopy (APC 0042)

For CY 2010, we proposed to continue to assign CPT codes 29806 (Arthroscopy, shoulder, surgical; capsulorrhaphy) and 29807 (Arthroscopy, shoulder, surgical; repair of slap lesion) to APC 0042 (Level II Arthroscopy), with a proposed payment rate of approximately \$3,279.

Comment: One commenter recommended that CMS reassign CPT codes 29806 and 29807 to APC 0052 (Level IV Musculoskeletal Procedures Except Hand and Foot), which had a proposed payment rate of approximately \$5,889. The commenter believed that these procedures are clinically similar to the other procedures in APC 0052.

Response: We continue to believe that CPT codes 29806 and 29807 are appropriately assigned to APC 0042 based on clinical and resource considerations. We note that most other shoulder arthroscopy procedures that are similar to CPT codes 29806 and 29807 are assigned to APC 0042, while most procedures assigned to APC 0052 are bone procedures that are not performed arthroscopically. Analysis of our claims data revealed that the median cost of CPT code 29806, based

on 161 single claims (out of 759 total claims), is approximately \$4,003, which is significantly lower than the median cost of approximately \$5,921 for APC 0052. Likewise, our claims data showed that the median cost of CPT code 29807, based on 199 single claims (out of 3,802 total claims), is approximately \$4,202, which is also significantly lower than the median cost for APC 0052. The CPT code-specific median costs of these two procedure codes fall within the range of median costs (approximately \$2,730 to \$4,592) of significant procedures that are also assigned to APC 0042 for CY 2010. Therefore, we believe that CPT codes 29806 and 29807 are most similar clinically and with respect to resource costs to other procedures assigned to APC 0042.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign CPT codes 29806 and 29807 to APC 0042, which has a final CY 2010 APC median cost of approximately \$3,261.

d. Fasciotomy Procedures (APC 0049)

For CY 2010, we proposed to continue to assign the following seven CPT codes for fasciotomy procedures to APC 0049 (Level I Musculoskeletal Procedures Except Hand and Foot): CPT code 25020 (Decompression fasciotomy, forearm and/or wrist, flexor or extensor compartment; without debridement of nonviable muscle and/or nerve); CPT code 27496 (Decompression fasciotomy, thigh and/or knee, one compartment (flexor or extensor or adductor)); CPT code 27498 (Decompression fasciotomy, thigh and/or knee, multiple compartments); CPT code 27499 (Decompression fasciotomy, thigh and/or knee, multiple compartments; with debridement of nonviable muscle and/or nerve); CPT code 27892 (Decompression fasciotomy, leg; anterior and/or lateral compartments only, with debridement of nonviable muscle and/or nerve); CPT code 27893 (Decompression fasciotomy, leg; posterior compartment(s) only, with debridement of nonviable muscle and/or nerve); and CPT code 27894 (Decompression fasciotomy, leg; anterior and/or lateral, and posterior compartment(s), with debridement of nonviable muscle and/or nerve). The CY 2010 proposed payment rate for APC 0049 was approximately \$1,490.

Comment: One commenter recommended that CMS reassign CPT codes 25020, 27496, 27498, 27599, 27892, 27893, and 27894 from APC 0049 to APC 0050 (Level II Musculoskeletal Procedures Except Hand and Foot) based on their clinical

and resource similarity to the other fasciotomy procedures proposed for assignment to APC 0050. Although the commenter recommended assignment of CPT code 27599 (Unlisted procedure, femur or knee) among its list of codes for assignment to APC 0050, we believe that the commenter may have intended to reference CPT code 27499 instead. CPT code 27499 describes a decompression fasciotomy on the thigh and/or knee and was proposed for assignment to APC 0049. CPT code 27599 was proposed for assignment to APC 0129 (Level I Closed Treatment Fracture Finger/Toe/Trunk) and does not describe a fasciotomy procedure.

Response: We reviewed the clinical characteristics associated with each of the seven fasciotomy procedures, and based on our analysis, we agree with the commenter's recommendation. We note that, while we have no or very limited hospital claims data for these services that reflect hospital costs, a number of other similar fasciotomy procedures are already assigned to APC 0050. Based on further analysis, we believe that CPT codes 25020, 27496, 27498, 27499, 27892, 27893, and 27894 are sufficiently similar to those other fasciotomy procedures to warrant reassignment to APC 0050.

After consideration of the public comment we received, for CY 2010, we are reassigning CPT codes 25020, 27496, 27498, 27499, 27892, 27893, and 27894 from APC 0049 to APC 0050, which has a final CY 2010 APC median cost of approximately \$2,122.

e. Fibula Repair (APC 0062)

For CY 2010, we proposed to continue to assign CPT code 27726 (Repair of fibula nonunion and/or malunion with internal fixation) to APC 0062 (Level I Treatment Fracture/Dislocation), with a proposed payment rate of approximately \$1,735.

Comment: One commenter recommended that CMS reassign CPT code 27726 from APC 0062 to APC 0063 (Level II Treatment Fracture/Dislocation) because the procedure is comparable in clinical and resource characteristics to CPT code 27760 (Closed treatment of medial malleolus fracture; without manipulation), which was proposed for assignment to APC 0063, with a proposed payment rate of approximately \$3,023. In particular, the commenter argued that repair of a fibular nonunion is similar clinically and with respect to resource costs to repair of a tibial nonunion and, therefore, the two procedures should be assigned to the same clinical APC. Although the commenter compared CPT code 27726 to CPT code 27760, we

believe that the commenter may have intended to reference CPT code 27720 (Repair of nonunion or malunion, tibia; without graft, (eg, compression technique)), which describes a repair of a tibial nonunion and was proposed for assignment to APC 0063, instead of CPT code 27760. CPT code 27760 describes a closed treatment of an ankle fracture and was proposed for assignment to APC 0129 (Level I closed Treatment Fracture Finger/Toe/Trunk).

Response: We reviewed the clinical characteristics and resource use associated with CPT code 27726, and based on our analysis, we agree with the commenter's recommendation. For CY 2010, our claims data showed a median cost of approximately \$3,486 for CPT code 27726, based on 59 single claims (of 121 total claims), which is significantly higher than the median cost of approximately \$1,726 for APC 0062. Further, our claims data showed that the median cost of CPT code 27726 is similar to that of APC 0063, which has an APC median cost of approximately \$3,037. In addition, CPT code 27726 clinically resembles CPT code 27720, which is also assigned to APC 0063.

After consideration of the public comment we received, for CY 2010, we are modifying our CY 2010 proposal and reassigning CPT code 27726 to APC 0063 for CY 2010, which has a final CY 2010 APC median cost of approximately \$3,037.

f. Forearm Orthopedic Procedures (APCs 0050, 0051, and 0052)

For CY 2010, we proposed to assign the 14 forearm fracture procedures listed in Table 25 below to APC 0050 (Level II Musculoskeletal Procedures Except Hand and Foot), APC 0051, (Level III Musculoskeletal Procedures Except Hand and Foot), or APC 0052 (Level IV Musculoskeletal Procedures Except Hand and Foot). The CY 2010 proposed payment rate for APCs 0050 was approximately \$2,135; for APC 0051, approximately \$3,156; and for APC 0052, approximately \$5,889.

Comment: One commenter recommended that CMS reassign six forearm fracture procedures to APC 0051. In particular, the commenter stated that CPT codes 25350, 25355, 25360, 25370, 25390, and 25400 describe forearm surgical procedures involving only one bone and the hospital resource costs for the procedures are similar to those of procedures assigned to APC 0051. In addition, the commenter suggested that CMS reassign both CPT codes 24400 and 24410 to APC 0051 because these procedures are similar in clinical

characteristics and resource costs to other procedures in APC 0051. Further, the commenter recommended that CPT codes 25365, 25375, 25393, 25405, 25415, and 25420 be reassigned to APC 0052 based on considerations of clinical and resource homogeneity.

Response: We reviewed the clinical characteristics and resource costs associated with each surgical procedure discussed by the commenter. Based on our analysis of hospital claims data and clinical review, we agree with the commenter's recommendation that CPT codes 24400, 24410, 25350, 25355, 25360, 25370, 25390, and 25400 should be assigned to APC 0051. We have very few hospital outpatient claims for these procedures upon which to estimate their hospital costs. We note that these procedures are all performed on only one forearm bone, either the radius or the ulna, and we believe they share significant clinical and resource characteristics with other procedures assigned to APC 0051. Therefore, we are reassigning CPT codes 24400, 24410,

25350, 25360, and 25390 to APC 0051 for CY 2010. As we proposed, we are continuing to assign CPT codes 25355, 25370, and 25400 to APC 0051 for CY 2010.

With regard to the procedures that were recommended for reassignment to APC 0052, we agree with the commenter's argument that CPT codes 25405, 25415, and 25420 have similar resource costs to other procedures already assigned to APC 0052. These procedures were assigned to APC 0052 for CY 2009 and, as we proposed, for CY 2010, we are continuing their assignment to APC 0052.

However, we do not agree with the commenter's recommendation to reassign CPT codes 25365, 25375, and 25393 to APC 0052. We have very few claims for these procedures from CY 2008, but we believe their clinical and resource characteristics are sufficiently similar to other procedures assigned to APC 0051 that they should all be assigned to APC 0051 for CY 2010. While we proposed to assign CPT codes

25375 and 25393 to APC 0051 for CY 2010, we proposed to assign CPT code 25365 to APC 0050. In this final rule with comment period, we are modifying the assignment of CPT code 25365 to APC 0051, where it will reside along with CPT codes 25375 and 25393.

After consideration of the public comment we received, we are finalizing our CY 2010 proposals, without modification, to continue to assign CPT codes 25355, 25370, 25375, and 25393, and 25400 to APC 0051, which has a final CY 2010 APC median cost of approximately \$3,111, and CPT codes 25405, 25415, and 25420 to APC 0052, which has a final CY 2010 APC median cost of approximately \$5,921. We are modifying our CY 2010 proposals and assigning CPT codes 24400, 24410, 25350, 25360, 25365, and 25390 to APC 0051, which has a final CY 2010 APC median cost of approximately \$3,111. Table 25 below lists the final APC assignments for the 14 forearm fracture procedures discussed in this section.

TABLE 25—CY 2010 APC ASSIGNMENT FOR CERTAIN FOREARM FRACTURE PROCEDURES

CY 2010 HCPCS code	CY 2010 Long descriptor	Proposed CY 2010 APC	Final CY 2010 APC
24400	Osteotomy, humerus, with or without internal fixation	0050	0051
24410	Multiple osteotomies with realignment on intramedullary rod, humeral shaft (sofield type procedure).	0050	0051
25350	Osteotomy, radius; distal third	0052	0051
25355	Osteotomy, radius; middle or proximal third	0051	0051
25360	Osteotomy; ulna	0050	0051
25365	Osteotomy; radius and ulna	0050	0051
25370	Multiple osteotomies, with realignment on intramedullary rod (sofield type procedure); radius or ulna.	0051	0051
25375	Multiple osteotomies, with realignment on intramedullary rod (sofield type procedure); radius and ulna.	0051	0051
25390	Osteoplasty, radius or ulna; shortening	0050	0051
25393	Osteoplasty, radius and ulna; lengthening with autograft	0051	0051
25400	Repair of nonunion or malunion, radius or ulna; without graft (eg, compression technique).	0051	0051
25405	Repair of nonunion or malunion, radius or ulna; with autograft (includes obtaining graft).	0052	0052
25415	Repair of nonunion or malunion, radius and ulna; without graft (eg, compression technique).	0052	0052
25420	Repair of nonunion or malunion, radius and ulna; with autograft (includes obtaining graft).	0052	0052

g. Low Energy Extracorporeal Shock Wave Therapy (Low Energy ESWT)

For CY 2010, we proposed to continue to assign CPT code 0019T (Extracorporeal shock wave involving musculoskeletal system, not otherwise specified, low energy) status indicator "A" (Services furnished to a hospital outpatient that are paid under a fee schedule or payment system other than OPSS).

Comment: One commenter urged CMS to assign CPT code 0019T status indicator "T" (Significant Procedure:

Multiple Reduction Applies), and to place the CPT code in an APC that pays appropriately. The commenter indicated that high energy ESWT, specifically CPT code 0101T (Extracorporeal shock wave involving musculoskeletal system, not otherwise specified, high energy), is assigned to APC 0050 (Level II Musculoskeletal Procedures Except Hand and Foot), with a proposed CY 2010 payment rate of approximately \$2,135. The commenter argued that both the low energy and high energy ESWT treat similar conditions and both use

Class III medical devices that are subject to the most stringent FDA approval process that restricts the sale of the device to by or on the order of a physician. Because of this similarity, the commenter urged CMS to be consistent in its payment policy and recommended that both CPT codes 0101T and 0019T be assigned the same status indicator to specify their separate payment under the OPSS.

Response: We do not agree that low energy ESWT is similar to high energy ESWT. High energy ESWT requires the

use of anesthesia during the procedure and usually involves only one treatment session. Alternatively, low energy ESWT does not require anesthesia and usually is furnished over several sessions. Because of the complexity of high energy ESWT, we believe that it is appropriate to pay for CPT code 0101T as a hospital outpatient service under the OPSS through APC 0050. However, CPT code 0019T is assigned status indicator "A" because it is designated as a "sometimes therapy" service to indicate that it is a therapy service when furnished by a therapist. When performed in the HOPD, we believe CPT code 0019T would be furnished as a therapy service paid under the MPFS and, therefore, the service is appropriately assigned status indicator "A" for hospital outpatient payment purposes. Regulation of the device by the FDA as a Class III medical device for sale by or on the order of a physician and the need for special training to use the technology for its approved use are not inconsistent with our considering CPT code 0019T to be a "sometimes therapy" service, that is, a therapy service when furnished by a therapist.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign CPT code 0019T to status indicator "A" for CY 2010.

h. Insertion of Posterior Spinous Process Distraction Device (APC 0052)

For CY 2009 (73 FR 68620), we reassigned CPT codes 0171T (Insertion of posterior spinous process distraction device (including necessary removal of bone or ligament for insertion and imaging guidance), lumbar, single level) and 0172T (Insertion of posterior spinous process distraction device (including necessary removal of bone or ligament for insertion and imaging guidance), lumbar, each additional level) from APC 0050 (Level II Musculoskeletal Procedures Except Hand and Foot) to APC 0052 (Level IV Musculoskeletal Procedures Except Hand and Foot). For CY 2007 and CY 2008, the device implanted in procedures described by CPT codes 0171T and 0172T, HCPCS code C1821 (Interspinous process distraction device (implantable)), was assigned pass-through payment status and, therefore, was paid separately at charges adjusted to cost. The period of pass-through payment for HCPCS code C1821 expired after December 31, 2008. According to our established methodology, the costs of devices no longer eligible for pass-through payments are packaged into the costs of the procedures with which the

devices are reported in the claims data used to set the payment rates for those procedures. Therefore, the costs of the implanted device identified by HCPCS code C1821 are packaged into the costs of CPT codes 0171T and 0172T beginning in CY 2009.

At the February 2009 meeting, the APC Panel heard a public presentation that recommended reassignment of CPT codes 0171T and 0172T from APC 0052 to APC 0425 (Level II Arthroplasty or Implantation with Prosthesis). The presenter believed that APC resource homogeneity would be improved if CPT codes 0171T and 0172T were reassigned to APC 0425. The presenter asserted, based on its analysis of CY 2007 claims data, that the median cost of CPT code 0171T was significantly higher than the median cost of APC 0052, while only slightly lower than the median cost of APC 0425. The presenter indicated that, while the median cost of APC 0052 was significantly higher than the median cost of device HCPCS code C1821, device costs are only one element of the overall procedure cost and other associated procedure costs are more than \$3,200. Regarding clinical homogeneity, the presenter stated that kyphoplasty is the only spine procedure currently assigned to APC 0052 other than CPT codes 0171T and 0172T. The presenter also claimed that 36 percent of claims for CPT code 0171T are reported without HCPCS code C1821, which identified a device that is always implanted in procedures reported with CPT codes 0171T and 0172T. The presenter requested reassignment of CPT codes 0171T and 0172T to APC 0425 because this APC is a device-dependent APC, and CPT codes 0171T and 0172T would then be subject to procedure-to-device claims processing edits.

The APC Panel recommended that CMS continue the assignment of CPT codes 0171T and 0172T to APC 0052 for CY 2010, institute procedure-to-device claims processing edits for HCPCS code C1821, and then reevaluate the APC assignments of CPT codes 0171T and 0172T in one year.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35305), we stated that under our existing policy, we generally do not identify any individual HCPCS codes as device-dependent codes under the OPSS. We create device edits, when appropriate, for procedures assigned to device-dependent APCs, where those APCs have been historically identified under the OPSS as having very high device costs. We noted in the CY 2009 OPSS/ASC final rule with comment period regarding APC 0052 (73 FR 68621) that we typically do not

implement procedure-to-device edits for an APC where there are not device HCPCS codes for all possible devices that could be used to perform a procedure that always requires a device, and the APC is not designated as a device-dependent APC. APC 0052 is not a device-dependent APC because a number of the procedures assigned to the APC do not require the use of implantable devices. Furthermore, in some cases, there may not be HCPCS codes that describe all devices that may be used to perform the procedures in APC 0052.

We examined the CY 2008 claims data available for the CY 2010 proposed rule to determine the frequency of billing CPT code 0171T (which is the main procedure code reported with HCPCS code C1821) with and without device HCPCS code C1821. CPT code 0172T is an add-on code to CPT code 0171T. We recognize that our single claims for CPT code 0172T may not be correctly coded claims and, therefore, our cost estimation for CPT code 0172T may not be accurate. Our analysis showed that the CY 2010 proposed rule median cost for CPT code 0171T was approximately \$7,717 based on over 800 single claims. The CY 2010 proposed rule claims data for CPT code 0171T revealed a median cost of approximately \$7,916 based on over 500 single claims with HCPCS code C1821, and a median cost of approximately \$7,387 based on approximately 300 single claims without HCPCS code C1821. Therefore, we concluded that the median cost of claims for CPT code 0171T reported with HCPCS code C1821 is similar to the median cost of claims for the procedure reported without HCPCS code C1821. We stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35305) that we have no reason to believe that those hospitals not reporting the device HCPCS code had failed to consider the cost of the device in charging for the procedure. Furthermore, claims for CPT code 0171T reported with HCPCS code C1821 accounted for about two-thirds of the single claims available for ratesetting. For the CY 2010 OPSS/ASC proposed rule, we concluded that the overall median cost of CPT code 0171T fell within an appropriate range of HCPCS code-specific median costs for those services proposed for CY 2010 assignment to APC 0052, which had a proposed APC median cost of approximately \$5,939 and no 2 times violation. Moreover, in the CY 2010 OPSS/ASC proposed rule (74 FR 35305), we indicated that we do not believe that procedure-to-device claims processing

edits are necessary to ensure accurate cost estimation for CPT code 0171T.

The CY 2010 OPPS/ASC proposed rule line-item median cost for HCPCS code C1821 was approximately \$4,625, while the CY 2010 OPPS/ASC proposed rule median cost of APC 0052 was approximately \$1,300 more than this device cost. We stated in the proposed rule (74 FR 35305) that previous estimates of procedure time presented to us at the time of the device pass-through application for the interspinous process distraction device described by HCPCS code C1821 were approximately 30 to 60 minutes of procedure time for the service currently described by CPT code 0171T. This is reasonably comparable to the typical procedure time for kyphoplasty described by CPT code 22523 (Percutaneous vertebral augmentation, including cavity creation (fracture reduction and bone biopsy included when performed) using mechanical device, one vertebral body, unilateral or bilateral cannulation (eg, kyphoplasty); thoracic) and CPT code 22524 (Percutaneous vertebral augmentation, including cavity creation (fracture reduction and bone biopsy included when performed) using mechanical device, one vertebral body, unilateral or bilateral cannulation (eg, kyphoplasty); lumbar), which are also assigned to APC 0052.

Because we reasoned that APC 0052 pays appropriately for the procedure cost of CPT codes 0171T and 0172T, we proposed to maintain the assignment of CPT codes 0171T and 0172T to APC 0052 for CY 2010 and not to implement device edits for these procedures. We proposed to accept one part of the APC Panel's recommendation regarding the continued assignment of CPT codes 0171T and 0172T to APC 0052, but we proposed to not accept the APC Panel's further recommendation to institute procedure-to-device edits for these services for CY 2010. As we do for all OPPS services, we stated that we would reevaluate the APC assignments of CPT codes 0171T and 0172T when additional claims data become available for CY 2011 ratesetting, in accordance with the final part of the APC Panel's recommendation for these procedures (74 FR 35305).

Comment: Some commenters recommended that CMS reassign CPT codes 0171T and 0172T from APC 0052 to APC 0425 for CY 2010, arguing that the resource costs associated with these procedures are more similar to the resource costs of procedures assigned to APC 0425 than to procedures assigned to APC 0052. One commenter noted, for example, that the median cost for CPT code 0171T is approximately 30 percent

higher than the median cost for APC 0052, but only two percent lower than the median cost for APC 0425. In response to CMS' observation in the CY 2010 OPPS/ASC proposed rule that the proposed median cost of APC 0052 was approximately \$1,300 more than the line-item median cost for HCPCS code C1821 of approximately \$4,625, the commenter pointed out that device costs are but one element of the overall procedure costs. The commenter presented data to demonstrate that the service costs associated with CPT code 0171T are greater than this \$1,300 difference. The commenter agreed that the 30 to 60 minute procedure time associated with CPT code 0171T that CMS noted in the proposed rule is reasonable, but argued that intraservice time should not be used as a sole basis for judging resource homogeneity because there is not a direct correlation between intraservice time and hospital costs.

The commenters also disagreed with CMS' assertion that the procedures described by CPT codes 0171T and 0172T are more similar clinically to procedures assigned to APC 0052 than to procedures assigned to APC 0425. One commenter argued that kyphoplasty is the only spine procedure assigned to APC 0052, and that, like all of the other procedures assigned to APC 0052, it does not involve the implantation of a device. The commenter acknowledged that, while CMS' statement of clinical similarity for APC 0052 is true to some extent, the procedure described by CPT code 0171T is more similar to procedures assigned to APC 0425 because it is orthopedic in nature and requires the use of a device that is classified as a prosthesis by the FDA.

Moreover, the commenter claimed that there are relevant precedents for reassignment of CPT codes 0171T and 0172T to APC 0425, such as CMS' proposed reassignment of CPT code 27446 (Arthroplasty, knee, condyle and plateau; medial OR lateral compartment) to APC 0425 for CY 2010. The commenter also argued that reassigning CPT 0171T and 0172T to device-dependent APC 0425, to which procedure-to-device edits apply, would help ensure that only correctly coded claims are used in ratesetting.

Response: We continue to believe that APC 0052 is an appropriate APC assignment for CPT codes 0171T and 0172T based on consideration of the procedures' clinical and resource characteristics. We do not agree with the commenters that the resource costs of providing the procedures described by CPT codes 0171T and 0172T are

substantially different from the resource costs of providing other procedures assigned to APC 0052 and that they should not be assigned to APC 0052, which has a final CY 2010 APC median cost of approximately \$5,921. Based on the CY 2008 claims data reviewed for this final rule with comment period, the final median costs of CPT codes 0171T and 0172T are approximately \$7,522 (based on 939 single claims) and approximately \$14,617 (based on 6 single claims), respectively. As we have noted previously (73 FR 68620), we recognize that our single claims for CPT code 0172T may not be correctly coded and, therefore, our cost estimation for CPT code 0172T may not be accurate. CPT code 0171T has the highest median cost of the significant procedures (defined as those procedures with a frequency of greater than 1,000 single claims or a frequency of greater than 99 and more than 2 percent of the single claims in the APC) assigned to APC 0052, while the lowest cost significant procedure has a median cost of approximately \$5,072. Therefore, the configuration of APC 0052 does not violate the 2 times rule. We continue to believe that, based on resource considerations, assignment to APC 0052 would provide appropriate payment for CPT codes 0171T and 0172T. We agree with the commenters that we should consider factors such as line-item median costs for devices and intraservice times as two data elements among several when we evaluate the clinical and resource homogeneity of APCs. In this case, we continue to believe that, as described in the proposed rule, both the line-item median cost for HCPCS code C1821 and the intraservice time for the procedure described by CPT code 0171T support our assessment that this procedure is similar in terms of resource utilization to other procedures assigned to APC 0052, consistent with the fact there is no 2 times violation within this APC.

We continue to believe the posterior spinous process distraction device procedures described by CPT codes 0171T and 0172T are clinically similar to other procedures, such as the kyphoplasty procedures, that are assigned to APC 0052. We disagree with the commenter that the kyphoplasty procedures described by CPT codes 22523 and 22524 do not involve the implantation of a device. Our definition of an implantable device includes surgically inserted or implanted devices that may not remain with the patient following the procedure, and thus may include expensive devices used in kyphoplasty such as expanders and

other single-use disposal devices used to create a cavity in the vertebral body. We note the code descriptor of kyphoplasty CPT code 22523 states, "using mechanical device." Based on a kyphoplasty New Technology APC application we received in CY 2004, the prices for these implantable devices are approximately \$3,000. Moreover, the kyphoplasty procedures are clinical substitutes for vertebroplasty procedures, such as the procedure described by CPT code 22520 (Percutaneous vertebroplasty, one vertebral body, unilateral or bilateral injection; thoracic) and are assigned to APC 0050 (Level II Musculoskeletal Procedures Except Hand and Foot). CPT code 22520 has a CY 2010 final rule median cost of approximately \$2,181, which is nearly \$3,800 less than the final rule median cost of approximately \$5,976 calculated for the kyphoplasty procedure described by CPT code 22523. This differential appears to be largely attributable to implantable device costs in kyphoplasty procedures. Therefore, we continue to believe that kyphoplasty and posterior spinous process distraction device procedures are clinically similar in that they are spinal procedures involving implantable devices. We note that there are no procedures involving the spine assigned to APC 0425.

We do not agree with the commenter that our reassignment of the knee arthroplasty procedure described by CPT code 27446 to APC 0425 serves as a precedent for the reassignment of CPT codes 0171T and 0172T to APC 0425. As discussed in section II.A.2.d.(1) of this final rule with comment period, we reassigned CPT code 27446 from APC 0681 (Knee Arthroplasty) to APC 0425 for CY 2010 in order to consolidate APC 0425 with APC 0681, in which CPT code 27446 was the only code. As noted in section II.A.2.d.(1) of this final rule with comment period, over the past several years, the median cost for CPT code 27446 has fluctuated due to a low volume of services being performed by a small number of providers in the HOPD, and to a single provider performing the majority of services. We believe that by reassigning CPT code 27446 to APC 0425 and deleting APC 0681, we can maintain greater stability from year to year in the payment rate for CPT code 27446. Therefore, we do not believe this is a similar situation to that of CPT codes 0171T and 0172T, as the commenter argued. Furthermore, we do not believe that implantation of an interspinous process distraction device, a minimally invasive procedure, is clinically comparable to a knee

replacement procedure that is performed on the majority of Medicare beneficiaries on a hospital inpatient basis. We also do not agree that we should reassign CPT codes 0171T and 0172T to APC 0425 in order to implement device edits for these procedures. As we described in the proposed rule (74 FR 35305), based upon analysis of CY 2010 proposed rule claims data for CPT code 0171T, we have no reason to believe that the minority of hospitals that do not bill HCPCS code C1821 along with CPT code 0171T are not already considering the costs of the interspinous process distraction device in charging for the procedure.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to continue to assign CPT codes 0171T and 0172T to APC 0052, which has a final CY 2010 APC median cost of approximately \$5,921.

7. Radiation Therapy Services

a. Proton Beam Therapy (APCs 0664 and 0667)

For CY 2010, we proposed to continue to assign CPT codes 77520 (Proton treatment delivery; simple, without compensation) and 77522 (Proton treatment delivery; simple, with compensation) to APC 0664 (Level I Proton Beam Radiation Therapy), which had a proposed payment rate of approximately \$713. We also proposed to continue to assign CPT codes 77523 (Proton treatment delivery; intermediate) and 77525 (Proton treatment delivery; complex) to APC 0667 (Level II Proton Beam Radiation Therapy), which had a proposed payment rate of approximately \$933.

Comment: Several commenters supported the proposed payment increases for the proton beam treatment CPT codes. The commenters cited a payment increase of 1.43 percent for CPT codes 77520 and 77522, and a payment increase of 11.02 percent for CPT codes 77523 and 77525.

Response: We appreciate the commenters' support for our proposals. The CY 2010 OPPS payment rates for CPT codes 77520, 77522, 77523, and 77525 are based on the APC median costs calculated from CY 2008 hospital claims data and the most current cost reports, according to the standard OPPS ratesetting methodology. We are confident that the observed costs in the claims data are representative of the costs of the proton beam therapy services provided in CY 2008 because

almost all of the claims are single claims that can be used for ratesetting.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to assign CPT codes 77520 and 77522 to APC 0664, with a final CY 2010 APC median cost of approximately \$934, and CPT codes 77523 and 77525 to APC 0667, with a final CY 2010 APC median cost of approximately \$1,221.

b. Stereotactic Radiosurgery (SRS) Treatment Delivery Services (APCs 0065, 0066, 0067, and 0127)

For CY 2010, we proposed to continue to assign CPT code 77371 (Radiation treatment delivery, stereotactic radiosurgery (SRS), complete course of treatment of cranial lesion(s) consisting of 1 session; multi-source Cobalt 60 based) to APC 0127 (Level IV Stereotactic Radiosurgery, MRgFUS, and MEG), with a proposed payment rate of approximately \$7,714.

We also proposed to continue to recognize for separate payment in CY 2010 four existing HCPCS G-codes that describe linear accelerator-based SRS treatment delivery services. Specifically, we proposed the following: to assign HCPCS code G0173 (Linear accelerator based stereotactic radiosurgery, complete course of therapy in one session) to APC 0067 (Level III Stereotactic Radiosurgery, MRgFUS, and MEG), with a proposed payment rate of approximately \$3,507; to assign HCPCS code G0251 (Linear accelerator-based stereotactic radiosurgery, delivery including collimator changes and custom plugging, fractionated treatment, all lesions, per session, maximum five sessions per course of treatment) to APC 0065 (Level I Stereotactic Radiosurgery, MRgFUS, and MEG), with a proposed payment rate of approximately \$894; to assign HCPCS code G0339 (Image-guided robotic linear accelerator-based stereotactic radiosurgery, complete course of therapy in one session or first session of fractionated treatment) to APC 0067, with a proposed payment rate of approximately \$3,507; and to assign HCPCS code G0340 (Image-guided robotic linear accelerator-based stereotactic radiosurgery, delivery including collimator changes and custom plugging, fractionated treatment, all lesions, per session, second through fifth sessions, maximum five sessions per course of treatment) to APC 0066 (Level II Stereotactic Radiosurgery, MRgFUS, and MEG), with a proposed payment rate of approximately \$2,505.

Further, we proposed to continue to assign CPT codes 77372 (Radiation treatment delivery, stereotactic

radiosurgery (SRS) (complete course of treatment of cerebral lesion(s) consisting of 1 session); linear accelerator based) and 77373 (Stereotactic body radiation therapy, treatment delivery, per fraction to 1 or more lesions, including image guidance, entire course not to exceed 5 fractions) status indicator "B" (Codes that are not recognized by OPPS when submitted on an outpatient hospital Part B bill type (12x and 13x)) under the OPPS, to indicate that these CPT codes are not payable under the OPPS.

Comment: Several commenters expressed concern about their belief that payment for HCPCS code G0173 and CPT code 77371 is based on the utilization of specific SRS equipment. The commenters stated that no clinical data exist to support the need for differential payment for linear accelerator-based and Cobalt-60 SRS procedures. The commenters further explained that current medical literature cites no difference in clinical effectiveness for the systems associated with linear accelerator-based and Cobalt-60 SRS procedures. One commenter provided an extensive bibliography of relevant peer-reviewed articles supporting this finding. The commenters recommended that CMS assign HCPCS code G0173 and CPT code 77371 to the same APC so that payment for both services would be the same. Specifically, the commenters suggested capping the payment rate for CPT code 77371 at the payment rate for HCPCS code G0173. One commenter added that, based on an internal analysis of CY 2007 claims data using the CY 2009 OPPS payment rates for CPT code 77371 and HCPCS code G0173, paying both procedures at the payment rate for HCPCS code G0173 would lead to Medicare savings of at least \$272 million over 10 years and about \$104 million over 5 years. The commenters encouraged CMS to consider this payment methodology and, thereby, pay for services appropriately regardless of the specific equipment used to deliver SRS treatment, especially as Medicare moves towards a value-based purchasing system.

Response: Analysis of our claims data shows that the median costs for linear accelerator-based and Cobalt-60 SRS procedures vary significantly. Since the creation of CPT code 77371, which was made effective January 1, 2007, our claims data has shown a median cost of more than approximately \$7,000 for this procedure. Based on data available for

CY 2010 ratesetting, our claims data showed a median cost of approximately \$7,277 for CPT code 77371 that is derived from 483 single claims (of 4,142 total claims), which is significantly higher than the median cost of approximately \$2,877 for HCPCS code G0173 that is based on 459 single claims (of 1,471 total claims). Likewise, for claims submitted for CY 2007, the data year used for CY 2009 ratesetting, our claims data showed a median cost of approximately \$7,470 based on 518 single claims (of 4,208 total claims) for CPT code 77371, which is much higher than the median cost of approximately \$3,523 for HCPCS code G0173, based on 528 single claims (of 1,616 total claims).

The OPPS is a prospective payment system, where APC payment rates are based on the relative costs of services as reported to us by hospitals according to the most recent claims and cost report data as described in section II.A. of this final rule with comment period. The 2 times rule specifies that the median cost of the highest cost item or service within a payment group may be no more than 2 times greater than the median cost of the lowest cost item or service within the same group. Based on application of the 2 times rule, we cannot assign HCPCS code G0173 and CPT code 77371 to the same APC. In addition, because hospitals continue to report very different costs for these services, we believe it is appropriate to maintain the assignment of these two codes to different payment groups for CY 2010. As a matter of payment policy, the OPPS does not set payment rates for services based on considerations of clinical effectiveness. Furthermore, in accordance with the statute, we budget neutralize payments under the OPPS each year in the annual update so that projected changes in spending for certain services are redistributed to payment for other services.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to continue to assign CPT code 77371 to APC 0127, which has a final CY 2010 APC median cost of approximately \$7,277, and to continue to assign HCPCS code G0173 to APC 0067, which has a final CY 2010 APC median cost of approximately \$3,539.

Comment: One commenter requested that CMS finalize the proposed APC and status indicator assignments for HCPCS codes G0173, G0251, G0339, G0340, 77372, and 77373 for CY 2010. The

commenter also recommended that CMS revise code descriptors of HCPCS code G0173, G0251, G0339, and G0340 for SRS, to distinguish between non-gantry and gantry-based SRS systems. Based on internal analysis, the commenter stated that, within the past year, there has been an increase in the OPPS volume of incorrectly coded claims. The commenter suggested specific code descriptor changes for the four revised HCPCS G-codes, as well as specific language changes to the SRS billing instructions in Chapter 4 of the Medicare Claims Processing Manual.

Response: These HCPCS G-codes for SRS have been in effect for several years and, based on questions brought to our attention by hospitals, we have no reason to believe that hospitals are confused about the reporting of these codes. Further, we see resource differences reflected in the median costs of the four HCPCS G-codes that are reasonably consistent with our expectations for different median costs for the services based on the current code descriptors. We believe it would be confusing to hospitals if we were to revise the code descriptors for HCPCS codes G0173, G0251, G0339, and G0340. Moreover, such a change could lead to instability in our median costs and inaccurate payments for some services. Therefore, we believe that modifying the G-code descriptors is not necessary for us to continue to provide appropriate payment for the services they describe. We also do not believe changes to our current billing instructions for SRS services in the Medicare Claims Processing Manual are necessary.

After consideration of the public comment we received, we are finalizing our CY 2010 proposals, without modification, to maintain the existing code descriptors for HCPCS codes G0173, G0251, G0339, and G0340 for linear accelerator-based SRS. In addition, we are finalizing our proposals, without modification, to continue to assign CPT codes 77372 and 77373 to status indicator "B" under the OPPS, and to continue to assign the four linear accelerator-based SRS HCPCS codes to the same APCs for CY 2010 as CY 2009, specifically APCs 0065, 0066, and 0067, with final CY 2010 APC median costs of approximately \$954, \$2,465, and \$3,539, respectively. Table 26 displays the final APC median costs for the SRS treatment delivery HCPCS codes and CPT code 77371.

TABLE 26—FINAL CY 2010 APC ASSIGNMENTS FOR ALL SRS TREATMENT DELIVERY SERVICES

CY 2010 HCPCS code	CY 2010 Short descriptor	Final CY 2010 SI	Final CY 2010 APC	Final CY 2010 approximate APC median cost
G0173	Linear acc stereo radsur com	S	0067	\$3,539
G0251	Linear acc based stero radio	S	0065	954
G0339	Robot lin-radsurg com, first	S	0067	3,539
G0340	Robt lin-radsurg fractx 2-5	S	0066	2,465
77371	SRS, multisource	S	0127	7,277

c. Clinical Brachytherapy (APCs 0312 and 0651)

For CY 2010, we did not propose any change to the HCPCS codes for assignment to APC 0312 (Radioelement Applications) or APC 0651 (Complex Interstitial Radiation Source Application). The proposed CY 2010 payment rates for these APCs were approximately \$298 and \$808, respectively.

Comment: Several commenters objected to the proposed reduction in the payment rate for brachytherapy services assigned to APC 0312 from approximately \$421 in CY 2009 to approximately \$298 in CY 2010, and the proposed reduction in the payment rate for APC 0651 from approximately \$847 in CY 2009 to approximately \$808 in CY

2010. The commenters believed these reductions in payment rates are the result of reduced numbers of single claims for the services assigned to the APCs, caused by the trimming of lines for which no payment was made on the claim. They objected to the use of only 2 percent of total claims or a 30 percent reduction in single claims for CPT code 77778 (Interstitial radiation source application; complex) that is assigned to APC 0651, and to the use of only 9 percent of total claims for CPT code 77776 (Interstitial radiation source application; simple) and 19 percent of total claims for CPT code 77777 (Interstitial radiation source application; intermediate) that are both assigned to APC 0312. The commenters speculated that the problem could be associated

with changes to the bypass list, trimming of unpaid lines, or other general problems with CMS' cost estimation methodology. They believed that, regardless of the source of the problem, CMS must establish appropriate and stable payment rates for these services to allow Medicare beneficiaries consistent access to brachytherapy procedures.

Response: The median cost for APC 0312 for CY 2010, calculated using final rule data, is approximately \$300. Our review of final rule claims data indicates that the reduction in median cost for APC 0312 from CY 2009 to CY 2010 appears to be caused by changes in the median costs for the HCPCS codes assigned to the APC that drive the median cost for the APC.

TABLE 27—MEDIAN COST AND FREQUENCY DATA FOR SERVICES ASSIGNED TO APC 0312* IN CY 2009 AND CY 2010

HCPCS code in APC 0312	Short descriptor	CY 2009 approximate median cost	CY 2009 frequency of single claims	CY 2009 percentages of single claims	CY 2009 total claims	CY 2010 approximate median cost	CY 2010 single claims	CY 2010 percentages of single claims	CY 2010 total claims
77776	Apply interstit radiat simpl.	\$119	23	6	149	\$522	9	4	104
77762	Apply intrcav radiat interm.	180	70	18	161	345	25	11	69
77763	Apply intrcav radiat compl.	507	131	34	352	345	112	48	250
77777	Apply interstit radiat inter.	608	7	2	51	300	11	5	54
77761	Apply intrcav radiat simple.	681	158	41	247	85	78	33	124
Totals			389	41	960		235	39	601

* Data exclude claims for CPT code 77799, which were not used in setting the APC median cost.

Specifically, in CY 2009, CPT codes 77761 and 77763 dominated APC 0312 and the APC median cost was approximately \$420. For CY 2010, CPT codes 77761 and 77763 continue to dominate APC 0312 but their HCPCS-specific median costs declined. Hence, the median cost for APC 0312 decreased to approximately \$300. We do not believe that the exclusion of the lines for which no payment was made was the controlling factor in the decline of the APC median cost. We excluded 97

lines (including one unlisted line that is not relevant) from the claims containing CPT codes assigned to APC 0312 before we split the claims into single claims. Therefore, it is not possible to know how many of the line-items we trimmed were on claims that might have become single claims that could be used for ratesetting purposes. The total frequency of HCPCS codes reported on claims used for CY 2010 ratesetting declined to 601 from 960 (before the line-item trim). Therefore, a reduction

in the number of single claims that are available for calculation of the median cost for the APC is to be expected because the universe of claims assigned to APC 0312 declined by more than one third. However, we note that the single claims used in the APC median calculation, as a percent of the total frequency, was 41 percent in CY 2009 and declined only minimally to 39 percent in CY 2010, notwithstanding the decrease in total frequency from CY 2009 to CY 2010 and the trim of 96

relevant lines of the 601 total claims for the codes used to set the APC median cost for APC 0312. We agree that the decline in the median costs for CPT codes 77761 and 77763 is notable. However, we know that, for CY 2007 (the year of the claims used for the CY 2009 OPPS), there were no CPT codes for the insertion of the needles and catheters used to apply brachytherapy sources interstitially to body areas other than the prostate. We believe it is possible that the costs of the needles and catheters may have been incorporated into the CY 2009 payment for some of the CPT codes assigned to APC 0312.

For CY 2008, the AMA's CPT Editorial Panel created CPT code 20555 (Placement of needles or catheters into muscle and/or soft tissue for subsequent interstitial radioelement application (at the time of or subsequent to the procedure)), and payment has been made for that CPT code through APC 0050 (Level II Musculoskeletal Procedures Except Hand and Foot) in CY 2008 and CY 2009. In the updated claims data used for this CY 2010 final rule with comment period, for services furnished in CY 2008, CPT code 20555 has a total frequency of 67 and a single claim frequency of 25. CPT code 20555 is assigned to APC 0050, which has a final CY 2010 median cost of approximately \$2,122. Because the needles and catheters must be placed before services reported by certain CPT codes assigned to APC 0312 can be performed, the hospital would receive not only the payment for APC 0312, but would also be paid for the placement of the needles and catheters or other devices, whether reported under CPT code 20555 or another code for placement of needles and catheters or other brachytherapy source delivery devices. Therefore, although the payment rate for APC 0312 has declined between CY 2009 and CY 2010, hospitals will commonly receive a separate payment for the placement of the needles and catheters or other devices that, when added to the payment for the application of the sources, will provide a robust payment for the service in its entirety.

The final CY 2010 median cost of APC 0651 is approximately \$885, compared to the median cost of approximately \$847 for CY 2009. We note that most claims that report CPT code 77778 are for low dose rate prostate brachytherapy that is paid through APC 8001 (LDR Prostate Brachytherapy Composite) rather than through APC 0651. Therefore, the total claim frequency for APC 0651 of 9,649 includes both the 7,742 claims that meet the criteria for

payment through APC 8001 and the 1,907 claims that meet the criteria for payment through APC 0651. For this final rule with comment period, we were able to use approximately 11 percent of the claims (206 of 1,907 total claims) that meet the criteria for payment through APC 0651 in the calculation of the median cost for APC 0651. Not only does the CY 2010 median cost for APC 0651 increase over the CY 2009 median cost, but when the separate payment for the placement of brachytherapy insertion devices is made, the full payment for the comprehensive service is substantial. For example, if CPT code 20555 was reported for placement of needles and catheters, the hospital would be paid for both one unit of APC 0651 (based on a CY 2010 median cost of approximately \$885) and one unit of APC 0050 (based on a CY 2010 median cost of approximately \$2,122).

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to calculate the median costs for APCs 0312 and 0651 according to the standard OPPS ratesetting methodology, applying the final bypass list and line-item trim as discussed in sections II.A.1. and II.A.2. of this final rule with comment period. The final CY 2010 median costs of APCs 0312 and 0651 are approximately \$300 and \$885, respectively. We believe that when hospitals fully report the services required for brachytherapy treatment, the combined OPPS payment for insertion of the source application devices and application of the sources themselves provides appropriate payment for the comprehensive service.

8. Other Services

a. Low Frequency, Non-Contact, Non-Thermal Ultrasound (APC 0013)

The CPT Editorial Panel created CPT code 0183T (Low frequency, non-contact, non-thermal ultrasound, including topical application(s), when performed, wound assessment, and instruction(s) for ongoing care, per day), effective January 1, 2008. Under the OPPS, we assigned CPT code 0183T to APC 0015 (Level III Debridement & Destruction) for CY 2008 and CY 2009. For CY 2009, APC 0015 has a payment rate of approximately \$100. Based upon our review of the first year of hospital claims data for CPT 0183T, for CY 2010 we proposed to reassign CPT code 0183T to APC 0013 (Level II Debridement & Destruction), with a proposed payment rate of approximately \$59.

Comment: Several commenters recommended that CMS continue to assign CPT code 0183T to APC 0015. The commenters asserted that the proposed payment for APC 0013 would not cover hospitals' costs for performing the procedure. One commenter stated that the single-use kit for the service costs \$55. Another commenter reported that the majority of hospitals with the highest utilization of CPT code 0183T either failed to report or underreported the packaged supply costs associated with CPT code 0183T. The commenter analyzed CMS' claims data according to hospitals' reporting of "packaged" supplies with CPT code 0183T and found that 52 percent of all single claims were from 5 hospitals, and that 4 of these 5 hospitals, representing 39 percent of single claims for CPT code 0183T used in ratesetting, reported \$0 or an insignificant (less than \$5) packaged supply cost. Moreover, the commenter stated that the analysis indicated that, overall, only one-third of the single claims for CPT code 0183T included any packaged costs, although costly supplies are required for hospitals to furnish the service. In addition, the commenter reported that it surveyed hospitals that provided the service and learned that those hospitals reported a median procedure cost of approximately \$153.

One commenter offered several reasons why hospitals might not report packaged supply costs with CPT code 0183T, including the fact that CPT code 0183T was a new CPT code in CY 2008, the year of claims data for the CY 2010 OPPS rates; hospitals' historical failure to consider supply costs in setting their procedure charges; the fact that relatively low cost supplies are often overlooked when hospitals charge for services; and the lack of a specific Level II HCPCS code to report a charge for the applicator kit. The commenter estimated that 32,000 procedures were furnished to Medicare beneficiaries in the HOPD in CY 2008, yet there were far fewer CY 2008 OPPS claims for the service. The commenter cited several examples of contractors providing instructions to report other CPT codes, such as CPT code 97602 (Removal of devitalized tissue from wound(s), non-selective debridement, without anesthesia (eg wet-to-moist dressings, enzymatic, abrasion), including topical application(s), wound assessment, and instruction(s) for ongoing care, per session), when providing the low frequency, non-contact, non-thermal ultrasound procedure. Another commenter argued that APC 0015 is the most clinically appropriate APC for CPT

code 0183T, and stated that if the service were reassigned to APC 0013 as proposed, it would be the only wound healing procedure and the only procedure requiring a single use disposable supply in APC 0013.

Response: We proposed to reassign CPT code 0183T to APC 0013 for CY 2010 based on clinical and resource considerations. The final CY 2010 median cost of CPT code 0183T is approximately \$77, based on 9,335 single claims. The final CY 2010 median cost of APC 0013 is approximately \$59, and the final CY 2010 final cost of APC 0015 is approximately \$103. The final CY 2010 HCPCS code-specific median costs of other significant services assigned to APC 0013 range from approximately \$46 to \$82; therefore, the \$77 final median cost of CPT code 0183T for CY 2010 is well within that range. While CY 2008 is the first year we have cost information from hospitals for the service, the large number of single claims provides a robust estimate of the service's cost based on claims from those hospitals that furnished the service in CY 2008. While the commenters were concerned that many claims did not include separate charges for the associated supplies, we have found that it is common for hospitals to consider the cost of necessary supplies when setting the procedure charge, rather than reporting a separate line-item charge for the associated supplies. Many supplies where payment is always packaged into procedure payments do not have specific Level II HCPCS codes under which to report the associated charges. Hospitals incorporate the charge for such supplies in the procedure charge or provide a charge on a separate line under an appropriate revenue code without a HCPCS code, and we package the costs from these uncoded line-items into payment for the associated procedure. Therefore, we have no reason to believe that our estimated cost for CPT code 0183T from CY 2008 claims data does not include the cost of the necessary supplies. The final CY 2010 median cost of CPT code 0183T is closer to the final CY 2010 median cost of APC 0013 than APC 0015. In fact, if we were to continue to assign CPT code 0183T to APC 0015 for CY 2010, APC 0015 would violate the 2 times rule. That is, if we maintained CPT code 0183T in APC 0015 for CY 2010 as requested by the commenters, it would be the significant procedure with the lowest median cost assigned to APC 0015. In turn, the median cost of approximately \$158 for the highest cost significant procedure, CPT code 11000 (Debridement of extensive eczematous

or infected skin; up to 1 of body surface), would be more than 2 times the median cost of CPT code 0183T, resulting in a 2 times violation in APC 0015. We note that the APC Panel heard several public presentations that addressed the proposed CY 2010 APC assignment of CPT code 0183T at the August 2009 meeting but made no recommendation regarding the CY 2010 assignment of the code. In particular, the APC Panel did not make a recommendation to us to maintain an APC configuration that would violate the 2 times rule and require that we except APC 0015 from the 2 times rule for CY 2010.

We also believe that APC 0013 is an appropriate APC assignment for CPT code 0183T based on clinical considerations. Other wound care services with similar median costs are assigned to APC 0013 for CY 2010, specifically CPT codes 97602 and 97605 (Negative pressure wound therapy (eg, vacuum assisted drainage collection), including topical application(s), wound assessment, and instruction(s) for ongoing care, per session; total wound(s) surface area less than 50 square centimeters).

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to reassign CPT 0183T from APC 0015 to APC 0013, with a final CY 2010 APC median cost of approximately \$59.

b. Skin Repair (APCs 0134 and 0135)

For CY 2010, we proposed to continue to assign the CPT skin repair codes for the application of Apligraf, Oasis, and Dermagraft skin substitutes to the same procedural APCs for CY 2010 as their CY 2009 assignments. Specifically, we proposed to continue to assign the Apligraf application CPT codes 15340 (Tissue cultured allogeneic skin substitute; first 25 sq cm or less) and 15341 (Tissue cultured allogeneic skin substitute; each additional 25 sq cm, or part thereof) to APC 0134 (Level II Skin Repair), with a proposed payment rate of approximately \$214. Likewise, we proposed to continue to assign the Dermagraft application CPT codes 15365 (Tissue cultured allogeneic dermal substitute, face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits; first 100 sq cm or less, or 1% of body area of infants and children) and 15366 (Tissue cultured allogeneic dermal substitute, face, scalp, eyelids, mouth, neck, ears, orbits, genitalia, hands, feet, and/or multiple digits; each additional 100 sq cm, or each additional 1% of body area of infants and children, or

part thereof) to APC 0134. We proposed to continue to assign the Oasis application CPT codes 15430 (Acellular xenograft implant; first 100 sq cm or less, or 1% of body area of infants and children) and 15431 (Acellular xenograft implant; each additional 100 sq cm, or each additional 1% of body area of infants and children, or part thereof) to APC 0135 (Level III Skin Repair), with a proposed payment rate of approximately \$297.

At the August 2009 meeting of the APC Panel, one public presenter requested that the APC Panel recommend that CMS reassign CPT codes 15340 and 15341 from APC 0134 to APC 0135. The presenter stated that the CY 2010 proposal to continue to assign both codes to APC 0134 would create a financial incentive favoring Dermagraft application. Specifically, the presenter explained that CPT instructions allow the separate reporting of the CPT codes for site preparation when Dermagraft is applied, while the CPT instructions for Apligraf application codes specify that site preparation cannot be separately reported. The presenter believed that this reporting difference and the resulting expected differences in the associated application procedure costs could be addressed by assigning the Apligraf application CPT codes to a higher paying APC than the Dermagraft application codes, instead of the same APC as CMS proposed for CY 2010. After discussion, the APC Panel requested that CMS provide data at the next APC Panel meeting on the frequency of primary and add-on CPT codes billed for Apligraf, Oasis, and Dermagraft application in order to assess the apparent variability in billing for the application of these products. In addition, the APC Panel requested median cost data for site preparation and debridement that may be separately reported in preparation for application of Dermagraft.

Comment: Several commenters supported the CY 2010 proposal to continue the CY 2009 APC assignments for the Apligraf, Dermagraft, and Oasis application CPT codes. One commenter argued that reassignment of the Apligraf application codes from APC 0134 to APC 0135 would create a financial incentive for hospitals to choose Apligraf instead of other products. Another commenter stated that the current APC assignments for all three sets of skin repair codes are appropriate based on an assessment of clinical homogeneity and resource costs.

Another commenter requested that CMS reassign the Apligraf application CPT codes 15340 and 15341 from APC

0134 to APC 0135 because of their similarity, from clinical and resource perspectives, to the Oasis application CPT codes 15430 and 15431 that are currently assigned to APC 0135. The commenter noted that none of these procedures allow separate reporting and payment of site preparation when performed. The commenter expressed concern that the variable APC assignments for similar procedures would create an unlevel playing field that would lead to financial incentives for hospitals to use one product rather than the other, as opposed to the most clinically appropriate product. Further, the commenter indicated that site preparation and debridement procedures are not paid separately when associated with Apligraf application, yet these site preparation services are paid separately when reported with Dermagraft application procedures that are assigned to the same APC as Apligraf application procedures. The commenter also requested that CMS not reassign the Oasis application CPT codes 15430 and 15431 from APC 0135 to APC 0134 because such a reassignment would inappropriately group skin repair procedures that incorporate site preparation with those that allow separate reporting and payment of that preparation.

Response: The current Apligraf, Oasis, and Dermagraft application CPT codes were made effective January 1, 2006. In the CY 2006 OPPS final rule with comment period (70 FR 68762), we assigned the Apligraf application CPT codes 15340 and 15341 and the Dermagraft application CPT codes 15365 and 15366 to the Level I Skin Repair APC (then designated as APC 0024 with a payment rate of approximately \$92). We assigned the Oasis application CPT codes 15430 and 15431 to the Level II Skin Repair APC (then designated as APC 0025 with a payment rate of approximately \$315) based on consideration of clinical and resource homogeneity.

For CY 2007 (71 FR 68054 through 68056), we assigned the three sets of skin repair CPT codes to the Level II Skin Repair APC (then designated as APC 0025) in response to comments received from the public regarding their clinical and expected resource similarity. However, for CY 2008, because of a 2 times violation in two of the four skin repair APCs that resulted from hospital claims data that were first available for these codes, we reconfigured the APC assignments for the Apligraf, Dermagraft, and Oasis application procedures. This reconfiguration resulted in our again differentiating the APC assignments for

the Oasis application CPT codes from the APC assignments for the Apligraf and Dermagraft application procedures, similar to the initial CY 2006 APC configuration. We also renumbered the Skin Repair APCs. We note that, for CY 2008, we made no change to the APC assignments for the Apligraf and Dermagraft application CPT codes, maintaining them in APC 0134, but we reassigned the Oasis application codes to APC 0135.

We retained these configurations for CY 2009 and, for CY 2010, we proposed to continue to assign these procedures to their CY 2009 APCs. We also proposed to pay separately for the Apligraf, Dermagraft, and Oasis products themselves in CY 2010. Analysis of our claims data for the application procedures revealed that the hospital resource costs associated with the Apligraf and Oasis application procedures are different. The median cost of the Apligraf application CPT code 15340 is approximately \$234, based on 13,551 single claims (of 17,534 total claims), and approximately \$186 for CPT code 15341, based on 1,789 single claims (of 4,424 total claims). For the Oasis application CPT code 15430, the median cost is approximately \$276 based on 12,807 single claims (of 14,723 total claims), and approximately \$261 for CPT code 15431 based on 150 single claims (of 293 total claims). These CPT code-specific median costs are consistent with the APC 0134 and APC 0135 median costs of approximately \$210 and \$296, respectively, where the different two sets of procedure codes are assigned.

The OPPS is a payment system that is based on the relativity of costs of procedures as reported to us by hospitals. Hospital costs, based on significant numbers of single claims, have been and continue to be consistently higher for the Oasis application procedures than for Apligraf or Dermagraft application procedures, despite the differences in CPT reporting instructions for Apligraf and Oasis application procedures in comparison with Dermagraft application procedures. We also note that the coverage areas for the Apligraf application codes are based on 25 square centimeter increments, whereas the Oasis and Dermagraft application codes are based on 100 square centimeter increments. While we are not sure of the contribution application of different products to different size wounds may have on hospital costs, we have no reason to believe that our high volume and consistent hospital claims data for these services do not accurately represent the costs of the procedures that have been

reported in accordance with their specific code descriptors since CY 2006.

Further, we do not agree that different APC assignments for similar skin repair procedures would create an unlevel playing field that would lead to financial incentives for hospitals to use one product rather than the other, as opposed to the most clinically appropriate product. Payments under the OPPS are based on the relative costs of services as reported to us by hospitals in claims and cost report data. In part, we assign services to APCs based on considerations of resource homogeneity, and hospital resources are reflected in the costs reported to us by hospitals. The skin repair CPT codes differ significantly from one another in terms of the other services that are bundled into them (such as site preparation) and in the coverage areas they describe. The specific Apligraf, Dermagraft, and Oasis application procedures have different median costs based on CY 2008 hospital claims that have led us to continue to assign them to different APCs for CY 2010, and we do not believe that appropriate payment for hospitals' costs for procedures provides incentives for hospitals to use one product instead of another. Instead, accurate payment based on the relative costs of services is an important principle of the OPPS, specifically intended to minimize any financial incentives for use of one product rather than the other in the case of similar procedures. We agree with the commenter that the choice of a patient's treatment should be based on clinical considerations, not financial incentives due to OPPS payment rates. We believe our final CY 2010 APC assignments for the Apligraf, Dermagraft, and Oasis application CPT codes are fully consistent with our interest in hospitals providing the most clinically appropriate treatments in an efficient manner.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to continue to assign the Apligraf and Dermagraft application CPT codes to APC 0134, which has a final CY 2010 APC median cost of approximately \$210, and to continue to assign the Oasis application CPT codes to APC 0135, which has a final CY 2010 APC median cost of approximately \$296. We note that when hospitals are performing these procedures, they also would report the Level II HCPCS codes that describe the biological products that are used with the Apligraf, Dermagraft, and Oasis application CPT codes, which are paid separately in CY 2010. Further, we are accepting the August 2009

recommendation of the APC Panel and will provide information at the winter 2010 APC Panel meeting on the frequency of primary and add-on CPT codes billed for Apligraf, Oasis, and Dermagraft application procedures, in addition to providing median cost data for site preparation and debridement that may be separately reported in preparation for application of Dermagraft.

c. Group Psychotherapy (APC 0325)

For CY 2010, we proposed to continue to assign CPT codes 90849 (Multiple-family group psychotherapy), 90853 (Group psychotherapy (other than of a multiple-family group)), and 90857 (Interactive group psychotherapy) to APC 0325 (Group Psychotherapy), with a proposed payment rate of approximately \$61, calculated according to the standard OPPS ratesetting methodology. In CY 2009, these three CPT codes also were the only codes assigned to APC 0325, with a payment rate of approximately \$65.

Comment: Several commenters expressed concern that the CY 2010 proposed payment rate for APC 0325 of approximately \$61 is 21 percent less than the CY 2006 payment rate for this APC, and 24 percent less than the CY 2004 payment rate for this APC. The commenters stated that the proposed payment rate would be insufficient to cover hospitals' costs for providing group mental health services and, as a result, would threaten beneficiary access to these services. Some commenters recommended that CMS increase the final CY 2010 payment rate for APC 0325 by approximately 17 percent, which the commenters calculated is the average increase from CY 2006 to CY 2010 for the other psychotherapy APCs, specifically APC 0322 (Brief Individual Psychotherapy), APC 0323 (Extended Individual Psychotherapy), and APC 0324 (Family Psychotherapy).

Response: As we have stated in the past regarding APC 0325 (72 FR 66739 and 73 FR 68627), we cannot speculate as to why the median cost of group psychotherapy services has decreased significantly since CY 2004. We again note that we have robust claims data for the CPT codes that map to APC 0325. Specifically, we were able to use more than 99 percent of the approximately 1.6 million claims submitted by hospitals to report group psychotherapy services. We set the payment rates for APC 0325 using our standard OPPS methodology based on relative costs from hospital outpatient claims. We have no reason to believe that our claims data, as reported by hospitals, do not accurately reflect

the hospital costs of group psychotherapy services. It would appear that the relative cost of providing these mental health services, in comparison with other HOPD services has decreased in recent years.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to calculate the payment rate for APC 0325 by applying our standard OPPS ratesetting methodology that relies on all single claims for all procedures assigned to the APC. The final CY 2010 APC median cost of APC 0325 is approximately \$59.

d. Portable X-Ray Services

Consistent with applicable requirements, hospitals may bill and be paid under the OPPS for diagnostic x-ray tests performed in locations other than HOPDs, such as a skilled nursing facility (SNF), if the patient is receiving the x-ray as a covered outpatient department service and not in the course of a Medicare-covered SNF stay. The charge for the x-ray (but not the transportation and set-up charges) is billed on a hospital outpatient claim. Medicare does not pay under the OPPS for transportation or set-up when the x-ray equipment is transported to another location where the x-ray is taken.

Comment: One commenter objected to the assignment of status indicator "B" (Codes that are not recognized by OPPS when submitted on an outpatient hospital Part B bill type (12X or 13X)) to HCPCS codes R0070 (Transportation of portable x-ray equipment and personnel to home or nursing home, per trip to facility or location, one patient seen); R0075 (Transportation of portable x-ray equipment and personnel to home or nursing home, per trip to facility or location, more than one patient seen); and Q0092 (Set up portable x-ray equipment) under the OPPS when a hospital transports and sets up a portable x-ray machine in a SNF or other nonhospital site of service to furnish an x-ray to a patient who is not in the course of a SNF stay that is covered by Medicare. The commenter indicated that to be paid for the transportation and set-up of the portable x-ray, the hospital must enroll as a supplier and bill the Medicare carrier or MAC on a HCFA 1500 claim for the transportation and set-up services, although the hospital may bill the fiscal intermediary or MAC on a UB-04 claim for the x-ray service itself. The commenter requested that CMS revise its billing instructions so that the transportation and set-up charges for portable x-ray services could be

reported on the same claim as the hospital's charge for the x-ray.

Response: In the case in which a patient receiving the portable x-ray service is not in a Medicare-covered SNF stay but a hospital furnishes the portable x-ray service in the SNF as a covered outpatient department service consistent with all applicable requirements, the HCPCS code and charge for the x-ray service (but not transportation and set-up charges) are billed to the fiscal intermediary or MAC. Payment is made under the OPPS for the x-ray service under such circumstances. The transportation and set-up of the portable x-ray are also covered services which are currently reported on the HCFA 1500 claim and are carrier-priced. We assign status indicator "B" to HCPCS codes R0070, R0075, and Q0092 because these services (transportation and set-up of the portable x-ray) are not paid under the OPPS and are rejected by the I/OCE if they are billed in an outpatient hospital bill type. We will explore whether it is feasible to revise the billing instructions to enable hospitals to bill for these transportation and set-up services on the same claim on which they report the charge for the x-ray service to which the transportation and set-up charges are ancillary. If we determine that it would be feasible and desirable to propose this change, we would propose to change the status indicators of these codes accordingly.

After consideration of the public comment we received, we are finalizing our CY 2010 proposals, without modification, to continue to assign the status indicator of "B" to HCPCS codes R0070, R0075, and Q0092. We will explore the feasibility of alternatives for billing and payment of these services that could reduce the hospital administrative burden associated with billing for the services.

e. Home Sleep Study Tests (APC 0213)

For CY 2010, we proposed to continue to assign Level II HCPCS codes G0398 (Home sleep study test (HST) with type II portable monitor, unattended; minimum of 7 channels: EEG, EOG, EMG, ECG/heart rate, airflow, respiratory effort and oxygen saturation), G0399 (Home sleep test (HST) with type III portable monitor, unattended; minimum of 4 channels: 2 respiratory movement/airflow, 1 ECG/heart rate and 1 oxygen saturation), and G0400 (Home sleep test (HST) with type IV portable monitor, unattended; minimum of 3 channels) to APC 0213 (Level I Extended EEG, Sleep, and Cardiovascular Studies), with a

proposed payment rate of approximately \$160.

Comment: One commenter urged CMS to pay appropriately for Level II HCPCS codes G0398, G0399, and G0400 to adequately cover the cost of devices used in performing these procedures. Specifically, the commenter stated that the acquisition costs for the devices used with these procedures are significant and vary between \$4,400 and \$16,500. The commenter argued that it was unreasonable for CMS to assign all three HCPCS G-codes to the same APC because the devices used for the procedures vary significantly in their costs and, therefore, payment at the same rate for all three services would violate the 2 times rule. The commenter urged CMS to review the proposed payment rates for HCPCS G-codes G0398, G0399, and G0400.

Response: As we explained in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68602), we created these three HCPCS G-codes to describe the various types of home sleep tests that Medicare determined could be used to allow for coverage of continuous positive airway pressure (CPAP) therapy based upon a diagnosis of obstructive sleep apnea (OSA) according to a home sleep study. We further explained that we decided to assign these HCPCS G-codes to an APC under the OPPS because we believe these diagnostic services may be provided by HOPDs to Medicare beneficiaries.

HCPCS codes G0398, G0399, and G0400 were made effective in March 2008. Analysis of our claims data from CY 2008 reveals that these services are not commonly performed in the hospital outpatient setting for Medicare beneficiaries. Our claims data show no single claims and only three total claims for HCPCS code G0398. The median cost of HCPCS code G0399 is approximately \$236 based on 12 single claims (of 13 total claims), and the median cost of HCPCS code G0400 is approximately \$80 based on 11 single claims (of 12 total claims). We believe it would be difficult to draw any conclusions about the resource differences among these three services based upon such limited claims data from a single year.

With regard to the commenter's concern about a violation of the 2 times rule, there is no 2 times violation in APC 0213 because none of the sleep study HCPCS G-codes are significant procedures in the APC. Generally, we review, on an annual basis, the items and services within an APC group to determine, with respect to comparability of the use of resources, if the median cost of the highest cost item

or service within an APC group is more than 2 times greater than the median cost of the lowest cost item or service within that same group, thereby assessing for 2 times rule violations. We make exceptions to the 2 times rule in unusual cases, such as low-volume items and services, and we only consider significant procedures for purposes of the 2 times assessment. We define significant procedures as those with a single claim frequency of greater than 1,000 or those with a frequency of greater than 99 and that constitute at least 2 percent of single claims in the APC. For APC 0213, our CY 2008 hospital outpatient claims used for CY 2010 ratesetting show that the median cost of the lowest cost significant service is approximately \$150 compared to approximately \$241 for the highest cost service. Based on our claims data, there is no 2 times violation in APC 0213.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign HCPCS codes G0398, G0399, and G0400 to APC 0213, which has a final CY 2010 APC median cost of approximately \$161.

IV. OPPS Payment for Devices

A. Pass-Through Payments for Devices

1. Expiration of Transitional Pass-Through Payments for Certain Devices

Section 1833(t)(6)(B)(iii) of the Act requires that, under the OPPS, a category of devices be eligible for transitional pass-through payments for at least 2, but not more than 3, years. This pass-through payment eligibility period begins with the first date on which transitional pass-through payments may be made for any medical device that is described by the category. We may establish a new device category for pass-through payment in any quarter. Under our established policy, we base the pass-through status expiration dates for the category codes on the date on which a category is in effect. The date on which a category is in effect is the first date on which pass-through payment may be made for any medical device that is described by such category. We propose and finalize the dates for expiration of pass-through status for device categories as part of the OPPS annual update.

We also have an established policy to package the costs of the devices no longer eligible for pass-through payments into the costs of the procedures with which the devices are reported in the claims data used to set the payment rates (67 FR 66763).

Brachytherapy sources, which are now separately paid in accordance with section 1833(t)(2)(H) of the Act, are an exception to this established policy.

There currently are no device categories eligible for pass-through payment, and there are no categories for which we proposed expiration of pass-through status. If we create new device categories for pass-through payment status during the remainder of CY 2009 or during CY 2010, we will propose future expiration dates in accordance with the statutory requirement that they be eligible for pass-through payments for at least 2, but not more than 3, years from the date on which pass-through payment for any medical device described by the category may first be made.

2. Provisions for Reducing Transitional Pass-Through Payments To Offset Costs Packaged Into APC Groups

a. Background

We have an established policy to estimate the portion of each APC payment rate that could reasonably be attributed to the cost of the associated devices that are eligible for pass-through payments (66 FR 59904). We deduct from the pass-through payments for identified device categories eligible for pass-through payments an amount that reflects the portion of the APC payment amount that we determine is associated with the cost of the device, defined as the device APC offset amount, as required by section 1833(t)(6)(D)(ii) of the Act. We have consistently employed an established methodology to estimate the portion of each APC payment rate that could reasonably be attributed to the cost of an associated device eligible for pass-through payment, using claims data from the period used for the most recent recalibration of the APC rates (72 FR 66751 through 66752). We establish and update the applicable device APC offset amounts for eligible pass-through device categories through the transmittals that implement the quarterly OPPS updates.

We currently have published a list of all procedural APCs with the CY 2009 portions (both percentages and dollar amounts) of the APC payment amounts that we determine are associated with the cost of devices, on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/01_overview.asp. The dollar amounts are used as the device APC offset amounts. In addition, in accordance with our established practice, the device APC offset amounts in a related APC are used in order to evaluate whether the cost of a device in an application for a new

device category for pass-through payment is not insignificant in relation to the APC payment amount for the service related to the category of devices, as specified in our regulations at § 419.66(d).

b. Final Policy

In the CY 2010 OPPI/ASC proposed rule (74 FR 35306), for CY 2010, we proposed to continue our established policies for calculating and setting the device APC offset amounts for each device category eligible for pass-through payment. We also proposed to continue to review each new device category on a case-by-case basis to determine whether device costs associated with the new category are already packaged into the existing APC structure. If device costs packaged into the existing APC structure are associated with the new category, we proposed to deduct the device APC offset amount from the pass-through payment for the device category. As stated earlier, these device APC offset amounts also would be used in order to evaluate whether the cost of a device in an application for a new device category for pass-through payment is not insignificant in relation to the APC payment amount for the service related to the category of devices (§ 419.66(d)).

In section V.A.4. of the CY 2010 OPPI/ASC proposed rule (74 FR 35311 through 35314), we proposed to specify that, beginning in CY 2010, the pass-through evaluation process and pass-through payment methodology for implantable biologicals, that are surgically inserted or implanted (through a surgical incision or a natural orifice) and that are newly approved for pass-through status beginning on or after January 1, 2010, would be the device pass-through process and payment methodology only. As a result of that proposal, we then proposed that, beginning in CY 2010, we would include implantable biologicals in our calculation of the device APC offset amounts. As of CY 2009, the costs of implantable biologicals not eligible for pass-through payment are packaged into the costs of the procedures in which they are implanted because nonpass-through implantable biologicals are not separately paid. We proposed to calculate and set any device APC offset amount for a new device pass-through category that includes a newly eligible implantable biological beginning in CY 2010 using the same methodology we have historically used to calculate and set device APC offset amounts for device categories eligible for pass-through payment (72 FR 66751 through 66752), with one modification. Because

implantable biologicals would be considered devices rather than drugs for purposes of pass-through evaluation and payment under this proposal for CY 2010, the device APC offset amounts would include the costs of implantable biologicals for the first time. We also proposed to utilize these revised device APC offset amounts to evaluate whether the cost of an implantable biological in an application for a new device category for pass-through payment is not insignificant in relation to the APC payment amount for the service related to the category of devices. Further, we proposed to no longer use the "policy-packaged" drug APC offset amounts for evaluating the cost significance of implantable biological pass-through applications under review and for setting the APC offset amounts that would apply to pass-through payment for those implantable biologicals, effective for new pass-through status determinations beginning in CY 2010. In addition, we proposed to update, on the CMS Web site at <http://www.cms.hhs.gov/HospitalOutpatientPPS>, the list of all procedural APCs with the final CY 2010 portions of the APC payment amounts that we determine are associated with the cost of devices so that this information is available for use by the public in developing potential CY 2010 device pass-through payment applications and by CMS in reviewing those applications.

Comment: One commenter noted that a significant consequence of paying for new implantable biologicals under the device pass-through payment methodology would be that the payment for the implantable biological would be reduced by the estimated cost of any predecessor devices included in the APC payment rate. The commenter believed that it is reasonable for CMS to reduce the payment for the pass-through implantable biological when the biological is used in lieu of a predecessor device whose cost is already incorporated into payment for the associated procedure. However, the commenter also stated that if the hospital implanted the predecessor device during the procedure in addition to the pass-through implantable biological, a reduction in the pass-through payment for the implantable biological by the predecessor device cost should not be taken.

Response: Concerning the commenter's request that we not take a reduction (that is, device APC offset) when both a predecessor device and an implantable biological that is on pass-through status are used in a procedure in the case of medical necessity, we note

that our standard policy when establishing a new device category for pass-through payment is to determine whether device costs associated with the new category are already packaged into the relevant existing clinical APC. If device costs packaged into the existing clinical APC are associated with the new pass-through device category and these predecessor devices would generally not be used when a device described by the new device category was implanted, we identify the device APC offset that would be deducted from the pass-through payment amount each time the new category is reported with the related clinical APC. We make determinations about the applicability of a device APC offset based on our overall clinical understanding of the device category and its associated procedures, rather than on a claim-by-claim basis for each different scenario. In the rare case where an implantable biological that is described by a device category with pass-through status was used in addition to a predecessor device in the performance of a procedure for which we had determined that a device APC offset was applicable, we would still apply the device APC offset to the pass-through payment for the implantable biological. With respect to a prospective payment system such as the OPPI, in some individual cases, payment exceeds the average cost; in other cases, payment is less than the average cost of an individual case. On balance, however, payment should approximate the relative cost of the average case, recognizing that, as a prospective payment system, the OPPI is a system of averages. We would not expect the scenario of implanting both a new implantable biological and the predecessor device described by the commenter to be common. If such a clinical scenario were common, we would determine that no device APC offset would apply to the new device category because the implantable biological was typically used in addition to the predecessor device in performing the associated procedure.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to reduce device pass-through payments based on device costs already included in the associated procedural APCs, when we determine that device costs associated with the new category are already packaged into the existing APC structure.

B. Adjustment to OPSS Payment for No Cost/Full Credit and Partial Credit Devices

1. Background

In recent years, there have been several field actions on and recalls of medical devices as a result of implantable device failures. In many of these cases, the manufacturers have offered devices without cost to the hospital or with credit for the device being replaced if the patient required a more expensive device. In order to ensure that payment rates for procedures involving devices reflect only the full costs of those devices, our standard ratesetting methodology for device-dependent APCs uses only claims that contain the correct device code for the procedure, do not contain token charges, and do not contain the "FB" modifier signifying that the device was furnished without cost or with a full credit. As discussed in section II.A.2.d.(1) of the CY 2010 OPSS/ASC proposed rule (74 FR 35267) and this final rule with comment period, we are further refining our standard ratesetting methodology for device-dependent APCs for CY 2010 by also excluding claims with the "FC" modifier signifying that the device was furnished with partial credit.

To ensure equitable payment when the hospital receives a device without cost or with full credit, in CY 2007 we implemented a policy to reduce the payment for specified device-dependent APCs by the estimated portion of the APC payment attributable to device costs (that is, the device offset) when the hospital receives a specified device at no cost or with full credit (71 FR 68071 through 68077). Hospitals are instructed to report no cost/full credit cases using the "FB" modifier on the line with the procedure code in which the no cost/full credit device is used. In cases in which the device is furnished without cost or with full credit, the hospital is instructed to report a token device charge of less than \$1.01. In cases in which the device being inserted is an upgrade (either of the same type of device or to a different type of device) with a full credit for the device being replaced, the hospital is instructed to report as the device charge the difference between its usual charge for the device being implanted and its usual charge for the device for which it received full credit. In CY 2008, we expanded this payment adjustment policy to include cases in which hospitals receive partial credit of 50 percent or more of the cost of a specified device. Hospitals are instructed to append the "FC" modifier to the

procedure code that reports the service provided to furnish the device when they receive a partial credit of 50 percent or more of the cost of the new device. We reduce the OPSS payment for the implantation procedure by 100 percent of the device offset for no cost/full credit cases when both a specified device code is present on the claim and the procedure code maps to a specified APC. Payment for the implantation procedure is reduced by 50 percent of the device offset for partial credit cases when both a specified device code is present on the claim and the procedure code maps to a specified APC. Beneficiary copayment is based on the reduced payment amount when either the "FB" or the "FC" modifier is billed and the procedure and device codes appear on the lists of procedures and devices to which this policy applies. We refer readers to the CY 2008 OPSS/ASC final rule with comment period for more background information on the "FB" and "FC" payment adjustment policies (72 FR 66743 through 66749).

2. APCs and Devices Subject to the Adjustment Policy

In the CY 2010 OPSS/ASC proposed rule (74 FR 35307), we proposed for CY 2010 to continue the policy of reducing OPSS payment for specified APCs by 100 percent of the device offset amount when a hospital furnishes a specified device without cost or with a full credit and by 50 percent of the device offset amount when the hospital receives partial credit in the amount of 50 percent or more of the cost for the specified device. Because the APC payments for the related services are specifically constructed to ensure that the full cost of the device is included in the payment, we stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35307) that we continue to believe it is appropriate to reduce the APC payment in cases in which the hospital receives a device without cost, with full credit, or with partial credit, in order to provide equitable payment in these cases. (We refer readers to section II.A.2.d.(1) of this final rule with comment period for a description of our standard ratesetting methodology for device-dependent APCs.) Moreover, the payment for these devices comprises a large part of the APC payment on which the beneficiary copayment is based, and we continue to believe it is equitable that the beneficiary cost sharing reflects the reduced costs in these cases.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35307), we also proposed to continue using the three criteria established in the CY 2007 OPSS/ASC final rule with comment period for

determining the APCs to which this policy applies (71 FR 68072 through 68077). Specifically, (1) all procedures assigned to the selected APCs must involve implantable devices that would be reported if device insertion procedures were performed; (2) the required devices must be surgically inserted or implanted devices that remain in the patient's body after the conclusion of the procedure (at least temporarily); and (3) the device offset amount must be significant, which, for purposes of this policy, is defined as exceeding 40 percent of the APC cost. We proposed to continue to restrict the devices to which the APC payment adjustment would apply to a specific set of costly devices to ensure that the adjustment would not be triggered by the implantation of an inexpensive device whose cost would not constitute a significant proportion of the total payment rate for an APC. We stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35307) that we continue to believe these criteria are appropriate because free devices and device credits are likely to be associated with particular cases only when the device must be reported on the claim and is of a type that is implanted and remains in the body when the beneficiary leaves the hospital. We believe that the reduction in payment is appropriate only when the cost of the device is a significant part of the total cost of the APC into which the device cost is packaged, and that the 40-percent threshold is a reasonable definition of a significant cost.

As indicated in the CY 2010 OPSS/ASC proposed rule (74 FR 35307), we examined the offset amounts calculated from the CY 2010 proposed rule data and the clinical characteristics of APCs to determine whether the APCs to which the no cost/full credit and partial credit device adjustment policy applies in CY 2009 continue to meet the criteria for CY 2010, and to determine whether other APCs to which the policy does not apply in CY 2009 would meet the criteria for CY 2010. Based on the CY 2008 claims data available for the CY 2010 proposed rule, we did not propose any changes to the APCs and devices to which this policy applies. Table 19 of the CY 2010 OPSS/ASC proposed rule (74 FR 35307 through 35308) listed the proposed APCs to which the payment adjustment policy for no cost/full credit and partial credit devices would apply in CY 2010 and displayed the proposed payment adjustment percentages for both no cost/full credit and partial credit circumstances. Table 20 of the CY 2010 OPSS/ASC proposed rule (74 FR

35308) listed the proposed devices to which this policy would apply in CY 2010. We stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35307) that we would update the lists of APCs and devices to which the no cost/full credit and partial credit device adjustment policy would apply in CY 2010, consistent with the three selection criteria discussed earlier in this section and based on the final CY 2008 claims data available for this CY 2010 OPPS/ASC final rule with comment period.

We did not receive any public comments on our CY 2010 proposal to continue the policy of reducing OPPS payment for specified APCs by 100 percent of the device offset amount when a hospital furnishes a specified device without cost or with a full credit and by 50 percent of the device offset amount when the hospital receives partial credit in the amount of 50 percent or more of the cost for the specified device. We also did not receive any public comments on our CY 2010 proposal to continue using the three criteria established in the CY 2007 OPPS/ASC final rule with comment

period for determining the APCs to which this policy applies (71 FR 68072 through 68077). Therefore, we are finalizing our CY 2010 proposals, without modification, to continue the established no cost/full credit and partial credit device adjustment policy. For CY 2010, OPPS payments for implantation procedures to which the “FB” modifier is appended are reduced by 100 percent of the device offset for no cost/full credit cases when both a device code listed in Table 29, below, is present on the claim and the procedure code maps to an APC listed in Table 28 below. OPPS payments for implantation procedures to which the “FC” modifier is appended are reduced by 50 percent of the device offset when both a device code listed in Table 29 is present on the claim and the procedure code maps to an APC listed in Table 28. Beneficiary copayment is based on the reduced amount when either the “FB” or “FC” modifier is billed and the procedure and device codes appear on the lists of procedures and devices to which this policy applies.

We are adding device HCPCS code L8680 (Implantable neurostimulator electrode, each) to the list of devices in Table 29 because we are changing the status indicator for this code from “B” (Codes that are not recognized by OPPS when submitted on an outpatient hospital Part B bill type (12x and 13x)) to “N” (Items and Services Packaged into APC Rates) for CY 2010, as reflected in Addendum B to this final rule with comment period. We are recognizing HCPCS code L8680 for payment purposes under the OPPS because it appropriately describes neurostimulator electrodes, and we typically try to recognize all valid HCPCS codes that hospitals may use to report items and services provided to hospital outpatients that are packaged or otherwise payable under the OPPS. This change in status indicator for HCPCS code L8680 for CY 2010 does not require hospitals to change their current billing practices in any way, but it does provide them with the flexibility to use this code if they choose to do so.

TABLE 28—APCS TO WHICH THE NO COST/FULL CREDIT AND PARTIAL CREDIT DEVICE ADJUSTMENT POLICY WILL APPLY

Final CY 2010 APC	CY 2010 APC title	Final CY 2010 device off-set percentage for no cost/full credit case	Final CY 2010 device off-set percentage for partial credit case
0039	Level I Implantation of Neurostimulator Generator	85	43
0040	Percutaneous Implantation of Neurostimulator Electrodes	58	29
0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electrodes.	64	32
0089	Insertion/Replacement of Permanent Pacemaker and Electrodes	72	36
0090	Insertion/Replacement of Pacemaker Pulse Generator	74	37
0106	Insertion/Replacement of Pacemaker Leads and/or Electrodes	44	22
0107	Insertion of Cardioverter-Defibrillator	89	44
0108	Insertion/Replacement/Repair of Cardioverter-Defibrillator Leads	88	44
0225	Implantation of Neurostimulator Electrodes, Cranial Nerve	73	37
0227	Implantation of Drug Infusion Device	83	41
0259	Level VII ENT Procedures	85	42
0315	Level II Implantation of Neurostimulator Generator	88	44
0385	Level I Prosthetic Urological Procedures	59	30
0386	Level II Prosthetic Urological Procedures	71	35
0418	Insertion of Left Ventricular Pacing Elect.	81	41
0425	Level II Arthroplasty or Implantation with Prosthesis	58	29
0648	Level IV Breast Surgery	48	24
0654	Insertion/Replacement of a permanent dual chamber pacemaker	75	37
0655	Insertion/Replacement/Conversion of a permanent dual chamber pacemaker.	75	37
0680	Insertion of Patient Activated Event Recorders	73	36

TABLE 29—DEVICES TO WHICH THE NO COST/FULL CREDIT AND PARTIAL CREDIT DEVICE ADJUSTMENT POLICY WILL APPLY

CY 2010 device HCPCS code	CY 2010 short descriptor
C1721	AICD, dual chamber.
C1722	AICD, single chamber.
C1728	Cath, brachytx seed adm.
C1764	Event recorder, cardiac.
C1767	Generator, neurostim, imp.
C1771	Rep dev, urinary, w/sling.
C1772	Infusion pump, programmable.
C1776	Joint device (implantable).
C1777	Lead, AICD, endo single coil.
C1778	Lead, neurostimulator.
C1779	Lead, pmkr, transvenous VDD.
C1785	Pmkr, dual, rate- resp.
C1786	Pmkr, single, rate- resp.
C1789	Prosthesis, breast, imp.
C1813	Prosthesis, penile, inflatab.
C1815	Pros, urinary sph, imp.
C1820	Generator, neuro rechg bat sys.
C1881	Dialysis access system.
C1882	AICD, other than sing/dual.
C1891	Infusion pump, non-prog, perm.
C1895	Lead, AICD, endo dual coil.
C1896	Lead, AICD, non sing/dual.
C1897	Lead, neurostim, test kit.
C1898	Lead, pmkr, other than trans.
C1899	Lead, pmkr/AICD combination.
C1900	Lead coronary venous.
C2619	Pmkr, dual, non rate- resp.
C2620	Pmkr, single, non rate- resp.
C2621	Pmkr, other than sing/dual.
C2622	Prosthesis, penile, non-inf.
C2626	Infusion pump, non-prog, temp.
C2631	Rep dev, urinary, w/o sling.
L8600	Implant breast silicone/eq.
L8614	Cochlear device/system.
L8680	Implt neurostim elctr each.
L8685	Implt nrostm pls gen sng rec.
L8686	Implt nrostm pls gen sng non.
L8687	Implt nrostm pls gen dua rec.
L8688	Implt nrostm pls gen dua non.
L8690	Aud osseo dev, int/ext comp.

V. OPPS Payment Changes for Drugs, Biologicals, and Radiopharmaceuticals

A. OPPS Transitional Pass-Through Payment for Additional Costs of Drugs, Biologicals, and Radiopharmaceuticals

1. Background

Section 1833(t)(6) of the Act provides for temporary additional payments or “transitional pass-through payments” for certain drugs and biological agents. As enacted by the Medicare, Medicaid, and SCHIP Balanced Budget Refinement

Act (BBRA) of 1999 (Pub. L. 106–113), this provision requires the Secretary to make additional payments to hospitals for current orphan drugs, as designated under section 526 of the Federal Food, Drug, and Cosmetic Act (Pub. L. 107–186); current drugs and biological agents and brachytherapy sources used for the treatment of cancer; and current radiopharmaceutical drugs and biological products. For those drugs and biological agents referred to as “current,” the transitional pass-through payment began on the first date the hospital OPPS was implemented.

Transitional pass-through payments also are provided for certain “new” drugs and biological agents that were not being paid for as an HOPD service as of December 31, 1996, and whose cost is “not insignificant” in relation to the OPPS payments for the procedures or services associated with the new drug or biological. For pass-through payment purposes, radiopharmaceuticals are included as “drugs.” Under the statute, transitional pass-through payments for a drug or biological described in section 1833(t)(6)(C)(i)(II) of the Act can be made for at least 2 years but not more than 3 years after the product’s first payment as a hospital outpatient service under Part B. The pass-through payment eligibility period is discussed in detail in section V.A.5. of this final rule with comment period. CY 2010 pass-through drugs and biologicals and their designated APCs are assigned status indicator “G” in Addenda A and B to this final rule with comment period.

Section 1833(t)(6)(D)(i) of the Act specifies that the pass-through payment amount, in the case of a drug or biological, is the amount by which the amount determined under section 1842(o) of the Act (or, if the drug or biological is covered under a competitive acquisition contract under section 1847B of the Act, an amount determined by the Secretary to be equal to the average price for the drug or biological for all competitive acquisition areas and the year established under such section as calculated and adjusted by the Secretary) for the drug or biological exceeds the portion of the otherwise applicable Medicare OPD fee schedule that the Secretary determines is associated with the drug or biological. This methodology for determining the pass-through payment amount is set forth in § 419.64 of the regulations, which specifies that the pass-through payment equals the amount determined under section 1842(o) of the Act minus the portion of the APC payment that CMS determines is associated with the drug or biological. Section 1847A of the Act establishes the use of the average

sales price (ASP) methodology as the basis for payment for drugs and biologicals described in section 1842(o)(1)(C) of the Act that are furnished on or after January 1, 2005. The ASP methodology, as applied under the OPPS, uses several sources of data as a basis for payment, including the ASP, wholesale acquisition cost (WAC), and average wholesale price (AWP). In this final rule with comment period, the term “ASP methodology” and “ASP-based” are inclusive of all data sources and methodologies described therein. Additional information on the ASP methodology can be found on the ASP Web site at: <http://www.cms.hhs.gov/McrPartBDrugAvgSalesPrice>.

As noted above, section 1833(t)(6)(D)(i) of the Act also states that if a drug or biological is covered under a competitive acquisition contract under section 1847B of the Act, the payment rate is equal to the average price for the drug or biological for all competitive acquisition areas and the year established as calculated and adjusted by the Secretary. Section 1847B of the Act establishes the payment methodology for Medicare Part B drugs and biologicals under the competitive acquisition program (CAP). The Part B drug CAP was implemented on July 1, 2006, and included approximately 190 of the most common Part B drugs provided in the physician’s office setting. As we noted in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68633), the Part B drug CAP program was suspended beginning in CY 2009 (Medicare Learning Network (MLN) Matters Special Edition 0833, available via the Web site: <http://www.medicare.gov>). Therefore, there is no effective Part B drug CAP rate for pass-through drugs and biologicals as of January 1, 2009. As we indicated in the CY 2010 OPPS/ASC proposed rule (74 FR 35309), if the program is reinstated during CY 2010 and Part B drug CAP rates become available, we would again use the Part B drug CAP rate for pass-through drugs and biologicals if they are a part of the Part B drug CAP program. Otherwise, we would continue to use the rate that would be paid in the physician’s office setting for drugs and biologicals with pass-through status. We note that the CY 2010 MPFS proposed rule (74 FR 33623 through 33633) included proposed changes to the operation of the Part B drug CAP program, including a proposed change in the frequency of CAP drug pricing updates. A discussion of the final CAP policies is available in the CY 2010 MPFS final rule with comment period.

For CYs 2005, 2006, and 2007, we estimated the OPPS pass-through

payment amount for drugs and biologicals to be zero based on our interpretation that the “otherwise applicable Medicare OPD fee schedule” amount was equivalent to the amount to be paid for pass-through drugs and biologicals under section 1842(o) of the Act (or section 1847B of the Act, if the drug or biological is covered under a competitive acquisition contract). We concluded for those years that the resulting difference between these two rates would be zero. For CYs 2008 and 2009, we estimated the OPPS pass-through payment amount for drugs and biologicals to be \$6.6 million and \$23.3 million, respectively. Our final OPPS pass-through payment estimate for drugs and biologicals in CY 2010 is \$35.5 million, which is discussed in section VI.B. of this final rule with comment period.

The pass-through application and review process for drugs and biologicals is explained on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/04_passthrough_payment.asp.

2. Drugs and Biologicals With Expiring Pass-Through Status in CY 2009

In the CY 2010 OPPS/ASC proposed rule (74 FR 35309), we proposed that the pass-through status of 6 drugs and biologicals would expire on December 31, 2009, as listed in Table 21 of the proposed rule (74 FR 35309 through 35310). These items were approved for pass-through status on or before January 1, 2008 and, therefore, all of these drugs and biologicals will have received OPPS pass-through payment for at least 2 years and no more than 3 years by December 31, 2009.

Two of the products with proposed expiring pass-through status for CY 2010 are biologicals that are solely surgically implanted according to their Food and Drug Administration approved indications. As discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68634), we package payment for those implantable biologicals that have expiring pass-through status into payment for the associated surgical procedure. In the CY 2010 OPPS/ASC proposed rule, we proposed to package payment for two products described by HCPCS codes C9354 (Acellular pericardial tissue matrix of non-human origin (Veritas), per square centimeter) and C9355 (Collagen nerve cuff (NeuroMatrix), per 0.5 centimeter length).

To date, for other nonpass-through biologicals paid under the OPPS that may sometimes be used as implantable devices, we have instructed hospitals, via Transmittal 1336, Change Request 5718, dated September 14, 2007, to not separately bill for the HCPCS codes for the products when using these items as implantable devices (including as a scaffold or an alternative to human or nonhuman connective tissue or mesh used in a graft) during surgical procedures. In such cases, we consider payment for the biological used as an implantable device in a specific clinical case to be included in payment for the surgical procedure.

As we established in the CY 2003 OPPS final rule with comment period (67 FR 66763), when the pass-through payment period for an implantable device ends, it is standard OPPS policy to package payment for the implantable device into payment for its associated surgical procedure. We consider nonpass-through implantable devices to be integral and supportive items and services for which packaged payment is most appropriate. According to our regulations at § 419.2(b), as a prospective payment system, the OPPS establishes a national payment rate that includes operating and capital-related costs that are directly related and integral to performing a procedure or furnishing a service on an outpatient basis including, but not limited to, implantable prosthetics, implantable durable medical equipment, and medical and surgical supplies. Therefore, when the period of nonbiological device pass-through payment ends, we package the costs of the devices no longer eligible for pass-through payment into the costs of the procedures with which the devices were reported in the claims data used to set the payment rates for the upcoming calendar year. As described in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68634), we believed that this policy to package payment for implantable devices that are integral to the performance of separately paid procedures should also apply to payment for implantable biologicals without pass-through status, when those biologicals function as implantable devices. As stated above, implantable biologicals may be used in place of other implantable nonbiological devices whose costs are already accounted for in the associated procedural APC payments for surgical

procedures. If we were to provide separate payment for these implantable biologicals without pass-through status, we would potentially be providing duplicate device payment, both through the packaged nonbiological device cost included in the surgical procedure's payment and separate biological payment. We indicated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68634) that we saw no basis for treating implantable biological and nonbiological devices without pass-through status differently for OPPS payment purposes because both are integral to and supportive of the separately paid surgical procedures in which either may be used.

With the exception of those groups of drugs and biologicals that are always packaged when they do not have pass-through status, specifically diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals, our standard methodology of providing payment for drugs and biologicals with expiring pass-through status in an upcoming calendar year is to determine the product's estimated per day cost and compare it with the OPPS drug packaging threshold for that calendar year (which is \$65 for CY 2010), as discussed further in section V.B.2. of this final rule with comment period. If the drug's or biological's estimated per day cost is less than or equal to the applicable OPPS drug packaging threshold, we would package payment for the drug or biological into the payment for the associated procedure in the upcoming calendar year. If the estimated per day cost of the drug or biological is greater than the OPPS drug packaging threshold, we would provide separate payment at the applicable relative ASP-based payment amount (which is at ASP+4 percent for CY 2010, as discussed further in section V.B.3. of this final rule with comment period). Section V.B.2.d. of this final rule with comment period discusses the packaging of all nonpass-through contrast agents, diagnostic radiopharmaceuticals, and implantable biologicals.

We did not receive any public comments on our proposal to expire certain drugs and biologicals from pass-through status, effective December 31, 2009. Therefore, we are finalizing our proposal, without modification, to expire the pass-through status of the six drugs and biologicals listed in Table 30 below, effective December 31, 2009.

TABLE 30—DRUGS AND BIOLOGICALS FOR WHICH PASS-THROUGH STATUS EXPIRES DECEMBER 31, 2009

CY 2009 HCPCS code	CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 SI	Final CY 2010 APC
C9354	C9354	Acellular pericardial tissue matrix of non-human origin (Veritas), per square centimeter.	N	N/A
C9355	C9355	Collagen nerve cuff (NeuroMatrix), per 0.5 centimeter length.	N	N/A
J1300	J1300	Injection, eculizumab, 10 mg	K	9236
J3488	J3488	Injection, zoledronic acid (Reclast), 1 mg	K	0951
J9261	J9261	Injection, nelarabine, 50 mg	K	0825
J9330	J9330	Injection, temsirolimus, 1 mg	K	1168

3. Drugs, Biologicals, and Radiopharmaceuticals With New or Continuing Pass-Through Status in CY 2010

In the CY 2010 OPPI/ASC proposed rule (74 FR 35310), we proposed to continue pass-through status in CY 2010 for 31 drugs and biologicals. These items, which were approved for pass-through status between April 1, 2008 and July 1, 2009, were listed in Table 22 of the proposed rule (74 FR 35310 through 35311). None of these products will have received OPPI pass-through payment for at least 2 years and no more than 3 years by December 31, 2009. The APCs and HCPCS codes for these drugs and biologicals were assigned status indicator “G” in Addenda A and B to the proposed rule.

Section 1833(t)(6)(D)(i) of the Act sets the amount of pass-through payment for pass-through drugs and biologicals (the pass-through payment amount) as the difference between the amount authorized under section 1842(o) of the Act (or, if the drug or biological is covered under a CAP under section 1847B of the Act, an amount determined by the Secretary equal to the average price for the drug or biological for all competitive acquisition areas and the year established under such section as calculated and adjusted by the Secretary) and the portion of the otherwise applicable OPD fee schedule that the Secretary determines is associated with the drug or biological. Payment for drugs and biologicals with pass-through status under the OPPI is currently made at the physician’s office payment rate of ASP+6 percent. We believe it is consistent with the statute to continue to provide payment for drugs and biologicals with pass-through status at a rate of ASP+6 percent in CY 2010, the amount that drugs and biologicals receive under section 1842(o) of the Act. Thus, for CY 2010, we proposed to pay for pass-through drugs and biologicals at ASP+6 percent, equivalent to the rate these drugs and biologicals would receive in the

physician’s office setting in CY 2010. The difference between ASP+4 percent that we proposed to pay for nonpass-through separately payable drugs under the CY 2010 OPPI and ASP+6 percent, therefore, would be the CY 2010 pass-through payment amount for these drugs and biologicals. In the case of pass-through contrast agents, diagnostic radiopharmaceuticals, and implantable biologicals, their pass-through payment amount would be equal to ASP+6 percent because, if not on pass-through status, payment for these products would be packaged into the associated procedures.

As discussed in more detail in section V.B.2.d. of this final rule with comment period, over the last 2 years, we implemented a policy whereby payment for all nonpass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals is packaged into payment for the associated procedure, and we proposed to continue the packaging of these items, regardless of their per day cost, in CY 2010. As stated earlier, pass-through payment is the difference between the amount authorized under section 1842(o) of the Act (or, if the drug or biological is covered under a CAP under section 1847B of the Act, an amount determined by the Secretary equal to the average price for the drug or biological for all competitive acquisition areas and the year established under such section as calculated and adjusted by the Secretary) and the portion of the otherwise applicable OPD fee schedule that the Secretary determines is associated with the drug or biological. Because payment for a drug that is either a diagnostic radiopharmaceutical or a contrast agent (identified as a “policy-packaged” drug, first described in the CY 2009 OPPI/ASC final rule with comment period (73 FR 68639)) or for an implantable biological (which we proposed to consider to be a device for all payment purposes beginning in CY 2010 as discussed in sections V.A.4. and V.B.2.d. of the CY 2010 OPPI/ASC

proposed rule (74 FR 35311 through 35314 and 74 FR 35323 through 35324) and this final rule with comment period) would otherwise be packaged if the product did not have pass-through status, we believe the otherwise applicable OPPI payment amount would be equal to the “policy-packaged” drug or device APC offset amount for the associated clinical APC in which the drug or biological is utilized. The calculation of the “policy-packaged” drug and device APC offset amounts are described in more detail in sections V.A.6.b. and IV.A.2. of this final rule with comment period, respectively. It follows that the copayment for the nonpass-through payment portion (the otherwise applicable fee schedule amount that we would also offset from payment for the drug or biological if a payment offset applies) of the total OPPI payment for those drugs and biologicals would, therefore, be accounted for in the copayment for the associated clinical APC in which the drug or biological is used. According to section 1833(t)(8)(E) of the Act, the amount of copayment associated with pass-through items is equal to the amount of copayment that would be applicable if the pass-through adjustment was not applied. Therefore, beginning in CY 2010, we proposed to set the associated copayment amount for pass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals that would otherwise be packaged if the item did not have pass-through status to zero. The separate OPPI payment to a hospital for the pass-through diagnostic radiopharmaceutical, contrast agent, or implantable biological, after taking into account any applicable payment offset for the item due to the device or “policy-packaged” APC offset policy, is the item’s pass-through payment, which is not subject to a copayment according to the statute. Therefore, we did not publish a copayment amount for these items in Addendum A and B to the proposed rule.

We also proposed to update pass-through payment rates on a quarterly basis on the CMS Web site during CY 2010 if later quarter ASP submissions (or more recent WAC or AWP information, as applicable) indicate that adjustments to the payment rates for these pass-through drugs or biologicals are necessary. If the Part B drug CAP is reinstated during CY 2010, and a drug or biological that has been granted pass-through status for CY 2010 becomes covered under the Part B drug CAP, we proposed to provide pass-through payment at the Part B drug CAP rate and to make the appropriate adjustments to the payment rates for these drugs and biologicals on a quarterly basis as appropriate.

As is our standard methodology, we annually review new permanent HCPCS codes and delete temporary HCPCS C-codes if an alternate permanent HCPCS code is available for purposes of OPSS billing and payment. For our CY 2010 review, we have determined that HCPCS code J2796 (Injection, romiplostim, 10 micrograms) describes the product reported under HCPCS code C9245 (Injection, romiplostim, 10 mcg); HCPCS code A9581 (Injection, gadoxetate disodium, 1 ml) describes the product reported under HCPCS code C9246 (Injection, gadoxetate disodium, per ml); HCPCS code A9582 (Iodine I-123 iobenguane, diagnostic, per study dose, up to 15 millicuries) describes the product reported under HCPCS code C9247 (Iobenguane, I-123, diagnostic, per study dose, up to 10 millicuries); HCPCS code J0718 (Injection, certolizumab pegol, 1 mg) describes the product reported under HCPCS code C9249 (Injection, certolizumab pegol, 1 mg); HCPCS code J0598 (Injection, C1 esterase inhibitor (human), 10 units) describes the product reported under HCPCS code C9251 (Injection, C1 esterase inhibitor (human), 10 units); HCPCS code J2562 (Injection, plerixafor, 1 mg) describes the product reported under HCPCS code C9252 (Injection, plerixafor, 1 mg); and HCPCS code J9328 (Injection, temozolomide, 1 mg) describes the product reported under HCPCS code C9253 (Injection, temozolomide, 1 mg). These new CY 2010 HCPCS codes are included in Table 31 below.

Comment: Several commenters supported CMS' proposal to provide payment at ASP+6 percent for drugs, biologicals, contrast agents, and radiopharmaceuticals that are granted pass-through status. Further, the commenters approved of the proposal to use the ASP methodology that would provide payment based on WAC if ASP information is not available, and

payment at 95 percent of AWP if WAC information is not available. Some commenters requested that CMS provide an additional payment for radiopharmaceuticals that are granted pass-through status because radiopharmaceuticals typically have higher overhead and pharmacy handling costs associated with their preparation than the overhead costs of other drugs and biologicals.

Response: As discussed above, the statutorily mandated pass-through payment equals the amount determined under section 1842(o) of the Act minus the portion of the APC payment that CMS determines is associated with the drug or biological. Therefore, the pass-through payment is determined by subtracting the otherwise applicable payment amount under the OPSS (determined to be ASP+4 percent for CY 2010) from the amount determined under section 1842(o) (ASP+6 percent).

For CY 2010, consistent with our CY 2009 policy for diagnostic radiopharmaceuticals, we proposed to provide payment for both diagnostic and therapeutic radiopharmaceuticals with pass-through status based on the ASP methodology. As stated above, the ASP methodology, as applied under the OPSS, uses several sources of data as a basis for payment, including the ASP, WAC if ASP is unavailable, and AWP if ASP and WAC are unavailable. For purposes of pass-through payment, we consider radiopharmaceuticals to be drugs under the OPSS and, therefore, if a diagnostic or therapeutic radiopharmaceutical receives pass-through status during CY 2010, we proposed to follow the standard ASP methodology to determine its pass-through payment rate under the OPSS. We have routinely provided a single payment for drugs, biologicals, and radiopharmaceuticals under the OPSS to account for the acquisition and pharmacy overhead costs, including compounding costs. We continue to believe that a single payment is appropriate for diagnostic radiopharmaceuticals with pass-through status in CY 2009, and that the payment rate of ASP+6 (or payment based on the ASP methodology) is appropriate to provide payment for both the radiopharmaceutical acquisition cost and any associated nuclear medicine handling and compounding costs. We refer readers to section V.B.5.b. of this final rule with comment period for further discussion of payment for radiopharmaceuticals based on ASP information submitted by manufacturers.

Comment: Several commenters expressed concern that a pass-through

payment period of possibly only 2 years discourages new product development, especially for radiopharmaceutical products. One commenter recommended providing pass-through payment for approved radiopharmaceuticals for a full 3-year time period to allow hospitals time to incorporate new products into their chargemasters and billing practices.

Response: The pass-through statute specifically allows for pass-through payment of drugs and biologicals to be made for at least 2 years, but no more than 3 years. We believe this period of payment facilitates dissemination of these new products into clinical practice and collection of hospital claims data reflective of their costs for future OPSS ratesetting. Our longstanding practice has been to provide pass-through payment for a period of 2 to 3 years, with expiration of pass-through status proposed and finalized through the annual rulemaking process. Each year when proposing to expire the pass-through status of certain drugs and biologicals, we examine our claims data for these products and we have generally seen no evidence that hospitals have not fully incorporated these items into their chargemasters based on the utilization and costs observed in our claims data. As discussed further in section V.A.5. of this final rule with comment period, we are making no operational changes to the drug and biological pass-through program for CY 2010 and plan to continue to expire pass-through status on an annual basis through rulemaking. Under this existing operational policy, which was generally supported by the commenters, because we begin pass-through payment on a quarterly basis that depends on when applications are submitted to us for consideration and we expire pass-through status only on an annual basis, there is no way to ensure that all pass-through drugs and biologicals receive pass-through payment for a full 3 years, while also providing pass-through payment for no more than 3 years as the statute requires. Therefore, we will continue to provide drug and biological pass-through payment for at least 2 years, but no more than 3 years, as required by the statute. We continue to receive numerous pass-through applications for drugs and biologicals for consideration each quarter, and we have no evidence that our current pass-through payment policies discourage new product development.

There is currently one diagnostic radiopharmaceutical, HCPCS code C9247 (Iodine I-123 iobenguane, diagnostic, per study dose, up to 15

millicuries), that has been granted pass-through status at the time of this final rule with comment period. We proposed to continue pass-through status for this diagnostic radiopharmaceutical as it would not have received at least 2 but not more than 3 years of pass-through payment by December 31, 2009. This is consistent with the OPPS provision that provides for at least 2 but not more than 3 years of pass-through payment for drugs and biologicals that are approved for pass-through payments.

We provide an opportunity through the annual OPPS/ASC rulemaking cycle for public comment on those drugs and biologicals that are proposed for expiration of pass-through payment at the end of the next calendar year. We have often received public comments related to our proposed expiration of pass-through status for particular drugs and biologicals, and we expect to continue to receive public comments regarding the proposed expiration of pass-through status for drugs and biologicals in the future. In this manner, we would address specific concerns about the pass-through payment period for individual drugs and biologicals in the future, including radiopharmaceuticals.

Comment: A few commenters supported the CY 2010 proposal to set the associated copayment amounts for pass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals that would otherwise be packaged if the product did not have pass-through status to zero. The commenters noted increased

beneficiary savings by setting the copayment amount to zero.

Response: We appreciate the commenters' support. As discussed in the CY 2010 OPPS/ASC proposed rule (74 FR 35311), we believe that for drugs and biologicals that are "policy-packaged," the copayment for the nonpass-through payment portion of the total OPPS payment for this subset of drugs and biologicals is accounted for in the copayment for the associated clinical APC in which the drug or biological is used. According to section 1833(t)(8)(E) of the Act, the amount of copayment associated with pass-through items is equal to the amount of copayment that would be applicable if the pass-through adjustment was not applied. Therefore, it is our belief that the amount should be zero for drugs and biologicals that are "policy-packaged," including diagnostic radiopharmaceuticals.

We did not receive any public comments on our proposal to update pass-through payment rates on a quarterly basis during CY 2010 if later quarter ASP submissions (or more recent WAC or AWP information, as applicable) indicate that adjustments to the payment rates for these pass-through drugs and biologicals are necessary.

After consideration of the public comments we received, we are finalizing our CY 2010 pass-through payment proposals, without modification. Specifically, we will provide pass-through payment in CY 2010 for those drugs, biologicals and radiopharmaceuticals listed in Table 31

below. Pass-through payment for drugs, biologicals, and radiopharmaceuticals granted pass-through status will be made at the payment rate indicated in section 1842(o) of the Act, that is, ASP+6 percent. If ASP data are not available, pass-through payment will be based on the OPPS ASP methodology—that is, payment at WAC+6 percent if ASP data are not available and payment at 95 percent of the pass-through radiopharmaceutical's most recent AWP if WAC information is not available. We will update pass-through payment rates on a quarterly basis during CY 2010 if later ASP submissions (or more recent WAC or AWP information, as applicable) indicate that adjustments to the payment rates for pass-through drugs and biologicals are necessary. We will set the associated copayment amount for pass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals that would otherwise be packaged if the item did not have pass-through status to zero. Finally, if a drug or biological that has been granted pass-through status for CY 2010 becomes covered under the Part B drug CAP if the program is reinstated, we will provide payment for Part B drugs that are granted pass-through status and are covered under the Part B drug CAP at the Part B drug CAP rate.

The drugs and biologicals that are continuing pass-through status for CY 2010 or that have been granted pass-through status as of January 2010 are displayed in Table 31 below.

TABLE 31—DRUGS AND BIOLOGICALS WITH PASS-THROUGH STATUS IN CY 2010

CY 2009 HCPCS code	CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 SI	Final CY 2010 APC
C9245	J2796	Injection, romiplostim, 10 micrograms	G	9245
C9246	A9581	Injection, gadoxetate disodium, 1 ml	G	9246
C9247	A9582	Iodine I-123 iobenguane, diagnostic, per study dose, up to 15 millicuries.	G	9247
	A9583	Injection, gadofosveset trisodium, 1 ml	G	1299
C9248	C9248	Injection, clevidipien butyrate, 1 mg	G	9248
C9249	J0718	Injection, certolizumab pegol, 1 mg	G	9249
C9250	C9250	Human plasma fibrin sealant, vapor-heated, solvent-detergent (Artiss), 2ml.	G	9250
C9251	J0598	Injection, C1 esterase inhibitor (human), 10 units	G	9251
C9252	J2562	Injection, plerixafor, 1 mg	G	9252
C9253	J9328	Injection, temozolomide, 1 mg	G	9253
	C9255	Injection, paliperidone palmitate, 1 mg	G	1300
	C9256	Injection, dexamethasone intravitreal implant, 0.1 mg	G	9256
C9356	C9356	Tendon, porous matrix of cross-linked collagen and glycosaminoglycan matrix (TenoGlide Tendon Protector Sheet), per square centimeter.	G	9356
C9358	C9358	Dermal substitute, native, non-denatured collagen, fetal bovine origin (SurgiMend Collagen Matrix), per 0.5 square centimeters.	G	9358
C9359	C9359	Porous purified collagen matrix bone void filler (Integra Mozaik Osteoconductive Scaffold Putty, Integra OS Osteoconductive Scaffold Putty), per 0.5 cc.	G	9359

TABLE 31—DRUGS AND BIOLOGICALS WITH PASS-THROUGH STATUS IN CY 2010—Continued

CY 2009 HCPCS code	CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 SI	Final CY 2010 APC
C9360	C9360	Dermal substitute, native, non-denatured collagen, neonatal bovine origin (SurgiMend Collagen Matrix), per 0.5 square centimeters.	G	9360
C9361	C9361	Collagen matrix nerve wrap (NeuroMend Collagen Nerve Wrap), per 0.5 centimeter length.	G	9361
C9362	C9362	Porous purified collagen matrix bone void filler (Integra Mozaik Osteoconductive Scaffold Strip), per 0.5 cc.	G	9362
C9363	C9363	Skin substitute, Integra Meshed Bilayer Wound Matrix, per square centimeter.	G	9363
C9364	C9364	Porcine implant, Permacol, per square centimeter	G	9364
J0641	J0641	Injection, levoleucovorin calcium, 0.5 mg	G	1236
J1267	J1267	Injection, doripenem, 10 mg	G	9241
J1453	J1453	Injection, fosaprepitant, 1 mg	G	9242
J1459	J1459	Injection, immune globulin (privigen), intravenous, non-lyophilized (e.g. liquid), 500 mg.	G	1214
J1571	J1571	Injection, hepatitis b immune globulin (hepagam b), intramuscular, 0.5 ml.	G	0946
J1573	J1573	Injection, hepatitis B immune globulin (Hepagam B), intravenous, 0.5ml.	G	1138
J1953	J1680	Injection, human fibrinogen concentrate, 100 mg	G	1290
J2785	J1953	Injection, levetiracetam, 10 mg	G	9238
J8705	J2785	Injection, regadenoson, 0.1 mg	G	9244
J9033	J8705	Topotecan, oral, 0.25 mg	G	1238
J9207	J9033	Injection, bendamustine hcl, 1 mg	G	9243
J9225	J9155	Injection, degarelix, 1 mg	G	1296
J9226	J9207	Injection, ixabepilone, 1 mg	G	9240
	J9225	Histrelin implant (vantas), 50 mg	G	1711
	J9226	Histrelin implant (supprelin la), 50 mg	G	1142
	Q0138	Injection, ferumoxylol, for treatment of iron deficiency anemia, 1 mg (non-esrd use).	G	1297
Q4114	Q4114	Dermal substitute, granulated cross-linked collagen and glycosaminoglycan matrix (Flowable Wound Matrix), 1 cc.	G	1251

4. Pass-Through Payment for Implantable Biologicals

a. Background

Section 1833(t)(6)(A)(iv) of the Act authorizes transitional pass-through payments for new medical devices, drugs, and biologicals, for those items where payment was not being made as a hospital outpatient service under Part B as of December 31, 1996, and whose cost is not insignificant in relation to the OPD fee schedule amount payable for the service (or group of services) involved. These pass-through payments are in addition to the usual APC payments for services in which the product is used. Coding and payment for drugs and biologicals with pass-through status are generally provided on a product-specific basis for a period of no less than 2 and no more than 3 years from the date pass-through payment is first made as discussed in section V.A.5. of this final rule with comment period, while coding and payment for devices with pass-through status are provided for categories of devices that may describe numerous products. The Act specifies that the duration of transitional pass-through payments for

devices must be no less than 2 and no more than 3 years from the first date on which payment is made for any medical device that is described by the category. Therefore, we utilize separate pass-through application and evaluation processes and criteria for drugs and biologicals and device categories because the statutory provisions are not the same for all items that may receive pass-through payment. These processes and the applicable evaluation criteria are available on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/04_passthrough_payment.asp#TopOfPage. The regulations that govern pass-through payment for drugs and biologicals are found in § 419.64 and those applicable to pass-through device categories are found in § 419.66.

Section 1833(t)(6)(D)(i) of the Act specifies that the pass-through payment amount, in the case of a drug or biological, is the amount by which the amount determined under section 1842(o) of the Act (or, if the drug or biological is covered under a competitive acquisition contract under section 1847B of the Act, an amount determined by the Secretary equal to the

average price for the drug or biological for all competitive acquisition areas and the year established under such section as calculated and adjusted by the Secretary) for the drug or biological exceeds the portion of the otherwise applicable Medicare OPD fee schedule that the Secretary determines is associated with the drug or biological. For the drugs and biologicals that would have otherwise been paid under the Part B drug CAP, because the Part B drug CAP has been suspended beginning January 1, 2009, pass-through payment for these drugs and biologicals is currently made at the physician's office payment rate of ASP+6 percent. In the case of diagnostic radiopharmaceuticals, where all products without pass-through status are packaged into payment for nuclear medicine procedures, the pass-through payment is reduced by an amount that reflects the diagnostic radiopharmaceutical portion of the APC payment amount for the associated nuclear medicine procedure (the "policy-packaged" drug APC offset) that we determine is associated with the cost of predecessor diagnostic radiopharmaceuticals. In the CY 2010 OPPS/ASC proposed rule (74 FR 35318),

we proposed a similar payment offset policy for contrast agents beginning in CY 2010, as discussed in section V.A.6.c. of the proposed rule, and we are finalizing this policy for CY 2010, as discussed in section V.A.6.c. of this final rule with comment period. Pass-through payment for a category of devices is made at the hospital's charge for the device, adjusted to cost by application of the hospital's CCR. If applicable, the device payment is reduced by an amount that reflects the portion of the APC payment amount for the associated surgical procedure that we determine is associated with the cost of the device, called the device APC offset and discussed further in section IV.A.2. of the proposed rule (74 FR 35306) and this final rule with comment period.

In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68633 through 68636), we finalized a policy to package payment for implantable biologicals without pass-through status that are surgically inserted or implanted (through a surgical incision or a natural orifice) into payment for the associated surgical procedure. Prior to our implementation of this policy for nonpass-through implantable biologicals, we adopted in the CY 2003 OPSS final rule with comment period (67 FR 66763) the current OPSS policy that packages payment for an implantable device into the associated surgical procedures when its pass-through payment period ends because payment for all implantable devices without pass-through status under the OPSS is packaged. We consider nonpass-through implantable devices to be integral and supportive items for which packaged payment is most appropriate. As we stated in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68634), we believe this policy to package payment for implantable devices that are integral to the performance of procedures paid separately through an APC payment should also apply to payment for implantable biologicals without pass-through status, when those biologicals function as implantable devices. Implantable biologicals may be used in place of other implantable nonbiological devices whose costs are already accounted for in the associated procedural APC payments for surgical procedures. We reasoned that if we were to provide separate payment for nonpass-through implantable biologicals, we would potentially be providing duplicate device payment, both through the packaged nonbiological device cost included in

the surgical procedure's payment and the separate biological payment.

In the CY 2009 OPSS/ASC final rule with comment period (73 FR 68634), we stated our belief that the three implantable biologicals with expiring pass-through status for CY 2009 differ from other biologicals paid under the OPSS in that they specifically always function as surgically implanted devices. We noted that both implantable nonbiological devices under the OPSS and the three biologicals with expiring pass-through status in CY 2009 are surgically inserted or implanted (including through a surgical incision or a natural orifice). These three biologicals are approved by the FDA as devices, and they are solely surgically implanted according to their FDA-approved indications. Furthermore, in some cases, these implantable biologicals can substitute for implantable nonbiological devices (such as for synthetic nerve conduits or synthetic mesh used in tendon repair).

For other nonpass-through biologicals paid under the OPSS that may sometimes be used as implantable devices, we have instructed hospitals, beginning via Transmittal 1336, Change Request 5718, dated September 14, 2007, to not separately bill the HCPCS codes for the products when using these items as implantable devices (including as a scaffold or an alternative to human or nonhuman connective tissue or mesh used in a graft) during surgical procedures. In such cases, we consider payment for the biological used as an implantable device in a specific clinical case to be included in payment for the surgical procedure. We stated that hospitals may include the charge for the biological in their charge for the procedure, report the charge on an uncoded revenue center line, or report the charge under a device HCPCS code, if one exists, so that the biological costs may be considered in future ratesetting for the associated surgical procedures.

Several commenters who responded to the CY 2009 OPSS/ASC proposed rule supported CMS' proposal to package payment for implantable biologicals without pass-through status into payment for the associated surgical procedure (73 FR 68635). One commenter also recommended that CMS treat biologicals that are always surgically implanted or inserted and have FDA device approval as devices for purposes of pass-through payment, rather than as drugs. The commenter observed that this would allow all implantable devices, biological and otherwise, to be subject to a single pass-through payment policy. The commenter concluded that this policy

change would provide consistency in billing and payment for these products functioning as implantable devices during their pass-through payment period, as well as after the expiration of pass-through status.

We finalized in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68635) our proposal to package payment for any nonpass-through biological that is surgically inserted or implanted (through a surgical incision or a natural orifice) into the payment for the associated surgical procedure, just as we package payment for all nonpass-through, implantable, nonbiological devices. As a result of this final policy, the three implantable biologicals with expiring pass-through status in CY 2009 were packaged and assigned status indicator "N" as of January 1, 2009. In addition, any new biologicals without pass-through status that are surgically inserted or implanted (through a surgical incision or a natural orifice) are also packaged beginning in CY 2009. Hospitals continue to report the HCPCS codes that describe biologicals that are always used as implantable devices on their claims, and we package the costs of those biologicals into the associated procedures, according to the standard OPSS ratesetting methodology that is described in section II.A.2. of the CY 2010 OPSS/ASC proposed rule (74 FR 35254 through 35267) and this final rule with comment period. Moreover, for nonpass-through biologicals that may sometimes be used as implantable devices, we continue to instruct hospitals to not bill separately for the HCPCS codes for the products when used as implantable devices. This reporting ensures that the costs of these products that may be, but are not always, used as implanted biologicals are appropriately packaged into payment for the associated implantation procedures when the products are used as implantable devices.

b. Policy for CY 2010

Some implantable biologicals are described by device category codes for expired pass-through categories, including HCPCS code C1781 (Mesh (implantable)), HCPCS code C1762 (Connective tissue, human), and HCPCS code C1763 (Connective tissue, non-human). All implantable devices described by the latter two categories are biologicals, while HCPCS code C1781 describes both implantable biological and nonbiological devices. Historically, these category codes included biological products that we approved for pass-through payment under the device pass-through process, initially when we paid for pass-through

devices on a brand-specific basis from CY 2000 through March 31, 2001, and later through the device categories described by HCPCS codes C1781, C1762, and C1763, which were developed effective April 1, 2001.

We believe that it is most appropriate for a product to be eligible for a single period of OPPS pass-through payment, rather than a period of device pass-through payment and a period of drug or biological pass-through payment. The limited timeframe for transitional pass-through payment ensures that new devices, drugs, and biologicals may receive special payment consideration under the OPPS for the first few years after their initial use, in order to allow sufficient time for their cost information to be reflected in hospital claims data and, therefore, to be available for OPPS ratesetting. After the pass-through payment period ends, like other existing services, we have cost information regarding these new products provided to us by hospitals from claims and cost report data. We then utilize that information when packaging the costs of the items (all devices, diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals, and other drugs with an estimated per day cost equal to or less than the annual drug packaging threshold) or paying separately for the products (drugs except contrast agents and diagnostic radiopharmaceuticals and also nonimplantable biologicals with estimated per day costs above the annual drug packaging threshold). Further, although implantable biologicals with pass-through status may substitute for nonpass-through implantable devices whose costs are packaged into procedural APC payments, our existing APC offset policies for the costs of predecessor items packaged into APC payment for the associated services do not apply to pass-through payment for biologicals. We note that the APC offset amount that would be most applicable to implantable biologicals, if we determine that an offset applies for a given APC, would be the device APC offset amount, based on their similarity of function to the implantable devices whose costs have been included in establishing the procedural APC payment, not the "policy-packaged" or "threshold-packaged" drug APC offset amounts that one would expect to apply to pass-through drugs and biologicals. Similarly, when we currently evaluate a pass-through implantable biological application for the cost significance of the product, our methodology utilizes the "policy-packaged" APC offset

amount to assess the candidate implantable biological, not the device APC offset amount that would be more reflective of the costs of predecessor devices related to the candidate implantable biological, such as those of device category HCPCS codes C1781, C1762, and C1763.

Many implantable biologicals, such as the three biologicals that expired from pass-through status after CY 2008, have FDA approval as devices. A number of other implantable biologicals with FDA approval as devices also have been approved for OPPS pass-through payment over the past several years, based on their product-specific pass-through applications as biologicals, not devices. Moreover, outside of the period of pass-through payment, the costs of implantable biologicals, like the costs of implantable devices, are now packaged into the cost of the procedure in which they are used. Implantable biologicals may be used in place of other implantable nonbiological devices whose costs are already accounted for in the associated procedural APC payments. Payment is made for nonpass-through implantable biologicals, like for devices, through the APC payment for the associated surgical procedure.

In view of these considerations, in the CY 2010 OPPS/ASC proposed rule (74 FR 35313), we proposed that the pass-through evaluation process and pass-through payment methodology for implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) and that are newly approved for pass-through status beginning on or after January 1, 2010, be the device pass-through process and payment methodology only. Given the shared payment methodologies for implantable biological and nonbiological devices during their nonpass-through payment periods, as well as their overlapping and sometimes identical clinical uses and their similar regulation by the FDA as devices, we believe that the most consistent pass-through payment policy for these different types of items that are surgically inserted or implanted and that may sometimes substitute for one another is to evaluate all such devices, both biological and nonbiological, only under the device pass-through process. As a result, implantable biologicals would no longer be eligible to submit biological pass-through applications and to receive biological pass-through payment at ASP+6 percent. While we understand that implantable biologicals have characteristics that result in their meeting the definitions of both devices and biologicals, we believe that

implantable biologicals are most similar to devices because of their required surgical insertion or implantation and that it would be appropriate to only evaluate them as devices because they share significant clinical similarity with implantable nonbiological devices. We refer readers to the CMS Web site specified previously in this section to view the device pass-through application requirements and review criteria that would apply to the evaluation of all implantable biologicals for pass-through status when their pass-through payment would begin on or after January 1, 2010.

However, those implantable biologicals that are surgically inserted or implanted (through a surgical incision or natural orifice) and that are receiving pass-through payment as biologicals prior to January 1, 2010, would continue to be considered pass-through biologicals for the duration of their period of pass-through payment. These products have already been evaluated for pass-through status based on their applications as biologicals and have been approved for pass-through status based on the established criteria for biological pass-through payment. We believe it would be most appropriate for them to complete their 2- to 3-year period of pass-through payment as biologicals in accordance with the pass-through payment policies that were applicable at the time their pass-through status was initially approved.

We note that, in conducting our pass-through review of implantable biologicals as devices beginning with CY 2010 pass-through payment, we would apply the portions of APC payment amounts associated with devices (that is, the device APC offset amounts) to assess the cost significance of the candidate implantable biologicals, as we do for other devices. The CY 2009 device APC offset amounts are posted on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/04_passthrough_payment.asp. The result of evaluating all implantable biological items only for device pass-through payment is that payment for implantable biologicals eligible for pass-through payment beginning on or after January 1, 2010, would be based on hospital charges adjusted to cost, rather than the ASP methodology that is applicable to pass-through drugs and biologicals. Treating implantable biologicals as devices for evaluation of pass-through payment eligibility and payment would result in their consistent treatment with respect to coding and payment during their pass-through and nonpass-through periods of

payment. This proposed policy would allow us to appropriately offset the pass-through payment for an implantable biological using the device APC offset amounts, which would incorporate the costs of predecessor devices (both biological and nonbiological) that are similar to the implantable biological item with pass-through status. Finally, this proposed policy would ensure that each implantable biological is eligible for OPPS pass-through payment for only one 2- to 3-year time period (as a device only, not as a biological), so that once OPPS claims data incorporate cost information for the implantable biological, the product would not be again eligible for OPPS pass-through payment in the future.

Further, because we proposed that the pass-through evaluation process for CY 2010 pass-through status approvals and pass-through payment methodology for implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) beginning in CY 2010 be the device pass-through process and payment methodology only, we also proposed to revise our regulations at §§ 419.64 and 419.66 to conform to this new policy. Specifically, we proposed to amend § 419.64 by adding a new paragraph (a)(4)(iii) and language under a new paragraph (c)(3) to exclude implantable biologicals from consideration for drug and biological pass-through payment. Furthermore, under proposed new paragraph (a)(4)(iv) of § 419.64, we proposed to specify the continued inclusion of implantable biologicals for which pass-through payment as a biological is made on or before December 31, 2009, as eligible for biological pass-through payment, consistent with our proposal to allow these products to complete their period of pass-through payment as biologicals.

Moreover, in light of our CY 2010 proposal that implantable biological applications for pass-through status beginning on or after January 1, 2010, would be considered only for device pass-through evaluation and payment, we stated in the proposed rule (74 FR 35314) that we believe it would also be appropriate to clarify the current example in § 419.66(b)(4)(iii) of the regulations regarding the exclusion of materials, for example, biological or synthetic materials, that may be used to replace human skin from device pass-through payment eligibility. While, by definition, implantable biologicals that are surgically implanted or inserted would not be biological materials that replace human skin, we proposed to more precisely state this in the regulations. Therefore, we proposed to

revise § 419.66(b)(4)(iii), which currently states that a device is not a material that may be used to replace human skin and provides an example of such a material as “a biological or synthetic material.” We proposed to revise § 419.66(b)(4)(iii) to specify that the biological materials be a “biological skin replacement material” rather than a “biological” and the synthetic materials be a “synthetic skin replacement material” rather than a “synthetic material” because we do not believe this example should refer to biologicals or synthetic materials that are used for purposes other than as a skin replacement material, given that the regulatory provision in § 419.66(b)(4)(iii) applies only to a material that may be used to replace human skin.

Comment: A few commenters requested that CMS continue to pay for all pass-through biologicals under the ASP methodology for drugs and nonimplantable biologicals, and not pay for new implantable biologicals eligible for pass-through payment based on charges adjusted to cost. One commenter believed that the ASP methodology is well understood by hospitals and Medicare contractors and asserted that some new implantable biologicals under development will cost several thousand dollars per procedure. Therefore, the commenter stated, many hospitals will be reluctant to mark up charges for these new implantable biologicals, thereby resulting in charge compression and an underestimate of the costs of biologicals. Furthermore, the commenter claimed that continued payment for pass-through implantable biologicals based on the ASP methodology would ensure consistent payment for new biologicals rather than variable payment based on hospitals' charging practices.

Response: Under our CY 2010 proposal to evaluate and pay for implantable biologicals under the device pass-through methodology, we would use the charges adjusted to cost payment methodology and apply a reduction to payment (that is, the device APC offset) for implantable biologicals eligible for pass-through payment beginning on or after January 1, 2010. Regarding the commenters' request that we continue the ASP payment methodology for pass-through implantable biologicals, we do not agree that payment under this methodology would be appropriate. Payment based on ASP for pass-through implantable biologicals would not provide the similar OPPS payment treatment of biological and nonbiological implantable devices that is our goal for

new devices. Given the shared payment methodologies for implantable biological and nonbiological devices during their nonpass-through payment periods, as well as their overlapping and sometimes identical clinical uses and their generally similar regulation by the FDA as devices, we believe that the most consistent pass-through payment policy for these different types of items that are surgically inserted or implanted and that may sometimes substitute for one another is to evaluate and pay for all such devices, both biological and nonbiological, only under the device pass-through process and payment methodology. As we stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35313), we believe that implantable biologicals are most similar to devices because of their required surgical insertion or implantation and that it would be appropriate to only evaluate them as devices because they share significant clinical similarity with implantable nonbiological devices. We note that we will continue pass-through payment under the ASP methodology for any implantable biological for which pass-through payment as a biological begins on or before December 31, 2009.

Comment: A few commenters supported the proposal to treat implantable biologicals and implanted devices the same regarding the pass-through eligibility criteria and payment methodology. Some commenters stated that payment for both implantable biological and nonbiological devices should be made on the same basis for items with both pass-through and nonpass-through status. One commenter asserted that the proposed treatment of implantable biologicals is consistent with CMS' policy to package the costs of implantable devices and would reinforce previous CMS instructions regarding the billing of biologicals when used as implanted devices.

Furthermore, another commenter also agreed with CMS' policy that separately payable HCPCS codes not be reported when biologicals that are sometimes implanted are surgically inserted during a procedure. The commenter urged CMS to continue educating providers about when HCPCS codes that describe biologicals that are sometimes implanted should be reported, including publishing a list of procedures with which the HCPCS codes for implantable biologicals would not typically be reported. The commenter encouraged CMS to publish “reverse” device-to-procedure edits for such procedures.

Response: We appreciate the commenters' support for our proposal. We agree that payment for both implantable biological and

nonbiological devices that may be substitutes for one another should be made on the same basis for items with both pass-through and nonpass-through status, that is, based on charges adjusted to cost while on pass-through status and packaged when not on pass-through status. Concerning the suggestion to publish a list of procedure codes with which the HCPCS codes for biologicals that are implanted would not typically be reported, we believe that creating and maintaining such a list would not be feasible because implantable biologicals may be used in a wide variety of surgical procedures. Moreover, creating and maintaining device-to-procedure edits for implantable biologicals also would not be feasible, given the broad array of surgical procedures in which such biologicals may be implanted.

Comment: One commenter requested that CMS delay the CY 2010 proposal to include implantable biologicals in the calculation of the device APC offset amounts. The commenter also recommended that CMS grandfather all implantable biological applications submitted under the drug and biological pass-through application process prior to the September 1, 2009 application filing deadline. The commenter noted that implantable biological applications submitted prior to September 1, 2009, could have received biological pass-through status if CMS had not proposed and finalized the policy to treat them as devices for pass-through purposes, beginning in CY 2010.

The commenter explained that two implantable biological products that are competitors to the product manufactured by the commenter currently have pass-through status as biologicals, and their pass-through status is proposed to continue for CY 2010. The commenter believed that treating implantable biologicals differently based on the date of their pass-through application would result in a competitive disadvantage for the product manufactured by the commenter.

Response: The commenter recommended delaying the packaging of implantable biologicals in calculating the device offset. As a practical matter, the packaging of nonpass-through implantable biologicals was proposed and finalized for CY 2009 (73 FR 68635) and was implemented beginning in CY 2009. Given our proposal to treat implantable biologicals as devices for pass-through purposes beginning in CY 2010 and our longstanding device APC offset policy for pass-through devices, we believe it is appropriate to consider the costs of implantable biologicals that are packaged in establishing the device

APC offset amounts under a policy that considers implantable biologicals to be devices for pass-through evaluation and payment purposes. We rely on the device APC offset amounts to account for the costs of all predecessor devices to a new device category when those predecessor devices are implanted in procedures assigned to an APC to which procedures associated with the new device category would be assigned, and the predecessor devices may now include implantable biologicals.

Concerning the commenter's request to grandfather all implantable biological applications submitted under the drug and biological pass-through application process prior to the September 1, 2009 application filing deadline, we believe it is important to adopt a consistent implantable biological pass-through policy for a full calendar year to provide appropriate payment under a single payment policy for that year and allow consistent use of our CY 2010 claims data for ratesetting in the future. The earliest an application filed for the September 1 deadline (applications are received and processed on a continual basis) could be considered for pass-through status is January 1 of the following year, in this case, CY 2010, as we have established and posted on the CMS Web site for pass-through applications at: http://www.cms.hhs.gov/HospitalOutpatientPPS/04_passthrough_payment.asp#TopOfPage. We do not believe it would be appropriate to implement pass-through evaluation and payment of implantable biologicals as devices later than the quarter beginning January 1, 2010. In order to meet the timeframes required by our claims processing systems, applications for drug and biological pass-through status received by the September 1, 2009 deadline for January 2010 payment have been evaluated based on the policy established in this final rule with comment period to evaluate implantable biologicals for device pass-through payment. We also note that when adopting any significant policy change under the OPSS with a specific effective date, we recognize that similar products or services may be treated differently because of the timing of their FDA approval, pass-through application submission, or other characteristics. Nevertheless, the rulemaking process provides significant opportunity for public notice and comment prior to such policy changes in order to ensure that we give full consideration to all issues and information related to proposals of new policy.

Comment: Several commenters recommended that both implantable

and nonimplantable biologicals approved by the FDA under a biologics license application (BLA) be evaluated for pass-through payment status under the drug pass-through evaluation process, and indicated their belief that Congress intended biologicals approved under BLAs to be paid under the specific OPSS statutory provisions that apply to specified covered outpatient drugs (SCODs), including the pass-through provisions. One commenter agreed that CMS should have similar payment methodologies for biological, nonbiological, and composite devices for fairness and consistency and recommended that CMS implement the proposed policy based on FDA approval status, specifically treating as devices for pass-through purposes only those implantable biologicals approved by the FDA as devices. The commenter claimed that CMS determined that several implantable devices that are currently treated as drugs or biologicals must be paid based on their product-specific ASP submissions because the requirement for combining drugs for the purpose of ASP is that the reference materials report them as clinical equivalents. The commenter reasoned that devices do not have equivalents identified in reference materials; therefore, those devices paid as drugs must always receive separate payment. The commenter also requested that CMS clarify when it will treat an implantable device as a biological for ASP payment.

One commenter suggested that CMS not use the device pass-through process for evaluating drugs or biologicals that are implanted using a device as merely a delivery vehicle, simply because the drug is administered through a device. The commenter recommended that CMS base its pass-through payment decision on the identity of the component that exerts the therapeutic effect of the combined product, either the biological component or the delivery vehicle, and provided as an example the practice of FDA's Office of Combination Products to assess combination products in development and assign their FDA regulation based on which component exerts the therapeutic effect claimed by the manufacturer. The commenter believed that there are clinical problems with using implantation to define whether a biological should be treated as a device because, for some drugs, implantation may always be the clinically superior route of administration. Another commenter claimed that some implantable biologicals meet the Act's definition of a biological under section 1861(t)(1) of

the Act even though they are approved by the FDA as devices.

Response: We proposed to evaluate implantable biologicals that function as and are substitutes for implantable devices, regardless of their category of FDA approval, as devices for OPPS payment purposes. We do not believe it is necessary to make our OPPS payment policies regarding implantable biologicals dependent on categories of FDA approval, the intent of which is to ensure the safety and effectiveness of medical products.

We do not agree with the commenters who asserted that Congress intended biologicals approved under BLAs to be paid under the specific OPPS statutory provisions that apply to SCODs, including the pass-through provisions. Moreover, Congress did not specify that we must pay for implantable biologicals as biologicals rather than devices, if they also meet our criteria for payment as a device. We believe that implantable biologicals meet the definitions of a device and a biological and that, for payment purposes, it is appropriate for us to consider implantable biologicals as implantable devices in all cases, not as biologicals. For example, beginning in CY 2009, we package the costs of implantable biologicals into the costs of the procedures in which they are used, as we do for implantable devices. Therefore, we do not believe that we must pay for implantable biologicals under our OPPS biological payment methodologies, rather than our device payment methodologies. Furthermore, because we consider implantable biologicals to be devices for payment purposes, any interpretation that a biological is unique in the context of the ASP payment methodology for biologicals would not apply. Thus, we disagree with the commenter's conclusion that implantable biologicals treated as devices must receive separate payment because devices do not have equivalents in reference materials, a concept applicable only to the requirements for combining biologicals for payment under the ASP methodology, because we consider these implantable biologicals to be devices under the OPPS, to which packaged payment outside of the pass-through payment period applies.

It is not our intention to consider biologicals under the device pass-through evaluation process and payment methodology when these products are merely administered through the implantation of a delivery system for the biological. Each implantable biological pass-through application for a combination product would be initially evaluated in such a

case to determine if the biological or device is the key therapeutic or diagnostic component, after which we would then determine whether to evaluate the item under the device or drug and biological pass-through process. If the key component of the candidate pass-through product is the biological and that biological is only implanted because it is administered through an implanted delivery system for the biological (that is, the biological itself is not functioning as an implantable device), we would evaluate the product under the drug and biological pass-through process. Conversely, if the key component of the candidate pass-through product is the biological and that biological is functioning as an implantable device or the key component of the product is the implantable delivery system for the biological, we would evaluate the product under the device pass-through process.

As we stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35313) and this final rule with comment period, while we understand that implantable biologicals have characteristics that result in their meeting the definitions of both devices and biologicals, we believe that biologicals are most similar to devices because of their required surgical insertion or implantation and that it would be appropriate to only evaluate them as devices because they share significant clinical similarity with implantable nonbiological devices. We do not believe that those implantable biologicals that meet the Act's definition of biological under section 1861(t)(1) necessarily must be evaluated and paid for under the OPPS drug and biological pass-through payment methodology, when they also meet the definition of a device for purposes of pass-through evaluation and payment.

Comment: One commenter requested that CMS clarify certain points regarding the proposal to evaluate and pay for implantable biologicals with pass-through status similarly to pass-through devices. The commenter requested that CMS designate that the types of biologicals that would be affected by the proposal would be connective tissue replacements that function as devices. The commenter also requested that CMS clarify that the proposed changes would apply to pass-through implantable biologicals and not to implantable drugs, and that CMS recognize that it would be inappropriate to treat implantable drugs as devices for pass-through purposes in the future.

Response: Our CY 2010 proposal was not limited to implantable biological connective tissue replacements, but

instead it applies to all implantable biologicals. For example, in the proposed rule (74 FR 35313), we cited expired device category HCPCS code C1781 (Mesh (implantable)) as describing implantable biologicals as well as implantable nonbiological devices, yet mesh need not necessarily function as a connective tissue replacement. We did not propose to treat implantable drugs as devices and, therefore, would not treat implantable drugs as devices for pass-through payment program purposes in CY 2010.

Comment: One commenter suggested that implantable biologicals should not be treated as devices, and observed that stakeholders have not had adequate time to consider the long-term implications of the CMS proposal. The commenter recommended that CMS not finalize the proposal at this time and hold a public meeting regarding the proposal.

Response: We believe that all stakeholders have had sufficient time to consider this proposal through the routine notice and comment rulemaking process. We received numerous public comments on our CY 2010 proposal and, while we are always open to meeting with stakeholders who would like to share their views with us, we do not believe a public meeting on this issue is needed.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, that the pass-through evaluation process and payment methodology for implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) and that are newly approved for pass-through status beginning on or after January 1, 2010, be the device pass-through process and payment methodology only. However, those implantable biologicals that are surgically inserted or implanted (through a surgical incision or natural orifice) and that are receiving pass-through payment as biologicals prior to January 1, 2010, would continue to be considered pass-through biologicals for the duration of their period of pass-through payment. As proposed, in conducting our pass-through review of implantable biologicals as devices beginning with CY 2010 pass-through payment, we will apply the portions of APC payment amounts associated with devices (that is, the device APC offset amounts) to assess the cost significance of the candidate implantable biologicals, as we do for other devices. Furthermore, we are finalizing our proposal to revise our regulations at §§ 419.64 and 419.66 to conform to this new policy.

Specifically, we are finalizing our proposal to amend § 419.64 by adding a new paragraph (a)(4)(iii) to exclude implantable biologicals from consideration for drug and biological pass-through payment. However, we note that, as discussed in section V.A.5. of this final rule with comment period, we are not finalizing our proposed addition of a new paragraph (c)(3) to § 419.64 and, therefore, we are not adopting our related proposed change to proposed paragraph (c)(3) that would have excluded implantable biologicals from consideration for drug and biological pass-through payment. Furthermore, we are adopting our proposed new paragraph (a)(4)(iv) of § 419.64, which specifies the continued inclusion of implantable biologicals for which pass-through payment as a biological is made on or before December 31, 2009, as eligible for biological pass-through payment. Finally, we are adopting our proposal stated above that clarifies the current example in § 419.66(b)(4)(iii) of the regulations regarding the exclusion of materials, for example, biological or synthetic materials, that may be used to replace human skin from device pass-through payment eligibility.

5. Definition of Pass-Through Payment Eligibility Period for New Drugs and Biologicals

Section 1833(t)(6) of the Act provides for transitional pass-through payments for medical devices, drugs, and biologicals. Section 1833(t)(6)(A) of the Act generally describes two groups of services—“current” and “new”—that are eligible for pass-through payments, depending, in part, on when they were first paid. One of the criteria for “new” drugs and biologicals to receive pass-through payments under section 1833(t)(6)(A)(iv)(I) of the Act is that payment for the item as an outpatient hospital service under Part B was not being made as of December 31, 1996. For those “new” drugs and biologicals, section 1833(t)(6)(C)(i)(II) of the Act specifies that there is a 2- to 3-year limitation on the pass-through period that begins on the first date on which payment is made under Part B for the drug or biological as an outpatient hospital service.

Section 419.64 of the regulations codifies the transitional pass-through payment provisions for drugs and biologicals. Section 419.64(a) describes the drugs and biologicals that are eligible for pass-through payments, essentially capturing the distinction between “new” and “current” services. Section 419.64(c)(2) provides that the pass-through payment eligibility period

for drugs and biologicals that fall into the “new” category begins on the date that CMS makes its first pass-through payment for the drug or biological.

In the CY 2010 OPPTS/ASC proposed rule (74 FR 35314), we noted that it had come to our attention that our pass-through payment eligibility period for “new” drugs and biologicals in § 419.64(c)(2) of the regulations might not most accurately reflect the statutory requirements of section 1833(t)(6)(C)(i)(II) of the Act. While our regulations indicate that the pass-through payment eligibility period for “new” drugs and biologicals begins on the first date on which pass-through payment is made for the item, section 1833(t)(6)(C)(i)(II) of the Act specifies that the pass-through period of 2 to 3 years for “new” drugs and biologicals begins on the first date on which payment is made under Part B for the drug or biological as an outpatient hospital service. In order to better reflect the statutory requirement for the pass-through period for a “new” drug or biological, in the CY 2010 OPPTS/ASC proposed rule (74 FR 35314), we proposed to revise paragraph (c)(2) of § 419.64 and add a new paragraph (c)(3) to § 419.64.

In order to conform the regulations to the statutory provisions, we proposed to change the start date of the pass-through payment eligibility period for a drug or biological from the first date on which pass-through payment is made to the date on which payment is first made for a drug or biological as an outpatient hospital service under Part B. Under this proposal, we needed to identify a first date of payment for a drug or biological as an outpatient hospital service under Part B. (Under our current policy, we had not established a start date for the eligibility period distinct from the beginning of pass-through payment because our current policy is to begin the pass-through payment eligibility period at the same time as we begin pass-through payment for the drug or biological.)

Due to the 2-year delay in the availability of claims data, under our CY 2010 proposal, we would not be able to identify an exact date of first payment for a drug or biological as an outpatient hospital service under Part B in order to determine the start date of the pass-through payment eligibility period until years after an application for pass-through payment for a “new” drug or biological has been submitted. At that later point in time, the pass-through payment eligibility period may be close to expiring, and the result of relying upon our claims data to evaluate an item for its eligibility for pass-through

status could result in a very short period of pass-through payment for the new drug or biological. Consequently, in the proposed rule, we stated our belief that it would be desirable to identify an appropriate and timely proxy for the date of first payment for the drug or biological as an outpatient hospital service under Part B. We proposed the date of first sale for a drug or biological in the United States following FDA approval as an appropriate proxy, as explained below, for the date on which the pass-through payment eligibility period would begin. We also noted that, in light of our CY 2010 proposal to treat implantable biologicals as medical devices for purposes of pass-through eligibility and payment under section 1833(t)(6) of the Act, described in section V.A.4. of the proposed rule (74 FR 35311 through 35314), these proposed revisions to the pass-through payment eligibility period for a drug or biological approved for pass-through payment beginning on or after January 1, 2010, would not apply to implantable biologicals, but rather only to nonimplantable biologicals.

We explained that the date of first sale of the drug or nonimplantable biological in the United States following FDA approval was an appropriate proxy for the first date of payment for the drug or nonimplantable biological as an outpatient hospital service under Part B for several reasons, including our expectation that Medicare beneficiaries would be among the first to use these drugs and nonimplantable biologicals. In addition, we currently rely on the date of first sale of a drug or biological in the United States following FDA approval under the ASP methodology and in the existing OPPTS pass-through payment eligibility determination. We stated that we did not believe that there is a more accurate and readily available proxy for the first date of payment for a drug or biological under Part B as an outpatient hospital service than the date of first sale of the drug or nonimplantable biological in the United States following FDA approval and that it was an accepted and available indicator of initial payment for the Medicare program.

For these reasons, we proposed that the date of first sale of a drug or nonimplantable biological in the United States following FDA approval would be the start date of the pass-through payment eligibility period for drugs or nonimplantable biologicals approved for pass-through payment beginning on or after January 1, 2010. We specified that our current policy—that the pass-through payment eligibility period of 2 to 3 years begins on the first date that

pass-through payment is made for the drug or biological—would apply only to drugs and biologicals approved for and receiving pass-through payment on or before December 31, 2009.

We currently implement new approvals of pass-through status for drugs and biologicals on a quarterly basis, and under our proposal for CY 2010, we stated that we would continue to implement these new approvals on a quarterly basis. We describe our quarterly process for reviewing and approving applications for drugs and biologicals to receive pass-through payment on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/04_passthrough_payment.asp. Interested parties may submit a complete application at any time. We typically review and make pass-through status approval decisions about complete applications for initiation of pass-through payment within 4 months of their submission and implement new pass-through status approvals on a quarterly basis through the next available OPPTS quarterly update. The CMS Web site provides a timeline showing the relationship between the date of submission of a complete application and the earliest date of pass-through payment that would result from approval of pass-through status for the drug or biological.

Under our current policy, the pass-through payment eligibility period and period of pass-through payment are the same. However, the pass-through payment eligibility period and the period of pass-through payment would not have been identical under our proposed policy. For our proposed policy, we identified both the pass-through payment eligibility period, as well as the period during which pass-through payment would be made, including the respective start and expiration dates of the pass-through payment eligibility period and the period of pass-through payment. We stated that the period of pass-through payment would coincide with the time period during which the drug or biological is designated as having pass-through status. (We note that being within the pass-through payment eligibility period alone does not qualify a “new” drug or biological for pass-through payment; the drug or biological must also meet the other requirements for pass-through payment, including a CMS determination that the cost of a drug or biological is not insignificant.) Under our proposal, the pass-through payment eligibility period would run for at least 2 years but no more than 3 years. We proposed to modify § 419.64

accordingly by adding new paragraph (c)(3) to state: “For a drug or nonimplantable biological described in paragraph (a)(4) of this section and approved for pass-through payment beginning on or after January 1, 2010—[the pass-through payment eligibility period begins on] the date of the first sale of the drug or nonimplantable biological in the United States after FDA approval.” Next, we proposed that pass-through payment itself would start on the first day of the calendar quarter following the calendar quarter during which the completed application was approved. We proposed to reflect this in regulation text, in proposed new § 419.64(c)(3), as follows. “Pass-through payment for the drug or nonimplantable biological begins on the first day of the hospital outpatient prospective payment system update following the update period during which the drug or nonimplantable biological was approved for pass-through status.” We noted that this start date for the period of pass-through payment would be specified in a letter to the applicant conveying pass-through status approval for the new drug or biological and would be the first day of the calendar quarter following the calendar quarter during which a complete pass-through application is approved by CMS for pass-through status.

Because the proposed revised definition of the pass-through payment eligibility period could have resulted in the eligibility period beginning well before application is made for pass-through payment for the drug or nonimplantable biological and could have resulted in a shorter period of pass-through payment for some drugs and biologicals than would be the case under our current policy, we also proposed to expire pass-through status for “new” drugs and biologicals on a quarterly basis. This proposal to expire the pass-through status of drugs and nonimplantable biologicals on a quarterly basis was a departure from our current policy for expiring the pass-through status of drugs and biologicals. Presently, we expire the pass-through status of drugs and biologicals at the end of the calendar year preceding the year of the applicable annual OPPTS update. Because our current pass-through payment eligibility period policy effectively aligns the start of pass-through payment with the beginning of the 2- to 3-year pass-through payment eligibility period, expiration of pass-through status on a calendar year basis affords those drugs and biologicals at least 2 but not more than 3 years of pass-through payment.

In addition to proposing to expire the pass-through status of “new” drugs and nonimplantable biologicals described by proposed new § 419.64(c)(3) on a quarterly basis, we also proposed to continue our established policy of determining whether a drug or biological would receive separate payment or packaged payment, after the expiration of the period of pass-through payment, on a calendar year basis through the annual OPPTS rulemaking process as described in section V.B.2. of the proposed rule (74 FR 35319 through 35321) and this final rule with comment period. Therefore, after the expiration of pass-through status of a “new” drug or biological in a given year’s calendar quarter, we proposed to continue to make separate payment through the end of that calendar year for those drugs and nonimplantable biologicals that would be subject to the drug packaging threshold when they did not have pass-through status at the applicable OPPTS payment rate for separately payable drugs and biologicals without pass-through status for that year, proposed to be ASP + 4 percent for CY 2010. (This proposal would exclude contrast agents and diagnostic radiopharmaceuticals for CY 2010, which would always be packaged when not on pass-through status.)

Comment: Several commenters disagreed with CMS’ proposal to change the pass-through payment eligibility period policy for new drugs and nonimplantable biologicals in CY 2010. Most of the commenters expressed concerns about separating the pass-through payment eligibility period from the period of pass-through payment, noting that delays that may occur between the date of the first sale of a drug in the United States and the date on which payment is first made under Part B would inevitably and inappropriately reduce the period of pass-through payment for new drugs. The commenters cited several examples, including a manufacturer’s delay in submitting a pass-through application after receiving FDA approval, the length of CMS’ pass-through review and approval process, delays in claim submissions and challenges associated with hospital billing for new services, and lags due to the resale process of a drug from a manufacturer to a wholesaler before the drug is available to the beneficiary. In addition, many commenters argued that non-Medicare beneficiaries, as opposed to Medicare beneficiaries, may be the first to receive a drug or biological, making the date of a drug’s first sale in the United States

after FDA approval irrelevant to the Medicare population.

One commenter acknowledged CMS' need to align the pass-through payment eligibility period policy with the statutory provisions. However, the commenter disagreed with CMS' proposal to use the date of the first sale in the United States following FDA approval as a proxy for the date on which payment is made under Part B. The commenter suggested that, considering all of the potential delays between the date of the first sale in the United States after FDA approval and the first date of payment under Part B as an outpatient hospital service, the date of the first sale in the United States after FDA approval is not a sufficiently precise proxy. The commenter suggested that CMS continue to use the current pass-through payment policy as a proxy for the first date on which payment is made under Part B, specifically the date that CMS first makes pass-through payment for a drug or biological, because it is the most accurate proxy. The commenter reasoned that establishing the date that CMS first makes pass-through payment for a drug or biological as a proxy for the first date on which payment is made under Part B as an outpatient hospital service is appropriate because the date of first pass-through payment would never predate the first payment under Part B as an outpatient hospital service, nor would it likely be made later than the date of first OPPS payment by an appreciable period of time. The commenter noted that, in general, manufacturers have an incentive to submit pass-through applications as quickly as possible and will do whatever they can to minimize any lag time between the date of first outpatient hospital payment and the availability of pass-through payments because pass-through status facilitates the product's introduction into the hospital outpatient setting.

Response: The commenter who urges us to adopt a different proxy than the one we proposed for the date of first payment under part B as an outpatient hospital service makes some very persuasive and compelling points. We have considered the merits and advantages of adopting the commenter's suggested proxy rather than the one we proposed, and we find that we agree with the commenter that the most appropriate policy is one that establishes the date that CMS makes its first pass-through payment for a drug or biological as the proxy for the first date on which payment is made under Part B as an outpatient hospital service. We believe that the date on which pass-

through payment is first made for a drug or nonimplantable biological is a more accurate proxy for the date on which payment is first made under Part B as an outpatient hospital service for several reasons. First, we agree with the commenter's points concerning the significant delays that may occur between the date of first sale of a drug or nonimplantable biological in the United States after FDA approval and the first date on which outpatient hospital payment is made under Part B. Such delays may result from numerous transactions in the drug distribution chain, initial use for non-Medicare patients with later diffusion to treatment of Medicare patients, delays in claims submission for new products without specific HCPCS codes, and established timeframes for Medicare processing payment of claims. All of these lags between the date of first sale and the date of first payment under Part B as an outpatient hospital service are cumulative and potentially significant. Therefore, adoption of the proposed proxy could, in some cases, lead to the start of the pass-through payment eligibility period substantially earlier than the start of the period of pass-through payment, thereby resulting in a reduction in the period of pass-through payment.

Second, we believe that utilizing the commenter's recommended proxy would eliminate the potential for delays between the proxy and the actual first date of payment under Part B as an outpatient hospital service, since the date of first pass-through payment would never predate the first payment under Part B as an outpatient hospital service. Although the first date of payment under Part B as an outpatient hospital service potentially could predate the date of first pass-through payment, it is also true that manufacturers have a significant incentive to submit pass-through applications as quickly as possible to minimize any lag between the date of first payment under the OPPS and the availability of pass-through payment. Pass-through payment can facilitate the availability of a product-specific HCPCS code for reporting its use and additional pass-through payment for the drug may allow beneficiaries access to the new drug in the HOPD. Therefore, in the rare circumstance that the date of first pass-through payment under the OPPS lags behind the first payment for the product under Part B as an outpatient hospital service, the delay is likely to be minimal. As a result, adopting this alternative date as a proxy would be unlikely to extend the pass-through

payment eligibility period beyond 2 to 3 years from the date of first payment under Part B as an outpatient hospital service as specified in the statute.

In addition, utilizing the date of first pass-through payment under the OPPS as a proxy for the date payment is first made for a product under Part B as an outpatient hospital service would afford drugs and nonimplantable biologicals at least a full 2 years of pass-through payment, whereas the proposed proxy might not have allowed for a full 2 years of pass-through payment in every case. Finally, using the date of first pass-through payment under the OPPS as the proxy for the date of first payment under Part B as an outpatient hospital service would not present an administrative burden to CMS or the public nor would it disrupt or change CMS' current operational practices. This administratively simple proxy would result in a continuation of the same smoothly functioning operational practices that CMS currently utilizes in determining pass-through payment for drugs and biologicals. Therefore, we are finalizing the date on which CMS makes its first pass-through payment as the proxy for the first date on which payment is made under Part B as an outpatient hospital service.

We note that, in the CY 2010 OPPS/ASC proposed rule (74 FR 35315 through 35317), we outlined CMS' pass-through payment policies for approving and expiring pass-through payment status for drugs and nonimplantable biologicals under the OPPS. In adopting the date on which CMS makes its first pass-through payment as a proxy for the first date on which payment is made under Part B as an outpatient hospital service and, therefore, as the start date for pass-through payment eligibility, we are not changing our current practices concerning application, approval, payment, and expiration of pass-through status for drugs and nonimplantable biologicals. In this regard, we will continue to accept applications as is currently described on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/04_passthrough_payment.asp. We will continue to begin pass-through payment on a quarterly basis through the next available OPPS quarterly update after the approval of a product's pass-through status. In addition, we will continue to expire pass-through status for drugs and nonimplantable biologicals on an annual basis through notice and comment rulemaking. Furthermore, our policy regarding the determination of packaging status after the pass-through status ends for a drug or biological, as discussed in section V.B.2. of this final

rule with comment period, remains the same. For those drugs with expiring pass-through status that are always packaged when they do not have pass-through status ("policy-packaged"), specifically diagnostic radiopharmaceuticals and contrast agents for CY 2010, we will package payment for these drugs once their pass-through status has expired. We discuss this policy in detail in section V.B.2.d. of this final rule with comment period.

Comment: One commenter argued that CMS' proposed proxy of the date of the first sale of a drug or biological in the United States following FDA approval was contradictory to section 1833(t)(6)(C)(i)(II) of the Act because that section references section 1833(t)(6)(A)(iv) of the Act, which defines a "new" drug or biological eligible for pass-through payment as being new after December 31, 1996, and as meeting the cost significance criteria. The commenter argued that a drug cannot be considered a pass-through drug until cost significance has been determined and that CMS would not determine cost significance until it qualifies a drug for pass-through status. Based on this assessment, the commenter argued that CMS should begin the pass-through payment period on the date CMS begins to treat the product as a pass-through drug, the first date of the pass-through payment period.

Response: We continue to believe that section 1833(t)(6)(C)(i)(II) of the Act requires the start date of the pass-through payment eligibility period for a drug or nonimplantable biological to begin on the date on which payment is first made for a drug or biological as an outpatient hospital service under Part B. As noted in the previous response, however, we are convinced by a commenter to adopt as the proxy for this date, the date on which CMS makes its first pass-through payment for the drug or nonimplantable biological.

Comment: Several commenters recommended that CMS continue to end pass-through status for drugs and nonimplantable biologicals on an annual basis, instead of ending pass-through status on a quarterly basis as CMS proposed. In the context of the specific proposal for the pass-through payment eligibility period, another commenter agreed with CMS' proposal to end pass-through status on a quarterly basis. Several other commenters argued that, because the proposal creates a delay between the beginning of the pass-through payment eligibility period and the period of pass-through payment, drugs and nonimplantable biologicals that are

approved for pass-through status should be given pass-through payment for the extent of the full 3-year pass-through eligibility period.

Response: Because we are adopting the date of first pass-through payment as the start of the pass-through payment eligibility period in this final rule with comment period, we will not change, as we proposed, the current operation of our drug and biological pass-through program. As is our current practice, we will continue to expire pass-through status for drugs and biologicals on an annual basis through notice and comment rulemaking. For example, if CMS receives a complete application for pass-through status for a drug on August 1, 2009, and approves the application for pass-through status for the January 1, 2010 OPPS quarterly update, the pass-through payment eligibility period would start on January 1, 2010. The pass-through payment period would extend for 2 but not more than 3 years, as is mandated by the statute, and we would propose to expire pass-through status for the drug on December 31, 2011 in the CY 2012 OPPS/ASC rulemaking process for January 1, 2012.

After consideration of the public comments we received, we are modifying our CY 2010 proposal and adopting the date of first pass-through payment for the drug or nonimplantable biological as the proxy for the first date on which payment for the product is made under Part B as an outpatient hospital service. Therefore, the 2- to 3-year pass-through payment eligibility period will start on the date of first pass-through payment and, consistent with our current policy, the pass-through payment eligibility period and the period of pass-through payment coincide. Finally, we will continue to expire the pass-through status of drugs and nonimplantable biologicals annually through the notice and comment rulemaking process.

Because our final policy reflects our current practice for implementing the pass-through eligibility and payment periods defined in section 1833(t)(6)(C)(i)(II) of the Act, we are not making any changes to § 419.64(c)(2), and we are not adding proposed new § 419.64(c)(3) to our regulations.

6. Provisions for Reducing Transitional Pass-Through Payments for Diagnostic Radiopharmaceuticals and Contrast Agents To Offset Costs Packaged Into APC Groups

a. Background

Prior to CY 2008, diagnostic radiopharmaceuticals and contrast agents were paid separately under the

OPPS if their mean per day costs were greater than the applicable year's drug packaging threshold. In CY 2008 (72 FR 66768), we began a policy of packaging payment for all nonpass-through diagnostic radiopharmaceuticals and contrast agents as ancillary and supportive items and services into their associated nuclear medicine procedures. Therefore, beginning in CY 2008, nonpass-through diagnostic radiopharmaceuticals and contrast agents were not subject to the annual OPPS drug packaging threshold to determine their packaged or separately payable payment status, and instead all nonpass-through diagnostic radiopharmaceuticals and contrast agents were packaged as a matter of policy. In the CY 2010 OPPS/ASC proposed rule (74 FR 35323), we proposed to continue to package payment for all nonpass-through diagnostic radiopharmaceuticals and contrast agents for CY 2010 as discussed in section V.B.2.d. of the proposed rule (74 FR 35323 through 35324).

b. Payment Offset Policy for Diagnostic Radiopharmaceuticals

As previously noted, radiopharmaceuticals are considered to be drugs for OPPS pass-through payment purposes. As described above, section 1833(t)(6)(D)(i) of the Act specifies that the transitional pass-through payment amount for pass-through drugs and biologicals is the difference between the amount paid under section 1842(o) (or the Part B drug CAP rate) and the otherwise applicable OPD fee schedule amount. There is currently one radiopharmaceutical with pass-through status under the OPPS, HCPCS code C9247 (Iobenguane, I-123, diagnostic, per study dose, up to 10 millicuries). HCPCS code C9247 was granted pass-through status beginning April 1, 2009 and will continue on pass-through status in CY 2010 under permanent HCPCS code A9582 (Iodine I-123 Iobenguane, diagnostic, per study dose, up to 15 millicuries). We currently apply the established radiopharmaceutical payment offset policy to pass-through payment for this product. As described earlier in section V.A.3. of this final rule with comment period, new pass-through diagnostic radiopharmaceuticals will be paid at ASP+6 percent, while those without ASP information will be paid at WAC+6 percent or, if WAC is not available, payment will be based on 95 percent of the product's most recently published AWP.

As a payment offset is necessary in order to provide an appropriate

transitional pass-through payment, we deduct from the payment for pass-through radiopharmaceuticals an amount that reflects the portion of the APC payment associated with predecessor radiopharmaceuticals in order to ensure no duplicate radiopharmaceutical payment is made. In CY 2009, we established a policy to estimate the portion of each APC payment rate that could reasonably be attributed to the cost of predecessor diagnostic radiopharmaceuticals when considering a new diagnostic radiopharmaceutical for pass-through payment (73 FR 68638 through 68641). Specifically, we utilize the “policy-packaged” drug offset fraction for APCs containing nuclear medicine procedures, calculated as 1 minus (the cost from single procedure claims in the APC after removing the cost for “policy-packaged” drugs divided by the cost from single procedure claims in the APC). We have previously defined “policy-packaged” drugs and biologicals as nonpass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals (73 FR 68639). In the CY 2010 OPPS/ASC proposed rule (74 FR 35323), we proposed for CY 2010 to redefine “policy-packaged” drugs as only nonpass-through diagnostic radiopharmaceuticals and contrast agents, as a result of the CY 2010 proposals discussed in sections V.A.4. and V.B.2.d. of the proposed rule (74 FR 35311 through 35314 and 74 FR 35323 through 35324) that would treat nonpass-through implantable

biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) and implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) with newly approved pass-through status beginning in CY 2010 or later as devices, rather than drugs. To determine the actual APC offset amount for pass-through diagnostic radiopharmaceuticals that takes into consideration the otherwise applicable OPPS payment amount, we multiply the “policy-packaged” drug offset fraction by the APC payment amount for the nuclear medicine procedure with which the pass-through diagnostic radiopharmaceutical is used and, accordingly, reduce the separate OPPS payment for the pass-through diagnostic radiopharmaceutical by this amount.

We will continue to post annually on the CMS Web site at <http://www.cms.hhs.gov/HospitalOutpatientPPS>, a file that contains the APC offset amounts that would be used for that year for purposes of both evaluating cost significance for candidate pass-through device categories and drugs and biologicals, including diagnostic radiopharmaceuticals, and establishing any appropriate APC offset amounts. Specifically, the file will continue to provide, for every OPPS clinical APC, the amounts and percentages of APC payment associated with packaged implantable devices, “policy-packaged” drugs, and “threshold-packaged” drugs and biologicals.

Table 23 of the proposed rule (74 FR 35318) displayed the proposed APCs to which nuclear medicine procedures would be assigned in CY 2010 and for which we expected that an APC offset could be applicable in the case of new diagnostic radiopharmaceuticals with pass-through status.

Comment: A few commenters supported the continuation of the pass-through diagnostic radiopharmaceutical offset policy for CY 2010.

Response: We continue to believe that a diagnostic radiopharmaceutical offset policy is necessary in order to ensure that duplicate payment is not made for diagnostic radiopharmaceuticals with pass-through status. We believe it is appropriate to remove the radiopharmaceutical payment amount that is already packaged into the payment for the associated nuclear medicine procedure when we provide pass-through payment for a diagnostic radiopharmaceutical with pass-through status.

Therefore, after consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to apply the diagnostic radiopharmaceutical offset policy to payment for pass-through diagnostic radiopharmaceuticals, as described above. Table 32 below displays the APCs to which nuclear medicine procedures are assigned in CY 2010 and for which we expect that an APC offset could be applicable in the case of diagnostic radiopharmaceuticals with pass-through status.

TABLE 32—APCs TO WHICH NUCLEAR MEDICINE PROCEDURES ARE ASSIGNED FOR CY 2010

CY 2010 APC	CY 2010 APC title
0307	Myocardial Positron Emission Tomography (PET) imaging.
0308	Non-Myocardial Positron Emission Tomography (PET) imaging.
0377	Level II Cardiac Imaging.
0378	Level II Pulmonary Imaging.
0389	Level I Non-imaging Nuclear Medicine.
0390	Level I Endocrine Imaging.
0391	Level II Endocrine Imaging.
0392	Level II Non-imaging Nuclear Medicine.
0393	Hematologic Processing & Studies.
0394	Hepatobiliary Imaging.
0395	GI Tract Imaging.
0396	Bone Imaging.
0397	Vascular Imaging.
0398	Level I Cardiac Imaging.
0400	Hematopoietic Imaging.
0401	Level I Pulmonary Imaging.
0402	Level II Nervous System Imaging.
0403	Level I Nervous System Imaging.
0404	Renal and Genitourinary Studies.
0406	Level I Tumor/Infection Imaging.
0408	Level III Tumor/Infection Imaging.
0414	Level II Tumor/Infection Imaging.

c. Payment Offset Policy for Contrast Agents

As described above, section 1833(t)(6)(D)(i) of the Act specifies that the transitional pass-through payment amount for pass-through drugs and biologicals is the difference between the amount paid under section 1842(o) (or the Part B drug CAP rate) and the otherwise applicable OPD fee schedule amount. There is currently one contrast agent with pass-through status under the OPSS, HCPCS code C9246 (Injection, gadoxetate disodium, per ml). HCPCS code C9246 was granted pass-through status beginning January 1, 2009, and will continue with pass-through status in CY 2010 under HCPCS code A9581 (Injection, gadoxetate disodium, 1 ml). As described earlier in section V.A.3. of this final rule with comment period, new pass-through contrast agents will be paid at ASP+6 percent, while those without ASP information would be paid at WAC+6 percent or, if WAC is not available, payment would be based on 95 percent of the product's most recently published AWP.

As discussed in the CY 2010 OPSS/ASC proposed rule (74 FR 35318), we believe that a payment offset, similar to the offset currently in place for pass-through devices and diagnostic radiopharmaceuticals, is necessary in order to provide an appropriate transitional pass-through payment for contrast agents because all of these items are packaged when they do not have pass-through status. In accordance with our standard offset methodology, in the CY 2010 OPSS/ASC proposed rule (74 FR 35318), we proposed to deduct from the payment for pass-through contrast agents an amount that reflects the portion of the APC payment associated with predecessor contrast agents in order to ensure no duplicate contrast agent payment is made.

In CY 2009, we established a policy to estimate the portion of each APC payment rate that could reasonably be attributed to the cost of predecessor diagnostic radiopharmaceuticals when considering a new diagnostic radiopharmaceutical for pass-through payment (73 FR 68638 through 68641). For CY 2010, we proposed to apply this same policy to contrast agents. Specifically, we proposed to utilize the "policy-packaged" drug offset fraction for clinical APCs calculated as 1 minus (the cost from single procedure claims in the APC after removing the cost for "policy-packaged" drugs divided by the cost from single procedure claims in the APC). As discussed above, while we have previously defined the "policy-

packaged" drugs and biologicals as nonpass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals (73 FR 68639), we proposed for CY 2010 to redefine "policy-packaged" drugs as only nonpass-through diagnostic radiopharmaceuticals and contrast agents, as a result of the CY 2010 proposal discussed in sections V.A.4. and V.B.2.d. of the proposed rule (74 FR 35311 through 35314 and 74 FR 35323 through 35324) that would treat all implantable biologicals as devices, rather than drugs. To determine the actual APC offset amount for pass-through contrast agents that takes into consideration the otherwise applicable OPSS payment amount, we proposed to multiply the "policy-packaged" drug offset fraction by the APC payment amount for the procedure with which the pass-through contrast agent is used and, accordingly, reduce the separate OPSS payment for the pass-through contrast agent by this amount.

We proposed to continue to post annually on the CMS Web site at <http://www.cms.hhs.gov/HospitalOutpatientPPS>, a file that contains the APC offset amounts that would be used for that year for purposes of both evaluating cost significance for candidate pass-through device categories and drugs and biologicals, including contrast agents, and establishing any appropriate APC offset amounts. Specifically, the file will continue to provide, for every OPSS clinical APC, the amounts and percentages of APC payment associated with packaged implantable devices, "policy-packaged" drugs, and "threshold-packaged" drugs and biologicals.

Comment: One commenter objected to the proposed offset policy for contrast agents, stating that an offset for new contrast agents granted pass-through status, combined with the packaging policy for all nonpass-through contrast agents, would discourage hospitals from providing contrast agents for financial reasons. The commenter argued that an offset policy is not necessary to avoid duplicate payment for pass-through contrast agents as the majority of older contrast agents have costs that are well below the \$65 OPSS drug packaging threshold and more expensive contrast agents would be eligible for pass-through status. Finally, the commenter believed that CMS does not have the appropriate contrast agent data available in order to calculate an offset amount for these products. Another commenter objected to CMS' proposed offset methodology for contrast agents and urged CMS to specify the APCs that

would be subject to an offset. Further, the commenter requested that CMS implement a contrast offset methodology that would be more similar to the offset methodology currently in place for pass-through devices and diagnostic radiopharmaceuticals.

Response: We have consistently implemented an offset policy for products receiving pass-through payment that would otherwise receive significant packaged payment if not for their pass-through status. An offset methodology ensures that we do not pay twice, first through a packaged payment included in the associated procedure payment and second through an individual separate payment, for the item with pass-through status. The potential for duplicate payment is higher for items such as contrast agents, diagnostic radiopharmaceuticals, and devices where the pass-through item typically substitutes for items that are otherwise always packaged. Furthermore, the potential magnitude of duplicate payment also is higher for these items because they are always packaged when they do not have pass-through status.

As discussed above, this offset policy appropriately provides for pass-through payment for the new product that represents the difference between the physician's office payment amount and the otherwise applicable OPD fee schedule amount, in the case of packaged contrast agents the "policy-packaged" drug APC offset amount, as specified by the statute. We note that the proposed contrast agent offset policy is virtually identical to the offset methodology currently in place for pass-through devices and diagnostic radiopharmaceuticals, consistent with the recommendation by one commenter that we adopt a similar policy for contrast agents. We believe that this methodology would pay appropriately for the cost of pass-through contrast agents and that hospitals should have no payment concerns when determining which contrast agent would be most clinically appropriate and efficient for a particular patient's study. Therefore, we do not believe that the application of a contrast agent offset methodology would discourage hospitals from using pass-through contrast agents insofar as providers determine they are necessary in the care of the patient.

As discussed above, we proposed to deduct from the payment for pass-through contrast agents an amount that reflects the portion of the APC payment associated with predecessor contrast agents in order to ensure no duplicate contrast agent payment is made. As

discussed above, we identified the “policy-packaged” drug APC offset amount as applicable to our offset policy because we have identified contrast agents as “policy-packaged” drugs in our claims data. To the extent that hospitals reported the HCPCS code for contrast agents when those drugs were administered during procedures, the contrast agent costs are included in our calculation of the “policy-packaged” drug APC offset amounts, and we believe that we have sufficient information regarding the costs of predecessor contrast agents to apply the resulting offset amounts to payment for pass-through contrast agents. To the extent hospitals did not report the use of contrast agents under specific HCPCS codes in CY 2008, we could not fully total the cost of contrast for a given imaging APC and we would underestimate an accurate “policy-packaged” drug APC offset amount. This unknown but potential bias would generally result in higher overall pass-through payment for a new contrast agent so any limitations of our current data on contrast agents for purposes of the offset would not inappropriately reduce pass-through payment for a new contrast agent.

We disagree with the commenter that an offset is unnecessary to avoid duplicate payment for contrast material. All nonpass-through contrast agents, regardless of their per day costs, are packaged into payments for the associated procedures. Therefore, OPSS payment for imaging and other procedures that currently utilize contrast agents already includes packaged payment for the necessary contrast agent. The observation that most contrast agents have per day costs below the \$65 threshold does not obviate the need for an offset policy for contrast agents with pass-through status. First, while the CY 2010 drug packaging threshold is low, \$65 as the per day cost, this cost may constitute a sizable percentage of a procedural APC’s median cost. Paying the full procedural APC amount plus the pass-through contrast agent payment of ASP+6 for an imaging scan with high volume could result in significant overpayment of the new contrast agent. Furthermore, a few contrast agents have per day costs above the \$65 drug packaging threshold, so that the amount of contrast agent cost represented in the “policy-packaged” drug amount of an APC median cost could be fairly substantial. Finally, unlike “threshold-packaged” drugs that are packaged based on the relationship of their per day cost to the \$65 drug packaging threshold, where the

packaged drug cost in a procedural APC may or may not represent predecessor drug costs and where multiple drugs may be administered in a single session paid under one procedural APC, contrast agents typically substitute for one another and hospitals rarely administer multiple contrast agents in the same session. Pass-through contrast agents are paid separately and are billed with procedures that already have costs of predecessor contrast agents packaged into the procedural APC payment, so duplicate contrast agent payment would result in the absence of an offset methodology.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35318), we proposed to utilize the “policy-packaged” drug offset fraction for procedural APCs calculated as 1 minus (the cost from single procedure claims in the APC after removing the cost for “policy-packaged” drugs divided by the cost from single procedure claims in the APC). To determine the actual APC offset amount for pass-through contrast agents that takes into consideration the otherwise applicable OPSS payment amount, we proposed to multiply the “policy-packaged” drug offset fraction by the APC payment amount for the procedure with which the pass-through contrast agent is used and, accordingly, reduce the separate OPSS payment for the pass-through contrast agent by this amount.

In response to the commenters’ concerns regarding our proposed methodology and request that we specify the APCs subject to the contrast agent offset policy, we reviewed the methodology and specifically examined the amount of contrast agent offsets associated with procedural APCs to determine which APCs, other than nuclear medicine APCs that contained the costs of diagnostic radiopharmaceuticals, included a significant “policy-packaged” drug amount in the APC payment. First, we excluded all APCs to which nuclear medicine procedures were assigned for CY 2010 from the APCs that would be subject to a contrast agent offset policy, reasoning that the “policy-packaged” drug costs associated with these APCs were for diagnostic radiopharmaceuticals. From a clinical perspective, there is very little overlap in the procedures that use contrast agents or diagnostic radiopharmaceuticals. Next, we reviewed the per day costs for all contrast agents with CY 2008 claims data and compared their aggregate, average per day cost to the “policy-packaged” drug amounts listed in the CY 2010 proposed rule APC offset file that was posted on the CMS Web site in

association with the CY 2010 OPSS/ASC proposed rule. When examining those APCs with “policy-packaged” drug amounts equal to or less than the 25th percentile of per day contrast agent cost (approximately \$22), we found that the majority of APCs with a “policy-packaged” drug offset amount other than zero but less than \$20 (our \$22 estimate rounded to the nearest \$5 increment) were generally APCs that were not likely to include procedures requiring significant use of contrast agents. We selected the 25th percentile of per day contrast agent cost to identify the majority of APCs with significant contrast agent cost because we believe that the 25th percentile is an appropriate threshold for representing significant contrast agent cost as it captures the lower bound of significant variation around the per day contrast agent cost. The interquartile range, the 25th to 75th percentile, is a typical descriptive statistic used to describe the variation in the center of a distribution. Further, the dollar value of the 25th percentile, \$22 was sufficiently high that we believed it would be worth establishing and implementing offset logic in our claims processing Pricer module. This allowed us to establish a meaningful threshold cost for application of a contrast agent offset policy that would identify APCs in which there is significant packaged contrast agent cost. Unlike the case of diagnostic radiopharmaceuticals, which are always administered during a limited number of nuclear medicine procedures so we are able to identify all APCs to which nuclear medicine procedures are assigned as those for which the diagnostic radiopharmaceutical offset policy would apply, contrast agents are utilized much more widely among procedures assigned to many OPSS APCs.

The APCs that we identified as below the threshold of \$20 included APC 0384 (GI Procedures with Stents) and APC 0427 (Level II Tube or Catheter Changes or Repositioning). As we would not expect contrast agents to generally be used in the procedures assigned to these APCs, we believe that implementing a threshold that would exclude these APCs from a contrast agent offset policy would be appropriate for administrative simplification of claims processing, while continuing to ensure no duplicate payment is made for pass-through contrast agents. Therefore, we have identified the APCs that would be subject to the contrast offset policy in CY 2010, within the scope of the criteria discussed above.

After consideration of the public comments we received, we are

finalizing a pass through contrast agent offset policy for CY 2010, with modification to specify the procedural APCs to which offsets for pass-through contrast agents would apply. Procedural APCs for which we expect a contrast agent offset could be applicable in the

case of a pass-through contrast agent have been identified as any procedural APC with a “policy-packaged” drug amount greater than \$20 that is not a nuclear medicine APC identified in Table 32 above, and these APCs are displayed in Table 33. For CY 2010,

when a contrast agent with pass-through status is billed with any procedural APC listed in Table 33, a specific offset based on the procedural APC will be applied to payment for the contrast agent to ensure that duplicate payment is not made for the contrast agent.

TABLE 33—APCs TO WHICH A CONTRAST AGENT OFFSET MAY BE APPLICABLE FOR CY 2010

CY 2010 APC	CY 2010 APC title
0080	Diagnostic Cardiac Catheterization.
0082	Coronary or Non-Coronary Atherectomy.
0083	Coronary or Non-Coronary Angioplasty and Percutaneous Valvuloplasty.
0093	Vascular Reconstruction/Fistula Repair without Device.
0104	Transcatheter Placement of Intracoronary Stents.
0128	Echocardiogram with Contrast.
0152	Level I Percutaneous Abdominal and Biliary Procedures.
0229	Transcatheter Placement of Intravascular Shunts.
0278	Diagnostic Urography.
0279	Level II Angiography and Venography.
0280	Level III Angiography and Venography.
0283	Computed Tomography with Contrast.
0284	Magnetic Resonance Imaging and Magnetic Resonance Angiography with Contrast.
0333	Computed Tomography without Contrast followed by Contrast.
0337	Magnetic Resonance Imaging and Magnetic Resonance Angiography without Contrast followed by Contrast.
0375	Ancillary Outpatient Services When Patient Expires.
0383	Cardiac Computed Tomographic Imaging.
0388	Discography.
0418	Insertion of Left Ventricular Pacing Elect.
0442	Dosimetric Drug Administration.
0653	Vascular Reconstruction/Fistula Repair with Device.
0656	Transcatheter Placement of Intracoronary Drug-Eluting Stents.
0662	CT Angiography.
0668	Level I Angiography and Venography.
8006	CT and CTA with Contrast Composite.
8008	MRI and MRA with Contrast Composite.

B. OPSS Payment for Drugs, Biologicals, and Radiopharmaceuticals Without Pass-Through Status

1. Background

Under the CY 2009 OPSS, we currently pay for drugs, biologicals, and radiopharmaceuticals that do not have pass-through status in one of two ways: packaged payment into the payment for the associated service; or separate payment (individual APCs). We explained in the April 7, 2000 OPSS final rule with comment period (65 FR 18450) that we generally package the cost of drugs and radiopharmaceuticals into the APC payment rate for the procedure or treatment with which the products are usually furnished. Hospitals do not receive separate payment for packaged items and supplies, and hospitals may not bill beneficiaries separately for any packaged items and supplies whose

costs are recognized and paid within the national OPSS payment rate for the associated procedure or service. (Transmittal A-01-133, issued on November 20, 2001, explains in greater detail the rules regarding separate payment for packaged services.)

Packaging costs into a single aggregate payment for a service, procedure, or episode-of-care is a fundamental principle that distinguishes a prospective payment system from a fee schedule. In general, packaging the costs of items and services into the payment for the primary procedure or service with which they are associated encourages hospital efficiencies and also enables hospitals to manage their resources with maximum flexibility.

Section 1833(t)(16)(B) of the Act, as added by section 621(a)(2) of Public Law 108-173, set the threshold for establishing separate APCs for drugs and biologicals at \$50 per

administration for CYs 2005 and 2006. Therefore, for CYs 2005 and 2006, we paid separately for drugs, biologicals, and radiopharmaceuticals whose per day cost exceeded \$50 and packaged the costs of drugs, biologicals, and radiopharmaceuticals whose per day cost was equal to or less than \$50 into the procedures with which they were billed. For CY 2007, the packaging threshold for drugs, biologicals, and radiopharmaceuticals that were not new and did not have pass-through status was established at \$55. For CYs 2008 and 2009, the packaging threshold for drugs, biologicals, and radiopharmaceuticals that were not new and do not have pass-through status was established at \$60. The methodology used to establish the \$55 threshold for CY 2007, the \$60 threshold for CYs 2008 and 2009, and our approach for CY 2010 are discussed in more detail in section

V.B.2.b. of this final rule with comment period.

2. Criteria for Packaging Payment for Drugs, Biologicals, and Radiopharmaceuticals

a. Background

As indicated in section V.B.1. of this final rule with comment period, in accordance with section 1833(t)(16)(B) of the Act, the threshold for establishing separate APCs for payment of drugs and biologicals was set to \$50 per administration during CYs 2005 and 2006. In CY 2007, we used the fourth quarter moving average Producer Price Index (PPI) levels for prescription preparations to trend the \$50 threshold forward from the third quarter of CY 2005 (when the Pub. L. 108–173 mandated threshold became effective) to the third quarter of CY 2007. We then rounded the resulting dollar amount to the nearest \$5 increment in order to determine the CY 2007 threshold amount of \$55. Using the same methodology as that used in CY 2007 (which is discussed in more detail in the CY 2007 OPPS/ASC final rule with comment period (71 FR 68085 through 68086)), we set the packaging threshold for establishing separate APCs for drugs and biologicals at \$60 for CYs 2008 and 2009.

Following the CY 2007 methodology, for the CY 2010 OPPS/ASC proposed rule we used updated fourth quarter moving average PPI levels to trend the \$50 threshold forward from the third quarter of CY 2005 to the third quarter of CY 2009 and again rounded the resulting dollar amount (\$65.07) to the nearest \$5 increment, which yielded a figure of \$65. In performing this calculation, we used the most up-to-date forecasted, quarterly PPI estimates from CMS' Office of the Actuary (OACT). As actual inflation for past quarters replaced forecasted amounts, the PPI estimates for prior quarters have been revised (compared with those used in the CY 2007 OPPS/ASC final rule with comment period) and were incorporated into our calculation. Based on the calculations described above, we proposed a packaging threshold for CY 2010 of \$65. (For a more detailed discussion of the OPPS drug packaging threshold and the use of the PPI for prescription drugs, we refer readers to the CY 2007 OPPS/ASC final rule with comment period (71 FR 68085 through 68086).)

b. Cost Threshold for Packaging of Payment for HCPCS Codes that Describe Certain Drugs, Nonimplantable Biologicals, and Therapeutic Radiopharmaceuticals (“Threshold-Packaged Drugs”)

To determine their proposed CY 2010 packaging status, for the CY 2010 OPPS/ASC proposed rule we calculated the per day cost of all drugs on a HCPCS code-specific basis (with the exception of those drugs and biologicals with multiple HCPCS codes that include different dosages as described in section V.B.2.c. of the CY 2010 OPPS/ASC proposed rule (74 FR 35321) and excluding diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals that we proposed to continue to package in CY 2010 as discussed in section V.B.2.d. of the CY 2010 OPPS/ASC proposed rule (74 FR 35323 through 35324) and this final rule with comment period), nonimplantable biologicals, and therapeutic radiopharmaceuticals (collectively called “threshold-packaged” drugs) that had a HCPCS code in CY 2008 and were paid (via packaged or separate payment) under the OPPS, using CY 2008 claims data processed before January 1, 2009. In order to calculate the per day costs for drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals to determine their proposed packaging status in CY 2010, we used the methodology that was described in detail in the CY 2006 OPPS proposed rule (70 FR 42723 through 42724) and finalized in the CY 2006 OPPS final rule with comment period (70 FR 68636 through 70 FR 68638).

To calculate the CY 2010 proposed rule per day costs, we used an estimated payment rate for each drug and nonimplantable biological HCPCS code of ASP+4 percent (which was the payment rate we proposed for separately payable drugs and nonimplantable biologicals in CY 2010, as discussed in more detail in section V.B.3.b. of the CY 2010 OPPS/ASC proposed rule (74 FR 35324 through 35326)). We used the manufacturer submitted ASP data from the fourth quarter of CY 2008 (data that were used for payment purposes in the physician's office setting, effective April 1, 2009) to determine the proposed rule per day cost.

As is our standard methodology, for CY 2010, we proposed to use payment rates based on the ASP data from the fourth quarter of CY 2008 for budget neutrality estimates, packaging determinations, impact analyses, and completion of Addenda A and B to the proposed rule because these were the

most recent data available for use at the time of development of the proposed rule. These data were also the basis for drug payments in the physician's office setting, effective April 1, 2009. For items that did not have an ASP-based payment rate, such as therapeutic radiopharmaceuticals, we used their mean unit cost derived from the CY 2008 hospital claims data to determine their proposed per day cost. We packaged items with a per day cost less than or equal to \$65 and identified items with a per day cost greater than \$65 as separately payable. Consistent with our past practice, we crosswalked historical OPPS claims data from the CY 2008 HCPCS codes that were reported to the CY 2009 HCPCS codes that we displayed in Addendum B to the proposed rule for payment in CY 2010.

Comment: Several commenters supported CMS' proposal to increase the packaging threshold to \$65 for CY 2010. However, the majority of commenters objected to the proposed increase to the OPPS packaging threshold.

A few commenters recommended that CMS consider either eliminating the drug packaging threshold and providing separate payment for all drugs with HCPCS codes or freezing the packaging threshold at \$60 for CY 2010. Some commenters objected to the use of a packaging threshold under the OPPS when one is not used for physician's office payment and believed that eliminating the drug packaging threshold would allow for parity in drug payment between the HOPD setting and the physician's office setting. These commenters expressed concern that the packaging threshold may impede beneficiary access to lower-cost packaged drugs in the HOPD setting. In addition, some commenters believed that eliminating the packaging threshold and paying separately for all drugs in the HOPD setting would allow a more accurate calculation of the separately payable payment amount for drugs (otherwise referred to as the ASP+X percent amount). Other commenters stated that CMS should not increase the drug packaging threshold because other changes in the drug payment ratesetting methodology were proposed. These commenters requested that CMS only change one aspect of the drug payment methodology at a time to allow for greater understanding of the impact of proposed changes to drug payment.

Response: As fully discussed in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66757 through 66758) and the CY 2009 OPPS/ASC final rule with comment period (73 FR 68643), we continue to believe that unpackaging payment for all drugs,

biologicals, and radiopharmaceuticals is inconsistent with the concept of a prospective payment system and that such a change could create an additional reporting burden for hospitals. The OPSS and the MPFS that applies to physician's office services are fundamentally different payment systems with essential differences in their payment policies and structures. Specifically, the OPSS is a prospective payment system, based on the concept of payment for groups of services that share clinical and resource characteristics. Payment is made under the OPSS according to prospectively established payment rates that are related to the relative costs of hospital resources for services. The MPFS is a fee schedule based on the relative value of each individual component of a service. Consistent with the MPFS approach, separate payment is made for each drug provided in the physician's office, but the OPSS packages payment for certain drugs into the associated procedure payments for the APC group. Given the fundamental differences between the MPFS payment mechanism and the OPSS payment mechanism, differences in the degrees of packaged payment and separate payment between these two systems are only to be expected. In general, we do not believe that our packaging methodology under the OPSS results in limited beneficiary access to drugs because packaging is a fundamental component of a prospective payment system that accounts for the cost of certain items and services in larger payment bundles, recognizing that some clinical cases may be more costly and others less costly but that, on average, OPSS payment is appropriate for the services provided.

We note that, in CYs 2005 and 2006, the statutorily mandated drug packaging threshold was set at \$50, and we continue to believe that it is appropriate to continue a modest drug packaging threshold for the CY 2010 OPSS for the reasons set forth below. As stated in the CY 2007 OPSS/ASC final rule with comment period (71 FR 68086), we believe that packaging certain items is a fundamental component of a prospective payment system, that packaging these items does not lead to beneficiary access issues and does not create a problematic site of service differential, that the packaging threshold is reasonable based on the initial establishment in law of a \$50 threshold for the CY 2005 OPSS, that updating the \$50 threshold is consistent with industry and government practices, and that the PPI for prescription preparations is an appropriate

mechanism to gauge Part B drug inflation. Therefore, because of our continued belief that packaging is a fundamental component of a prospective payment system that contributes to important flexibility and efficiency in the delivery of high quality hospital outpatient services, we are not adopting the commenters' recommendations to pay separately for all drugs, biologicals, and radiopharmaceuticals for CY 2010 or to eliminate or to freeze the packaging threshold at \$60.

Finally, we believe that our continued application of the methodology initially adopted in CY 2007 to update the drug packaging threshold does not inhibit our ability to propose additional changes to the nonpass-through drug payment methodology under the OPSS. We note that for the past several years, we have made a number of proposals to revise our drug payment methodology, while continuing to implement our established methodology for annually updating the drug packaging threshold. While we have not finalized any of these previous proposals, we have consistently applied the methodology described above to update the drug packaging threshold while examining a variety of alternatives for determining payment for separately payable drugs without pass-through status.

Comment: One commenter to the CY 2009 OPSS/ASC final rule with comment period noted that HCPCS code J3300 (Injection, triamcinolone acetonide, preservative free, 1 mg) should not be packaged as established in the final rule with comment period because the per day cost of this drug is over the CY 2009 OPSS drug packaging threshold of \$60 per day.

Response: While the payment for HCPCS code J3300 was adopted on an interim final basis as packaged for CY 2009 (status indicator "N"), upon receipt of this public comment we reviewed our calculation and released a correction notice changing the status indicator to "K" for CY 2009 (74 FR 4343). In addition, we discussed this status indicator change in the April 2009 OPSS quarterly update CR (Transmittal 1702, CR 6416, dated March 13, 2009).

Comment: One commenter stated that HCPCS code J3473 (Injection, hyaluronidase, recombinant, 1 USP unit) was incorrectly assigned status indicator "N" in the CY 2010 OPSS/ASC proposed rule. The commenter argued that coding errors resulted in hospital claims data indicating that per day costs of HCPCS code J3473 is below the drug packaging threshold for CY 2010. The commenter

explained that a variety of HCPCS codes and various dosage descriptors for similar products contributed to hospital coding errors, and that the product described by HCPCS code J3473 is only sold in a single use vial of 150 units, with an ASP that exceeds the CY 2010 packaging threshold. The commenter noted that this concern had been raised with the CMS HCPCS Workgroup but a request for a new HCPCS code descriptor was denied.

Response: HCPCS code J3473 expired from pass-through status on December 31, 2008, and was paid separately in CY 2009 because the estimated per day cost, using updated final rule claims data from CY 2007, showed that the per day cost of this drug exceeded the CY 2009 drug packaging threshold. For CY 2010, we proposed to package HCPCS code J3473 as the estimated per day cost did not exceed the proposed CY 2010 drug packaging threshold. The OPSS relies on hospital claims data in order to determine payment rates. For drugs and biologicals, we rely upon hospital claims data, in part, to determine the estimated per day cost we use in our annual packaging determination. In addition, the concern about discrepancies between HCPCS code descriptors for similar products is under the purview of the CMS HCPCS Workgroup, the sole creator and maintainer of HCPCS codes and their descriptors. We remind hospitals through each OPSS quarterly update CR that when billing for drugs, biologicals, and radiopharmaceuticals, they should make certain that the reported units of service of the reported HCPCS code are consistent with the quantity of the drug, biological, or radiopharmaceutical that was used in the care of the patient. Therefore, we expect that the data that we receive on hospital claims accurately reflect the services that were provided to the beneficiary.

As is our standard methodology, we used updated claims data and ASP rates to make final packaging determinations for CY 2010. For HCPCS code J3473, our CY 2008 claims data showed approximately 2,100 days and 226,800 units from 37 providers. While this drug was not commonly used in CY 2008, we have no reason to believe that the estimated per day cost of HCPCS code J3473 of approximately \$57, based on our methodology described above as applied to claims from a modest number of providers, is not reflective of the per day cost to hospitals for furnishing the drug. Therefore, we have determined that the per day cost of HCPCS code J3473 does not exceed the \$65 packaging threshold for drugs and

biologicals and payment for HCPCS code J3473 is packaged in CY 2010.

For purposes of this final rule with comment period, we again followed the CY 2007 methodology for CY 2010 and used updated fourth quarter moving average PPI levels to trend the \$50 threshold forward from the third quarter of CY 2005 to the third quarter of CY 2010 and again rounded the resulting dollar amount (\$66.55) to the nearest \$5 increment, which yielded a figure of \$65. In performing this calculation, we used the most up-to-date forecasted, quarterly PPI estimates from CMS' OACT.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue use of the established methodology for annually updating the OPPS packaging threshold for drugs and biologicals by the PPI for prescription drugs. The final CY 2010 drug packaging threshold is \$65, calculated according to the threshold update methodology that we have applied since CY 2007.

In CY 2005 (69 FR 65779 through 65780), we implemented a policy that exempted the oral and injectable forms of 5-HT3 antiemetic products from our packaging policy, providing separate payment for these drugs regardless of their estimated per day costs through CY 2009. There are currently seven Level II HCPCS codes for 5-HT3 antiemetics that describe four different drugs, specifically dolasetron mesylate, granisetron hydrochloride, ondansetron hydrochloride, and palonosetron hydrochloride. Each of these drugs, except palonosetron hydrochloride, is available in both injectable and oral forms, so seven HCPCS codes are available to describe the four drugs in all of their forms. As of 2008, both ondansetron hydrochloride and granisetron hydrochloride were available in generic versions. We have now paid separately for all 5-HT3 antiemetics for 5 years under a policy that exempts these products from the drug packaging methodology. While we continue to believe that use of these antiemetics is an integral part of an anticancer treatment regimen and that OPPS claims data demonstrate their increasingly common hospital outpatient utilization, in the CY 2010 OPPS/ASC proposed rule (74 FR 35320), we indicated that we no longer believe that a specific exemption to our standard drug payment methodology is necessary for CY 2010 to ensure access to the most appropriate antiemetic product for Medicare beneficiaries.

We analyzed historical hospital outpatient claims data for the seven

5-HT3 antiemetic products that have been subject to this packaging exemption, and we found that HCPCS code J2405 (Injection, ondansetron hydrochloride, per 1 mg) was the dominant product used in the hospital outpatient setting both before and after the adoption of our 5-HT3 packaging exemption in CY 2005. Prior to this packaging exemption, payment for HCPCS code J2405 was packaged in CY 2004. HCPCS code J2405 was modestly costly relative to the other 5-HT3 antiemetics in CY 2004, but its per day cost still fell below the applicable packaging threshold of \$50. Since CY 2005, the injectable form of ondansetron hydrochloride has experienced a significant change in its pricing structure as generic versions of the drug have become available, including a steady decline in its estimated per day cost. Notwithstanding this change in price, we have observed continued growth in its OPPS utilization. For CY 2008, HCPCS code J2405 was the least costly of the seven 5-HT3 antiemetics, with an estimated per day cost of only approximately \$1 in CY 2008 (based on July 2008 ASP information), yet we observed that it constituted 88 percent of all treatment days of 5-HT3 antiemetics in the CY 2008 OPPS claims data. Using April 2009 ASP information for the CY 2010 proposed rule, we estimated a per day cost of only approximately \$1 for HCPCS code J2405. For the five modestly priced 5-HT3 antiemetics, we estimated CY 2010 per day costs between approximately \$7 and \$50, while we estimated a per day cost for the most costly 5-HT3 antiemetic, J2469 (Injection, palonosetron hcl, 25 mcg), of \$174 per day. In light of an anticipated relatively constant pricing structure for these drugs in CY 2010, combined with our experience that prescribing patterns for these 5-HT3 antiemetics are not very sensitive to changes in price, we did not believe that continuing to exempt these drugs from our standard OPPS drug packaging methodology was appropriate for CY 2010. Therefore, for CY 2010, because we proposed to no longer exempt the 5-HT3 antiemetic products from our standard packaging methodology, we proposed to package payment for all of the 5-HT3 antiemetics except palonosetron hydrochloride, consistent with their estimated per day costs from CY 2008 claims data.

At the August 2009 meeting of the APC Panel, the APC Panel recommended that when CMS changes the dollar amount of the drug packaging threshold and determines that some drugs within a single therapeutic class

fall on either side of the packaging threshold, CMS consider packaging all of the drugs within that class on the basis of feedback from providers, the APC Panel, and stakeholders. Our response to this recommendation is included in our response to comments below.

Comment: The majority of commenters opposed the proposal to no longer continue to exempt the oral and injectable forms of 5-HT3 antiemetics from packaging, thereby packaging all but one 5-HT3 antiemetic. Many commenters requested that CMS continue to exempt all 5-HT3 antiemetics from the packaging methodology in order to preserve access to these products. The commenters expressed concern that hospitals may choose to only provide the separately payable antiemetic instead of the antiemetic that is most beneficial for the beneficiary. One commenter requested that CMS not finalize the CY 2010 proposal to apply the packaging threshold to 5-HT3 antiemetics until more information is available on the impact of packaging these products and to avoid unintended consequences, such as changes in prescribing practices, which may result from this policy.

However, several commenters expressed support for the proposed payment for 5-HT3 antiemetic products in the HOPD for CY 2010. These commenters stated that the majority of the products would be packaged under the proposal, and that would lead to reduced beneficiary copayments. The commenters offered their support due to the availability of lower-cost generic versions of some of the products and CMS' data analysis. The commenters also noted that the single product that would be paid separately under the proposal, HCPCS code J2469 (Injection, palonosetron hcl, 25 mcg), has unique properties that indicate separate payment would be appropriate.

Response: We continue to believe that use of these antiemetics is an integral part of an anticancer treatment regimen and that OPPS claims data demonstrate their increasingly common hospital outpatient utilization. As discussed above, our analysis for the CY 2010 OPPS/ASC proposed rule (74 FR 35320 through 35321) found that the most frequently used 5-HT3 antiemetic constituted 88 percent of all treatment days, and had an estimated per day cost of approximately \$1 in CY 2008. The per day costs of other 5-HT3 antiemetics with per day costs below the CY 2010 drug packaging threshold of \$65 (as discussed above) ranged from \$8 to \$51 per day. The single 5-HT3 antiemetic with a per day cost that exceeded the

CY 2010 drug packaging threshold is HCPCS code J2469. As stated in the proposed rule (74 FR 35320), we no longer believe that a specific exemption to our standard drug payment methodology is necessary for CY 2010 to ensure access to the most appropriate antiemetic product for Medicare beneficiaries. We believe that our analysis, along with the historical stability in prescribing patterns and the availability of generic alternatives for

several of these products, allows us to discontinue our policy of specifically exempting these products from the OPPS drug packaging threshold. After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to apply the CY 2010 drug packaging threshold to all 5-HT3 antiemetics. We expect that packaging will encourage hospitals to use the most cost-efficient 5-HT3

antiemetic that is clinically appropriate. We also anticipate that hospitals will continue to provide care that is aligned with the best interests of the patient. We do not believe that our CY 2010 policy to apply the drug packaging threshold to 5-HT3 antiemetics will limit beneficiaries' ability to receive clinically appropriate drugs and biologicals. The final CY 2010 OPPS status indicators for 5-HT3 antiemetics are listed in Table 34 below.

TABLE 34—FINAL CY 2010 STATUS INDICATORS FOR 5-HT3 ANTIEMETICS

CY 2010 HCPCS code	CY 2010 long descriptor	CY 2010 SI
J1260	Injection, dolasetron mesylate, 10 mg	N
J1626	Injection, granisetron hydrochloride, 100 mcg	N
J2405	Injection, ondansetron hydrochloride, per 1 mg	N
J2469	Injection, palonosetron hcl, 25 mcg	K
Q0166	Granisetron HCL, 1 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 24-hour dosage regimen	N
Q0179	Ondansetron HCL 8 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0180	Dolasetron mesylate, 100 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 24-hour dosage regimen	N

Comment: One commenter suggested that CMS institute a packaging threshold exemption for antineoplastic agents and other anticancer therapeutic agents. The commenter believed that anticancer agents, as a class, are not appropriate for packaging because of the toxicity, side effects, interactions with other drugs, and level of patient specificity associated with these therapies. The commenter requested that CMS not apply the drug packaging threshold for anticancer agents and any product that is typically used in chemotherapy supportive care regimens. Instead, the commenter requested that CMS provide separate payment for all of these products in CY 2009.

In addition, several commenters requested that CMS apply the same principle to other groups of drugs in order to equalize payment methodologies across drugs in the same clinical category. One commenter suggested that CMS institute a similar policy for anticoagulant therapies provided in the HOPD. The commenter noted that in the group of anticoagulant therapies, the majority are packaged while one drug is paid separately. The commenter was concerned that these different payment methodologies provide hospitals an incentive to use the separately payable drug, although the commenter noted that treatments are not interchangeable and that benefits vary by patient.

Response: As we discussed in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66757) and the CY 2009 OPPS/ASC final rule with comment period (73 FR 68643), as we continue to explore the possibility of additional encounter-based or episode-based payment in future years, we may consider additional options for packaging drug payment in the future. For example, a higher drug packaging threshold could eliminate existing disparities in payment methodologies for other drug groups and provide similar methods of payment across items in a group. Nevertheless, as discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68643), while we may be interested in alternative threshold methodologies for future ratesetting purposes, we realize that there are existing situations where drugs in a particular category vary in their payment treatment under the OPPS, with some drugs packaged and others separately paid.

We continue to believe the challenges associated with categorizing drugs to assess them for differences in their OPPS payment methodologies are significant, and we are not convinced that ensuring the same payment treatment for all drugs in any particular drug category is essential at this time. Therefore, we do not believe that it would be appropriate at this time to take any additional steps to ensure that all

drugs in a specific category, including anticoagulants and antineoplastic agents, are all separately paid (or, alternatively, all packaged), as requested by some commenters.

While some commenters requested that we seek feedback from interested stakeholders when the packaging threshold creates a payment methodology disparity between drugs within a single therapeutic class, we note that we provide an opportunity through the annual OPPS/ASC rulemaking cycle for public comment on the proposed packaging status of drugs and biologicals for the next calendar year. Further, we regularly accept meeting requests from interested providers and stakeholders on a variety of issues, and we address APC Panel recommendations in our annual proposed and final rules. We have often received public comments related to our proposed packaging status for particular drugs and biologicals, and we expect to continue to receive public comments regarding the proposed packaging status for drugs and biologicals in the future. In this manner, we would address specific concerns about the proposed packaging status for individual drugs and biologicals in the future, including those within a single therapeutic class where some drugs may be proposed to be packaged while others are proposed to be separately payable. While we have not defined classes of drugs that may or

may not be affected by the packaging threshold, we are accepting the APC Panel recommendation to continue to seek feedback on the proposed packaging status of all drugs under the OPSS through the annual rulemaking process. However, implicit in the APC Panel's recommendation is that we consider packaging all drugs within a therapeutic class and, as described above, we have not defined classes of drugs for consideration in the context of proposed changes to the annual drug packaging threshold.

In summary, after consideration of the public comments we received, we are finalizing our proposed CY 2010 treatment of 5-HT₃ antiemetics as follows. We are finalizing, without modification, our proposal to apply the drug packaging methodology to all 5-HT₃ antiemetics for CY 2010. In addition, we are not providing any exceptions to the standard drug packaging methodology for any class of drugs, including anticoagulants and anticancer therapies, for CY 2010. Finally, we are accepting the APC Panel recommendation to continue to consider feedback from providers, the APC Panel, and stakeholders when finalizing the packaging status of drugs and biologicals.

Having specified our standard drug packaging methodology for all drugs and biologicals makes no exceptions for different drugs and biologicals in the same therapeutic class for CY 2010, we must adopt final packaging determinations for CY 2010 for each drug and biological for this final rule with comment period. Our policy during previous cycles of the OPSS has been to use updated ASP and claims data to make final determinations of the packaging status of HCPCS codes for drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals for the final rule with comment period. We note that it is also our policy to make an annual packaging determination for a HCPCS code only when we develop the OPSS/ASC final rule for the update year. Only HCPCS codes that are identified as separately payable in the final rule with comment period are subject to quarterly updates. For our calculation of per day costs of HCPCS codes for drugs and nonimplantable biologicals in the CY 2010 OPSS/ASC final rule with comment period, we proposed to use ASP data from the first quarter of CY 2009, which is the basis for calculating payment rates for drugs and biologicals in the physician's office setting using the ASP methodology, effective July 1, 2009, along with updated hospital claims data from CY 2008. We note that we also used these

data for budget neutrality estimates and impact analyses for this CY 2010 OPSS/ASC final rule with comment period. Payment rates for HCPCS codes for separately payable drugs and nonimplantable biologicals included in Addenda A and B to this final rule with comment period are based on ASP data from the second quarter of CY 2009, which are the basis for calculating payment rates for drugs and biologicals in the physician's office setting using the ASP methodology, effective October 1, 2009. These rates would then be updated in the January 2010 OPSS update, based on the most recent ASP data to be used for physician's office and OPSS payment as of January 1, 2010. For items that do not currently have an ASP-based payment rate, we recalculated their mean unit cost from all of the CY 2008 claims data and updated cost report information available for the CY 2010 final rule with comment period to determine their final per day cost.

Consequently, the packaging status of some HCPCS codes for drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals in this CY 2010 OPSS/ASC final rule with comment period using the updated data may be different from the same drug HCPCS code's packaging status determined based on the data used for the proposed rule. Under such circumstances, in the CY 2010 OPSS/ASC proposed rule (74 FR 35320), we proposed to continue the established policies initially adopted for the CY 2005 OPSS (69 FR 65780) in order to more equitably pay for those drugs whose median cost fluctuates relative to the CY 2010 OPSS drug packaging threshold and the drug's payment status (packaged or separately payable) in CY 2009. Specifically, we proposed for CY 2010 to apply the following policies to these HCPCS codes for drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals whose relationship to the \$65 drug packaging threshold changes based on the final updated data:

- HCPCS codes for drugs and nonimplantable biologicals that were paid separately in CY 2009 and that were proposed for separate payment in CY 2010, and then have per day costs equal to or less than \$65, based on the updated ASPs and hospital claims data used for the CY 2010 final rule with comment period, would continue to receive separate payment in CY 2010.

- HCPCS codes for drugs and nonimplantable biologicals that were packaged in CY 2009 and that were proposed for separate payment in CY 2010, and then have per day costs equal

to or less than \$65, based on the updated ASPs and hospital claims data used for the CY 2010 final rule with comment period, would remain packaged in CY 2010.

- HCPCS codes for drugs and nonimplantable biologicals for which we proposed packaged payment in CY 2010 but then have per day costs greater than \$65, based on the updated ASPs and hospital claims data used for the CY 2010 final rule with comment period, would receive separate payment in CY 2010.

We did not receive any public comments on our proposal to apply the established policies initially adopted for the CY 2005 OPSS (69 FR 65780) in order to more equitably pay for those drugs whose median cost fluctuates relative to the CY 2010 OPSS drug packaging threshold and the drug's payment status (packaged or separately payable) in CY 2009. Therefore, we are finalizing our proposal, without modification, for CY 2010.

We note that HCPCS codes J1652 (Injection, fondaparinux sodium, 0.5 mg), J2430 (Injection, pamidronate disodium, per 30 mg); J7191 (Factor viii (antihemophilic factor (porcine)), per i.u.), J9165 (Injection, diethylstilbestrol diphosphate, 250 mg), and J9209 (Injection, mesna, 200 mg) were all paid separately in CY 2009 and were proposed for separate payment in CY 2010 but had final per day costs of less than the \$65 drug packaging threshold, based on the updated ASPs and the CY 2008 hospital claims data available for this CY 2010 final rule with comment period. Therefore, HCPCS codes J1652, J2430, J7191, J9165 and J9209 will continue to be paid separately in CY 2010 according to the established methodology set forth above.

In addition, we proposed to provide separate payment for HCPCS codes J2670 (Injection, tolazoline HCL, up to 25 mg) and J3320 (Injection, spectinomycin dihydrochloride, up to 2 gm) in CY 2010, although their payment was packaged in CY 2009. Using updated ASPs and the CY 2008 hospital claims data available for this final rule with comment period, HCPCS codes J2670 and J3320 now have per day costs less than \$65. In accordance with our established policy for such cases, for CY 2010 we will package payment for HCPCS codes J2670 and J3320.

Finally, we proposed to package HCPCS code Q2004 (Irrigation solution for treatment of bladder calculi, for example renacidin, per 500 ml) for CY 2010. Using updated ASPs and the CY 2008 hospital claims data available for this final rule with comment period, HCPCS code Q2004 now has a per day

cost greater than \$65. In accordance with our established policy for such cases, for CY 2010 we will pay for HCPCS code Q2004 separately.

c. Packaging Determination for HCPCS Codes That Describe the Same Drug or Biological But Different Dosages

In the CY 2008 OPPS/ASC final rule with comment period (72 FR 66776), we began recognizing, for OPPS payment purposes, multiple HCPCS codes reporting different dosages for the same covered Part B drugs or biologicals in order to reduce hospitals' administrative burden by permitting them to report all HCPCS codes for drugs and biologicals. In general, prior to CY 2008, the OPPS recognized for payment only the HCPCS code that described the lowest dosage of a drug or biological. We extended this recognition to multiple HCPCS codes for several other drugs under the CY 2009 OPPS (73 FR 68665). During CYs 2008 and 2009, we applied a policy that assigned the status indicator of the previously recognized HCPCS code to the associated newly recognized code(s), reflecting the new code(s)' packaged or separately payable status. In the CY 2008 OPPS/ASC final rule with comment period (72 FR 66775), we explained that once claims data were available for these previously unrecognized HCPCS codes, we would determine the packaging status and resulting status indicator for each HCPCS code according to the general, established HCPCS code-specific methodology for determining a code's packaging status for a given update year. However, we also stated that we planned to closely follow our claims data to ensure that our annual packaging determinations for the different HCPCS codes describing the same drug or biological did not create inappropriate payment incentives for hospitals to report certain HCPCS codes instead of others.

CY 2008 is the first year of claims data for the HCPCS codes describing different dosages of the same drug or biological that were newly recognized in CY 2008. Applying our standard HCPCS code-specific packaging determination methodology as described in the CY 2010 OPPS/ASC proposed rule (74 FR 35321 through 35323), we found that our CY 2008 claims data would result in several different packaging determinations for different codes describing the same drug or biological. Furthermore, our claims data included few units and days for a number of these newly recognized HCPCS codes, resulting in our concern that these data reflected claims from only a small number of hospitals, even though the

drug or biological itself may be reported by many other hospitals under the most common HCPCS code. We were concerned about proposing different packaging determinations for multiple HCPCS codes for the same drug or biological driven by different costs associated with the varying dosages of the same drug or biological and a small number of claims for the less common dosages that are not representative of the costs of all hospitals billing for the drug or biological. This is especially true when the general policy of the current CMS HCPCS Workgroup is to establish a single HCPCS code for a drug or biological, with a dosage that would allow accurate reporting of a patient dose for all anticipated clinical uses of the drug or biological.

Based on these findings from our first available claims data for the newly recognized HCPCS codes, in the CY 2010 OPPS/ASC proposed rule (74 FR 35321 through 35323) we explained that we believe that adopting our standard HCPCS code-specific packaging determinations for these codes could lead to payment incentives for hospitals to report certain HCPCS codes instead of others, particularly because we do not currently require hospitals to report all drug and biological HCPCS codes under the OPPS in consideration of our previous policy that generally recognized only the lowest dosage HCPCS code for a drug or biological for OPPS payment. Therefore, for CY 2010 we proposed to make packaging determinations on a drug-specific basis, rather than a HCPCS code-specific basis, for those HCPCS codes that describe the same drug or biological but different dosages. To identify all HCPCS codes for drugs and biologicals to which this proposed policy would apply, we first included the drugs and biologicals with multiple HCPCS codes that we newly recognized for payment in CY 2008 and CY 2009. We then reviewed all of the remaining drug and biological HCPCS codes to identify other drugs and biologicals for which longstanding OPPS policy recognized for payment multiple HCPCS codes for different dosages of the same drug or biological, so that our CY 2010 proposal would apply to the packaging determinations for these drugs and biologicals and their associated HCPCS codes. All of the drug and biological HCPCS codes that we proposed to be subject to this drug-specific packaging determination methodology were listed in Table 24 of the proposed rule (74 FR 35321 through 35323).

In order to propose a packaging determination that is consistent across all HCPCS codes that describe different

dosages of the same drug or biological, we aggregated both our CY 2008 claims data and our pricing information at ASP+4 percent across all of the HCPCS codes that describe each distinct drug or biological in order to determine the mean units per day of the drug or biological in terms of the HCPCS code with the lowest dosage descriptor. We then multiplied the weighted average ASP+4 percent payment amount across all dosage levels of a specific drug or biological by the estimated units per day for all HCPCS codes that describe each drug or biological from our claims data to determine the estimated per day cost of each drug or biological at less than or equal to \$65 (whereupon all HCPCS codes for the same drug or biological would be packaged) or greater than \$65 (whereupon all HCPCS codes for the same drug or biological would be separately payable). The proposed packaging status of each drug and biological HCPCS code to which this methodology would apply was displayed in Table 24 of the proposed rule (74 FR 35321 through 35323).

Comment: Several commenters supported the proposal to make packaging determinations on a drug-specific basis rather than a HCPCS code-specific basis for drugs with multiple HCPCS codes describing different dosages.

Response: We continue to believe that adopting the standard HCPCS code-specific packaging determinations for these codes could lead to payment incentives for hospitals to report certain HCPCS codes for drugs instead of others. Making packaging determinations on a drug-specific basis eliminates these incentives and allows hospitals flexibility in choosing to report all HCPCS codes for different dosages of the same drug or only the lowest dosage HCPCS code.

Therefore, after consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to make a single packaging determination for a drug, rather than an individual HCPCS code, when a drug has multiple HCPCS codes describing different dosages. For CY 2010, we have aggregated both our CY 2008 claims data and our pricing information at ASP+4 percent across all of the HCPCS codes that describe each distinct drug or biological in order to determine the mean units per day of the drug or biological in terms of the HCPCS code with the lowest dosage descriptor. We then multiplied the weighted average ASP+4 percent payment amount across all dosage levels of a specific drug or biological by the estimated units per day for all HCPCS codes that

describe each drug or biological from our claims data to determine the estimated per day cost of each drug or biological at less than or equal to \$65 (whereupon all HCPCS codes for the same drug or biological would be packaged) or greater than \$65 (whereupon all HCPCS codes for the same drug or biological would be separately payable). The final CY 2010 packaging status of each drug and biological HCPCS code to which this methodology applies is displayed in Table 35 below.

We note that new HCPCS code Q2024 (Injection, bevacizumab, 0.25 mg) was implemented effective in October 2009 and represents a different dosage descriptor for the same drug described by HCPCS code J9035 (injection, bevacizumab, 10 mg). Further, HCPCS code Q2024 has been replaced with HCPCS code C9257 (Injection, bevacizumab, 0.25 mg), effective January 1, 2010. In accordance with our CY 2010 policy to make a single packaging determination for a single drug, we are applying the methodology described above to bevacizumab and are assigning the applicable bevacizumab HCPCS codes the same packaging status for CY 2010. HCPCS codes C9257 and J9035 are included in Table 35 below.

In addition, HCPCS codes J0530 (Injection, penicillin g benzathine and penicillin g procaine, up to 600,000 units); J0540 (Injection, penicillin g benzathine and penicillin g procaine, up to 1,200,000 units); and J0550 (Injection, penicillin g benzathine and penicillin g procaine, up to 2,400,000 units), have been replaced with HCPCS code J0559 (injection, penicillin G benzathine and penicillin G procaine, 2500 units) for CY 2010. While we had proposed to treat HCPCS codes J0530, J0540 and J0550 as drugs with multiple HCPCS codes and multiple dosage descriptors via the methodology finalized above, this is no longer necessary as there is a single code for this product in CY 2010. In order to make a packaging determination for new HCPCS code J0559, we used updated hospital claims data from HCPCS codes J0530, J0540 and J0550 and ASP pricing information to determine the estimated per day cost for the drug as described above. Because the estimated per day cost was less than our CY 2010 packaging threshold of \$65, we assigned status indicator "N" to HCPCS code J0559 for CY 2010. We note that HCPCS codes J0530, J0540, and J0550 are not displayed in Table 35 below because there is only a single HCPCS code for the drug in CY 2010.

Finally, HCPCS codes J7502 (Cyclosporine, oral, 100 mg) and J7515 (Cyclosporine, oral, 25 mg) were proposed to be packaged in CY 2010 based on the methodology discussed above for drugs with multiple HCPCS codes with different dosage descriptors. As is our standard methodology, we use updated final rule data and updated ASP rates for purposes of this final rule with comment period to calculate per day estimates for final packaging determinations. Using this updated data and the multiple HCPCS code methodology discussed above, the per day cost of the drug described by HCPCS codes J7502 and J7515 would exceed the packaging threshold for CY 2010. Therefore, in accordance with the policy that was finalized in section V.B.2.b. above for HCPCS codes for drugs and nonimplantable biologicals for which we proposed packaged payment in CY 2010 but then have per day costs greater than \$65, based on the updated ASPs and hospital claims data available for the CY 2010 final rule with comment period, HCPCS codes J7502 and J7515 are separately payable in CY 2010.

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TABLE 35.—HCPCS CODES TO WHICH THE CY 2010 DRUG-SPECIFIC PACKAGING DETERMINATION METHODOLOGY APPLIES

CY 2010 HCPCS Code	CY 2010 Long Descriptor	Final CY 2010 SI
J0560	Injection, penicillin g benzathine, up to 600,000 units	N
J0570	Injection, penicillin g benzathine, up to 1,200,000 units	N
J0580	Injection, penicillin g benzathine, up to 2,400,000 units	N
J0970	Injection, estradiol valerate, up to 40 mg	N
J1020	Injection, methylprednisolone acetate, 20 mg	N
J1030	Injection, methylprednisolone acetate, 40 mg	N
J1040	Injection, methylprednisolone acetate, 80 mg	N
J1070	Injection, testosterone cypionate, up to 100 mg	N
J1080	Injection, testosterone cypionate, 1 cc, 200 mg	N
J1380	Injection, estradiol valerate, up to 10 mg	N
J1390	Injection, estradiol valerate, up to 20 mg	N
J1440	Injection, filgrastim (g-csf), 300 mcg	K
J1441	Injection, filgrastim (g-csf), 480 mcg	K
J1460	Injection, gamma globulin, intramuscular, 1 cc	K
J1470	Injection, gamma globulin, intramuscular 2 cc	K
J1480	Injection, gamma globulin, intramuscular 3 cc	K
J1490	Injection, gamma globulin, intramuscular 4 cc	K
J1500	Injection, gamma globulin, intramuscular 5 cc	K
J1510	Injection, gamma globulin, intramuscular 6 cc	K
J1520	Injection, gamma globulin, intramuscular 7 cc	K
J1530	Injection, gamma globulin, intramuscular 8 cc	K
J1540	Injection, gamma globulin, intramuscular 9 cc	K
J1550	Injection, gamma globulin, intramuscular 10 cc	K

CY 2010 HCPCS Code	CY 2010 Long Descriptor	Final CY 2010 SI
J1560	Injection, gamma globulin, intramuscular over 10 cc	K
J1642	Injection, heparin sodium, (heparin lock flush), per 10 units	N
J1644	Injection, heparin sodium, per 1000 units	N
J1840	Injection, kanamycin sulfate, up to 500 mg	N
J1850	Injection, kanamycin sulfate, up to 75 mg	N
J2270	Injection, morphine sulfate, up to 10 mg	N
J2271	Injection, morphine sulfate, 100mg	N
J2320	Injection, nandrolone decanoate, up to 50 mg	K
J2321	Injection, nandrolone decanoate, up to 100 mg	K
J2322	Injection, nandrolone decanoate, up to 200 mg	K
J2788	Injection, rho d immune globulin, human, minidose, 50 micrograms (250 i.u.)	K
J2790	Injection, rho d immune globulin, human, full dose, 300 micrograms (1500 i.u.)	K
J2920	Injection, methylprednisolone sodium succinate, up to 40 mg	N
J2930	Injection, methylprednisolone sodium succinate, up to 125 mg	N
J3120	Injection, testosterone enanthate, up to 100 mg	N
J3130	Injection, testosterone enanthate, up to 200 mg	N
J3471	Injection, hyaluronidase, ovine, preservative free, per 1 usp unit (up to 999 usp units)	N
J3472	Injection, hyaluronidase, ovine, preservative free, per 1000 usp units	N
J7030	Infusion, normal saline solution , 1000 cc	N
J7040	Infusion, normal saline solution, sterile (500 ml=1 unit)	N
J7050	Infusion, normal saline solution , 250 cc	N
J7502	Cyclosporine, oral, 100 mg	K
J7515	Cyclosporine, oral, 25 mg	K
J8520	Capecitabine, oral, 150 mg	K
J8521	Capecitabine, oral, 500 mg	K
J9035	Injection, bevacizumab, 10 mg	K
C9257	Injection, bevacizumab, 0.25 mg	K
J9060	Cisplatin, powder or solution, per 10 mg	N
J9062	Cisplatin, 50 mg	N
J9070	Cyclophosphamide, 100 mg	N
J9080	Cyclophosphamide, 200 mg	N
J9090	Cyclophosphamide, 500 mg	N
J9091	Injection, cyclophosphamide, 1.0 gram	N

CY 2010 HCPCS Code	CY 2010 Long Descriptor	Final CY 2010 SI
J9092	Cyclophosphamide, 2.0 gram	N
J9093	Cyclophosphamide, lyophilized, 100 mg	N
J9094	Cyclophosphamide, lyophilized, 200 mg	N
J9095	Cyclophosphamide, lyophilized, 500 mg	N
J9096	Cyclophosphamide, lyophilized, 1g	N
J9097	Cyclophosphamide, lyophilized, 2g	N
J9100	Injection, cytarabine, 100 mg	N
J9110	Injection, cytarabine, 500 mg	N
J9130	Dacarbazine, 100 mg	N
J9140	Injection, dacarbazine, 200 mg	N
J9250	Methotrexate sodium, 5 mg	N
J9260	Methotrexate sodium, 50 mg	N
J9280	Mitomycin, 5 mg	K
J9290	Mitomycin, 20 mg	K
J9291	Mitomycin, 40 mg	K
J9370	Vincristine sulfate, 1 mg	N
J9375	Vincristine sulfate, 2 mg	N
J9380	Vincristine sulfate, 5 mg	N
Q0164	Prochlorperazine maleate, 5 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0165	Prochlorperazine maleate, 10 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0167	Dronabinol, 2.5 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0168	Dronabinol, 5 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N

CY 2010 HCPCS Code	CY 2010 Long Descriptor	Final CY 2010 SI
Q0169	Promethazine hydrochloride, 12.5 mg, oral, FDA approved prescription anti-emetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0170	Promethazine hydrochloride, 25 mg, oral, FDA approved prescription anti-emetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0171	Chlorpromazine hydrochloride, 10 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0172	Chlorpromazine hydrochloride, 25 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0175	Perphenazine, 4 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0176	Perphenazine, 8 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0177	Hydroxyzine pamoate, 25 mg, oral, FDA approved prescription anti-emetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N
Q0178	Hydroxyzine pamoate, 50 mg, oral, FDA approved prescription antiemetic, for use as a complete therapeutic substitute for an IV antiemetic at the time of chemotherapy treatment, not to exceed a 48-hour dosage regimen	N

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d. Packaging of Payment for Diagnostic Radiopharmaceuticals, Contrast Agents, and Implantable Biologicals ("Policy-Packaged" Drugs and Devices)

Prior to CY 2008, the methodology of calculating a product's estimated per day cost and comparing it to the annual OPPS drug packaging threshold was used to determine the packaging status

of drugs, biologicals, and radiopharmaceuticals under the OPPS (except for our CYs 2005 through 2009 exemption for 5-HT₃ antiemetics). However, as established in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66766 through 66768), we began packaging payment for all diagnostic radiopharmaceuticals and contrast agents into the payment for the associated procedure, regardless of their

per day costs. In addition, in CY 2009 we adopted a policy that packaged the payment for nonpass-through implantable biologicals into payment for the associated surgical procedure on the claim (73 FR 68633 through 68636). We refer to diagnostic radiopharmaceuticals and contrast agents collectively as "policy-packaged" drugs and to implantable biologicals as devices because we proposed to treat

implantable biologicals as devices for all OPSS payment purposes beginning in CY 2010.

According to our regulations at § 419.2(b), as a prospective payment system, the OPSS establishes a national payment rate that includes operating and capital-related costs that are directly related and integral to performing a procedure or furnishing a service on an outpatient basis including, but not limited to, implantable prosthetics, implantable durable medical equipment, and medical and surgical supplies. Packaging costs into a single aggregate payment for a service, encounter, or episode-of-care is a fundamental principle that distinguishes a prospective payment system from a fee schedule. In general, packaging the costs of items and services into the payment for the primary procedure or service with which they are associated encourages hospital efficiencies and also enables hospitals to manage their resources with maximum flexibility.

Prior to CY 2008, we noted that the proportion of drugs, biologicals, and radiopharmaceuticals that were separately paid under the OPSS had increased in recent years, a pattern that we also observed for procedural services under the OPSS. Our final CY 2008 policy that packaged payment for all nonpass-through diagnostic radiopharmaceuticals and contrast agents, regardless of their per day costs, contributed significantly to expanding the size of the OPSS payment bundles and is consistent with the principles of a prospective payment system.

We believe that packaging the payment for diagnostic radiopharmaceuticals and contrast agents into the payment for their associated procedures continues to be appropriate for CY 2010. As discussed in more detail in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68645 through 68649), we presented several reasons supporting our initial policy to package payment of diagnostic radiopharmaceuticals and contrast agents into their associated procedures on a claim. Specifically, we stated that we believed packaging was appropriate because: (1) The statutory requirement that we must pay separately for drugs and biologicals for which the per day cost exceeds \$50 under section 1833(t)(16)(B) of the Act has expired; (2) we believe that diagnostic radiopharmaceuticals and contrast agents function effectively as supplies that enable the provision of an independent service; and (3) section 1833(t)(14)(A)(iii) of the Act requires that payment for specified covered

outpatient drugs (SCODs) be set prospectively based on a measure of average hospital acquisition cost. For these reasons, we believed that our proposal to continue to treat diagnostic radiopharmaceuticals and contrast agents differently from other SCODs was appropriate for CY 2010. Therefore, in the CY 2010 OPSS/ASC proposed rule (74 FR 35323), we proposed to continue packaging payment for all contrast agents and diagnostic radiopharmaceuticals, collectively referred to as "policy-packaged" drugs, regardless of their per day costs, for CY 2010.

For more information on how we proposed to set CY 2010 payment rates for nuclear medicine procedures in which diagnostic radiopharmaceuticals are used and echocardiography services provided with and without contrast agents, we refer readers to sections II.A.2.d.(5) (74 FR 35276) and (4) (74 FR 35269), respectively, of the proposed rule and this final rule with comment period.

In CY 2009 (73 FR 68634), we began packaging the payment for all nonpass-through implantable biologicals into payment for the associated surgical procedure. Because implantable biologicals may sometimes substitute for nonbiological devices, we noted that if we were to provide separate payment for implantable biologicals without pass-through status, we would potentially be providing duplicate device payment, both through the packaged nonbiological device cost already included in the surgical procedure's payment and separate biological payment. We concluded that we saw no basis for treating implantable biological and nonbiological devices without pass-through status differently for OPSS payment purposes because both are integral to and supportive of the separately paid surgical procedures in which either may be used. Therefore, in CY 2009, we adopted a final policy to package payment for all nonpass-through implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice), like our longstanding policy that packages payment for all implantable nonbiological devices without pass-through status.

For CY 2010, we continue to believe that the policy to package payment for implantable devices that are integral to the performance of separately paid procedures should also apply to payment for all implantable biologicals without pass-through status, when those biologicals function as implantable devices. Therefore, in the CY 2010 OPSS/ASC proposed rule (74 FR 35323),

we proposed to continue to package payment for nonpass-through implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) into the body, referred to as devices, in CY 2010. In accordance with this proposal, two of the products with expiring pass-through status for CY 2010 are biologicals that are solely surgically implanted according to their FDA-approved indications. These products are described by HCPCS codes C9354 (Acellular pericardial tissue matrix of non-human origin (Veritas), per square centimeter) and C9355 (Collagen nerve cuff (NeuroMatrix), per 0.5 centimeter length). Like the three implantable biologicals with expiring pass-through status in CY 2009 that were discussed in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68633 through 68634), we believe that the two biologicals specified above with expiring pass-through status for CY 2010 differ from other biologicals paid under the OPSS in that they specifically function as surgically implanted devices. As a result of the proposed CY 2010 packaged payment methodology for all nonpass-through implantable biologicals, we proposed to package payment for HCPCS codes C9354 and C9355 and assign them status indicator "N" for CY 2010. In addition, any new biologicals without pass-through status that are surgically inserted or implanted (through a surgical incision or a natural orifice) would be packaged in CY 2010. Moreover, for nonpass-through biologicals that may sometimes be used as implantable devices, we continue to instruct hospitals to not bill separately for the HCPCS codes for the products when used as implantable devices. This reporting ensures that the costs of these products that may be, but are not always, used as implanted biologicals are appropriately packaged into payment for the associated implantation procedures.

Comment: Several commenters objected to CMS' proposal to package payment for all diagnostic radiopharmaceuticals and contrast agents in CY 2010. A number of commenters stated that diagnostic radiopharmaceuticals and contrast agents with per day costs over the proposed OPSS drug packaging threshold are defined as SCODs and, therefore, should be assigned separate APC payments. In particular, the commenters questioned CMS' authority to classify groups of drugs, such as diagnostic radiopharmaceuticals and contrast agents, and implement packaging and payment policies that do

not reflect their status as SCODs. Some commenters stressed that hospitals consider radiopharmaceuticals to be drugs, rather than supplies, and that these products are not interchangeable for patients receiving specific nuclear medicine scans. The commenters recommended that diagnostic radiopharmaceuticals should be subject to the same per day cost drug packaging threshold that applies to other drugs, in order to determine whether their payment would be packaged or made separately.

In addition, the commenters objected to the proposal to package payment for diagnostic radiopharmaceuticals and contrast agents because, as SCODs, the commenters believed these products were required by statute to be paid at average acquisition cost. The commenters explained that, when several different diagnostic radiopharmaceuticals or contrast agents may be used for a particular procedure, the costs of those diagnostic radiopharmaceuticals or contrast agents are averaged together and added to the cost for the procedure in order to determine the payment rate for the associated procedural APC. Therefore, the commenters argued that the amount added to the procedure cost through packaging, representing the cost of the diagnostic radiopharmaceutical or contrast agent, did not reflect the average acquisition cost of any one particular item but, rather, reflected the average cost of whatever items may have been used with that particular procedure.

Finally, one commenter requested clarification on when CMS treats an implantable device as a biological for payment purposes.

Response: As discussed in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66766), the CY 2009 OPPS/ASC final rule with comment period (73 FR 68645) and the CY 2010 OPPS/ASC proposed rule (74 FR 35323), we continue to believe diagnostic radiopharmaceuticals and contrast agents are different from other drugs and biologicals for several reasons. We note that the statutorily required OPPS drug packaging threshold has expired, and we continue to believe that diagnostic radiopharmaceuticals and contrast agents are always ancillary and supportive to an independent service, rather than serving themselves as the therapeutic modality. We packaged their payment in CYs 2008 and 2009 as ancillary and supportive services in order to provide incentives for greater efficiency and to provide hospitals with additional flexibility in managing their

resources. In order for payment to be packaged, it is not necessary that all products be interchangeable in every case, and we recognize that in some cases hospitals may utilize higher cost products and in some cases lower cost products, taking into consideration the clinical needs of the patient and efficiency incentives. While we recognize this variability from case to case, on average under a prospective payment system we expect payment to pay appropriately for the services furnished. In the past, we have classified different groups of drugs for specific payment purposes, as evidenced by our CY 2005 through CY 2009 policy regarding 5-HT₃ antiemetics and their exemption from the drug packaging threshold. We note that we treat diagnostic radiopharmaceuticals and contrast agents as “policy-packaged” drugs because our policy is to package payment for all of the products in the category.

In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68634), we also began packaging the payment for all nonpass-through implantable biologicals into payment for the associated surgical procedure because we consider these products to always be ancillary and supportive to independent services, just like implantable nonbiological devices that are always packaged. Therefore, we currently package payment for nonpass-through implantable biologicals, also known as devices, that are surgically inserted or implanted (through a surgical incision or a natural orifice) into the body. As we stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35324), we continue to believe that payment should be packaged for nonpass-through implantable biologicals for CY 2010.

Although our final CY 2009 policy that we are continuing for CY 2010, as discussed below, packages payment for all diagnostic radiopharmaceuticals, contrast agents, and nonpass-through implantable biologicals into the payment for their associated procedures, we are continuing to provide payment for these items in CY 2010 based on a proxy for average acquisition cost just as we did in CY 2009. We continue to believe that the line-item estimated cost for a diagnostic radiopharmaceutical, contrast agent, or nonpass-through implantable biological in our claims data is a reasonable approximation of average acquisition and preparation and handling costs for diagnostic radiopharmaceuticals, contrast agents, and nonpass-through implantable biologicals, respectively. As we discussed in the CY 2009 OPPS/ASC

final rule with comment period (73 FR 68645), we believe that hospitals have adapted to the CY 2006 coding changes for radiopharmaceuticals and responded to our instructions to include charges for radiopharmaceutical handling in their charges for the radiopharmaceutical products. Further, because the standard OPPS packaging methodology packages the total estimated cost of each radiopharmaceutical, contrast agent, or nonimplantable biological on each claim (including the full range of costs observed on the claims) with the cost of associated procedures for ratesetting, this packaging approach is consistent with considering the average cost for radiopharmaceuticals, contrast agents, or nonpass-through implantable biologicals, rather than the median cost. In addition, as we noted in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68646), these drugs, biologicals, or radiopharmaceuticals for which we have not established a separate APC and, therefore, for which payment would be packaged rather than separately provided under the OPPS, could be considered to not be SCODs. Similarly, drugs and biologicals with per day costs of less than \$65 in CY 2010 that are packaged and for which a separate APC has not been established also would not be SCODs. This reading is consistent with our final payment policy whereby we package payment for diagnostic radiopharmaceuticals, contrast agents, and nonpass-through implantable biologicals and provide payment for these products through payment for their associated procedures.

Comment: Several commenters disagreed with the proposal to distinguish between diagnostic and therapeutic radiopharmaceuticals for payment purposes under the OPPS. The commenters noted that CMS’ identification of HCPCS codes A9542 (Indium In-111 ibritumomabixetan, diagnostic, per study dose, up to 5 millicuries) and A9544 (Iodine I-131 tositumomab, diagnostic, per study dose) as diagnostic radiopharmaceuticals was inappropriate because these radiopharmaceuticals function as dosimetric radiopharmaceuticals, and they both have higher than average costs associated with their acquisition and significant compounding costs in comparison with other nuclear medicine imaging agents. A few commenters explained that these radiopharmaceutical products are used as part of a therapeutic regimen and, therefore, should be considered therapeutic for OPPS payment purposes.

Response: As discussed above, and in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66641), the CY 2009 OPPS/ASC final rule with comment period (73 FR 68645) and the CY 2010 OPPS/ASC proposed rule (74 FR 35323), we classified each radiopharmaceutical into one of two groups according to whether its long descriptor contained the term “diagnostic” or “therapeutic.” HCPCS codes A9542 and A9544 both contain the term “diagnostic” in their long code descriptors. Therefore, according to our established methodology, we continue to classify them as diagnostic for the purposes of CY 2010 OPPS payment. While we understand that these items are provided in conjunction with additional supplies, imaging tests, and therapeutic radiopharmaceuticals for patients already diagnosed with cancer, we continue to believe that the purpose of administering the products described by HCPCS codes A9542 and A9544 is diagnostic in nature. As we first stated in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66641), we continue to believe that HCPCS codes A9542 and A9544 are diagnostic radiopharmaceuticals. While they are not used to diagnose disease, they are used to determine whether future therapeutic services would be beneficial to the patient and to determine how to proceed with therapy. While a group of associated services may be considered a therapeutic regimen by some commenters, HCPCS codes A9542 and A9544 are provided in conjunction with a series of nuclear medicine imaging scans. Many nuclear medicine studies using diagnostic radiopharmaceuticals are provided to patients who already have an established diagnosis. We continue to consider HCPCS codes A9542 and A9544 to be diagnostic because these items are provided for the purpose of a diagnostic imaging procedure and are used to identify the proper dose of the therapeutic agent to be provided at a later time.

Comment: Some commenters recommended using the ASP methodology to package payment for nonpass-through diagnostic radiopharmaceuticals, noting that it would be inconsistent for CMS to provide payment for diagnostic radiopharmaceuticals that have pass-through status based on the ASP methodology, and then, after the diagnostic radiopharmaceutical’s pass-through payment status expires, package the costs included in historical hospital claims data, rather than using the ASP methodology to pay for the product. The commenters believed that the ASP

methodology would be more reflective of actual diagnostic radiopharmaceutical costs and would not be subject to the billing inconsistencies that are present in hospital claims data. Therefore, the commenters concluded that it would be illogical to transition from an accurate methodology to estimate hospital costs (such as the ASP methodology) to a less accurate methodology (based on hospital claims data) once a product is packaged after its pass-through payment expires.

Response: While we understand the commenters’ request for the continued use of ASP data for purposes of packaging costs after a diagnostic radiopharmaceutical’s pass-through payment period has ended, based on their belief that ASP data are more accurate than hospital claims data, we continue to believe that hospitals have the ability to identify and set charges for any new diagnostic radiopharmaceutical product accurately during its 2 to 3 year pass-through time period while the product has the potential to be paid based on ASP. Packaging hospital costs based on hospital claims data is how the costs of all packaged items are factored into payment rates for associated procedures under the OPPS. We believe that the costs reported on claims, as determined by hospitals, are the most appropriate representation of the costs of diagnostic radiopharmaceuticals that should be packaged into payment for the associated nuclear medicine procedures.

We recognize that radiopharmaceuticals are specialized products that have unique costs associated with them. However, we believe that the costs are reflected in the charges that hospitals set for them and in the Medicare cost report where the full costs and charges associated with the services are reported. Therefore, the packaged costs of diagnostic radiopharmaceuticals are calculated like any other OPPS costs and packaged into the cost of the nuclear medicine service to which they are ancillary and supportive. This methodology is the basis for the payment of nuclear medicine procedures in the same way that other packaged costs contribute to the payment rates for the services to which they are an integral part.

Comment: Some commenters believed that packaging payment for diagnostic radiopharmaceuticals and nonpass-through implantable biologicals would undermine the clinical and resource homogeneity in the various procedural APCs, resulting in 2 times violations.

Response: As we stated in the CY 2009 OPPS/ASC final rule with

comment period (73 FR 68647), we agree that packaging the costs of ancillary and supportive services into the cost of an independent service can change the median cost for that service and could result in 2 times violations. However, we disagree that we should refrain from packaging payment for ancillary and supportive items into the payment for the service in which they are used in order to prevent the occurrence of 2 times violations. Instead, we believe that we should reconfigure APCs when necessary to resolve 2 times violations where they occur. As we discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68647), because we have traditionally paid for a service package under the OPPS as represented by a HCPCS code for the major procedure that is assigned to an APC group for payment, we assess the applicability of the 2 times rule to services at the HCPCS code level, not at a more specific level based on the individual diagnostic radiopharmaceuticals or nonpass-through implantable biologicals that may be utilized in a service reported with a single HCPCS code. Furthermore, if the use of a very expensive diagnostic radiopharmaceutical in a clinical scenario causes a specific procedure to be much more expensive for the hospital than the APC payment, we consider such a case to be the natural consequence of a prospective payment system that anticipates that some cases will be more costly and others less costly than the procedure payment. This same logic would apply to situations in which a nonpass-through implantable biological is implanted in a surgical procedure and results in an increase in a procedure’s cost to the hospital for an individual case. In addition, very high cost cases could be eligible for outlier payment. As we note elsewhere in this final rule with comment period, decisions about packaging and bundling payment involve a balance between ensuring some separate payment for individual services and establishing incentives for efficiency through larger units of payment. In the case of diagnostic radiopharmaceuticals and nonpass-through implantable biologicals, these products are part of the OPPS payment package for the procedures in which they are used. We refer readers to section II.A.d.(5) of this final rule with comment period for a discussion of payment for nuclear medicine procedures.

Comment: One commenter recommended that CMS provide separate payment for all diagnostic radiopharmaceuticals with a median per

day cost greater than \$200. The commenter believes that this recommendation is most consistent with the APC Panel's recommendation to CMS at the September 2007 APC Advisory Panel meeting.

Response: At the September 2007 APC Panel meeting, the APC Panel recommended that CMS package radiopharmaceuticals with a median per day cost of less than \$200 but pay separately for radiopharmaceuticals with a per day cost of \$200 or more. In the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66638), we did not accept the APC Panel's recommendation, citing an inability to determine an empirical basis for paying separately for radiopharmaceuticals with a per day cost in excess of \$200. Instead, as proposed, for CY 2008 we finalized the packaging of payment for all diagnostic radiopharmaceuticals. We continue to believe that diagnostic radiopharmaceuticals are ancillary and supportive to the nuclear medicine procedures in which they are used and that their costs should be packaged into the primary procedures with which they are associated. We do not believe it would be appropriate to set a cost threshold for packaging diagnostic radiopharmaceuticals because, regardless of their per day cost, they are always supportive of an independent procedure that is the basis for administration of the diagnostic radiopharmaceutical.

After consideration of the public comments we received, we are finalizing our CY 2010 proposals, without modification, to continue to package payment for all nonpass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals that are surgically inserted or implanted into the body, regardless of their per day costs. Given the inherent function of contrast agents and diagnostic radiopharmaceuticals as ancillary and supportive to the performance of an independent procedure and the similar functions of implantable biological and nonbiological devices, we continue to view the packaging of payment for contrast agents, diagnostic radiopharmaceuticals, and implantable biologicals as a logical expansion of packaging payment for drugs and biologicals. In addition, as we initially established in the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66768), we are finalizing our proposal to continue to identify diagnostic radiopharmaceuticals specifically as those Level II HCPCS codes that include the term "diagnostic" along with a radiopharmaceutical in their long code

descriptors, and therapeutic radiopharmaceuticals as those Level II HCPCS codes that include the terms "therapeutic" along with a radiopharmaceutical in their long code descriptors. For more information on how we set CY 2010 payment rates for nuclear medicine procedures in which diagnostic radiopharmaceuticals are used and echocardiography services provided with and without contrast agents, we refer readers to section II.A.2.d (5) and (4), respectively, of this final rule with comment period.

3. Payment for Drugs and Biologicals Without Pass-Through Status That Are Not Packaged

a. Payment for Specified Covered Outpatient Drugs (SCODs) and Other Separately Payable and Packaged Drugs and Biologicals

Section 1833(t)(14) of the Act defines certain separately payable radiopharmaceuticals, drugs, and biologicals and mandates specific payments for these items. Under section 1833(t)(14)(B)(i) of the Act, a "specified covered outpatient drug" is a covered outpatient drug, as defined in section 1927(k)(2) of the Act, for which a separate APC has been established and that either is a radiopharmaceutical agent or is a drug or biological for which payment was made on a pass-through basis on or before December 31, 2002.

Under section 1833(t)(14)(B)(ii) of the Act, certain drugs and biologicals are designated as exceptions and are not included in the definition of "specified covered outpatient drugs," known as SCODs. These exceptions are—

- A drug or biological for which payment is first made on or after January 1, 2003, under the transitional pass-through payment provision in section 1833(t)(6) of the Act.
- A drug or biological for which a temporary HCPCS code has not been assigned.
- During CYs 2004 and 2005, an orphan drug (as designated by the Secretary).

Section 1833(t)(14)(A)(iii) of the Act requires that payment for SCODs in CY 2006 and subsequent years be equal to the average acquisition cost for the drug for that year as determined by the Secretary, subject to any adjustment for overhead costs and taking into account the hospital acquisition cost survey data collected by the Government Accountability Office (GAO) in CYs 2004 and 2005. If hospital acquisition cost data are not available, the law requires that payment be equal to payment rates established under the methodology described in section

1842(o), section 1847A, or section 1847B of the Act, as calculated and adjusted by the Secretary as necessary.

Section 1833(t)(14)(E) of the Act provides for an adjustment in OPPTS payment rates for overhead and related expenses, such as pharmacy services and handling costs. Section 1833(t)(14)(E)(i) of the Act required MedPAC to study pharmacy overhead and to make recommendations to the Secretary regarding whether, and if so how, a payment adjustment should be made to compensate hospitals for them. Section 1833(t)(14)(E)(ii) of the Act authorizes the Secretary to adjust the weights for ambulatory procedure classifications for SCODs to take into account the findings of the MedPAC study.

In the CY 2006 OPPTS proposed rule (70 FR 42728), we discussed the June 2005 report by MedPAC regarding pharmacy overhead costs in HOPDs and summarized the findings of that study:

- Handling costs for drugs, biologicals, and radiopharmaceuticals administered in the HOPD are not insignificant;
- Little information is available about the magnitude of pharmacy overhead costs;
- Hospitals set charges for drugs, biologicals, and radiopharmaceuticals at levels that reflect their respective handling costs; and
- Hospitals vary considerably in their likelihood of providing services which utilize drugs, biologicals, or radiopharmaceuticals with different handling costs.

As a result of these findings, MedPAC developed seven drug categories for pharmacy and nuclear medicine handling costs based on the estimated level of hospital resources used to prepare the products (70 FR 42729). Associated with these categories were two recommendations for accurate payment of pharmacy overhead under the OPPTS.

1. CMS should establish separate, budget neutral payments to cover the costs hospitals incur for handling separately payable drugs, biologicals, and radiopharmaceuticals.

2. CMS should define a set of handling fee APCs that group drugs, biologicals, and radiopharmaceuticals based on attributes of the products that affect handling costs; CMS should instruct hospitals to submit charges for these APCs and base payment rates for the handling fee APCs on submitted charges reduced to costs.

In response to the MedPAC findings, in the CY 2006 OPPTS proposed rule (70 FR 42729), we discussed our belief that, because of the varied handling resources

required to prepare different forms of drugs, it would be impossible to exclusively and appropriately assign a drug to a certain overhead category that would apply to all hospital outpatient uses of the drug. Therefore, our CY 2006 OPPS proposal included a proposal to establish three distinct Level II HCPCS C-codes and three corresponding APCs for drug handling categories to differentiate overhead costs for drugs and biologicals (70 FR 42730). We also proposed: (1) To combine several overhead categories recommended by MedPAC; (2) to establish three drug handling categories, as we believed that larger groups would minimize the number of drugs that may fit into more than one category and would lessen any undesirable payment policy incentives to utilize particular forms of drugs or specific preparation methods; (3) to collect hospital charges for these HCPCS C-codes for 2 years; and (4) to ultimately base payment for the corresponding drug handling APCs on CY 2006 claims data available for the CY 2008 OPPS.

In the CY 2006 OPPS final rule with comment period (70 FR 68659 through 68665), we discussed the public comments we received on our proposal regarding pharmacy overhead. The overwhelming majority of commenters did not support our proposal and urged us not to finalize this policy, as it would be administratively burdensome for hospitals to establish charges for HCPCS codes for pharmacy overhead and to report them. Therefore, we did not finalize this proposal for CY 2006. Instead, we established payment for separately payable drugs and biologicals at ASP+6 percent, which we calculated by comparing the estimated aggregate cost of separately payable drugs and biologicals in our claims data to the estimated aggregate ASP dollars for separately payable drugs and biologicals, using the ASP as a proxy for average acquisition cost (70 FR 68642). Hereinafter, we refer to this methodology as our standard drug payment methodology. We concluded that payment for drugs and biologicals and pharmacy overhead at a combined ASP+6 percent rate would serve as the best proxy for the combined acquisition and overhead costs of each of these products.

In the CY 2007 OPPS/ASC final rule with comment period (71 FR 68091), we finalized our proposed policy to provide a single payment of ASP+6 percent for the hospital's acquisition cost for the drug or biological and all associated pharmacy overhead and handling costs. The ASP+6 percent rate that we finalized was higher than the equivalent average ASP-based amount calculated

from claims of ASP+4 percent according to our standard drug payment methodology, but we adopted payment at ASP+6 percent for stability while we continued to examine the issue of the costs of pharmacy overhead in the HOPD.

In the CY 2008 OPPS/ASC proposed rule (72 FR 42735), in response to ongoing discussions with interested parties, we proposed to continue our methodology of providing a combined payment rate for drug and biological acquisition and pharmacy overhead costs. We also proposed to instruct hospitals to remove the pharmacy overhead charge for both packaged and separately payable drugs and biologicals from the charge for the drug or biological and report the pharmacy overhead charge on an uncoded revenue code line on the claim. We believed that this would provide us with an avenue for collecting pharmacy handling cost data specific to drugs in order to package the overhead costs of these items into the associated procedures, most likely drug administration services. Similar to the public response to our CY 2006 pharmacy overhead proposal, the overwhelming majority of commenters did not support our CY 2008 proposal and urged us to not finalize this policy (72 FR 66761). At its September 2007 meeting, the APC Panel recommended that hospitals not be required to separately report charges for pharmacy overhead and handling and that payment for overhead be included as part of drug payment. The APC Panel also recommended that CMS continue to evaluate alternative methods to standardize the capture of pharmacy overhead costs in a manner that is simple to implement at the organizational level (72 FR 66761). Because of concerns expressed by the APC Panel and public commenters, we did not finalize the proposal to instruct hospitals to separately report pharmacy overhead charges for CY 2008. Instead, in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66763), we finalized a policy of providing payment for separately payable drugs and biologicals and their pharmacy overhead at ASP+5 percent as a transition from their CY 2007 payment of ASP+6 percent to payment based on the equivalent average ASP-based payment rate calculated from hospital claims according to our standard drug payment methodology, which was ASP+3 percent for the CY 2008 OPPS/ASC final rule with comment period. Hospitals continued to include charges for pharmacy overhead costs in the line-

item charges for the associated drugs reported on claims.

For CY 2009, we proposed to pay separately payable drugs and biologicals at ASP+4 percent, including both SCODs and other drugs without CY 2009 OPPS pass-through status, based on our standard drug payment methodology, and we also proposed to split the "Drugs Charged to Patients" cost center into two cost centers: One for drugs with high pharmacy overhead costs and one for drugs with low pharmacy overhead costs (73 FR 41492). We noted that we expected that CCRs from the proposed new cost centers would be available in 2 to 3 years to refine OPPS drug cost estimates by accounting for differential hospital markup practices for drugs with high and low overhead costs. After consideration of the public comments received and the APC Panel recommendations, we finalized a CY 2009 policy (73 FR 68659) to provide payment for separately payable nonpass-through drugs and biologicals based on costs calculated from hospital claims at a 1-year transitional rate of ASP+4 percent, in the context of an equivalent average ASP-based payment rate of ASP+2 percent calculated according to our standard drug payment methodology from the final rule claims and cost report data. We did not finalize our proposal to split the single standard "Drugs Charged to Patients" cost center into two cost centers largely due to concerns raised to us by hospitals about the associated administrative burden. Instead, we indicated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68659) that we would continue to explore other potential approaches to improve our drug cost estimation methodology, thereby increasing payment accuracy for separately payable drugs and biologicals.

In response to the CMS proposals for the CY 2008 and CY 2009 OPPS, a group of pharmacy stakeholders (hereinafter referred to as the pharmacy stakeholders), including some cancer hospitals, some pharmaceutical manufacturers, and some hospital and professional associations, commented that CMS should pay an acquisition cost of ASP+6 percent for separately payable drugs, should substitute ASP+6 percent for the packaged cost of all packaged drugs and biologicals on procedure claims, and should redistribute the difference between the aggregate estimated packaged drug cost in claims and payment for all drugs, including packaged drugs at ASP+6 percent, as separate pharmacy overhead payments for separately payable drugs. They

indicated that this approach would preserve the aggregate drug cost observed in the claims data, while significantly increasing payment accuracy for individual drugs and procedures using packaged drugs. Their suggested approach would provide a separate overhead payment for each separately payable drug or biological at one of three different levels, depending on the pharmacy stakeholders' assessment of the complexity of pharmacy handling associated with each specific drug or biological (73 FR 68651 through 68652). Each separately payable drug or biological HCPCS code would be assigned to one of the three overhead categories, and the separate pharmacy overhead payment applicable to the category would be made when each of the separately payable drugs or biologicals was paid.

At the February 2009 APC Panel meeting, the APC Panel recommended that CMS pay for the acquisition cost of all separately payable drugs at no less than ASP+6 percent. The APC Panel also recommended that CMS package payment at ASP+6 percent on claims for all drugs that are not separately payable and use the difference between these rates and CMS' cost derived from charges to create a pool to provide more appropriate payment for pharmacy service costs and that CMS pay for pharmacy services costs using this pool, applying a tiered approach to payments based on some objective criteria related to the pharmacy resources required for groups of drugs. The APC Panel further recommended that, if CMS does not implement the drug payment recommendations specified above, CMS should exclude data from hospitals that participate in the 340B Federal drug pricing program from its ratesetting calculations for drugs and CMS should pay 340B hospitals in the same manner as it pays non-340B hospitals. Hospitals that participate in the 340B program are generally hospitals that serve a disproportionate share of low-income patients and receive disproportionate share payments under the IPPS. These facilities may acquire outpatient drugs and biologicals at prices that are substantially below ASP because the 340B program requires drug manufacturers to provide outpatient drugs to eligible entities at a reduced price and these reduced price sales are not included in the ASP submissions of manufacturers to Medicare. Public presenters at the February 2009 APC Panel meeting emphasized that the purpose of the 340B Federal drug pricing program is to ensure access to drugs for low-income patients by

supplementing the higher cost of providing care to low-income patients born by hospitals serving a disproportionate share of these patients. The agenda, recommendations, and report from the February 2009 APC Panel meeting are posted on the CMS Web site at: <http://www.cms.hhs.gov/FACA>. We respond to these APC Panel recommendations in our discussion of the proposed CY 2010 policy that follows.

b. Payment Policy

Section 1833(t)(14)(A)(iii) of the Act, as described above, continues to be applicable to determining payments for SCODs for CY 2010. This provision requires that payment for SCODs be equal to the average acquisition cost for the drug for that year as determined by the Secretary, subject to any adjustment for overhead costs and taking into account the hospital acquisition cost survey data collected by the GAO in CYs 2004 and 2005. If hospital acquisition cost data are not available, the law requires that payment be equal to payment rates established under the methodology described in section 1842(o), section 1847A, or section 1847B of the Act, as calculated and adjusted by the Secretary as necessary. In addition, section 1833(t)(14)(E)(ii) of the Act authorizes the Secretary to adjust APC weights to take into account the 2005 MedPAC report relating to overhead and related expenses, such as pharmacy services and handling costs. Since CY 2006, when we first adopted our standard methodology of paying for separately payable drugs and biologicals based on the equivalent average ASP-based payment rate calculated from claims and cost report data, we have applied this methodology to payment for all separately payable drugs and biologicals without pass-through status, both SCODs and other drugs and biologicals that do not meet the statutory definition of SCODs. We have seen no reason to distinguish SCODs from these other separately payable drugs and biologicals, and under our standard drug payment methodology, we have used ASP data and costs estimated from charges on hospital claims data as a proxy for the average hospital acquisition cost that the statute requires for payment of SCODs and to provide payment for the associated pharmacy overhead cost.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35332), we proposed to redistribute between one-third and one-half of the difference between the aggregate claims cost for coded packaged drugs and biologicals with an ASP and ASP dollars for those products,

which resulted in our proposal to pay for the acquisition and pharmacy overhead costs of separately payable drugs and biologicals that did not have pass-through payment status at ASP+4 percent. Based on the rationale described in the CY 2010 OPPS/ASC proposed rule (74 FR 35326 through 35333), we believed that approximately \$150 million of the estimated \$395 million total in pharmacy overhead cost included in our claims data for coded packaged drugs and biologicals with an ASP above the aggregate ASP dollars of these packaged products should be attributed to separately payable drugs and biologicals to provide an adjustment for the pharmacy overhead costs of these separately payable products. As a result, we also proposed to reduce the cost of these packaged drugs and biologicals that is included in the payment for procedural APCs to offset the \$150 million adjustment to payment for separately payable drugs and biologicals. In addition, we proposed that any redistribution of pharmacy overhead cost that may arise from CY 2010 final rule data would occur only from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals, thereby maintaining the estimated total cost of drugs and biologicals.

Using our CY 2010 proposed rule data, and applying our longstanding methodology for calculating the total cost of separately payable drugs and biologicals in our claims compared to the ASP dollars for the same drugs and biologicals, without applying the proposed overhead cost redistribution, we determined that the estimated aggregate cost of separately payable drugs and biologicals (status indicators "K" and "G"), including acquisition and pharmacy overhead costs, was equivalent to ASP-2 percent. Therefore, under our standard drug payment methodology, we would pay for separately payable drugs and biologicals at ASP-2 percent for CY 2010, their equivalent average ASP-based payment rate. We also determined that the estimated aggregate cost of coded packaged drugs and biologicals with an ASP (status indicator "N"), including acquisition and pharmacy overhead costs, was equivalent to ASP+247 percent. We found that the estimated aggregate cost for all coded drugs and biologicals (status indicators "N," "K," and "G"), including acquisition and pharmacy overhead costs, was equivalent to ASP+13 percent. For a detailed explanation of our standard process for these calculations, we refer

readers to the CY 2006 OPPTS proposed rule (70 FR 42725).

TABLE 36—STANDARD DRUG PAYMENT METHODOLOGY USING CY 2010 OPPTS PROPOSED RULE DATA: ASP+X CALCULATION

	Total ASP dollars for drugs and biologicals in claims data (in millions)*	Total cost of drugs and biologicals in claims data (in millions)**	Ratio of cost to ASP	ASP+X percent
Coded Packaged Drugs and Biologicals with an ASP	\$160	\$555	3.47	ASP+247
Separately Payable Drugs and Biologicals	2,589	2,539	0.98	ASP-2
All Coded Drugs and Biologicals	2,749	3,094	1.13	ASP+13

*Total April 2009 ASP dollars (ASP multiplied by drug or biological units in CY 2008 claims) for drugs and biologicals with a HCPCS code and ASP information.

**Total cost in the CY 2008 claims data for drugs and biologicals with a HCPCS code and April 2009 ASP information.

In the proposed rule, we recognized that there may be concern over whether the actual full cost (acquisition and pharmacy overhead) of separately payable drugs and biologicals could be 2 percent less than ASP for these products (74 FR 35327), although we did not have ASP information specifically for their sales to hospitals. Similarly, we acknowledged that a full cost (acquisition and pharmacy overhead) of ASP+247 percent for coded packaged drugs and biologicals with an ASP seemed relatively high. When we subtracted the total ASP dollars for packaged drugs and biologicals with a reported ASP amount in the CY 2008 claims data (\$160 million), our proxy for their acquisition cost, from the total cost of packaged drugs and biologicals in the same claims (\$555 million), we found that the difference, which we viewed as the pharmacy overhead cost currently attributed to packaged drugs and biologicals was \$395 million. While we had no way of assessing whether this current distribution of overhead cost to coded packaged drugs and biologicals with an ASP was appropriate, we acknowledged that the established method of converting billed charges to costs had the potential to “compress” the calculated costs to some degree. Further, we recognized that the attribution of pharmacy overhead costs to packaged or separately payable drugs and biologicals through our standard drug payment methodology of a combined payment for acquisition and pharmacy overhead costs depends, in part, on the treatment of all drugs and biologicals each year under our annual drug packaging threshold. Changes to the packaging threshold may result in changes to payment for the overhead cost of drugs and biologicals that do not reflect actual changes in hospital pharmacy overhead cost for those products. For these reasons, we believed that some portion, but not all, of the

\$395 million in total overhead cost that is associated with coded packaged drugs and biologicals with an ASP based on our standard drug payment methodology should, at least for CY 2010, be attributed to separately payable drugs and biologicals. Although we believed that for CY 2010 it would be prudent to redistribute some pharmacy overhead cost between coded packaged drugs and biologicals with an ASP at ASP+247 percent and separately payable drugs and biologicals at ASP-2 percent that would result from our standard drug payment methodology, the amount of overhead cost redistribution that would be appropriate between the packaged and separately payable drugs and biologicals in a payment system that is fundamentally based on averages was not fully evident. Pharmacy overhead cost includes, but is not limited to, some costs of indirect overhead that are shared by all hospital items and services, such as administrative and general costs, capital costs, staff benefits, and other facility costs. With regard to these indirect overhead costs, the amount of indirect overhead cost that is attributable to an inexpensive (typically packaged) drug is the same in dollar value as the amount of indirect overhead cost that is attributable to an extremely costly drug (typically separately payable). Hence, the indirect overhead costs that are common to all drugs and biologicals have no relationship to the cost of an individual drug or biological, or to the complexity of the handling, preparation, or storage of that individual drug or biological. Therefore, we believed that the indirect overhead cost alone for an inexpensive drug or biological could be far in excess of the ASP for that inexpensive product.

Layered on these indirect overhead costs are the pharmacy overhead direct costs of staff, supplies, and equipment that are directly attributable only to the

storage, handling, preparation, and distribution of drugs and biologicals and which do vary, sometimes considerably, depending upon the drug being furnished. As we describe above, in its June 2005 Report to Congress, MedPAC found that drugs can be categorized into seven different categories based on the handling costs (that is, the direct costs) incurred (70 FR 42729). Similarly, the pharmacy stakeholders, whose suggested approach the APC Panel recommended that we accept for CY 2010, identified three categories of pharmacy overhead complexity with variable costs, to which they assigned individual drugs and biologicals for purposes of implementing their recommended redistribution of the difference between aggregate dollars for all drugs and biologicals at ASP+6 percent and aggregate cost for all drugs and biologicals in the claims data as additional pharmacy overhead payments.

In the CY 2010 OPPTS/ASC proposed rule (74 FR 35328), we acknowledged that the observed combined payment for acquisition and pharmacy overhead costs of ASP-2 percent for separately payable drugs and biologicals may be too low and ASP+247 percent for coded packaged drugs and biologicals with an ASP in the CY 2010 claims data may be too high. However, we stated our belief that the pharmacy stakeholders’ recommendation to set packaged drug and biologicals dollars to ASP+6 percent was inappropriate given our understanding that an equal allocation of indirect overhead costs among packaged and separately payable drugs and biologicals would lead to a higher observed ASP+X percent than ASP+6 percent for packaged drugs and biologicals. As discussed above, the indirect overhead costs that are common to all drugs and biologicals have no relationship to the cost of an individual drug or biological, or to the complexity

of the handling, preparation, or storage of that individual drug or biological. Therefore, we stated our belief that the indirect overhead cost alone for an inexpensive drug or biological which would be packaged could be far in excess of the ASP for that inexpensive product. In contrast, we would expect that the indirect overhead cost alone for an expensive drug or biological which would be separately paid could be far less than the ASP for that expensive product.

Therefore, as discussed in the proposed rule, we believed that some middle ground would represent the most accurate redistribution of pharmacy overhead cost. We stated that the assumption that approximately one-third to one-half of the total pharmacy overhead cost currently associated with coded packaged drugs and biologicals with an ASP was a function of both charge compression and our choice of

an annual drug packaging threshold offered a more appropriate allocation of drug and biological cost to separately payable drugs and biologicals. One-third of the \$395 million of pharmacy overhead cost associated with coded packaged drugs and biologicals with an ASP was \$132 million, whereas one-half was \$198 million. Within the one-third to one-half parameters, we proposed that reallocating \$150 million in drug and biological cost observed in the claims data from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals for CY 2010 would more appropriately distribute pharmacy overhead cost among packaged and separately payable drugs and biologicals than either of the two other options, that is, paying for separately payable drugs and biologicals at ASP-2 percent according to our standard drug payment methodology or adopting the pharmacy stakeholders'

recommendation. We stated that if we attributed \$150 million in additional cost to the payment for the drugs and biologicals for which we proposed to pay separately for the CY 2010 OPPS, we determined a payment rate for separately payable drugs and biologicals of ASP+4 percent as displayed in Table 26 of the proposed rule (74 FR 35328) that is reprinted below as Table 37. Thus, we proposed a pharmacy overhead adjustment for separately payable drugs and biologicals in CY 2010 that would result in their payment at ASP+4 percent. We proposed to accomplish this adjustment by redistributing one-third to one-half of the pharmacy overhead cost of coded packaged drugs and biologicals with an ASP (\$150 million), which represented a reduction in cost of coded packaged drug and biologicals with an ASP in the CY 2010 proposed rule claims data of 27 percent.

TABLE 37—CY 2010 PROPOSED PHARMACY OVERHEAD ADJUSTMENT PAYMENT METHODOLOGY FOR SEPARATELY PAYABLE AND PACKAGED DRUGS AND BIOLOGICALS

	Total ASP dollars for drugs and biologicals in claims data (in millions)*	Total cost of drugs and biologicals in claims Data After Adjustment (in millions)**	Ratio of cost to ASP (column C/column B)	ASP+X percent
Coded Packaged Drugs and Biologicals with an ASP	\$160	\$405	2.53	ASP+153
Separately Payable Drugs and Biologicals	2,589	2,689	1.04	ASP+4
All Coded Drugs and Biologicals	2,749	3,094	1.13	ASP+13

*Total April 2009 ASP dollars (ASP multiplied by drug or biological units in CY 2008 claims) for drugs and biologicals with a HCPCS code and ASP information.

**Total cost in the CY 2008 claims data for drugs and biologicals with a HCPCS code and April 2009 ASP information.

Comment: Many commenters agreed with CMS that it was unlikely that the full cost (acquisition and pharmacy overhead) of separately payable drugs and biologicals could be 2 percent less than ASP for these products, and that the full cost (acquisition and pharmacy overhead) of packaged drugs could be 247 percent of ASP.

Response: We continue to find that the results of our standard drug payment methodology are unlikely to accurately reflect the full cost of acquisition and pharmacy overhead for separately payable drugs and biologicals and packaged drugs and biologicals due to hospital charging practices and our use of an annual drug packaging threshold. Using our CY 2010 final rule

data, and applying our longstanding methodology for calculating the total cost of separately payable drugs and biologicals in our claims compared to the ASP dollars for the same drugs and biologicals and without applying the proposed overhead cost redistribution, we determined that the estimated aggregate cost of separately payable drugs and biologicals (status indicators "K" and "G"), including acquisition and pharmacy overhead costs, is equivalent to ASP-3 percent (compared to ASP-2 percent as presented in the proposed rule). Therefore, under our standard drug payment methodology, we would pay for separately payable drugs and biologicals at ASP-3 percent for CY 2010, their equivalent average ASP-

based payment rate. We also determined that the estimated aggregate cost of coded packaged drugs and biologicals with an ASP (status indicator "N"), including acquisition and pharmacy overhead costs, is equivalent to ASP+259 percent (compared to ASP+247 percent as presented in the proposed rule). We found that the estimated aggregate cost for all coded drugs and biologicals (status indicators "N," "K," and "G), including acquisition and pharmacy overhead costs, is equivalent to ASP+11 percent (compared to ASP+13 percent as presented in the proposed rule). These values are shown in Table 38 below.

TABLE 38—PROPOSED AND FINAL ASP+X VALUES FOR ALL CODED DRUGS AND BIOLOGICALS WITH AN ASP, CODED PACKAGED DRUGS AND BIOLOGICALS WITH AN ASP, AND SEPARATELY PAYABLE DRUGS AND BIOLOGICALS

	ASP+X for all coded drugs and biologicals with an ASP	ASP+X for coded packaged drugs and biologicals with an ASP	ASP+X for separately payable drugs and biologicals
CY 2010 Proposed Rule*	ASP+13	ASP+247	ASP-2
CY 2010 Final Rule**	ASP+11	ASP+258	ASP-3

*Based on CY 2010 proposed rule claims data and April 2009 ASPs.
 **Based on CY 2010 final rule claims data and July 2009 ASPs.

Comment: Some commenters agreed with CMS’ assertion that packaged drugs and biologicals typically have an aggregate absolute pharmacy overhead cost that exceeds the acquisition cost of the packaged drugs and biologicals. One commenter claimed that ASP+6 percent would be insufficient to accurately account for the acquisition and pharmacy overhead costs of packaged drugs. In addition, a few commenters recommended that CMS modify the pharmacy stakeholders’ approach by packaging the cost of drugs and biologicals at ASP plus 100 percent, rather than the stakeholder’s recommended amount of ASP+6 percent.

Response: We continue to be concerned with a methodology that would package the cost of all packaged drugs and biologicals at ASP+6 percent. As stated in our proposal, we have no data specific to the overhead costs of these drugs and biologicals and, therefore, we cannot determine with any certainty an ASP+X value relating specifically to their costs for acquisition and pharmacy overhead. While we appreciate the recommendation of the commenters to package payment for the acquisition and pharmacy overhead costs of inexpensive drugs below the drug packaging threshold at ASP plus 100 percent, we cannot verify that using ASP plus 100 percent would result in an accurate estimate of acquisition and pharmacy overhead costs of packaged drugs and biologicals.

Comment: One commenter noted that a key CMS assumption behind the standard methodology for calculating the ASP+X percent payment rate, and the redistribution methodology by implication, is that the average overhead cost for drugs and biologicals ultimately must appear in the revenue producing cost center 5600 “Drugs Charged to Patients,” along with the acquisition cost of drugs and biologicals. The commenter acknowledged that CMS’ standard drug payment methodology relies on appropriate allocation of pharmacy overhead cost in order to

derive an accurate payment amount for separately payable versus packaged drugs and biologicals. The commenter specifically cited weak cost reporting instructions for the “Drugs Charged to Patients” cost center and cost center 1600 “Pharmacy” and questioned whether pharmacy overhead cost adequately and appropriately appears in the “Drugs Charged to Patients” cost center. Specifically, the commenter noted that costs accumulated in the “Drugs Charged to Patients” cost center remain in that cost center, but asserted that costs accumulated in the 1600 “Pharmacy” cost center would be allocated across revenue producing cost centers on the basis of costed requisitions. The commenter asked a series of specific questions about how costs are accumulated in both the “Pharmacy” and “Drugs Charged to Patients” cost centers, including (1) what costs hospitals actually report in the “Pharmacy” cost center versus the “Drugs Charged to Patients” cost center; (2) which revenue producing cost centers have costed requisitions that would receive “Pharmacy” cost center overhead costs, that is, do hospitals account for contrast agent costs under radiology revenue producing cost centers; (3) how much of “Pharmacy” cost center overhead costs is allocated to “Drugs Charged to Patients;” and (4) when would hospitals not account for the cost of a drug in the “Drugs Charged to Patients” cost center.

Response: We acknowledge that the CCR for the “Drugs Charged to Patients” cost center reflects the average acquisition cost for drugs and biologicals reported in that cost center, as well as the average pharmacy overhead cost for those drugs and biologicals. In addition, use of this CCR to estimate costs from charges on claims has the potential to “compress” the pharmacy overhead costs for expensive drugs and biologicals, where hospitals differentially distribute pharmacy overhead among their charges for drugs and biologicals by marking up the charges for expensive drugs and

biologicals proportionally less than inexpensive drugs and biologicals. We have stated that combining this compression with a packaging threshold may lead us to underestimate the pharmacy overhead costs of separately payable drugs and biologicals and overestimate the overhead costs of packaged drugs and biologicals.

At a minimum, the CCR for the cost center “Drugs Charged to Patients” that CMS uses to estimate costs from charges for drugs and biologicals in our claims data should consist of charges for all drugs and biologicals separately charged to patients and the related costs for those separately chargeable drugs and biologicals. A hospital would not include the charge and cost for a drug or biological in the “Drugs Charged to Patients” cost center if the hospital did not separately charge the drug or biological to a patient. The identification of costs for the “Drugs Charged to Patients” cost center can occur in several ways.

First, we generally believe that the indirect costs that are common to all drugs and biologicals, including administrative and general costs, capital costs, staff benefits, and other facility costs, and the more direct costs of handling, preparation, and storage are accumulated as total pharmacy operation costs in cost center 1600 “Pharmacy.” Second, hospitals can choose to treat the acquisition cost of their drugs and biologicals in several ways. Frequently, hospitals accumulate and report the acquisition costs of drugs and biologicals (costed requisitions) directly in the most appropriate revenue producing cost center on Worksheet A of the cost report. We expect that, the majority of the time, hospitals accrue the largest acquisition cost of drugs and biologicals in the “Drugs Charged to Patients” cost center, which is specific to these items. However, hospitals may also account for the acquisition cost of unique drugs and biologicals, such as contrast agents, in other revenue producing cost centers such as the radiology cost centers, when the

contrast agents are not separately charged to the patient. Assignment of acquisition cost to a different revenue producing cost center can only occur if the drug or biological was not separately charged to a patient. Although the commenter suggested that contrast agents may appear in other cost centers, our claims data demonstrate a significant volume of contrast agents reported under a pharmacy revenue code. Therefore, we believe that hospitals largely are charging patients specifically for contrast agents and accounting for these costs and charges in the "Drugs Charged to Patients" cost center. The hospital would then allocate the total overhead costs from the "Pharmacy" general services cost center to all revenue producing cost centers that have costed requisitions for drugs and biologicals on Worksheet B-1. In this circumstance, a large proportion of the total cost of the "Pharmacy" cost center would be allocated to the "Drugs Charged to Patients" cost center, assuming a concentration of costed requisitions for drugs and biologicals in the "Drugs Charged to Patients" cost center. The total pharmacy cost being allocated is an aggregation that commingles the overhead costs of a variety of drugs and biologicals. The resulting CCR for the "Drugs Charged to Patients" cost center should reflect both the average acquisition cost of drugs and biologicals, including those that are expensive and inexpensive, in that cost center and the average pharmacy overhead cost apportioned to that cost center.

Hospitals also may include the acquisition cost of drugs and biologicals directly in the "Pharmacy" general services cost center and reclassify this cost to revenue producing cost centers, including "Drugs Charged to Patients," before allocating the total cost of the "Pharmacy" cost center, which would have an effect similar to directly reporting the cost of drugs in the revenue producing cost centers on Worksheet A, as discussed above. In this situation, overhead cost from the "Pharmacy" cost center would be allocated to each of the revenue producing cost centers on the basis of costed requisitions. Some hospitals include the acquisition cost of drugs and biologicals directly in the "Pharmacy" cost center but do not reclassify this cost to the appropriate revenue producing cost center on Worksheet A, and instead allocate those costs on Worksheet B-1 together with the overhead cost of the pharmacy using costed requisitions. Regardless of which method described above that the

provider uses, the resulting CCR for the "Drugs Charged to Patients" cost center should reflect the average acquisition cost for drugs and biologicals in that cost center and the average pharmacy overhead apportioned to that cost center. Our redistribution methodology acknowledges that relying on a single CCR has the potential to "compress" overhead costs and that combining this compression with a packaging threshold leads us to underestimate the overhead costs of separately payable drugs and biologicals and overestimate the overhead costs of packaged drugs and biologicals.

As we discussed in our proposal, we did not propose to redistribute pharmacy overhead cost from packaged to separately payable drugs and biologicals utilizing a methodology that would provide a separate pharmacy overhead payment for each separately payable drug and biological based on its pharmacy complexity. The OPSS is a prospective payment system that provides payment for groups of services and we believe that it is important, at a minimum, to maintain the current size of the OPSS payment bundles, in order to encourage efficiency in the hospital outpatient setting. As we stated in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66613), we believe it is important that the OPSS create incentives for hospitals to provide only necessary, high quality care and to provide that care as efficiently as possible. We have considered in recent years how we could increase packaging under the OPSS in a manner that would create incentives for efficiency while providing hospitals with flexibility to provide care in the most appropriate way for each Medicare beneficiary. Hospitals have repeatedly explained that they consider the acquisition and pharmacy overhead costs of drugs in setting their charges for drugs, and we have continued to provide a single payment for the acquisition and pharmacy overhead costs of separately payable drugs and biologicals under the OPSS consistent with this hospital charging practice. While we have worked to develop, and are now implementing, a refined payment methodology for drugs and biologicals for the CY 2010 OPSS that we believe will pay more accurately for the pharmacy overhead cost of packaged and separately payable drugs and biologicals, we do not believe it would be appropriate to unbundle the current single combined payment for the acquisition and overhead costs of a separately payable drug into two

distinct payments, a drug payment and a pharmacy overhead payment. Furthermore, we note that section 1833(t)(14)(E)(ii) of the Act specifically authorizes the Secretary to adjust the APC payment weights for SCODs to take into account the recommendations of MedPAC on pharmacy overhead costs. We believe our CY 2010 approach that will adjust the APC payment for separately payable drugs and biologicals to more accurately pay for their associated pharmacy overhead cost, rather than provide a separate payment for a drug's pharmacy overhead cost each time the product is separately paid, is consistent with this statutory provision. Therefore, as we proposed, we are continuing to make a single bundled payment for the acquisition and pharmacy overhead costs of separately payable drugs and biologicals under the CY 2010 OPSS, an approach we believe both continues to encourage hospital efficiencies in the provision of drugs and biologicals to Medicare beneficiaries in the hospital outpatient setting and improves payment accuracy for the acquisition and pharmacy overhead costs of drugs and biologicals.

To confirm the portion of the \$395 million in estimated pharmacy overhead cost associated with coded packaged drugs and biologicals with an ASP that should be attributable to separately payable drugs and biologicals for the proposed rule, we used information from a variety of sources in order to corroborate the appropriateness of our policy to redistribute between one-third and one-half of the difference (\$150 million) between the aggregate claims cost for packaged drugs and biologicals and ASP dollars for the same drugs and biologicals to separately payable drugs and biologicals. In the CY 2010 OPSS/ASC proposed rule (74 FR 35330 through 35331), we presented two separate analyses which confirmed that our proposed redistribution of \$150 million in pharmacy overhead cost associated with the cost of packaged drugs and biologicals was appropriate.

We began the analytic exercise with three fundamental assumptions. The first assumption was that the hospital acquisition cost of separately payable drugs and biologicals, on average, is not less than 100 percent of ASP. We believed that this assumption was valid because we have been told that hospitals pay a range of prices for the same drug or biological. Some hospitals may be able to take advantage of volume and group purchasing to achieve significant discounts for certain drugs and biologicals, but other hospitals may pay more than average for drugs and biologicals because of their low volume

usage or a hospital's remote geographic location. Further, hospitals often serve as community care resources so they must provide drugs and biologicals to meet the needs of all of the patients who present to their facilities for care. The amounts and nature of those drugs and biologicals may vary significantly and unpredictably over time, particularly for smaller hospitals, due to changing availability of other care settings in their communities, such as physicians' offices, or due to emergencies, and this variability may constrain hospitals' ability to purchase all necessary quantities of certain drugs and biologicals based on best price contractual agreements negotiated in advance. Hence, we believed that the ASP was likely a fair estimate of hospitals' average acquisition cost of drugs and biologicals in general, excluding direct and indirect overhead costs.

The second assumption was that coded packaged drugs and biologicals with an ASP, as a group, typically have an aggregate absolute pharmacy overhead cost (direct and indirect) that exceeds the acquisition cost of the packaged drugs and biologicals, as measured by ASP. We believed that this assumption was appropriate because packaged drugs and biologicals carry the same absolute amount of indirect overhead cost per drug or biological administered as separately payable drugs and biologicals and because many packaged drugs and biologicals have extremely low ASPs but some of the same direct costs (for example, recordkeeping, storage, safety precautions, and disposal requirements) as separately payable drugs and biologicals. Our proposed rule claims data showed that the weighted average ASP for the coded drugs and biologicals with an ASP that we proposed to package for CY 2010 was approximately \$7 per day per packaged drug or biological, and we believed that it was a reasonable assumption that the full pharmacy overhead cost for a drug or biological (direct and indirect) equals or exceeds that amount.

Our final assumption was that, on average, the pharmacy overhead cost of separately payable drugs and biologicals, as a group, was not greater than the acquisition cost of the separately payable drugs and biologicals. We believed that this assumption is appropriate because separately payable drugs and biologicals carry the same absolute amount of indirect pharmacy overhead cost per drug or biological administered as packaged drugs and biologicals. While we have been told by MedPAC and the

pharmacy stakeholders that separately payable drugs and biologicals generally have direct pharmacy overhead costs that are significantly higher than the direct overhead costs of packaged drugs and biologicals, we do not believe that they exceed the acquisition cost of separately payable drugs and biologicals. The weighted average ASP for the drugs and biologicals in our proposed rule claims data that we proposed for separate payment for CY 2010 was approximately \$954 per day per separately payable drug or biological. We believed that the full pharmacy overhead cost for a separately payable drug or biological would not, on average, exceed the weighted average per day ASP. Hence, we believed these last two assumptions about the relationship of ASP to full pharmacy overhead cost (direct and indirect) for packaged and separately payable drugs and biologicals were appropriate for purposes of these analyses.

Having made these assumptions, for the proposed rule, we reduced the \$395 million in estimated pharmacy overhead cost that exceeded the ASP dollars for coded packaged drugs and biologicals with an ASP (their average acquisition cost) by \$50 million. Fifty million dollars in additional cost was necessary to raise the estimated cost calculated for separately payable drugs and biologicals from hospital claims data from 98 percent of ASP to 100 percent of ASP, in order to reach our estimate of the average hospital acquisition cost of separately payable drugs and biologicals of ASP. This left \$345 million in estimated residual pharmacy overhead cost that continued to be associated with packaged drugs and biologicals. We stated our belief that a portion of this cost was associated with coded packaged drugs and biologicals with an ASP in our claims data, both due to charge compression and our choice of an annual drug packaging threshold, and would continue to be less accurately associated with packaged drugs and biologicals were we not to engage in further redistribution of that portion of this residual pharmacy overhead cost of packaged drugs and biologicals.

We then performed two analyses using information provided by the MedPAC Report (June 2005 Report to Congress) and by the pharmacy stakeholders (February 2009 presentation to the APC Panel and other meetings with CMS) that we applied to our proposed rule claims data to estimate the amount of residual pharmacy overhead cost associated with packaged drugs and biologicals that should more accurately be attributed to

separately payable drugs and biologicals. To perform these analyses, we used proposed rule claims data only for those drugs and biologicals described by HCPCS codes that met the following criteria:

- The proposed CY 2010 OPPS status indicator for the HCPCS code was "G" for pass-through drugs and biologicals (excluding pass-through radiopharmaceuticals), "K" for separately payable drugs and biologicals that do not have pass-through status, or "N" for packaged drugs and biologicals, where the packaged status of these nonpass-through drugs and biologicals was determined by an estimate of cost per day based on ASP+4 percent;
- April 2009 pricing information based on the ASP methodology (other than mean cost from claims data) was available for the HCPCS code, and we would use the ASP methodology to pay for the HCPCS code if it had a status indicator of "K" or "G"; and
- CY 2008 OPPS claims data included claims for the HCPCS code or an equivalent predecessor code.

We first converted six of the seven categories that MedPAC recommended for reporting pharmacy overhead costs to three CMS categories (low, medium, and high), as we had proposed for the CY 2006 OPPS (70 FR 42729 through 42730); the seventh MedPAC category was not pertinent for this exercise because it is for the overhead cost attributable to radiopharmaceuticals. The CMS categories are defined as: Low (Orals); medium (Injection/Sterile Preparation; Single IV Solution/Sterile Preparation; Compounded Reconstituted IV Preparations); and high (Specialty IV or Agents requiring special handling in order to preserve their therapeutic value; Cytotoxic Agents in all formulations requiring personal protective equipment). We then derived a relative overhead weight for each of the three CMS categories by averaging the overhead weights for the six pertinent MedPAC categories. These averages were not weighted. The derived relative overhead weights for the CMS categories were as follows: Low = 1.00 (corresponding to MedPAC Category 1); medium = 3.61 (corresponding to MedPAC Categories 1, 2, and 3); and high = 11.11 (corresponding to MedPAC categories 5 and 6).

We also calculated a relative overhead weight for each of the three categories of pharmacy overhead complexity that were provided by the pharmacy stakeholders, using the different fixed dollar amounts that these stakeholders recommended that CMS pay for pharmacy overhead costs if we were to

make such payments for “all drugs” (packaged and separately payable). The pharmacy stakeholders’ categories are defined as: Low (Dispense without manipulation: e.g., oral drugs, pre-filled syringes); medium (Injectable drug with one step manipulation: e.g., simple injections); and high (Multiple step injectable products and chemotherapy that require safety considerations). The pharmacy stakeholders’ relative overhead weights were as follows: Low = 1; medium = 2.67; and high = 5.50.

Using the pharmacy stakeholders’ overhead categories (low, medium, and high) and incorporating the pharmacy stakeholders’ assignments of specific drugs and biologicals to levels of pharmacy complexity that they previously provided to CMS, we then assigned the remaining HCPCS codes for drugs and biologicals (approximately 50

percent of all drug and biological HCPCS codes with an associated ASP) based on our understanding of the characteristics of the categories. Similarly, we assigned all drug and biological HCPCS codes to the CMS categories created from the MedPAC groups for the derived relative overhead weights based on the definitions of those categories. Although the subsequent analytic processes were identical, we performed these analyses separately using the derived CMS overhead category weights (results are in Table 39) and using the pharmacy stakeholders’ overhead category weights (results are in Table 40).

Specifically, for the proposed rule we assigned the overhead weights to each drug and biological in the set of drugs and biologicals qualifying for this exercise. We then calculated a per unit

overhead cost by dividing the total relative weight for all drugs and biologicals in this exercise (low, medium, and high) into the residual pharmacy overhead cost from packaged drugs and biologicals of \$345 million. Using the relative weights for each scenario, we estimated the exact per unit pharmacy overhead cost reallocation for each low, medium, and high pharmacy overhead category. We then added this payment amount to ASP for each drug and biological and reassessed the amount of total claims cost for separately payable and packaged drugs and biologicals and calculated our standard ratio of aggregate claims cost to aggregate ASP dollars for separately payable and packaged drugs and biologicals. The results of these analyses are reprinted in Tables 39 and 40 below.

TABLE 39—ESTIMATED REDISTRIBUTION OF PHARMACY OVERHEAD COSTS USING RELATIVE WEIGHTS DERIVED FROM MEDPAC PHARMACY OVERHEAD CATEGORIES AND CY 2010 OPps PROPOSED RULE DATA

	Total ASP dollars for drugs and biologicals in claims data (in millions)*	Total cost of drugs and biologicals in claims data after adjustment (in millions)**	Ratio of cost to ASP (column C/ column B)	ASP+X Percent
Coded Packaged Drugs and Biologicals with an ASP	\$160	\$390	2.44	ASP+144
Separately Payable Drugs and Biologicals	2,589	2,704	1.04	ASP+4
All Coded Drugs and Biologicals	2,749	3,094	1.13	ASP+13

* Total April 2009 ASP dollars (ASP multiplied by drug or biological units in CY 2008 claims) for drugs and biologicals with a HCPCS code and ASP information.

** Total cost in the CY 2008 claims data after adjustment for drugs and biologicals with a HCPCS code and April 2009 ASP information.

TABLE 40—ESTIMATED REDISTRIBUTION OF PHARMACY OVERHEAD COST USING RELATIVE WEIGHTS CALCULATED FROM PHARMACY STAKEHOLDERS RECOMMENDED PHARMACY OVERHEAD PAYMENT LEVELS AND CY 2010 PROPOSED RULE DATA

	Total ASP dollars for drugs and biologicals in claims data (in millions)*	Total cost of drugs and biologicals in claims data after adjustment (in millions)**	Ratio of cost to ASP (column C/ column B)	ASP+X Percent
Coded Packaged Drugs and Biologicals with an ASP	\$160	\$402	2.51	ASP+151
Separately Payable Drugs and Biologicals	2,589	2,692	1.04	ASP+4
All Coded Drugs and Biologicals	2,749	3,094	1.13	ASP+13

* Total April 2009 ASP dollars (ASP multiplied by drug units in CY 2008 claims) for drugs with a HCPCS code and ASP information.

** Total cost in the CY 2008 claims data after adjustment for drugs with a HCPCS code and April 2009 ASP information.

As shown in Tables 39 and 40, the ratio of adjusted cost in the claims data for separately payable drugs and biologicals to ASP increased compared to the value derived from our standard methodology and declined for packaged drugs and biologicals with an associated ASP compared to the value calculated according to our standard drug payment methodology as shown in Table 41. Specifically, for the proposed rule under

our standard methodology without adjustment of the pharmacy overhead cost currently attributed to packaged drugs and biologicals with an associated ASP, we would have made packaged payment at ASP+247 percent. Using the CMS overhead weights, this value declined to ASP+144 percent and using the pharmacy stakeholders’ overhead weights, it declined to ASP+151 percent.

Under our standard drug payment methodology, without adjustment of the pharmacy overhead cost currently attributed to separately payable drugs and biologicals, we estimated for the proposed rule that separately payable drugs and biologicals would be paid at ASP–2 percent. Assuming a base average acquisition cost for all drugs and biologicals of ASP and using the CMS overhead weights to redistribute

the residual \$345 million in pharmacy overhead cost associated with coded packaged drugs and biologicals with an ASP in the claims data, this value increased to ASP+4 percent, and using the pharmacy stakeholders' overhead weights to redistribute the residual \$345 million in pharmacy overhead cost, this value also increased to ASP+4 percent.

Based on these analyses, for the proposed rule, we estimated that we would redistribute \$165 million in pharmacy overhead cost from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals by setting the average acquisition cost for all drugs and biologicals to ASP and using the CMS overhead weights, and we would redistribute \$153 million in pharmacy overhead cost from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals by setting the average acquisition cost for all drugs and biologicals to ASP and using the pharmacy stakeholders' overhead weights. These observed outcomes were consistent with our CY 2010 proposal to redistribute between one-third and one-half of the \$395 million of pharmacy overhead cost currently associated with packaged drugs and biologicals with an ASP to separately payable drugs and biologicals. These values were also consistent with the \$150 million we proposed to redistribute from the cost of coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals for CY 2010, which would represent a reduction in the cost of packaged drugs and biologicals of 27 percent.

After we performed these analyses but prior to display of the CY 2010 OPPS/ASC proposed rule, the pharmacy stakeholders provided us with updated assignments of CY 2009 drug HCPCS codes to their recommended levels of pharmacy complexity. We then assigned the remaining HCPCS codes for drugs and biologicals that the pharmacy stakeholders had not assigned based on our understanding of the characteristics of their categories. We recalibrated our model to incorporate the updated information. We observed no substantive changes in our findings, with the revised overhead category assignments redistributing \$159 million from packaged to separately payable drugs and biologicals and resulting in an ASP+X percentage of ASP+4 percent for separately payable drugs and biologicals and ASP+148 percent for packaged drugs and biologicals with an ASP.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35331), we indicated that

these analyses based on our synthesis of existing data and information from a variety of sources supported the appropriateness of a redistribution of the magnitude we proposed for CY 2010. We believed that our analyses of the claims data using the CMS relative overhead weights derived from the 2005 MedPAC pharmacy overhead study and using the pharmacy overhead category payments, levels of complexity, and assignments of drugs provided by the pharmacy stakeholders (where available), confirmed that payment for separately payable drugs and biologicals at ASP+4 percent would represent a reasonable aggregate adjustment for the pharmacy overhead cost of these separately payable drugs and biologicals, compared to the payment that would result from the standard drug payment methodology. We stated our belief that payment for separately payable drugs at ASP+4 percent would ensure that hospitals are paid appropriately for the average hospital acquisition cost and the pharmacy overhead cost that our analyses show would be appropriately redistributed from the estimated cost of overhead associated with drugs and biologicals with an ASP that we proposed to package for CY 2010.

Our proposal for CY 2010 relied upon the premise of providing a pharmacy overhead adjustment to payment for separately payable drugs by redistributing calculated pharmacy overhead cost from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals. Therefore, regardless of the payment level that the CY 2010 OPPS/ASC final rule with comment period claims and cost report data and July 2009 ASP data ultimately suggested, we believed that any redistributed amount of pharmacy overhead cost should be removed from the estimated cost of pharmacy overhead associated with coded packaged drugs and biologicals with an ASP. We proposed to redistribute pharmacy overhead cost within the estimated total amount of acquisition and overhead cost for all drugs and biologicals with an ASP that has been reported to us by hospitals by making a pharmacy overhead adjustment to payment for separately payable drugs and biologicals that is based upon a partial redistribution of the pharmacy overhead cost of coded packaged drugs and biologicals with an ASP. As described previously in this section, we proposed that any redistribution of pharmacy overhead cost that may arise from CY 2010 final rule data would occur only from some

drugs and biologicals to other drugs and biologicals, thereby maintaining the estimated total cost of drugs and biologicals in our claims data (no redistribution of cost would occur from other services to drugs and biologicals or vice versa). While there is some evidence that relatively more pharmacy overhead cost should be associated with separately payable drugs and biologicals and less pharmacy overhead cost should be associated with packaged drugs and biologicals in order to improve payment accuracy, we concluded that the recent RTI report on the OPPS' hospital-specific CCR methodology ("Refining Cost to Charge Ratios for Calculating APC and DRG Relative Payment Weights," July 2008 final report), the June 2005 MedPAC study of hospital outpatient pharmacy overhead costs, and our claims analyses discussed in the proposed rule presented no evidence that the total cost of drugs and biologicals (including acquisition and overhead costs) is understated in the claims data that we use to model the upcoming prospective payment year in relation to the costs of other services paid under the OPPS. Therefore, to improve the distribution of pharmacy overhead cost within the total estimated cost for all drugs and biologicals, without adversely affecting the relativity of payment weights for all services paid under the OPPS, we reasoned that it would be most appropriate to redistribute pharmacy overhead cost only within the total estimated cost of coded packaged and separately payable drugs and biologicals. By redistributing pharmacy overhead cost only within the total estimated cost of packaged and separately payable drugs and biologicals, we would maintain a constant total cost of drugs and biologicals under the OPPS as reported to us by hospitals, without redistributing cost from other OPPS services to the cost of drugs and biologicals under the budget neutral OPPS.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35332), we indicated that while we agree conceptually with the APC Panel that a redistribution of pharmacy overhead cost in our claims data from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals is appropriate, we did not accept the APC Panel's February 2009 recommendation that CMS pay for the acquisition cost of all separately payable drugs at no less than ASP+6 percent because, as we discussed previously in this section, our analyses of claims data indicated that appropriate payment for the acquisition

and pharmacy overhead costs of separately payable drugs would be ASP+4 percent. We also did not accept the APC Panel's February 2009 recommendation that CMS package the cost of packaged drugs at ASP+6 percent, use the difference between this cost and CMS' cost derived from charges to provide more appropriate payment for pharmacy services costs, and pay for pharmacy services using this amount by applying a tiered approach to payments based on criteria related to the pharmacy resources required for groups of drugs. We believed that the recommendation to package the cost of packaged drugs at ASP+6 percent would underpay for the pharmacy overhead cost of packaged drugs, which we expected would be higher in relation to ASP than the pharmacy overhead cost of separately payable drugs. Further, as discussed earlier in this section, because the OPSS is a prospective payment system that relies on payment for groups of services to encourage hospital efficiencies, we did not believe payment for pharmacy overhead costs that is separate from the OPSS payment for the acquisition costs of drugs would be appropriate.

The APC Panel further recommended that, if CMS did not adopt a methodology consistent with their recommendations summarized above, CMS should exclude data from hospitals that participate in the 340B program from its ratesetting calculations for drugs and that CMS should pay 340B hospitals in the same manner as it pays non-340B hospitals. In the proposed rule, we did not accept the APC Panel's recommendation that CMS propose to exclude data from hospitals that participate in the 340B program from its ratesetting calculations for drugs. For CY 2010, we note that we proposed a drug payment methodology that partially resembled the methodology recommended by the APC Panel because the proposal incorporated a redistribution of pharmacy overhead cost from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals. However, excluding data from hospitals that participate in the 340B program from our ASP+X calculation, but paying those hospitals at that derived payment amount, would effectively redistribute payment to drugs and biologicals from payment for other services under the OPSS, and we did not believe this redistribution would be appropriate. In our CY 2010 proposal, we did accept the APC Panel's February 2009 recommendation that CMS propose to pay 340B hospitals in the same manner

as non-340B hospitals are paid. Commenters on the CY 2009 OPSS/ASC final rule with comment period were generally opposed to differential payment for hospitals based on their 340B participation status, and we did not believe it would be appropriate to exclude claims from this subset of hospitals in the context of our CY 2010 proposal to pay all hospitals at the same rate for separately payable drugs and biologicals.

Moreover, as discussed above, while we did not propose to adopt the APC Panel's specific recommended methodology to redistribute pharmacy overhead cost that would otherwise be paid through payment for packaged drugs and biologicals, our proposed CY 2010 pharmacy adjustment methodology that would result in the payment of separately payable drugs and biologicals at ASP+4 percent incorporated a more limited redistribution of pharmacy overhead cost for coded packaged drugs and biologicals with an ASP (ASP is necessary to calculate an overhead amount) that would preserve the aggregate drug cost in the claims, a result consistent with the APC Panel's recommendations. Therefore, we believed that it would be appropriate to propose to pay 340B hospitals at the same rates that we are proposing to pay non-340B hospitals, and we proposed to include the claims and cost report data for 340B hospitals in the data we had used for our analyses in order to calculate the payment rates for drugs and biologicals and other services for the CY 2010 OPSS.

In conclusion, we proposed for CY 2010 to redistribute between one-third and one-half of the difference between the aggregate claims cost for coded packaged drugs and biologicals with an ASP and ASP dollars for those products, which resulted in proposed payment for the acquisition and pharmacy overhead costs of separately payable drugs and biologicals that do not have pass-through payment status of ASP+4 percent. This payment amount reflected an APC drug payment adjustment for pharmacy overhead cost. To accomplish this payment adjustment, we also proposed to reduce the cost of coded packaged drugs and biologicals with an ASP that was incorporated into the payment for procedural APCs by the amount of pharmacy overhead cost that was redistributed from these packaged drugs and biologicals to the payment for separately payable drugs and biologicals. The proposal was based on the proposed redistribution of \$150 million (through a 27 percent reduction in the cost of coded packaged drug and

biologicals with an ASP), between one-third and one-half of the pharmacy overhead cost (the cost above ASP) of coded packaged drugs and biologicals with an ASP in hospital outpatient claims, to the cost of separately payable drugs and biologicals, preserving the aggregate cost of all drugs and biologicals observed in the most recent claims and cost report data available for the proposed rule. We further proposed that the claims data for 340B hospitals be included in the calculation of payment for drugs and biologicals under the CY 2010 OPSS, and that 340B hospitals would be paid the same amounts for separately payable drugs and biologicals as hospitals that do not participate in the 340B program. Finally, we proposed that, in accordance with our standard drug payment methodology, the estimated payments for separately payable drugs and biologicals would be taken into account in the calculation of the weight scaler that would apply to the relative weights for all procedural services (but would not apply to separately payable drugs and biologicals) paid under the OPSS, as required by section 1833(t)(14)(H) of the Act.

At the August 2009 meeting of the APC Panel, the APC Panel recommended that CMS pay for all separately payable drugs at a rate of ASP+6 percent. The APC Panel recommended that CMS redistribute costs from packaged drugs to separately payable drugs as outlined in the CY 2010 OPSS/ASC proposed rule. Further, the APC Panel recommended that CMS analyze the impact on different classes of hospitals of payment at ASP+6 percent for separately payable drugs compared with CY 2009 payment at ASP+4 percent. In addition, the APC Panel requested that CMS provide an impact analysis of payment for separately payable drugs at ASP+6 percent on payment rates for other services that use packaged drugs compared with CY 2009 payment at ASP+4 percent. Finally, the APC Panel recommended that CMS and stakeholders continue to refine their analysis of payment for drugs, biologicals, and radiopharmaceuticals to assess the infrastructure costs associated with the preparation and handling of these products. Our responses to these recommendations are included in our responses to comments below.

Comment: Several commenters, including MedPAC, generally agreed with CMS' proposal to redistribute pharmacy overhead cost from packaged to separately payable drugs and biologicals. The commenters appreciated that the proposed

methodology would not pose an administrative burden to hospitals.

However, many commenters disagreed with CMS' calculation of the total estimated pharmacy overhead cost of \$395 million in the claims data associated with packaged drugs that resulted in the proposed redistribution of \$150 million, which was between one-half and one-third of this overhead cost. The commenters stated that CMS' estimate of \$395 million was too low to represent the aggregate pharmacy overhead cost of all packaged drugs and biologicals, resulting in an underestimate of how much overhead cost should be redistributed to separately payable drugs and biologicals and, therefore, a proposed payment rate for separately payable drugs and biologicals that was too low. They explained that, although CMS allows flexibility in hospital charging practices to account for drug and biological cost on hospital claims, CMS' proposed rule calculation did not take hospital charging practices for packaged drugs and biologicals into account.

Specifically, in CMS' estimation of the cost of packaged drugs and biologicals, the commenters pointed out that CMS omitted costs from claims data for drugs and biologicals that either do not have a HCPCS code or do not have a reported ASP, including those costs reported under a pharmacy revenue code line without a drug or biological HCPCS code due to hospital choice or claims processing requirements. The commenters argued that these uncoded packaged drug and biological costs represent a substantial portion of aggregate packaged drug and biological cost under the OPSS.

Some commenters estimated the additional packaged pharmacy overhead cost attributable to these uncoded drugs and biologicals to be nearly \$560 million. The commenters asserted that hospitals mark up the costs of drugs and biologicals reported on claims under pharmacy revenue code lines without HCPCS codes similarly to packaged drugs and biologicals reported with HCPCS codes. Several commenters provided analyses to support their contention that the costs of uncoded pharmacy revenue code lines reflect mostly packaged drug and biological costs, and that when hospitals do not report packaged drugs and biologicals with HCPCS codes, they report uncoded pharmacy revenue code lines instead for those drugs and biologicals. The commenters concluded that a significant percentage of the uncoded costs reported under pharmacy revenue code lines is pharmacy overhead cost disproportionately attributed to

packaged drugs and biologicals due to the tendency of our established methodology of converting billed charges to costs to "compress" the calculated costs to some degree and recognizing that our choice of an annual drug packaging threshold contributes to the magnitude of the ASP+X percent payment rate resulting from our standard drug payment methodology.

In order to address these concerns, the commenters recommended that CMS redistribute the pharmacy overhead cost attributed to uncoded cost reported under pharmacy revenue code lines to the cost of separately payable drugs and biologicals. Some commenters argued that, because they believe the costs on these uncoded pharmacy revenue code lines largely are for packaged drugs and biologicals with HCPCS codes, CMS could accurately assume the same proportional amount of ASP and mark up as for packaged drugs and biologicals with a HCPCS code and derive a simulated pharmacy overhead amount. Therefore, they suggested that one-third to one-half of residual pharmacy overhead cost associated with these uncoded pharmacy revenue code lines, which they estimate to total approximately \$560 million, should be redistributed to the cost of separately payable drugs and biologicals.

In addition, several commenters expressed concern that hospitals may not be billing packaged drugs and biologicals with HCPCS codes appropriately, resulting in uncoded costs reported under pharmacy revenue code lines, and that this contributed to the low estimate of pharmacy overhead costs included in the proposed rule. The commenters stated that a review of the OPSS claims data found variations in how hospitals are reporting drugs and biologicals with HCPCS codes under pharmacy revenue codes. The commenters stated that some hospitals are inappropriately assigning costs for drugs and biologicals with HCPCS codes to revenue code 0250 (Pharmacy (also see 063x, an extension of 025x); General Classification), rather than revenue code 0636 (Pharmacy—Extension of 025x; Drugs Requiring Detailed coding (a)). They speculated about a variety of reasons why more HCPCS-coding for packaged drug and biological cost was not available to CMS for proposed rule estimate purposes: (1) Hospitals may have reported their packaged drugs with revenue code 0250 and the associated charges and units with no HCPCS codes because HCPCS codes are not required to be reported for packaged drugs and biologicals; (2) the associated HCPCS code may not have printed on the claim because of provider billing system

settings; or (3) Medicare contractors may have instructed hospitals not to report HCPCS codes under revenue code 0250. As a result, the commenters believed that CMS' derived pharmacy overhead cost estimate for packaged drugs and biologicals based only on the cost of packaged drugs and biologicals with a HCPCS code and an ASP in the proposed rule were inaccurately low. In order to provide complete drug information for future years, they requested that CMS instruct hospitals to bill for drugs and biologicals with HCPCS codes under revenue code 0636.

Finally, some commenters expressed frustration that CMS did not provide information on the assignment of every drug and biological HCPCS code with a status indicator of "K," "G," or "N" to one of three categories of pharmacy overhead complexity in the analyses that CMS presented to validate the proposed redistribution of pharmacy overhead cost from packaged drugs and biologicals with an ASP to separately payable drugs and biologicals. In light of this omission, the commenters recommended that CMS redistribute the larger one-half portion of the one-third to one-half of the proposed pharmacy overhead cost to accurately account for all pharmacy costs represented in the HOPD.

Response: We proposed to reallocate approximately \$150 million in pharmacy overhead cost from coded packaged drugs and biologicals with an ASP to separately payable drugs and biologicals, representing a middle ground between the one-third to one-half of the total pharmacy overhead cost associated with this set of packaged drugs and biologicals. We agree with the commenters that we did not include uncoded drug and biological costs reported under pharmacy revenue code lines in our proposed rule estimate of the pharmacy overhead costs of packaged drugs and biologicals. We also agree with the commenters that costs on uncoded pharmacy revenue code lines represent OPSS drug and biological cost. The commenters suggested that we assume the same relationship between total claim cost and ASP for the uncoded drug and biological costs in our claims data as we observe for coded packaged drugs and biologicals with an ASP, and then redistribute one-third of the assumed, associated pharmacy overhead cost to the cost of separately payable drugs and biologicals. We were interested to review the analyses provided by some commenters that used statistical techniques to compare the uncoded drug and biological costs to the costs of packaged drugs and biologicals with HCPCS codes but, at this time, we

cannot be certain that the assumptions suggested by the commenters would represent an accurate portion of the uncoded drug and biological cost attributable to acquisition cost versus pharmacy overhead cost. In the proposed rule, we stated that a premise of our redistribution model was our assumption that the associated aggregate ASP for packaged drugs and biologicals was a proxy for acquisition cost for this group of drugs and biologicals (74 FR 35327). Our proposed methodology identified the difference between this proxy for acquisition cost and the cost of the same coded packaged drugs and biologicals with an ASP in our claims data as pharmacy overhead cost, and it was one-third to one-half of that pharmacy overhead cost (\$150 million) that we specifically proposed to redistribute from coded packaged drugs and biologicals with an ASP to the cost of separately payable drugs and biologicals.

As shown in Table 41, we determined that the estimated aggregate cost of separately payable drugs and biologicals, including acquisition and pharmacy overhead costs, is equivalent to ASP-3 percent for this final rule with comment period. A redistribution of \$150 million from the pharmacy overhead cost of coded packaged drugs and biologicals with an ASP (one-third of that pharmacy overhead cost from final rule data) to the cost of separately payable drugs and biologicals would result in payment for separately payable drugs and biologicals at ASP+2 percent for CY 2010. If we were to assume the same relationship between total claim cost and ASP for the uncoded drug and biological cost in our claims data as we observe for coded packaged drugs and biologicals with an ASP as recommended by some commenters, and if we were then to redistribute one-third of the assumed, associated pharmacy overhead cost (\$150 million) of uncoded drug and biological cost to the cost of separately payable drugs and biologicals, the result would be payment for separately payable drugs and biologicals at ASP+7 percent. The total cost redistribution to separately payable drugs and biologicals in this case would be \$300 million, \$150 million from the cost of coded packaged drugs and biologicals with an ASP and \$150 million from the cost of uncoded packaged drugs and biologicals.

We understand that our proposal for a redistribution of any drug and biological cost from packaged to separately payable drugs and biologicals already is not our usual OPSS cost estimation methodology, which uses the estimated cost from claims and cost

report data as reported to us by hospitals for an item or service to calculate a relative weight for that service, or in the case of drugs and biologicals, an ASP+X percent under our standard drug payment methodology. We made this redistribution proposal because we were concerned that by not redistributing pharmacy overhead cost from packaged to separately payable drugs and biologicals, an underpayment of separately payable drugs and biologicals at ASP-2 percent (ASP-3 percent based on final rule claims data) could result. We remain concerned that the redistribution of \$150 million from the pharmacy overhead cost of coded packaged drugs and biologicals with an ASP to payment for separately payable drugs and biologicals, which would provide payment at ASP+2 percent, also could result in underpayment of separately payable drugs and biologicals. We are also troubled, however, that payment for separately payable drug and biologicals at ASP+7 percent resulting from an assumption that the uncoded drug and biological cost resembles the coded packaged cost of drugs and biologicals with an ASP could result in a potential payment overestimation. As noted above, we cannot be certain that we know what portion of the uncoded drug and biological cost is acquisition cost versus pharmacy overhead cost. Therefore, we are not willing to make even broader assumptions about the magnitude of ASP for uncoded drug and biological cost in claims or layer any other assumptions on the proposed methodology that would further significantly redistribute costs as reported to us by hospitals within the framework of the OPSS ratesetting methodology.

While we are not making sweeping assumptions that this uncoded packaged drug and biological cost includes a pharmacy overhead amount comparable to that of coded packaged drugs and biologicals with an ASP, we do acknowledge that there must be some pharmacy overhead cost associated with these uncoded packaged drugs and biologicals that was not accounted for in our initial estimate of the pharmacy overhead cost of packaged drugs and biologicals because we expect that hospitals would have attributed some pharmacy overhead cost to these products through their mark-up practices. Therefore, while we further examine the issue of pharmacy overhead costs and while hospitals examine administrative changes that could result in their submission of more

accurate data to us as described below, we believe that the adoption of a transitional payment rate of ASP+4 percent based on a pharmacy overhead adjustment methodology for CY 2010 would base OPSS payment upon the best available proxy for the average acquisition and pharmacy overhead costs of separately payable drugs and biologicals. We note that payment for separately payable drugs and biologicals at ASP+4 percent falls within the range of ASP-3 percent, that would result from no pharmacy overhead cost redistribution from packaged to separately payable drugs and biologicals, to ASP+7 percent, that would result from redistribution of pharmacy overhead cost based on expansive assumptions about the nature of uncoded packaged drug and biological cost. We proposed payment for separately payable drugs and biologicals at ASP+4 percent for CY 2010, and our final CY 2010 transitional payment rate is consistent with this amount. We are confident that ASP+4 percent will provide appropriate payment for separately payable drugs and biologicals in CY 2010, noting that this payment is consistent with our payment in CY 2009. We are not aware of any current access problems for Medicare beneficiaries to drugs and biologicals in the HOPD based on our CY 2009 OPSS payment for separately payable drugs and biologicals at this rate.

Specifically, for CY 2010, to acknowledge the uncoded drug and biological cost without making significant further assumptions about the amount of pharmacy overhead cost associated with the drugs and biologicals captured by this cost and to pay separately payable drugs and biologicals at ASP+4 percent, we believe it currently would be appropriate to reallocate \$50 million of the total uncoded drug and biological cost in order to represent the pharmacy overhead cost of uncoded packaged drugs and biologicals that should be appropriately associated with the cost of separately payable drugs and biologicals. We believe that our proposal to reallocate \$150 million of cost from coded packaged drugs and biologicals with an ASP, or one-third of the pharmacy overhead cost of these products based upon the claims data available for this CY 2010 final rule, to separately payable drugs and biologicals continues to be appropriate. The commenters generally supported the one-third to one-half redistribution estimate. While some commenters requested a reallocation of one-half of

the pharmacy overhead cost of packaged drugs and biologicals to separately payable drugs and biologicals, as already discussed, we do not believe there is a compelling reason to reallocate that amount. We note that the reallocation of \$50 million or 8 percent of the total cost of uncoded packaged drugs and biologicals assumes that whatever pharmacy overhead cost is not accurately associated with uncoded packaged drugs and biologicals, it would not be less than 8 percent of total uncoded drug and biological cost. This is intentionally a conservative estimate, as compared with the case of coded packaged drugs and biologicals with an ASP and for which we have a specific pharmacy overhead cost estimate in relationship to their known ASPs where the reallocation of \$150 million constitutes 24 percent of the total cost of the coded packaged drugs and biologicals with an ASP. As stated earlier, we are unwilling to make sweeping assumptions that uncoded packaged drug and biological cost includes a pharmacy overhead amount comparable to that of coded packaged drugs and biologicals with an ASP. We are confident that this conservative

estimate of \$50 million for redistribution from the cost of uncoded packaged drugs and biologicals to separately payable drugs and biologicals, as opposed to the \$150 million redistribution that could result from broad assumptions about the ASPs of these uncoded drugs and biologicals, is an appropriate amount for CY 2010 in light of our uncertainty about the relationship between ASP and pharmacy overhead cost for the uncoded drugs and biologicals.

In summary, with a redistribution of a total of \$200 million, \$150 million from the pharmacy overhead cost of coded packaged drugs and biologicals with an ASP as we proposed and \$50 million from the cost of uncoded packaged drugs and biologicals for which we cannot estimate a more specific pharmacy overhead cost at this time, to separately payable drugs and biologicals, the final CY 2010 transitional payment rate for separately payable drugs and biologicals is ASP+4 percent based on the final pharmacy overhead adjustment methodology.

In response to commenters' frustration that we did not provide information on our assignment of every

drug or biological HCPCS code to one of three categories of pharmacy overhead complexity in the analyses that we presented to validate the proposed redistribution methodology, we did not base our proposed redistribution amount on these analyses. We explicitly made a proposal to redistribute \$150 million in estimated pharmacy overhead cost associated with coded packaged drugs and biologicals with an ASP, between one-third and one-half of the estimated pharmacy overhead cost. Although we did not provide the precise assignment of drugs and biologicals to the various categories, we did describe each set of pharmacy overhead complexity categories in the proposed rule and our methodology for redistributing pharmacy overhead cost under each scenario. In addition, we posted a clarification to this discussion for the public replicating our models on August 6, 2009 during the comment period.

Table 41 displays the final pharmacy overhead adjustment methodology for separately payable and packaged drugs and biologicals under the CY 2010 OPPS.

TABLE 41—CY 2010 FINAL RULE—PHARMACY OVERHEAD ADJUSTMENT PAYMENT METHODOLOGY FOR SEPARATELY PAYABLE AND PACKAGED DRUGS AND BIOLOGICALS

	Total ASP dollars for drugs and biologicals in claims data (in millions)*	Total cost of drugs and biologicals in claims data after adjustment (in millions)**	Ratio of cost to ASP (column C/ column B)	ASP+X Percent
Uncoded Packaged Drugs and Biologicals	Unknown	\$606	N/A	N/A
Coded Packaged Drugs and Biologicals with an ASP	172	466	2.71	ASP+171
Separately Payable Drugs and Biologicals with an ASP	2,972	3,039	1.04	ASP+4
All Coded Drugs and Biologicals with an ASP	3,144	3,505	1.11	ASP+11

* Total July 2009 ASP dollars (ASP multiplied by drug or biological units in CY 2008 claims) for drugs and biologicals with a HCPCS code and ASP information.

** Total cost in the CY 2008 claims data for drugs and biologicals.

We note that hospitals currently have a variety of ways to bill for drugs and biologicals that are not separately paid. They may report the charges for the HCPCS code separately on a line, and if the HCPCS code has a status indicator of "N," no separate payment is made for the drug or biological, but the reported charge information is available to use for future ratesetting. Provided that information for the ASP pricing methodology was available for the drug or biological HCPCS code, we included drug or biological cost estimated from charges for claims described by this scenario in our estimation of total pharmacy overhead costs of coded packaged drugs and biologicals with an

ASP for CY 2010 because we could identify these drugs and biologicals, estimate their cost from charges in CY 2008 claims data, and use their ASP pricing information. Another option available to hospitals billing for packaged drugs and biologicals is to incorporate the charge for the drug or biological in the charge for the procedure. We are unable to identify the cost estimated from charges as drug or biological cost because the procedures are not reported under a pharmacy revenue code line and, therefore, these packaged drug and biological costs were not included in our estimate of the total pharmacy overhead cost of packaged drugs and biologicals. The final way for

hospitals to bill for packaged drugs and biologicals is to include charges for these items under a pharmacy revenue code line, specifically revenue code 0250, without a HCPCS code, and it is an additional \$50 million from this uncoded cost of packaged drugs and biologicals that we have redistributed to the cost of separately payable drugs and biologicals in our final CY 2010 pharmacy overhead adjustment payment methodology for drugs and biologicals.

We have adopted this pharmacy overhead adjustment payment methodology for CY 2010 only after 3 distinct attempts over the 4 prior years to garner more accuracy in both the

claim drug or biological charge and Medicare hospital cost report data as submitted to us by hospitals in an effort to show consideration for the significant hospital administrative burden that the commenters cited in response to each proposal that, in turn, precluded further refinement of our data collection efforts. In light of our commitment to using hospital data as reported to us by hospitals to set OPPS payment rates, we believe that it would be inappropriate to assume that the costs reported under uncoded pharmacy revenue code lines are for the same drugs and biologicals, with the same ASPs, as the costs of packaged drugs and biologicals reported with HCPCS codes. We acknowledge that the pharmacy overhead cost associated with drug and biological costs reported under uncoded pharmacy revenue code lines were not included in the proposed rule estimate of total pharmacy overhead cost of coded packaged drugs and biologicals with an ASP. In response to the concerns of commenters, we have considered only a small percentage of this uncoded drug and biological cost to be misallocated pharmacy overhead cost that is appropriate for redistribution in our final CY 2010 methodology. We cannot be certain that the amount of uncoded pharmacy overhead cost is as high as some commenters suggested, that hospitals mark up these uncoded drugs and biologicals in the same way as packaged drugs and biologicals with HCPCS codes, or that significant volume for these uncoded drugs and biologicals might not warrant allocating a greater percentage of fixed pharmacy overhead cost to these drugs and biologicals. If hospitals truly desire significantly greater OPPS payment accuracy for separately payable drugs and biologicals, it is clear that hospitals will need to assume some burden in submitting more accurate data to us. In addition, we will continue to examine the issue of pharmacy overhead costs as we work to refine our transitional payment methodology for separately payable drugs and biologicals for future years.

CMS' longstanding policy is to refrain from instructing hospitals on charging practices for services under most revenue codes. We believe that this allows hospital flexibility in billing systems and provides the necessary autonomy for hospitals to manage the many variations that are possible when creating a hospital chargemaster for multiple payers. While we do not require hospitals to use revenue code 0636 (Pharmacy-Extension of 025x; Drugs Requiring Detailed coding (a))

when billing for drugs and biologicals that have HCPCS codes, whether they are separately payable or packaged, we believe that a practice of billing all drugs and biologicals with HCPCS codes under revenue code 0636 would be consistent with NUBC billing guidelines and would provide us with the most complete and detailed information for ratesetting. We note that we make packaging determinations for drugs annually based on cost information reported under HCPCS codes, so the OPPS ratesetting is best served when hospitals report charges for all items and services that have HCPCS codes under those HCPCS codes, whether or not payment for the items and services is packaged or not. As already discussed, it is our standard ratesetting methodology to rely on hospital cost and charge information as it is reported to us through the claims data. More complete data from hospitals on which drugs were provided for a specific episode would help improve payment accuracy for separately payable drugs in the future, and we encourage hospitals to change their reporting practices if they are not already reporting HCPCS codes for all drugs furnished, if specific codes are available.

Comment: Several commenters objected to the proposed redistribution methodology as it decreased payments for procedural APCs with high packaged drug costs included in their payment rates.

One commenter disagreed with the proposed redistribution methodology and its effect on the imaging procedure APCs. The commenter argued that because all contrast agents without pass-through status are packaged, regardless of an individual agent's relationship to the annual drug packaging threshold, imaging procedure APCs should be exempt from the proposed pharmacy cost redistribution methodology. If imaging procedures are not exempted from the redistribution, the commenter contended that these procedures would be disproportionately affected because the spectrum of contrast costs are currently represented as packaged costs within the imaging procedure APCs.

Similarly, another commenter requested that CMS exempt nuclear medicine procedures from the redistribution methodology. Again, the commenter stated that as all diagnostic radiopharmaceuticals are packaged, regardless of their estimated per day costs, their overhead costs are all represented in the nuclear medicine APCs and a redistribution would disproportionately affect these services.

Response: We agree that packaging all contrast agents into associated imaging

procedures results in the inclusion of payment for both expensive and relatively inexpensive contrast agents in the payment for the associated imaging procedures. While the commenters contended that this policy thereby incorporates all contrast agents with different hospital mark up practices in a single packaged payment methodology and, therefore, should not be subject to the cost redistribution, we believe that contrast agents are contributing to the overall charge compression for all drugs and biologicals that is the specific target of our redistribution methodology. When examining CY 2008 claims data for the final rule, we observed that hospitals typically billed costs for contrast agents under a pharmacy revenue code (025X (Pharmacy), 026X (IV Therapy), or 063X (Pharmacy—Extension of 025X)). We believe that in almost all cases, hospitals capture the costs and charges for pharmacy revenue codes in the cost center 5600 "Drugs Charged to Patients," and this is the cost center that we use to estimate costs from charges for the pharmacy revenue codes in our claims data each year. We make the revenue code-to-cost center crosswalk that we use to match Medicare hospital cost report information with claims data continually available for inspection and comment on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS>. The proposed methodology of redistributing pharmacy overhead cost from packaged drugs and biologicals to separately payable drugs and biologicals was a proposal to address charge compression observed within this specific cost center that captures the vast majority of costs and charges for drugs and biologicals billed on hospital outpatient claims. Therefore, as most hospitals billing contrast agents with pharmacy revenue codes are associating the contrast agent costs with the cost center 5600 "Drugs Charged to Patients," we believe it is appropriate to redistribute cost from contrast agents to separately payable drugs and biologicals under our final CY 2010 pharmacy overhead cost redistribution methodology.

The commenter also suggested that it would be inappropriate to redistribute cost from contrast agents because, as discussed in V.B.2.d. of this final rule with comment period, it has been OPPS policy to package payment for all contrast agents since CY 2008. The proposed methodology for redistributing pharmacy overhead cost from packaged drugs and biologicals to separately payable drugs and biologicals was not only a proposal to address charge

compression, but specifically a proposal to address charge compression in light of our adoption of a specific drug packaging threshold, which is \$65 for CY 2010. The argument that it would, therefore, be inappropriate to redistribute cost from contrast agents could have merit if there was a sizable amount of aggregate cost for contrast agents with per day costs greater than the drug packaging threshold of \$65. In that case, it could be argued that the compression in cost estimates for expensive contrast agents (those with per day costs greater than the \$65 packaging threshold) created by estimating costs for those agents by applying the CCR for the single cost center 5600 "Drugs Charged to Patients" to expensive contrast agents' charges would be offset by the overestimation of costs for inexpensive contrast agents (those with per day costs less than the \$65 packaging threshold) created by application of the same single CCR to inexpensive contrast agents' charges, assuming that hospitals apply a lower markup to expensive contrast agents and a higher markup to inexpensive contrast agents. If the mix of expensive and inexpensive contrast agents resembled the mix of expensive and inexpensive drugs generally captured in the cost center 5600 "Drugs Charged to Patients," the use of a single CCR would accurately estimate total cost of contrast agents in aggregate. Because all contrast agents not receiving pass-through payment are packaged, packaging an accurate aggregate cost estimate for contrast agents could argue against redistributing cost from packaged contrast agents to separately payable drugs and biologicals. However, in our CY 2010 final rule claims data, we observed only 3 percent of total contrast agent cost associated with those contrast agents that have per day costs above \$65.

In conclusion, both because contrast agents are billed under pharmacy revenue codes and accounted for in the cost center 5600 "Drugs Charged to Patients" and because the per day cost of almost all contrast agents falls under the CY 2010 packaging threshold of \$65, we believe the estimated cost of contrast agents contains a disproportionate amount of pharmacy overhead cost and that it is appropriate to include them in our final CY 2010 redistribution methodology.

While we believe that contrast agents are commonly billed under pharmacy revenue codes and that hospitals largely account for the cost of contrast agents under the cost center 5600 on their Medicare hospital cost report, we did not observe that hospitals apply the

same practice for diagnostic radiopharmaceuticals. After reviewing our claims data, we found that the majority of diagnostic radiopharmaceuticals are billed under revenue code 0343 (Nuclear Medicine; Diagnostic Radiopharmaceuticals). As specified in our revenue code-to-cost center crosswalk, we believe hospitals largely account for the costs and charges associated with revenue code 0343 in a nonstandard cost center for Diagnostic Nuclear Medicine or the cost center 4100 "Radiology-Diagnostic." Because the redistribution of pharmacy overhead cost from packaged drugs and biologicals to separately payable drugs and biologicals is intended to specifically address charge compression in the pharmacy cost center, in light of the above information, we excluded the costs of both diagnostic and therapeutic radiopharmaceuticals from our estimate of total drug and biological cost in the claims data for the final CY 2010 redistribution methodology. As a result, the final payment rates for nuclear medicine procedures that incorporate the costs of packaged diagnostic radiopharmaceuticals are not impacted by the final redistribution methodology.

Comment: Several commenters cited methodological concerns about the approach CMS used to calculate the proposed equivalent average ASP-based payment amount for separately payable drugs and biologicals. In addition, the commenters expressed concern that CMS' cost estimation methodology is very sensitive to changes in the underlying data and assumptions. Citing these concerns, some commenters requested payment at ASP+6 percent for parity with physician's office payment rates for drugs, arguing that hospital costs for acquisition and associated pharmacy overhead would be at least as high, if not significantly greater, than the physician's office costs.

Some commenters noted that the statute requires drug cost surveys for payment purposes for SCODs under the OPSS, and that the most recent survey available is outdated as it was performed in CY 2004 by the GAO. The commenters stated that the statute specifically requires survey data as the basis for hospital acquisition costs in order to provide a more appropriate payment methodology for drugs and biologicals, instead of costs calculated from claims data. They concluded that, by not performing a survey and by not paying for drugs and biologicals at the physician's office rate, CMS is not in compliance with the statute. The commenters acknowledged that drug cost surveys are difficult to perform. However, they asserted that, in order to

comply with the requirements of the statute, either a survey should be performed or payment should be made at ASP+6 percent. Other commenters cited the methodological concerns that are described below regarding the proposed proxy for average acquisition cost based upon claims data, and stated that until CMS is able to adequately address these concerns, CMS should implement payment at ASP+6 percent pursuant to section 1833(t)(14)(A)(iii)(I).

The commenters' first methodological concern is that CMS compared cost estimates from different points in time to develop payment rates for drugs and biologicals. Specifically, several commenters noted that for the proposed rule, CMS used ASP data from the fourth quarter of CY 2008, which is the basis for calculating payment rates for drugs and biologicals in the physician's office setting using the ASP methodology effective April 1, 2009, along with hospital claims data from CY 2008 to determine the relative ASP amount for CY 2010 under CMS' proposed drug payment methodology. The commenters requested that CMS use an alternative ASP file for the final rule calculation of ASP+X to better align ASP data with hospital claims and cost report data. The commenters stated that CMS compared hospital claims data from throughout CY 2008 with costs estimated from charges that include pharmacy overhead cost, with ASP data as a proxy for acquisition cost representing drug sales in the fourth quarter of CY 2008, well after the time hospitals would have purchased most of their drugs for administration in CY 2008. As an alternative, the commenters requested that CMS use an earlier ASP file that is more representative of the costs to hospitals when they purchase drugs for the claims year. Specifically, some commenters requested that CMS use the July 1, 2008 ASP file that represents sales from the first quarter of CY 2008 when comparing CY 2008 hospital claims data to ASP data to determine an ASP+X amount.

Second, many commenters reiterated concerns that when CMS applies a single CCR to adjust charges to costs for these drugs and biologicals, charge compression leads to misallocation of the pharmacy overhead costs associated with high and low cost drugs and biologicals during ratesetting. The commenters noted that hospitals disproportionately mark up their charges for low cost drugs and biologicals to account for pharmacy overhead costs. They indicated that while the aggregate charges for inexpensive and expensive drugs may include the total pharmacy overhead

costs of the hospital, the charges for individual drugs and biologicals do not represent the specific acquisition and pharmacy overhead costs of that particular drug or biological. The commenters explained that hospitals apply proportionately smaller markups to higher cost items and proportionately larger markups to lower cost items. The commenters argued that by using only separately payable drugs in the calculation of the equivalent average ASP-based amount, the pharmacy overhead costs associated with these separately payable drugs that are disproportionately included in the charges for packaged drugs are not factored into the calculation, resulting in an artificially low ASP add-on percentage. Therefore, some commenters suggested using the costs of both packaged drugs and separately payable drugs when calculating the equivalent average ASP-based payment amount for separately payable drugs, as they argued that this would provide a more accurate ASP-based payment amount for separately payable drugs. As an alternative, the commenters recommended that CMS eliminate the drug packaging threshold and provide separate payment for all Part B drugs under the OPSS at an ASP+X percent amount calculated from the cost of all drugs with HCPCS codes.

Finally, several commenters noted that CMS included, in the calculation of the costs of separately payable drugs and biologicals, OPSS claims data from hospitals that receive Federal discounts on drug prices under the 340B drug pricing program. The commenters pointed out that hospital participation in the 340B program had grown substantially over the past few years, and they believed that the costs from these hospitals now constituted a significant proportion of hospital drug costs on CY 2008 OPSS claims. The commenters stated that including 340B hospital claims data when comparing aggregate hospital costs based on claims data to ASP rates contributed to an artificially low equivalent average ASP-based payment rate because ASP data specifically exclude drugs sales under the 340B program.

Response: As discussed above, the provision in section 1833(t)(14)(A)(iii) of the Act continues to be applicable to determining payments for SCODs for CY 2010. This provision requires that payment for SCODs be equal to the average acquisition cost for the drug for that year (which, at the option of the Secretary, may vary by hospital group) as determined by the Secretary, subject to any adjustment for overhead costs and taking into account the hospital

acquisition cost survey data under section 1833(t)(14)(D) of the Act, or if hospital acquisition cost data are not available, then the average price for the drug in the year established under section 1842(o), 1847A, or 1847B of the Act, as the case may be, as calculated and adjusted by the Secretary as necessary for purposes of section 1833(t)(14) of the Act. In the CY 2006 OPSS final rule with comment period (70 FR 68640), we compared hospital drug cost data that were available to us at the time, specifically: (1) data from the GAO survey; (2) hospital claims data from CY 2004; and (3) ASP information. In addition, we discussed our methodology for comparing these data that represented different timeframes from 2004 to 2006. As a result of our analysis comparing these three sources, we concluded that, on average, the costs from hospital claims data representing SCODs were roughly equivalent to payment at ASP+6 percent. Therefore, we finalized a policy that used our hospital claims data as a proxy for average hospital acquisition cost and provided payment for separately payable drugs that do not have pass-through status at ASP+6 percent for CY 2006 (70 FR 68639 through 68642). While the commenters are correct that the statute allows for the use of the methodology described in section 1842(o), section 1847A or section 1847B of the Act, as calculated and adjusted by the Secretary as necessary, this is only when hospital acquisition cost data are not available. We believe that we have established both our hospital claims data and ASP data as an appropriate proxy for average hospital acquisition cost, taking the GAO survey information into account for the base year (70 FR 68641). Many of the drugs and biologicals covered under the OPSS are provided a majority of the time in the hospitals setting, and the ASP information we collect would be an adequate proxy for hospital acquisition cost. Further, as already discussed, the commenters have not disputed the accuracy of the total drug and biological cost estimated in our claims data, only the estimated cost of separately payable drugs and biologicals. While we have not yet performed hospital drug acquisition cost surveys similar to the GAO survey, we note that the statute only calls for "periodic" surveys. Therefore, we disagree that we are not complying with the statute by not performing a survey and not paying at the physician's office rate. We note, however, that we are considering the possibility of such a survey at some point in the future. Therefore, we do not

believe that it is appropriate at this time to provide payment at anything other than average acquisition cost, with a redistribution for pharmacy overhead, based on the drug and biological costs observed in hospital claims data and pricing information observed in ASP data. We disagree with the commenters who believe that our redistribution methodology is not an appropriate proxy for average hospital acquisition cost, with an adjustment for pharmacy overhead cost. We have no basis for providing payment for separately payable drugs at the physician's office rate in the face of an appropriate proxy for average hospital acquisition cost, as described in detail below.

In response to the commenters who claimed that hospital costs for drug acquisition and associated pharmacy overhead would be at least as high, if not significantly greater, than the physician's office costs, we have no information that would confirm this statement. ASP information is only available for all sales of drugs and biologicals, so we cannot compare hospital and physician's office acquisition costs for individual drugs and biologicals or in aggregate. While our final CY 2010 pharmacy overhead adjustment methodology for payment of separately payable drugs and biologicals relies on the assumption that ASP is a fair estimate of hospitals' average acquisition cost of drugs and biologicals in general, we expect that drug acquisition costs could vary across settings and clinical cases. In some cases hospital drug acquisition costs could be lower than the costs to physicians' offices, based on high volume purchasing agreements, and in other cases hospital acquisition costs could be greater, based on their need for emergency purchases outside of negotiated contracts with preestablished best rates. We also expect that pharmacy overhead costs could vary across hospital and physician's office settings, based on the drugs and biologicals administered in those settings. Many hospitals provide a range of drugs and biologicals, including those with high and low pharmacy overhead costs, whereas physicians' offices may be more likely to provide drugs and biologicals with typically high (or low) pharmacy overhead costs. This unknown variability in drug acquisition and pharmacy overhead costs across settings means that we cannot conclude that the acquisition and pharmacy overhead costs of drugs and biologicals in the HOPD are greater or less than the physician's office costs. Finally, the ASP-based payment rate for drugs

furnished in physicians' offices is specified in the statute as ASP+6 percent, whereas the OPSS payment is based on average hospital acquisition cost and associated pharmacy overhead cost. Therefore, we do not believe that comparisons of OPSS drug payment rates with physician's office payment rates suggesting that parity is necessary are applicable to determining appropriate payment rates for separately payable drugs and biologicals under the OPSS.

For our calculation of per day costs of HCPCS codes for drugs and nonimplantable biologicals in the CY 2010 OPSS/ASC final rule with comment period, we proposed to use ASP data from the first quarter of CY 2009, which is the basis for calculating payment rates for drugs and biologicals in the physician's office setting using the ASP methodology effective July 1, 2009, along with updated hospital claims data from CY 2008 (74 FR 35320). We also proposed to use these data for budget neutrality estimates and impact analyses for this CY 2010 OPSS/ASC final rule with comment period. Payment rates for HCPCS codes for separately payable drugs and nonimplantable biologicals included in Addenda A and B to this final rule with comment period are based on ASP data from the second quarter of CY 2009, which are the basis for calculating payment rates for drugs and biologicals in the physician's office setting using the ASP methodology effective October 1, 2009.

Since implementing the ASP+X methodology in CY 2006, we have used the most recently available data to establish our relative ASP payment rate for the upcoming year, consistent with our overall policy of updating the OPSS of using the most recent claims and cost report data. For the CY 2010 final rule, this results in using July 2009 ASP payment rates (based on first quarter CY 2009 sales), CY 2008 hospital claims data, and the most recently available hospital cost reports (in the majority of cases cost reports beginning in CY 2007). As we have noted in previous years, the relative ASP+X amount is likely to change from the proposed rule to the final rule as a result of updated ASP data, hospital claims data, and updated hospital cost reports. If we were to introduce significant error into our ASP+X percent calculation by not aligning all pricing and cost data to a single period of time, we would consider changing the ASP data that we use. However, we believe that if we were to use an ASP file from CY 2008, which commenters claim would more accurately represent hospital costs

associated with procuring drugs and biologicals for that claims year, we would need to offset any increases in the relative ASP amount resulting from the use of a different ASP file with a deflation adjustment for each hospital's CCRs for cost center 5600 "Drugs Charged to Patients" in order to simulate costs from claim charges in the claim year. We make comparable CCR deflation estimates when we set our fixed dollar eligibility threshold for outlier payments described in section II.F. of this final rule with comment period. Because over recent years hospital charges have typically grown faster than costs, we would expect such an adjustment to reduce estimated costs in our claims data. Therefore, we are continuing our current policy of using the most recently available claims, cost report, and ASP data when performing our ASP+X calculation under the final CY 2010 redistribution methodology in order to set payment rates for separately payable drugs and biologicals.

For CY 2010, we again attempted to address the issue of charge compression by proposing a methodology that reallocates pharmacy overhead costs from packaged drugs and biologicals to separately payable drugs and biologicals. We have made several proposals in the past to identify pharmacy overhead costs and address charge compression in the pharmacy revenue center. For the CY 2006 OPSS, we proposed to establish three distinct Level II HCPCS C-codes and three corresponding APCs for drug handling categories to differentiate overhead costs for drugs and biologicals (70 FR 42730). In the CY 2008 OPSS/ASC proposed rule (72 FR 42735), we proposed to instruct hospitals to remove the pharmacy overhead charge for both packaged and separately payable drugs and biologicals from the charge for the drug or biological and report the pharmacy overhead charge on an uncoded revenue code line on the claim. We believed that this would provide us with an avenue for collecting pharmacy handling cost data specific to drugs in order to package the overhead costs of these items into the associated procedures, most likely drug administration services. For CY 2009, we proposed to split the "Drugs Charged to Patients" cost center into two cost centers: one for drugs with high pharmacy overhead costs and one for drugs with low pharmacy overhead costs (73 FR 41492). We noted that we expected that CCRs from the proposed new cost centers would be available in 2 to 3 years to refine OPSS drug cost estimates by accounting for differential

hospital markup practices for drugs with high and low overhead costs. However, we did not finalize any of these proposals due to concerns from the hospital community that these proposals would create an overwhelming burden on hospitals and staff. We have once again proposed to address the issue of charge compression, in this case without requiring any changes to current hospital reporting practices.

It has been our policy, since CY 2006, to only use separately payable drugs and biologicals in the calculation of the equivalent average ASP-based payment amount under the OPSS. We do not include packaged drugs and biologicals in this standard analysis because cost data for these items are already accounted for within the APC ratesetting process through the median cost calculation methodology discussed in section II.A.2. of this final rule with comment period. To include the costs of coded packaged drugs and biologicals in both our APC ratesetting process (for associated procedures present on the same claim) and in our ratesetting process to establish an equivalent average ASP-based payment amount for separately payable drugs and biologicals would give these data disproportionate emphasis in the OPSS system by skewing our analyses, as the costs of these packaged items would be, in effect, counted twice. Accordingly, we are not adopting the suggestion from commenters that we include all packaged and separately payable drugs and biologicals when establishing an equivalent average ASP-based rate to provide payment for the hospital acquisition and pharmacy handling costs of drugs and biologicals. However, we remind commenters that because the costs of packaged drugs and biologicals, including their pharmacy overhead costs, are packaged into the payments for the procedures in which they are administered, the OPSS provides payment for both the drugs and the associated pharmacy overhead costs through the applicable procedural APC payments. Furthermore, we disagree with the commenters who recommend that we should pay separately for all drugs and biologicals with HCPCS codes. We continue to believe that packaging is a fundamental component of a prospective payment system that contributes to important flexibility and efficiency in the delivery of high quality hospital outpatient services and, therefore, we believe it is appropriate to maintain a modest drug packaging threshold that packages the costs of

inexpensive drugs into payment for the associated procedures.

Moreover, we continue to believe that excluding data from hospitals that participate in the 340B program from our ASP+X calculation, and paying those hospitals at that derived payment amount, would inappropriately redistribute payment to drugs and biologicals from payment for other services under the OPSS. The ASP-equivalent cost of drugs under the OPSS that would be calculated only from claims data for non-340B hospitals would likely be higher than the cost of all drugs from our aggregate claims for all hospitals. To set drug payment rates for all hospitals based on a subset of hospital cost data, determined only from claims data for non-340B hospitals would increase the final APC payment weights for drugs in a manner that does not reflect the drug costs of all hospitals, although all hospitals, including 340B hospitals, would be paid at these rates for drugs. Furthermore, as a consequence of the statutory requirement for budget neutrality, increasing the payment weights for drugs by excluding 340B hospital claims would reduce the relative payment weights for other services in a manner that does not reflect the procedural costs of all hospitals relative to the drug costs of all hospitals, thereby distorting the relativity of payment weights for services based on hospital costs. Many commenters on the CY 2009 OPSS/ASC final rule with comment period were generally opposed to differential payment for hospitals based on their 340B participation status, and we do not believe it would be appropriate to exclude claims from this subset of hospitals in the context of a CY 2010 drug and biological payment policy that pays all hospitals at the same rate for separately payable drugs and biologicals.

Therefore, for CY 2010, we are finalizing our proposed payment for separately payable drugs and biologicals, with modification. We are redistributing \$200 million from the cost of packaged drugs and biologicals to separately payable drugs and biologicals. The \$200 million consists of \$150 million (one-third of the pharmacy overhead cost) from the coded packaged drug and biological cost for drugs and biologicals with an ASP and \$50 million from the packaged drug and biological cost for drugs and biologicals without an ASP. To model this policy for the CY 2010 final rule with comment period, we reduced the cost of coded packaged drugs and biologicals with an ASP by 24 percent (based on final rule data; reduction was 27 percent based on

proposed rule data) and the cost of packaged drugs and biologicals without a HCPCS code or an ASP by 8 percent when we calculated the median cost of the CY 2010 procedural APCs. This redistribution results in payment for separately payable drugs and biologicals at a transitional rate of ASP+4 percent for CY 2010 under a pharmacy overhead adjustment methodology. We are, therefore, not accepting the August 2009 recommendation of the APC Panel to pay for separately payable drugs and biologicals at ASP+6 percent. Furthermore, because we are finalizing payment of separately payable drugs at APS+4 percent, we are not accepting the August 2009 APC Panel recommendations to analyze the impact of ASP+6 percent payment on different classes or for services that use packaged drugs compared with payment at ASP+4 percent. We are accepting the recommendation of the APC Panel to continue to refine our analyses of payment for drugs, biologicals, and radiopharmaceuticals and continue to welcome information and analyses from the public regarding pharmacy overhead costs.

Comment: One commenter requested that CMS create a HCPCS J-code for tositumomab, currently provided under a radioimmunotherapy regimen and billed as part of HCPCS code G3001 (Administration and supply of tositumomab, 450 mg). The commenter argued that because tositumomab is listed in compendia, is approved by the FDA as part of the BEXXAR® regimen, and has its own National Drug Code (NDC) number, it should be recognized as a drug and, therefore, be paid as other drugs are paid under the OPSS methodology, instead of having a payment rate determined by hospital claims data. The commenter suggested that a payment rate could be established using the ASP methodology.

Response: We have consistently noted that unlabeled tositumomab is not approved as either a drug or a radiopharmaceutical, but it is a supply that is required as part of the radioimmunotherapy treatment regimen (November 18, 2008 OPSS/ASC final rule with comment period (73 FR 68658); November 27, 2007 OPSS/ASC final rule with comment period for CY 2008 (72 FR 66765); November 10, 2005 OPSS final rule with comment period for CY 2006 (70 FR 68654); November 7, 2003 OPSS final rule with comment period for CY 2004 (68 FR 63443)). We do not make separate payment for supplies used in services provided under the OPSS. Payments for necessary supplies are packaged into payments for the separately payable services provided

by the hospital. Specifically, administration of unlabeled tositumomab is a complete service that qualifies for separate payment under its own clinical APC. This complete service is currently described by HCPCS code G3001. Therefore, we do not agree with the commenter's recommendation that we should assign a separate HCPCS code to the supply of unlabeled tositumomab. Rather, we will continue to make separate payment for the administration of tositumomab, and payment for the supply of unlabeled tositumomab is packaged into the administration payment.

We note that separately payable drug and biological payment rates listed in Addenda A and B to this final rule with comment period, which illustrate the final CY 2010 payment of ASP+4 percent for separately payable nonpass-through drugs and nonimplantable biologicals and ASP+6 percent for pass-through drugs and biologicals, reflect either ASP information that is the basis for calculating payment rates for drugs and biologicals in the physician's office setting effective October 1, 2009 or mean unit cost from CY 2008 claims data and updated cost report information available for this final rule with comment period. In general, these published payment rates are not reflective of actual January 2010 payment rates. This is because payment rates for drugs and biologicals with ASP information for January 2010 will be determined through the standard quarterly process where ASP data submitted by manufacturers for the third quarter of 2009 (July 1, 2009 through September 30, 2009) are used to set the payment rates that are released for the quarter beginning in January 2010 near the end of December 2009. In addition, payment rates for drugs and biologicals for which there was no ASP information available for October 2009 payment and, therefore, these products would be paid based on mean unit cost in CY 2010 based on available information at the time of this final rule with comment period, may have ASP information available for payment for the quarter beginning in January 2010. These drugs and biologicals would then be priced based on their newly available ASP information. Finally, there may be drugs and biologicals that have ASP information available for this final rule with comment period (reflecting October 2009 ASP data) that do not have ASP information available for the quarter beginning in January 2010. These drugs and biologicals would then be paid based on mean unit cost data derived from CY 2008 hospital claims.

Therefore, the payment rates listed in Addenda A and B to this final rule with comment period are not for January 2010 payment purposes and are only illustrative of the CY 2010 OPPS payment methodology using the most recently available information at the time of this final rule with comment period.

4. Payment for Blood Clotting Factors

For CY 2009, we are providing payment for blood clotting factors under the OPPS at ASP+4 percent, plus an additional payment for the furnishing fee. We note that when blood clotting factors are provided in physicians' offices under Medicare Part B and in other Medicare settings, a furnishing fee is also applied to the payment. The CY 2009 updated furnishing fee is \$0.164 per unit.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35333), for CY 2010, we proposed to pay for blood clotting factors at ASP+4 percent, consistent with our proposed payment policy for other nonpass-through separately payable drugs and biologicals, and to continue our policy for payment of the furnishing fee using an updated amount. Because the furnishing fee update is based on the percentage increase in the Consumer Price Index (CPI) for medical care for the 12-month period ending with June of the previous year and the Bureau of Labor Statistics releases the applicable CPI data after the MPFS and OPPS/ASC proposed rules are published, we were not able to include the actual updated furnishing fee in this proposed rule and we also are not able to include the actual updated furnishing fee in this final rule with comment period. Therefore, in accordance with our policy as finalized in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66765), we will announce the actual figure for the percent change in the applicable CPI and the updated furnishing fee calculated based on that figure through applicable program instructions and posting on the CMS Web site at: <http://www.cms.hhs.gov/McrPartBDrugAvgSalesPrice/>.

Comment: One commenter expressed support for CMS' proposal to continue to apply the furnishing fee for blood clotting factors provided in the OPD. The commenter stated that the furnishing fee helps ensure patient access to blood clotting factors by increasing the payment rate for these items. Another commenter disagreed with CMS' proposed payment rate of ASP+4 percent for blood clotting factors in CY 2010, even with the furnishing fee add-on. The commenters stated that ASP+4 percent was inadequate for all

drugs and biologicals, and is especially inappropriate for blood clotting factors. Finally, one commenter supported the payment of blood clotting factors at the same rate that applies to other nonpass-through separately payable drugs and biologicals in the HOPD.

Response: We continue to believe that applying the furnishing fee for blood clotting factors is appropriate for CY 2010. However, we see no compelling reason to provide payment for blood clotting factors under a different methodology for OPPS purposes at this time. We believe that the payment rate of ASP+4 percent that we are finalizing for payment of all nonpass-through separately payable drugs and biologicals in CY 2010 is appropriate. In addition, we believe that it continues to be appropriate to pay a furnishing fee for blood clotting factors under the OPPS, as is done in the physician's office setting and the inpatient hospital setting.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to provide payment for blood clotting factors under the same methodology as other separately payable drugs and biologicals under the OPPS and to continue paying an updated furnishing fee.

5. Payment for Therapeutic Radiopharmaceuticals

a. Background

Section 303(h) of Public Law 108-173 exempted radiopharmaceuticals from ASP pricing in the physician's office setting. Beginning in the CY 2005 OPPS final rule with comment period, we have exempted radiopharmaceutical manufacturers from reporting ASP data for payment purposes under the OPPS. (For more information, we refer readers to the CY 2005 OPPS final rule with comment period (69 FR 65811) and the CY 2006 OPPS final rule with comment period (70 FR 68655).) Consequently, we did not have ASP data for radiopharmaceuticals for consideration for previous years' OPPS ratesetting. In accordance with section 1833(t)(14)(B)(i)(I) of the Act, we have classified radiopharmaceuticals under the OPPS as SCODs. As such, we have paid for radiopharmaceuticals at average acquisition cost as determined by the Secretary and subject to any adjustment for overhead costs. Radiopharmaceuticals also are subject to the policies affecting all similarly classified OPPS drugs and biologicals, such as pass-through payment for diagnostic and therapeutic radiopharmaceuticals and individual

packaging determinations for therapeutic radiopharmaceuticals, discussed earlier in this proposed rule.

For CYs 2006 and 2007, we used mean unit cost data from hospital claims to determine each radiopharmaceutical's packaging status and implemented a temporary policy to pay for separately payable radiopharmaceuticals based on the hospital's charge for each radiopharmaceutical adjusted to cost using the hospital's overall CCR. In addition, in the CY 2006 OPPS final rule with comment period (70 FR 68654), we instructed hospitals to include charges for radiopharmaceutical handling in their charges for the radiopharmaceutical products so these costs would be reflected in the CY 2008 ratesetting process. The methodology of providing separate radiopharmaceutical payment based on charges adjusted to cost through application of an individual hospital's overall CCR for CYs 2006 and 2007 was finalized as an interim proxy for average acquisition cost because of the unique circumstances associated with providing radiopharmaceutical products to Medicare beneficiaries. The single OPPS payment represented Medicare payment for both the acquisition cost of the radiopharmaceutical and its associated handling costs.

During the CY 2006 and CY 2007 rulemaking processes, we encouraged hospitals and radiopharmaceutical stakeholders to assist us in developing a viable long-term prospective payment methodology for these products under the OPPS. As reiterated in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66766), we were pleased to note that we had many discussions with interested parties regarding the availability and limitations of radiopharmaceutical cost data.

In considering payment options for therapeutic radiopharmaceuticals for CY 2008, we examined several alternatives that we discussed in the CY 2008 OPPS/ASC proposed rule (72 FR 42738 through 42739) and CY 2008 OPPS/ASC final rule with comment period (72 FR 66769 through 66770). After considering the options and the public comments received, we finalized a CY 2008 methodology to provide prospective payment for therapeutic radiopharmaceuticals (defined as those Level II HCPCS codes that include the term "therapeutic" along with a radiopharmaceutical in their long code descriptors) using mean costs derived from the CY 2006 claims data, where the costs were determined using our standard methodology of applying hospital-specific departmental CCRs to

radiopharmaceutical charges, defaulting to hospital-specific overall CCRs only if appropriate departmental CCRs were unavailable (72 FR 66772). In addition, we finalized a policy to package payment for all diagnostic radiopharmaceuticals (defined as those Level II HCPCS codes that include the term “diagnostic” along with a radiopharmaceutical in their long code descriptors) for CY 2008. As discussed in the CY 2008 OPPS/ASC proposed rule (72 FR 42739), we believed that adopting prospective payment for therapeutic radiopharmaceuticals based on historical hospital claims data was appropriate because it served as our most accurate available proxy for the average hospital acquisition cost of separately payable therapeutic radiopharmaceuticals. In addition, we noted that we have found that our general prospective payment methodology based on historical hospital claims data results in more consistent, predictable, and equitable payment amounts across hospitals and likely provides incentives to hospitals for efficiently and economically providing these outpatient services.

Prior to the implementation of the final CY 2008 methodology of providing a prospective payment for therapeutic radiopharmaceuticals, section 106(b) of Public Law 110–173 was enacted on December 29, 2007 specifying payment for therapeutic radiopharmaceuticals based on individual hospital charges adjusted to cost. Therefore, hospitals continued to receive payment for therapeutic radiopharmaceuticals by applying the hospital-specific overall CCR to each hospital’s charge for a therapeutic radiopharmaceutical from January 1, 2008, through June 30, 2008. As we stated in the CY 2009 OPPS/ASC proposed rule (73 FR 41493), thereafter, the OPPS would provide payment for separately payable therapeutic radiopharmaceuticals on a prospective basis, with payment rates based upon mean costs from hospital claims data as set forth in the CY 2008 OPPS/ASC final rule with comment period, unless otherwise required by law.

Following issuance of the CY 2009 OPPS/ASC proposed rule, section 142 of Public Law 110–275 amended section 1833(t)(16)(C) of the Act, as amended by section 106(a) of Public Law 110–173, to further extend the payment period for therapeutic radiopharmaceuticals based on hospital’s charges adjusted to cost through December 31, 2009. Therefore, we are continuing to pay hospitals for therapeutic radiopharmaceuticals at charges adjusted to cost through the end of CY 2009.

b. Payment Policy

Since the start of the temporary cost-based payment methodology for radiopharmaceuticals in CY 2006, we have met with several interested parties on a number of occasions regarding payment under the OPPS for radiopharmaceuticals and have received numerous different suggestions from these stakeholders regarding payment methodologies that we could employ for future use under the OPPS.

In the CY 2008 OPPS/ASC final rule with comment period (72 FR 66771), we solicited comments requesting interested parties to provide information related to whether the existing ASP methodology could be used to establish payment for specific therapeutic radiopharmaceuticals under the OPPS. Similar to the recommendations we received during the CY 2008 OPPS/ASC proposed rule comment period (72 FR 66770), we received several suggestions regarding the establishment of an OPPS-specific methodology for radiopharmaceutical payment that would be similar to the ASP methodology, without following the established ASP procedures referenced at section 1847A of the Act and implemented through rulemaking. Some commenters recommended using external data submitted by a variety of sources other than manufacturers. Along this line, commenters suggested gathering information from nuclear pharmacies using methodologies with a variety of names such as Nuclear Pharmacy Calculated Invoiced Price (Averaged) (CIP) and Calculated Pharmacy Sales Price (CPSP). Other commenters recommended that CMS base payment for certain radiopharmaceuticals on manufacturer-reported ASP.

As noted in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66771), a ratesetting approach based on external data would be administratively burdensome for us because we would be required to collect, process, and review external information to ensure that it was valid, reliable, and representative of a diverse group of hospitals so that it could be used to establish rates for all hospitals. However, we specifically requested additional comments regarding the use of the existing ASP reporting structure for therapeutic radiopharmaceuticals as this established methodology was already used for payment of other drugs provided in the hospital outpatient setting (72 FR 66771). While we received several recommendations from commenters on the CY 2008 OPPS/ASC final rule with comment period regarding payment of

therapeutic radiopharmaceuticals based on estimated costs provided by manufacturers or other parties, we believe that the use of external data for payment of therapeutic radiopharmaceuticals should only be adopted if those external data are subject to the same well-established regulatory framework as the ASP data currently used for payment of separately payable drugs and biologicals under the OPPS. We have previously indicated that nondevice external data used for setting payment rates should be representative of a diverse group of hospitals both by location and type, and should also identify the relevant data sources. We do not believe that external therapeutic radiopharmaceutical cost data voluntarily provided outside of the established ASP methodology, either by manufacturers or nuclear pharmacies, would generally satisfy these criteria that are minimum standards for setting OPPS payment rates.

We received public comments on the CY 2008 OPPS/ASC final rule with comment period from certain radiopharmaceutical manufacturers who indicated that the standard ASP methodology could be used for payment of certain therapeutic radiopharmaceutical products. Specifically, these manufacturers expressed interest in providing ASP data for their therapeutic radiopharmaceutical products as a basis for payment under the OPPS.

In the CY 2009 OPPS/ASC proposed rule (73 FR 41495), we proposed to allow manufacturers to submit ASP information for any separately payable therapeutic radiopharmaceutical for payment purposes under the OPPS. If ASP information was not submitted or appropriately certified by the manufacturer for a given calendar year quarter, then for that quarter we proposed to provide prospective payment based on the therapeutic radiopharmaceuticals mean cost from hospital claims data. However, as stated above, section 142 of Public Law 110–275 amended section 1833(t)(16)(C) of the Act, as amended by section 106(a) of Public Law 110–173, to further extend the payment period for therapeutic radiopharmaceuticals based on hospital’s charges adjusted to cost through December 31, 2009, so we did not finalize this proposal. We note that, in response to our proposed therapeutic radiopharmaceutical payment methodology for CY 2009, we received a number of public comments that were supportive of the proposal for future years.

At the February 2009 meeting of the APC Panel, the APC Panel

recommended that CMS use the ASP methodology to pay for therapeutic radiopharmaceuticals and, where ASP data are not available, to pay based on mean costs from claims data for CY 2010. We accepted this recommendation, and for CY 2010, we proposed to allow manufacturers to submit ASP information for any separately payable therapeutic radiopharmaceutical in order for therapeutic radiopharmaceuticals to be paid based on ASP beginning in CY 2010 under the OPPTS (74 FR 35334 through 35336). Similar to our CY 2009 proposal, for CY 2010, we did not propose to compel manufacturers to submit ASP information. Furthermore, as discussed in the CY 2009 OPPTS/ASC proposed rule (73 FR 41495), we stated that the ASP data submitted would need to be provided for a patient-specific dose, or patient-ready form, of the therapeutic radiopharmaceutical in order to properly calculate the ASP amount for a given HCPCS code. In addition, in those instances where there is more than one manufacturer of a particular therapeutic radiopharmaceutical, we noted that all manufacturers would need to submit ASP information in order for payment to be made on an ASP basis. We specifically requested public comment on the development of a crosswalk, similar to the NDC/HCPCS crosswalk for separately payable drugs and biologicals posted on the CMS Web site at: http://www.cms.hhs.gov/McrPartBDrugAvgSalesPrice/01a1_2009aspfiles.asp#TopOfPage, for use for therapeutic radiopharmaceuticals.

Comment: Several commenters requested guidance on how costs associated with the manufacturing, compounding, and preparation of therapeutic radiopharmaceuticals should be reported when submitting ASP. The commenters stated that there are several activities that may take place at a variety of locations in order to prepare a radiopharmaceutical for patient administration. These services range in complexity from activities typical to any hospital pharmacy, such as drawing up a therapeutic radiopharmaceutical into a syringe and ensuring proper disposal of wasted product, to more complex processing such as preparing the therapeutic radiopharmaceutical itself by radiolabeling a cold kit (nonradioactive compound or complex that is combined with a radioisotope and results in a radiopharmaceutical) supplied by the manufacturer using a radioisotope supplied by the manufacturer or another

source. As CMS requested ASP information from separately payable radiopharmaceutical manufacturers in the form of a patient-specific dose, or patient-ready form, several commenters argued that the ASP amount reported to CMS should reflect all costs associated with these additional activities or items, such as the radioisotope and radiolabeling processes, needed to provide a patient-ready dose of a radiopharmaceutical. Several commenters pointed out that, based on current business practices, a single manufacturer might be unable to report ASP data for a therapeutic radiopharmaceutical described by a HCPCS code that incorporated all of these costs, and others were concerned that CMS intended for the manufacturer to collect and report the costs of the activities of other entities, such as freestanding radiopharmacies, that would not typically be reflected in the radiopharmaceutical manufacturer's sales price.

Response: In the following response, we discuss our expectations for manufacturer submission of ASP data to CMS to set OPPTS payment rates for separately payable therapeutic radiopharmaceuticals. This methodology also would apply to manufacturers submitting ASP data for diagnostic and therapeutic radiopharmaceuticals with pass-through status. As discussed in section V.A. of this final rule with comment period, we would use any submitted ASP information for separately payable diagnostic and therapeutic radiopharmaceuticals with pass-through status to establish a payment rate of ASP+6 percent, consistent with our policy for pass-through payment of drugs and biologicals. We note that ASP submissions for radiopharmaceutical payment under the OPPTS would need to meet all of the existing regulatory and subregulatory requirements of the ASP reporting process under sections 1847A and 1927(b)(3) of the Act, except as otherwise specified in this final rule with comment period.

For CY 2010, when reporting an ASP for a separately payable radiopharmaceutical, we expect that the ASP data reported by a manufacturer would be representative of the item(s) sold by the manufacturer. We used the term "patient-ready" in our proposed rule to ensure that ASP data submitted for OPPTS payment purposes for separately payable radiopharmaceuticals reflect the costs of all the component materials of the finished radiopharmaceutical product. We expect that the ASP data would represent the sales price of all of the

component materials of the finished radiopharmaceutical product sold by the manufacturer in terms that reflect the applicable HCPCS code descriptor, such as "treatment dose" or "millicurie." We understand that manufacturers of separately payable radiopharmaceuticals produce radiopharmaceuticals that require a variety of processing steps in order to finalize the product for administration to a beneficiary. For example, some radiopharmaceuticals are the combined product of a cold kit produced by one manufacturer, which is then radiolabeled with a radioisotope provided by a freestanding or hospital nuclear pharmacy. At a minimum, to be used for separate OPPTS radiopharmaceutical payment, the ASP data reported by a manufacturer must represent sales of all of the component materials associated with the radiopharmaceutical. In the context of radiopharmaceuticals used in the HOPD, we would expect that the component materials would include at least the cold kit and the radioisotope needed to radiolabel the cold kit in order to make the radiopharmaceutical.

With regard to additional processing steps, we believe manufacturers of radiopharmaceuticals could include in their calculations of ASP for OPPTS payment purposes in addition to the prices for the component materials, the portion of the sales price attributable to the production of the manufactured product as it is sold by the manufacturer reporting ASP data.

Radiopharmaceuticals are unique in that they require a radioisotope in addition to the cold kit and, at a minimum, they require radiolabeling the cold kit in order to produce a final radiopharmaceutical product. We note that manufacturers have the discretion to determine the form of the final product that is sold, and that the manufacturing process may include processing of the component materials in a variety of ways. To the extent that the price includes processing steps that are a service performed by the manufacturer to produce a radiopharmaceutical, we believe that ASP data submitted for purposes of calculating OPPTS payment for radiopharmaceuticals can appropriately capture those additional processing costs.

However, we do not believe that all processing steps that may be needed to prepare the separately payable radiopharmaceutical for administration to the beneficiary must be included in the submitted ASP data in order for the OPPTS to use manufacturer-reported ASP data as the basis for

radiopharmaceutical payment. We expect that the costs of any further processing of the radiopharmaceutical component materials after the manufacturer's sales, which could include radiolabeling when a manufacturer only sells the component materials or could consist of additional preparation besides radiolabeling, would not be included in the ASP data submitted by the manufacturer. However, these processing costs would be paid under the OPSS through the single ASP+4 percent payment rate for nonpass-through, separately payable therapeutic radiopharmaceuticals, in the same way that the OPSS currently pays for the acquisition and pharmacy overhead costs of separately payable drugs and biologicals through this single payment. We do not believe that it would be appropriate to include in the ASP for OPSS purposes the radiopharmaceutical processing services performed in a freestanding radiopharmacy or hospital pharmacy to prepare a final product or its component materials for patient administration after the manufactured product is sold by the manufacturer reporting ASP. In this case, the combined OPSS ASP+4 percent CY 2010 payment for the acquisition and pharmacy overhead and handling costs of the separately payable therapeutic radiopharmaceuticals would pay for these additional processing activities.

To be sufficient for purposes of calculating the OPSS payment, all radiopharmaceutical ASP submissions must meet the existing regulatory and subregulatory requirements of the ASP submission process under sections 1847A and 1927(b)(3) of the Act, except as otherwise specified in this final rule with comment period. In particular, we believe the "bona fide service fee" test in the ASP regulations is instructive here, and we would expect manufacturers to apply the "bona fide service fee" test to determine whether service fees it pays to another entity are "bona fide service fees." We believe the "bona fide service fee" test can be used in the OPSS ASP context to determine whether a fee that the manufacturer pays to a radiopharmacy for performing a service on behalf of the radiopharmaceutical manufacturer could be excluded from the ASP calculation—that is, it would not be considered a price concession that otherwise would reduce the ASP. The definition of a "bona fide service fee" is included in the ASP regulations (§ 414.802), which defines these fees as "fees paid by a manufacturer to an entity, that represent fair market value

for a bona fide, itemized service actually performed on behalf of the manufacturer that the manufacturer would otherwise perform (or contract for) in the absence of the service arrangement, and that are not passed on in whole or in part to a client or customer of an entity whether or not the entity takes title to the drug." In the context of the ASP calculation under section 1847A of the Act and its implementing regulations, "bona fide service fees" are not considered price concessions that must be deducted from the ASP. Similarly, we believe that for OPSS purposes, fees that are paid by the manufacturer that meet the "bona fide service fee" test would not reduce the ASP. Thus, a radiopharmaceutical manufacturer could contract with an entity, consistent with these regulations, to perform certain steps in the radiopharmaceutical manufacturing process that the manufacturer would otherwise perform itself in order to make the final radiopharmaceutical product, and the fees paid to the entity could qualify as a "bona fide service fee" that would be included in the ASP calculation for OPSS purposes. In light of the necessity of radiolabeling to the production of radiopharmaceuticals, we further believe that for OPSS purposes, the manufacturer's purchase of the radioisotope and payment for radiolabeling the cold kit could be factored into the manufacturer's price for the finished product, and if the fees the manufacturer paid meet the "bona fide service fee" test, they would not need to be netted against the price for purposes of calculating the manufacturer's ASP for the therapeutic radiopharmaceutical. Thus, in effect, for OPSS purposes, if a manufacturer chooses to contract for or purchase these items or services, fees for these bona fide services could be included in the manufacturer-reported ASP. We fully expect that the manufacturer would comply with the ASP regulations and, in particular, would factor these fees into the ASP only if the prices of the services performed by the radiopharmacy are fair market value, and the fees are not passed on to any purchasers of the separately payable radiopharmaceutical.

In summary, a patient-specific dose or patient-ready form in the context of OPSS ASP submission for radiopharmaceutical payment means that the ASP reflecting manufacturer sales must represent sales of all of the component materials for the radiopharmaceutical, including a minimum of a cold kit and a radioisotope, and be reported in terms that reflect the applicable HCPCS code

descriptor, such as "treatment dose" or "millicurie." The ASP would not necessarily take into account the preparation of the final form of the therapeutic radiopharmaceutical for patient administration, including radiolabeling, which may be conducted by the manufacturer, freestanding radiopharmacy, hospital pharmacy, or other entity. With respect to the latter, fees paid by the manufacturer for these services would be excluded from the ASP calculation (that is, would not be considered price concessions that reduce the ASP), only if they are "bona fide service fees" as defined in the regulations governing ASP. Thus, if the manufacturer pays a "bona fide service" fee for the services of the freestanding radiopharmacy, hospital pharmacy, or other entity, and reflects that fee in its price for the radiopharmaceutical, the amount of the "bona fide service fee" would be taken into account in the reported ASP data. However, manufacturers are not required to pay for the preparation of a radiopharmaceutical (including radiolabeling) in a freestanding radiopharmacy, hospital pharmacy, or other entity after sale of all of the component materials, and in that case, the cost of those services would not be reflected in the ASP data submitted to CMS. Manufacturers should submit ASP data for a separately payable radiopharmaceutical that incorporates prices for sales of all of the component materials by the manufacturer. Any additional costs associated with the preparation of the radiopharmaceutical for administration to a beneficiary after the manufacturer's sale of the component materials and any processing that the manufacturer conducts would be paid through the single OPSS ASP+4 percent payment for the acquisition and pharmacy overhead and handling costs of the nonpass-through, separately payable therapeutic radiopharmaceutical.

Comment: Several commenters expressed support for CMS's proposal to make prospective payment for therapeutic radiopharmaceuticals using the ASP methodology in CY 2010 to pay the same ASP+4 percent payment rate that CMS proposed for separately payable nonpass-through drugs and nonimplantable biologicals. A few commenters suggested that CMS consider alternatives to the percentage-based add-on to ASP inherent in the single combined payment for acquisition and handling costs of ASP+4 percent to better account for the more intensive handling that radiopharmaceuticals require. The

commenters recommended a variety of options, including a fixed add-on payment comparable to the complexity of handling involved, a consignment ASP method that would account for the costs associated with the handling of a radiopharmaceutical, the preparation of an invoice that would give a standard drug percentage for handling charges, or the establishment of some other separate payment mechanism to capture the costs of radiolabeling. One commenter suggested that CMS could create a Level II HCPCS code for hospitals to report their charges for radiolabeling conducted by a radiopharmacy, and hospital cost information developed from these charges could be used to establish a separate payment for radiolabeling services.

Response: We appreciate the commenters' support for our proposal to make prospective payment for nonpass-through, separately payable therapeutic radiopharmaceuticals at the same ASP+4 percent payment rate for a "patient-ready" dose of a radiopharmaceutical that we establish for separately payable drugs and nonimplantable biologicals. In our response to the previous comment, we established our interpretation of "patient-ready" for purposes of the OPSS to mean the ASP, reported in terms that reflect the applicable HCPCS code descriptor, for all component materials of the radiopharmaceutical and any additional processing, including radiolabeling, that is reflected in the price the manufacturer charges for the radiopharmaceutical so long as the fees paid for such additional processing meet the "bona fide service fee" test under the regulations implementing section 1847A of the Act. We explicitly note that because radiopharmaceuticals uniquely require radiolabeling of their component materials, we believe that for purposes of OPSS ASP reporting, radiolabeling could constitute a bona fide service on behalf of the manufacturer, and the fees for which could meet the "bona fide service fee" test. Given our position on radiolabeling, we similarly believe that significant processing costs associated with handling a radiopharmaceutical may be reflected in the prices used to calculate the manufacturer's ASP data for OPSS purposes. As noted above, the combined single payment for nonpass-through, separately payable therapeutic radiopharmaceutical acquisition and overhead costs embodied in the ASP+4 percent payment rate for CY 2010 would address any other processing after the sale by the manufacturer, and we believe this payment is sufficient for

these additional handling costs borne by the hospital. Under this interpretation of "patient-ready" dose, we do not believe that making an additional payment for more intensive handling costs is necessary.

We also do not believe that establishing a separate Level II HCPCS code to exclusively capture radiopharmaceutical handling costs is an appropriate approach when we have not adopted such an approach to capture the pharmacy overhead costs of other drugs and biologicals, which also may be substantial in some cases. We have heard from hospitals previously on the issue of separately reporting charges for pharmacy handling costs of drugs. In the CY 2007 OPSS/ASC final rule with comment period (71 FR 68090), we discussed our efforts to create a set of Level II HCPCS codes that hospitals would be able to use to indicate the relative resource levels of pharmacy handling involved in preparing a reported drug, biological, or radiopharmaceutical for administration. This methodology would have allowed us to begin collecting data on pharmacy overhead costs for possible use in future ratesetting calculations, yet we did not finalize this proposal due to the overwhelming response from the hospital community citing the tremendous administrative burden separately reporting these pharmacy handling codes and charges would have placed on hospital resources. We continue to believe that hospitals would likely view such an approach for radiopharmaceuticals alone as burdensome.

Comment: Several commenters responded to CMS' request for public comments on the development of a crosswalk similar to the NDC/HCPCS crosswalk for separately payable drugs and biologicals. These commenters support a NDC/HCPCS "crosswalk" to allow ASP to be utilized.

Response: We appreciate the commenter's support for implementing a "crosswalk" for use for separately payable radiopharmaceuticals. We believe that an NDC/HCPCS crosswalk for nonpass-through, separately payable therapeutic radiopharmaceuticals and pass-through diagnostic and therapeutic radiopharmaceuticals that is similar to the crosswalk for separately payable drugs and biologicals is appropriate, and we will, therefore, work to develop and implement the appropriate NDC/HCPCS crosswalk for separately payable radiopharmaceuticals.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35335), we stated that we continue to believe that the use of ASP information for OPSS payment would

provide an opportunity to improve payment accuracy for separately payable radiopharmaceuticals by applying an established methodology that has already been successfully implemented under the OPSS for other separately payable drugs and biologicals. As is the case with other drugs and biologicals subject to ASP reporting under section 1847A of the Act, we stated that in order for a separately payable radiopharmaceutical to receive OPSS payment based on ASP beginning January 1, 2010, we would need to receive ASP information from the manufacturer no later than November 2, 2009 that would reflect separately payable radiopharmaceutical sales in the third quarter of CY 2009 (July 1, 2009 through September 30, 2009). Our normal deadline for January submission is November 1, but because November 1 falls on a Sunday, the ASP submission deadline for January 2010 payment is November 2, 2009. We stated that these data would not be available for publication in the CY 2010 OPSS/ASC final rule with comment period but would be included in the January 2010 OPSS quarterly release that would update the payment rates for separately payable drugs, biologicals, and therapeutic radiopharmaceuticals based on the most recent ASP data, consistent with our customary practice over the past 4 years when we have used the ASP methodology for payment of separately payable drugs and biologicals under the OPSS. In addition, we proposed to receive information from radiopharmaceutical manufacturers that would allow us to calculate a unit dose cost estimate based on the applicable HCPCS code for the separately payable radiopharmaceutical.

In the CY 2010 OPSS/ASC proposed rule (74 FR 35335), we acknowledged that we realized that not all therapeutic radiopharmaceutical manufacturers may be willing or able to submit ASP information for a variety of reasons. We proposed to provide payment at the OPSS ASP rate if ASP information is available for a given calendar year quarter or, if ASP information is not available, we proposed to provide payment based on the most recent hospital mean unit cost data that we have available. We indicated our belief that both methodologies represent an appropriate and adequate proxy for average hospital acquisition cost and associated handling costs for these products. Therefore, if ASP information for the appropriate period of sales related to payment in any CY 2010 quarter was not available, we proposed to rely on the CY 2008 mean unit cost

data derived from hospital claims to set the payment rates for therapeutic radiopharmaceuticals. We noted that this is not the usual OPPS process that relies on alternative data sources, such as WAC or AWP, when ASP information is temporarily unavailable, prior to defaulting to the mean unit cost from hospital claims data. We proposed a methodology specific to nonpass-through, separately payable therapeutic radiopharmaceuticals where we would immediately default to the mean unit cost from hospital claims data if sufficient ASP data were not available because we did not propose to require therapeutic radiopharmaceutical manufacturers to report ASP data at this time. We indicated that we did not believe that WAC or AWP is an appropriate proxy to provide OPPS payment for average therapeutic radiopharmaceutical acquisition cost and associated handling costs when manufacturers are not required to submit ASP data. Payment based on WAC or AWP under the established OPPS ASP methodology for payment of separately payable drugs and biologicals is usually temporary for a calendar quarter until a manufacturer is able to submit the required ASP data in accordance with the quarterly ASP submission timeframes for reporting under section 1847A of the Act. However, we were concerned that because ASP reporting for OPPS payment of separately payable therapeutic radiopharmaceuticals would not be required for CY 2010, a manufacturer's choice to not submit ASP could result in payment for a separately payable therapeutic radiopharmaceutical based on WAC or AWP for a full year, a result which we believed would be inappropriate. Therefore, for separately payable therapeutic radiopharmaceutical payment under the OPPS, we proposed that the OPPS ASP methodology would pay based on ASP, with payment based on mean unit cost from OPPS claims data if ASP data were not available for a calendar quarter.

Recognizing that we may need to utilize mean unit cost data to pay for nonpass-through, separately payable therapeutic radiopharmaceuticals in CY 2010 if ASP data are not submitted, for the CY 2010 proposed rule we evaluated the mean unit cost information in the CY 2010 claims data for all therapeutic radiopharmaceuticals. We noticed that we had numerous claims with service units greater than one for HCPCS code A9543 (Yttrium Y-90 ibritumomab tiuxetan, therapeutic, per treatment dose, up to 40 millicuries) and A9545

(Iodine I-131 tositumomab, therapeutic, per treatment dose), when the long descriptors for these therapeutic radiopharmaceuticals clearly indicate "per treatment dose" and, therefore, we expected the service units on every claim to be one. In contrast, the other six therapeutic radiopharmaceuticals that would be separately payable in CY 2010 all include "per millicurie" in their HCPCS code descriptors, so reporting multiple service units for those items could be appropriate. We did not believe that hospitals billing more than one unit of HCPCS code A9543 or A9545 on a claim were correctly reporting these products and, therefore, we believed that these claims were incorrectly coded. Although we do not normally examine hospital reporting patterns for individual services, pricing an individual item, such as a therapeutic radiopharmaceutical with low volume, may argue for more aggressive trimming to remove inaccurate claims. Therefore, we removed all claims with units greater than one for these two therapeutic radiopharmaceuticals before estimating their mean unit costs. Because we did not have ASP data for therapeutic radiopharmaceuticals that were used for payment in April 2009, the proposed payment rates included in Addenda A and B to the proposed rule were based on mean costs from historical hospital claims data available for the proposed rule, subject to the additional trimming of incorrectly coded claims for HCPCS codes A9543 and A9545 as described above.

Similar to the ASP process already in place for separately payable drugs and biologicals under the OPPS, we proposed to update ASP data for therapeutic radiopharmaceuticals through our quarterly process as updates become available. In addition, we proposed to assess the availability of ASP data for therapeutic radiopharmaceuticals quarterly, and if ASP data become available midyear, we proposed to transition to the next available quarter to ASP-based payment. For example, if ASP data are not available for the quarter beginning January 2010 (that is, ASP information reflective of third quarter CY 2009 sales are not submitted in November 2009), then the next opportunity to begin payment based on ASP data for a therapeutic radiopharmaceutical would be April 2010 if ASP data reflective of fourth quarter CY 2009 sales were submitted in February 2010.

Comment: Many commenters agreed with CMS' proposal to permit, but not require, radiopharmaceutical manufacturers to submit ASP data. One

commenter encouraged CMS to obtain data from all therapeutic radiopharmaceutical manufacturers across all therapeutic radiopharmaceuticals, not just a few. Several commenters expressed concern over the proposed immediate collection of ASP data from manufacturers for the January 2010 OPPS quarterly update. They stated that manufacturers would need an adequate amount of time to submit ASP data for the third quarter of CY 2009 (July 1, 2009 through September 30, 2009).

A few commenters recommended that CMS establish a transition period of 6 months or longer to provide more time for manufacturers to compile and submit ASP data. Another commenter recommended that CMS accept therapeutic radiopharmaceutical ASP data 30 days after finalizing and publishing CMS' CY 2010 OPPS/ASC final rule. This would extend the deadline for which ASP data would be submitted for the January 2010 OPPS quarterly update from the usual November 2, 2009 ASP submission deadline to November 30, 2009.

During a transition period, several commenters recommended that CMS continue its current policy of paying for therapeutic radiopharmaceuticals at charges adjusted to cost, as opposed to the proposed default of mean unit cost derived from claims data. A number of commenters requested open dialogue with CMS on what a manufacturer would need to submit to accurately report ASP for a "patient-ready" radiopharmaceutical dose.

Response: We appreciate the commenters' support for our proposal to pay for nonpass-through, separately payable therapeutic radiopharmaceuticals in CY 2010 using the ASP methodology. We proposed to allow manufacturers to submit ASP information for any nonpass-through, separately payable therapeutic radiopharmaceutical in order to establish an ASP+4 percent payment rate under the OPPS for the therapeutic radiopharmaceutical beginning in CY 2010. Consistent with our authority to collect data in order to determine payment amounts, we intend to collect ASP data for separately payable nonpass-through and pass-through therapeutic radiopharmaceuticals (and diagnostic radiopharmaceuticals with pass-through status), adopting the same submission requirements as we do for drugs and biologicals under section 1847A of the Act and the corresponding regulations, except as otherwise specified in this final rule with comment period for specific OPPS purposes. As we stated in the CY 2010

OPPS/ASC proposed rule (74 CR 35335), we continue to believe that the use of ASP information for OPPS payment would provide an opportunity to improve payment accuracy for separately payable radiopharmaceuticals by applying an established methodology that has already been successfully implemented under the OPPS to set prospective payment rates for other separately payable drugs and biologicals.

In recognizing the potential burden involved in reporting ASP data and our belief in the accuracy of prospective payment rates based on claims data, we did not propose to require manufacturers to submit ASP information. Although one commenter suggested that we collect ASP from all manufacturers of therapeutic radiopharmaceuticals, we continue to believe that the challenges involved in reporting ASP for radiopharmaceuticals, given the variety of manufacturing processes, are significant in some cases and, therefore, that payment based on mean unit cost from historical hospital claims data offers the best proxy for average hospital acquisition cost and associated handling costs for a radiopharmaceutical in the absence of ASP. We continue to believe that we should allow, but not require, manufacturers to submit ASP information for therapeutic radiopharmaceuticals. If ASP information is unavailable for a therapeutic radiopharmaceutical, meaning if a manufacturer is not willing or not able to submit ASP information, we will provide payment based on the mean unit cost of the product that is applicable to payment rates for the year the nonpass-through therapeutic radiopharmaceutical is administered. We continue to believe that both methodologies represent an appropriate proxy for average hospital acquisition cost and associated handling costs for nonpass-through, separately payable therapeutic radiopharmaceuticals. We expect manufacturers to submit ASP data for all component materials and any "bona fide service fees" that are reflected in the price for additional processing to produce the separately payable radiopharmaceutical, in an aggregated form in per millicurie or per dosage unit that matches the HCPCS code descriptor for that radiopharmaceutical. We note that the separately payable, nonpass-through therapeutic radiopharmaceutical payment rates listed in Addenda A and B to this final rule with comment period are the mean unit costs from CY 2008 hospital claims data, subject to the

additional trimming of incorrectly coded claims for HCPCS codes A9543 and A9545 as proposed and described above, that we would use for payment of a separately payable radiopharmaceutical if ASP information from the manufacturer were not submitted for the product for the applicable OPPS payment quarter.

For CY 2010, we are not implementing a transition period of payment at charges adjusted to cost for therapeutic radiopharmaceuticals. We note that section 1833(t)(16)(C) of the Act continues the payment period for therapeutic radiopharmaceuticals based on a hospital's charges adjusted to cost through December 31, 2009, and this requirement expires beginning CY 2010. We believe it would not be consistent with the statutory expiration of the charges-adjusted-to-cost payment methodology to continue payment using this approach for any portion of CY 2010. For manufacturers that cannot initially submit ASP data, we believe that mean cost payment for a nonpass-through, separately payable therapeutic radiopharmaceutical provides our best proxy estimate of average hospital acquisition cost and associated handling costs and implements prospective payment. In examining the CY 2008 claims data, aggregate therapeutic radiopharmaceutical payment at mean unit cost would be comparable to payment at charges adjusted to cost in CY 2010 assuming no charge inflation between CY 2008 and CY 2010, and we observe deflation in per unit charges for some therapeutic radiopharmaceuticals between CY 2007 and CY 2008 claims. Finally, because we proposed to update payment based on ASP submissions on a quarterly basis, manufacturers would not need to wait one year to be paid based on ASP but could work toward submitting ASP data for April 2010 payment if they were unable to provide data for January 2010 payment.

We recognize that the timeframe for submitting ASP information by November 2, 2009, to begin ASP-based payment on January 1, 2010 is extremely close to the display date of this final rule with comment period. While we expect that most manufacturers interested in reporting ASP for their therapeutic radiopharmaceutical already have begun the process of compiling that data given that we have proposed ASP-based payment under the OPPS for nonpass-through, separately payable therapeutic radiopharmaceuticals for 2 years in a row, we understand that manufacturers will not have had the opportunity to consider our discussion in this final rule with comment period that clarifies the

term "patient-ready" in their preparation of ASP data for OPPS purposes. As suggested by the commenters, we recognize that some manufacturers may need to discuss with us how to report ASP for a "patient-ready" dose of their particular radiopharmaceutical. We encourage manufacturers with questions regarding their submissions to contact us, especially if they intend to submit by November 2, 2009. Manufacturers can contact us immediately by sending an email to the OPPS mailbox: OutpatientPPS@cms.hhs.gov. We will be monitoring this mailbox closely. We will provide any assistance that we can within the confines of the ASP quarterly production schedule to facilitate accurate and timely reporting of ASP and payment based on ASP as early as possible. To further our commitment to helping manufacturers submit ASP data in a timely fashion, we also intend to post guidance on the definition of "patient-ready" dose for reporting radiopharmaceutical ASP for OPPS use, and on how manufacturers should compile and submit ASP data for that dose on the CMS Web site at: http://www.cms.hhs.gov/HospitalOutpatientPPS/05_OPSPGuidance.asp#TopOfPage, at about the time that this final rule with comment period goes in display at the **Federal Register**.

Comment: One commenter expressed concern over CMS proposal to pay for therapeutic radiopharmaceuticals using the ASP methodology for CY 2010. The commenter stated that the methodology established in the proposal, to pay for therapeutic radiopharmaceuticals under the ASP methodology and if ASP is unavailable to make payment based upon the most recent hospital mean unit cost data that CMS has available, would not provide accurate data and, therefore, would not pay accurately for therapeutic radiopharmaceuticals. The commenter was skeptical that manufacturers would submit ASP data in a timely and accurate manner, because the commenter believed manufacturers have little incentive to do so. The commenter recommended that CMS base payment on hospital invoice data in order to provide accurate payment.

Response: We disagree with the commenter and continue to believe that providing payment for therapeutic radiopharmaceuticals based on ASP or mean unit cost if ASP information is not available would provide appropriate payment for these products. We acknowledge in the proposed rule (74 FR 35335) that some manufacturers may be unable or unwilling to submit ASP

data for the CY 2010 January OPPS quarterly update and we, therefore, proposed to make payment based on the most recent hospital mean unit cost data that we have available for therapeutic radiopharmaceuticals if ASP is not available. Many other commenters, including radiopharmaceutical manufacturers, stated that manufacturers have a significant incentive to submit ASP information because they believe payment based on the default of mean unit cost would not be most reflective of the cost to hospitals to acquire these products. Furthermore, some therapeutic radiopharmaceutical manufacturers already submit ASP data for separately payable drugs and biologicals and, therefore, are familiar with the submission process, including its timing and other requirements. As we stated previously, we continue to believe that the use of ASP information for OPPS payment would provide an opportunity to improve payment accuracy for nonpass-through, separately payable therapeutic radiopharmaceuticals by applying an established methodology that has already been successfully implemented under the OPPS for other separately payable drugs and biologicals. The OPPS has relied upon ASP information as an accurate method for providing payment for drugs and biologicals for several years.

Comment: One commenter requested that CMS not include payment rates based on mean unit cost for therapeutic radiopharmaceuticals in the Addenda to the CY 2010 OPPS/ASC final rule with comment period when a manufacturer intends to report ASP information for CY 2010. The commenter offered this recommendation in order to avoid having other payers that utilize Medicare payment rates adopt these payment rates that the commenter believes will never be those paid to hospitals in CY 2010.

Response: We believe that payment at mean unit cost would appropriately pay for the average hospital acquisition cost and associated handling costs of nonpass-through, separately payable therapeutic radiopharmaceuticals if ASP data are not available. Therefore, we have included the mean unit cost amounts for nonpass-through, separately payable therapeutic radiopharmaceuticals in Addenda A and B to this CY 2010 OPPS/ASC final rule with comment period that would be used for payment in any CY 2010 calendar quarter for which ASP information is not submitted by the product's manufacturer(s). Inclusion of mean unit cost for all nonpass-through, separately payable therapeutic

radiopharmaceuticals in the addenda is unique to CY 2010, because we have no ASP information available for these products based on their payment in CY 2009. For future years, based on our usual final rule addenda publication policy, we note that if a radiopharmaceutical manufacturer has submitted ASP for OPPS payment in October 2010, as long as our CY 2011 payment methodology for nonpass-through, separately payable therapeutic radiopharmaceuticals relies on ASP, we would publish a payment rate in the CY 2011 OPPS/ASC final rule with comment period that reflects the radiopharmaceutical's third quarter CY 2010 ASP information.

We follow this final rule addenda publication policy based on our general expectation that drugs and biologicals with ASP information available for payment in the fourth quarter of CY 2009 will have ASP information available for payment in the first quarter of CY 2010. Therefore, we believe that posting illustrative CY 2010 payment rates for drugs and biologicals based on the October 2009 ASP information, rather than mean unit cost, provides a better illustration of the likely payment rates for these product in January 2010. In the event that we have no ASP information for a therapeutic radiopharmaceutical or any other drug or biological paid based on the ASP methodology for any quarter of CY 2010, the applicable mean unit cost for payment of the product in CY 2010 is available on the CMS web site in the OPPS drug median file that is posted as supporting information for this final rule with comment period at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/HORD/list.asp#TopOfPage>.

Comment: One commenter requested that CMS establish a temporary Level II HCPCS C-code, effective January 1, 2010, to be used in CY 2010 to report the product currently described by HCPCS code A9605 (Samarium Sm-153 Lexidronam, per 50 millicuries). The commenter recommended that CMS replace the current "per 50 millicuries" in the code descriptor with "per treatment dose" in the HCPCS C-code descriptor. The commenter stated that the current code descriptor for HCPCS code A9605 is problematic for coding and ASP reporting purposes because of the decay in radiopharmaceutical radioactivity. Under the existing code descriptor of "per 50 millicuries," while the manufacturer would report ASP for the HCPCS code based on the radioactivity level of the product at the time of sale, the hospital would report units of the HCPCS code based on the

dose administered to the patient at a later point in time, and there would be a mismatch between the reported price and the dose actually administered to the patient. The commenter concluded that reporting the sales and administration of this product on a "per treatment dose" basis would allow ASP information to be aligned with payment for the product under the OPPS, taking into consideration radioactive decay over time since administration would always occur after the manufacturer's sale of the product.

Response: We understand the concerns raised by the commenter in regards to the current code descriptor for A9605. In response to these concerns, CMS' HCPCS Workgroup has decided to create a new HCPCS code, A9604 Samarium SM-153 lexidronam, therapeutic, per treatment dose, up to 150 millicuries), effective January 1, 2010, and delete existing HCPCS code A9605. This new code should facilitate alignment between ASP reporting by the manufacturer and hospital reporting of administration on a "per treatment dose" basis. We note that the default payment rate for HCPCS code A9604 included in Addendum A and B to this final rule with comment period is based on the per-day mean unit cost of HCPCS code A9605. We believe that the CY 2008 hospital per-day cost of HCPCS code A9605 reflects the cost of a single treatment dose and, therefore, it is the mean per-day cost that we will use for payment of new HCPCS code A9604 in CY 2010 if ASP information is not available.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to pay all nonpass-through, separately payable therapeutic radiopharmaceuticals at ASP+4 percent based on ASP information, if available, for a "patient-ready" dose beginning on January 1, 2010, and updated on a quarterly basis for products for which manufacturers report ASP data. We are defining a "patient-ready" dose for OPPS purposes as including all component materials of the radiopharmaceutical, at a minimum, and any other processing the manufacturer requires to produce the radiopharmaceutical that it sells that are reflected in the sales price, including radiolabeling, as long as any fees paid for such processing done on behalf of the manufacturer meet the definition of "bona fide service fees" under § 414.802. We also are finalizing our CY 2010 proposal, without modification, to base nonpass-through, separately payable therapeutic radiopharmaceutical payment on mean

unit cost derived from CY 2008 claims data when ASP pricing is not available. The nonpass-through therapeutic radiopharmaceuticals that are separately payable in CY 2010 are identified in Table 42 below. The CY 2010 payment rates for these products included in

Addenda A and B to this final rule with comment period are based on mean unit cost. Moreover, we note that, should ASP be submitted timely for January 2010 OPPS payment, according to our usual process for updating the payment rates for separately payable drugs and

biologicals on a quarterly basis if updated ASP information is available, these payment rates will be updated through the January 2010 OPPS quarterly release.

TABLE 42—CY 2010 NONPASS-THROUGH, SEPARATELY PAYABLE THERAPEUTIC RADIOPHARMACEUTICALS

CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 APC	Final CY 2010 SI
A9517	Iodine I-131 sodium iodide capsule(s), therapeutic, per millicurie	1064	K
A9530	Iodine I-131 sodium iodide solution, therapeutic, per millicurie	1150	K
A9543	Yttrium Y-90 ibritumomab tiuxetan, therapeutic, per treatment dose, up to 40 millicuries.	1643	K
A9545	Iodine I-131 tositumomab, therapeutic, per treatment dose	1645	K
A9563	Sodium phosphate P-32, therapeutic, per millicurie	1675	K
A9564	Chronic phosphate P-32 suspension, therapeutic, per millicurie	1676	K
A9600	Strontium Sr-89 chloride, therapeutic, per millicurie	0701	K
A9604	Samarium SM-153 leixidronam, therapeutic, per treatment dose, up to 150 millicuries.	1295	K

6. Payment for Nonpass-Through Drugs, Biologicals, and Radiopharmaceuticals With HCPCS Codes, but Without OPPS Hospital Claims Data

Public Law 108-173 does not address the OPPS payment in CY 2005 and after for drugs, biologicals, and radiopharmaceuticals that have assigned HCPCS codes, but that do not have a reference AWP or approval for payment as pass-through drugs or biologicals. Because there is no statutory provision that dictated payment for such drugs, biologicals, and radiopharmaceuticals in CY 2005, and because we had no hospital claims data to use in establishing a payment rate for them, we investigated several payment options for CY 2005 and discussed them in detail in the CY 2005 OPPS final rule with comment period (69 FR 65797 through 65799).

For CYs 2005 to 2007, we implemented a policy to provide separate payment for new drugs, biologicals, and radiopharmaceuticals with HCPCS codes (specifically those new drug, biological, and radiopharmaceutical HCPCS codes in each of those calendar years that did not crosswalk to predecessor HCPCS codes) but which did not have pass-through status, at a rate that was equivalent to the payment they received in the physician's office setting, established in accordance with the ASP methodology for drugs and biologicals, and based on charges adjusted to cost for radiopharmaceuticals. For CYs 2008 and 2009, we finalized a policy to provide payment for new drugs (excluding contrast agents) and biologicals (excluding implantable biologicals for

CY 2009) with HCPCS codes, but which did not have pass-through status and were without OPPS hospital claims data, at ASP+5 percent and ASP+4 percent, respectively, consistent with the final OPPS payment methodology for other separately payable drugs and biologicals. New therapeutic radiopharmaceuticals were paid at charges adjusted to cost based on the statutory requirement for CY 2008 and CY 2009 and payment for new diagnostic radiopharmaceuticals was packaged in both years. In the CY 2010 OPPS/ASC proposed rule (74 FR 35336 through 35337), for CY 2010, we proposed to continue the CY 2009 payment methodology for new drugs (excluding contrast agents) and nonimplantable biologicals and extend the methodology to payment for new therapeutic radiopharmaceuticals, when their period of payment at charges adjusted to cost no longer would apply. Therefore, for CY 2010, we proposed to provide payment for new drugs (excluding contrast agents), nonimplantable biologicals, and therapeutic radiopharmaceuticals with HCPCS codes (those new CY 2010 drug (excluding contrast agents), nonimplantable biological, and therapeutic radiopharmaceutical HCPCS codes that do not crosswalk to CY 2009 HCPCS codes), but which do not have pass-through status and are without OPPS hospital claims data, at ASP+4 percent, consistent with the proposed CY 2010 payment methodology for other separately payable nonpass-through drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals. We believed this proposed policy would ensure that new nonpass-through drugs,

nonimplantable biologicals, and therapeutic radiopharmaceuticals would be treated like other drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals under the OPPS, unless they are granted pass-through status. Only if they are pass-through drugs, nonimplantable biologicals, or therapeutic radiopharmaceuticals would they receive a different payment for CY 2010, generally equivalent to the payment these drugs and biologicals would receive in the physician's office setting, consistent with the requirements of the statute. We proposed to continue packaging payment for all new nonpass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals with HCPCS codes (those new CY 2010 diagnostic radiopharmaceutical, contrast agent, and implantable biological HCPCS codes that do not crosswalk to predecessor HCPCS codes), consistent with the proposed packaging of all existing nonpass-through diagnostic radiopharmaceuticals, contrast agents and implantable biologicals, as discussed in more detail in the proposed rule (74 FR 35323 through 35324). In accordance with the OPPS ASP methodology, in the absence of ASP data, in the CY 2010 OPPS/ASC proposed rule (74 FR 35336), for CY 2010, we proposed to continue the policy we implemented beginning in CY 2005 of using the WAC for the product to establish the initial payment rate for new nonpass-through drugs and biologicals with HCPCS codes, but which are without OPPS claims data. However, we note that if the WAC is

also unavailable, we would make payment at 95 percent of the product's most recent AWP. We also proposed to assign status indicator "K" to HCPCS codes for new drugs and nonimplantable biologicals without OPPS claims data and for which we have not granted pass-through status. We further noted that, with respect to new items for which we do not have ASP data, once their ASP data become available in later quarter submissions, their payment rates under the OPPS would be adjusted so that the rates would be based on the ASP methodology and set to the finalized ASP-based amount (proposed for CY 2010 at ASP+4 percent) for items that have not been granted pass-through status.

We did not receive any public comments specific to these proposals. While commenters, in general, objected to payment for drugs and biologicals at ASP+4 percent, these comments were not specific to new drugs and biologicals with HCPCS codes but without OPPS claims data. Further, we summarize the general public comments on payment for separately payable drugs and provide our responses in section V.B.3.b. of this final rule with comment period. In addition, commenters on the CY 2010 OPPS/ASC proposed rule objected to packaging payment for diagnostic radiopharmaceuticals and contrast agents in general, but these comments were not directed to new diagnostic radiopharmaceuticals or contrast agents with HCPCS codes but without OPPS claims data. We summarize these comments and provide our responses in section V.A.2.d. of this final rule with comment period.

Therefore, we are finalizing our CY 2010 proposals, without modification, as follows: Payment for new drugs (excluding contrast agents), nonimplantable biologicals, and therapeutic radiopharmaceuticals with HCPCS codes, but which do not have pass-through status and are without OPPS hospital claims data, will be made at ASP+4 percent for CY 2010. In cases where ASP information is not available, payment will be made using WAC, and if WAC is also unavailable payment will be made at 95 percent of the most recent AWP. Further, payment for all new nonpass-through diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals with HCPCS codes but without OPPS claims data will be packaged for CY 2010. Finally, we are assigning status indicator "K" to HCPCS codes for new drugs and nonimplantable biologicals without OPPS claims data and for

which we have not granted pass-through status in CY 2010.

For CY 2010, we also proposed to base payment for new therapeutic radiopharmaceuticals with HCPCS codes as of January 1, 2010, but which do not have pass-through status, on the WACs for these products if ASP data for these therapeutic radiopharmaceuticals are not available. If the WACs are also unavailable, we proposed to make payment for new therapeutic radiopharmaceuticals at 95 percent of their most recent AWP's because we would not have mean costs from hospital claims data upon which to base payment. Analogous to new drugs and biologicals, we proposed to assign status indicator "K" to HCPCS codes for new therapeutic radiopharmaceuticals for which we have not granted pass-through status.

We did not receive any public comments specific to our proposal for new therapeutic radiopharmaceuticals with HCPCS codes but without pass-through status. However, commenters on the CY 2010 OPPS/ASC proposed rule were supportive of the ASP methodology, in general, for payment for therapeutic radiopharmaceuticals in the HOPD, and we are finalizing an ASP payment methodology for separately payable therapeutic radiopharmaceuticals for CY 2010 as discussed in section V.B.5. of this final rule with comment period.

Therefore, we are finalizing our CY 2010 proposals, without modification, to provide payment for new therapeutic radiopharmaceuticals with HCPCS codes but without pass-through status, if ASP information is not available, based on WAC. If WAC information is also unavailable, we will make payment for new therapeutic radiopharmaceuticals at 95 percent of their most recent AWP. In addition, we are assigning status indicator "K" to HCPCS codes for new therapeutic radiopharmaceuticals in CY 2010 that do not have pass-through status.

Consistent with other ASP-based payments, for CY 2010, we proposed to announce any changes to the payment amounts for new drugs and biologicals in the CY 2010 OPPS/ASC final rule with comment period and also on a quarterly basis on the CMS Web site during CY 2010 if later quarter ASP submissions (or more recent WACs or AWP's) indicate that changes to the payment rates for these drugs and biologicals are necessary. The payment rates for new therapeutic radiopharmaceuticals would also be changed accordingly, based on later quarter ASP submissions. We note that the new CY 2010 HCPCS codes for

drugs, biologicals, and therapeutic radiopharmaceuticals were not available at the time of development of the proposed rule. However, they are included in Addendum B to this CY 2010 OPPS/ASC final rule with comment period. They are assigned comment indicator "NI" in Addendum B to reflect that their interim final OPPS treatment is open to public comment on the CY 2010 OPPS/ASC final rule with comment period.

We did not receive any public comments on our proposal to announce, via the CMS Web site, any changes to the OPPS payment amounts for new drugs and biologicals on a quarterly basis. Therefore, we are finalizing our proposals and will update payment rates for new drugs, biologicals, and therapeutic radiopharmaceuticals, as necessary, in association with our quarterly update process and provide this information on the CMS Web site.

There are several nonpass-through drugs and biologicals that were payable in CY 2008 and/or CY 2009 for which we did not have any CY 2008 hospital claims data available for the proposed rule and for which there were no other HCPCS codes that describe different doses of the same drug but for which we did have pricing information for the ASP methodology. In the CY 2010 OPPS/ASC proposed rule (74 FR 35337), we noted that there are currently no therapeutic radiopharmaceuticals in this category. In order to determine the packaging status of these products for CY 2010, we calculated an estimate of the per day cost of each of these items by multiplying the payment rate for each product based on ASP+4 percent, similar to other nonpass-through drugs and biologicals paid separately under the OPPS, by an estimated average number of units of each product that would typically be furnished to a patient during one administration in the hospital outpatient setting. We proposed to package items for which we estimated the per administration cost to be less than or equal to \$65, which is the general packaging threshold that we proposed for drugs, nonimplantable biologicals, and therapeutic radiopharmaceuticals in CY 2010. We proposed to pay separately for items with an estimated per day cost greater than \$65 (with the exception of diagnostic radiopharmaceuticals, contrast agents and implantable biologicals, which we proposed to continue to package regardless of cost, as discussed in more detail in the proposed rule (74 FR 35323 through 35324)) in CY 2010. We proposed that the CY 2010 payment for separately payable items without CY 2008 claims

data would be ASP+4 percent, similar to payment for other separately payable nonpass-through drugs and biologicals under the OPPS. In accordance with the ASP methodology used in the physician's office setting, in the absence of ASP data, we proposed to use the WAC for the product to establish the initial payment rate. However, we note that if the WAC is also unavailable, we would make payment at 95 percent of the most recent AWP available.

We did not receive any public comments on our proposal to use

estimated per day costs for these drugs and biologicals or on the resulting packaging status of these drugs and biologicals. Therefore, we are finalizing our CY 2010 proposal, without, modification to use the estimated number of units per day included in Table 43 below to determine estimated per day costs for the corresponding drugs and biologicals for CY 2010. Further, we are finalizing our proposal to package those drugs with an estimated per day cost less than or equal to \$65 and to provide separate payment

for those drugs and biologicals with estimated per day costs over \$65 for CY 2010. For those drugs and biologicals that we determined to be separately payable in CY 2010, payment will be made at ASP+4 percent. If ASP information is not available, payment will be based on WAC or 95 percent of the most recently published AWP if WAC is not available. The final estimated units per day and status indicators for these items are displayed in Table 43 below.

TABLE 43—DRUGS AND BIOLOGICALS WITHOUT CY 2008 CLAIMS DATA

CY 2010 HCPCS code	CY 2010 long descriptor	Estimated average number of units per administration	Final CY 2010 SI	Final CY 2010 APC
90681	Rotavirus vaccine, human, attenuated, 2 dose schedule, live, for oral use.	1	K	1239
90696	Diphtheria, tetanus toxoids, acellular pertussis vaccine and poliovirus vaccine, inactivated (DTaP-IPV), when administered to children 4 through 6 years of age, for intramuscular use.	1	N
J0364	Injection, apomorphine hydrochloride, 1 mg	12	N
J2724	Injection, protein c concentrate, intravenous, human, 10 iu	2240	K	1139
J3355	Injection, urofollitropin, 75 IU	2	K	1741
J9215	Injection, interferon, alfa-n3, (human leukocyte derived), 250,000 iu	5	K	0865

Finally, there were eight drugs and biologicals, shown in Table 31 of the CY 2010 OPPS/ASC proposed rule (74 FR 35337), that were payable in CY 2008, but for which we lacked CY 2008 claims data and any other pricing information for the ASP methodology for the CY 2010 OPPS/ASC proposed rule. In CY 2009, for similar items without CY 2007 claims data and without pricing information for the ASP methodology, we stated that we were unable to determine their per day cost and we packaged these items for the year, assigning these items status indicator "N."

For CY 2010, we proposed to change the status indicator for eight drugs and biologicals shown in Table 31 of the CY 2010 OPPS/ASC proposed rule (74 FR 35337) to status indicator "E" (Not paid by Medicare when submitted on outpatient claims (any outpatient bill type)) as we understood that these drugs and biologicals are not currently sold or have been identified as obsolete. In

addition, we proposed to provide separate payment for these drugs and biologicals if pricing information reflecting recent sales becomes available mid-year in CY 2010 for the ASP methodology. If pricing information becomes available, we would assign the products status indicator "K" and pay for them separately for the remainder of CY 2010.

We did not receive any public comments on our proposal to change the status indicators for drugs and biologicals without claims data or pricing information for the ASP methodology. Therefore, we are finalizing our CY 2010 proposal, without modification, to assign status indicator "E" to these drugs and biologicals. As we have used updated claims data and ASP pricing information for this final rule with comment period, we have newly identified for this final rule with comment period CPT codes 90393 (Vaccinia immune globulin, human, for

intramuscular use); 90477 (Adenovirus vaccine, type 7, live, for oral use); 90644 (Meningococcal conjugate vaccine, serogroups C & Y and Hemophilus influenza b vaccine, tetanus toxoid conjugate (Hib-MenCY-TT), 4-dose schedule, when administered to children 2–15 months of age, for intramuscular use); and 90670 (Pneumococcal conjugate vaccine, 13 valent, for intramuscular use) as lacking CY 2008 claims data and any other pricing information for the ASP methodology. Therefore, in addition to the HCPCS codes we proposed to assign status indicator "E" for CY 2010 on this basis in the proposed rule, we are assigning status indicator "E" to CPT codes 90393, 90477, 90644 and 90670 for CY 2010. All drugs and biologicals without CY 2008 hospital claims data and data based on the ASP methodology that are assigned status indicator "E" on this basis at the time of this final rule with comment period for CY 2010 are displayed in Table 44 below.

TABLE 44—DRUGS AND BIOLOGICALS WITHOUT CY 2008 CLAIMS DATA AND WITHOUT PRICING INFORMATION FOR THE ASP METHODOLOGY

CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 SI
90296	Diphtheria antitoxin, equine, any route	E
90393	Vaccinia immune globulin, human, for intramuscular use	E

TABLE 44—DRUGS AND BIOLOGICALS WITHOUT CY 2008 CLAIMS DATA AND WITHOUT PRICING INFORMATION FOR THE ASP METHODOLOGY—Continued

CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 SI
90477	Adenovirus vaccine, type 7, live, for oral use	E
90581	Anthrax vaccine, for subcutaneous use	E
90644	Meningococcal conjugate vaccine, serogroups C & Y and Hemophilus influenza b vaccine, tetanus toxoid conjugate (Hib-MenCY-TT), 4-dose schedule, when administered to children 2–15 months of age, for intramuscular use.	E
90670	Pneumococcal conjugate vaccine, 13 valent, for intramuscular use	E
90727	Plague vaccine, for intramuscular use	E
J0128	Injection, abarelix, 10 mg	E
J0350	Injection, anistreplase, per 30 units	E
J0395	Injection, arbutamine hcl, 1 mg	E
J1452	Injection, fomivirsen sodium, intraocular, 1.65 mg	E
J2460	Injection, oxytetracycline HCL, up to 50 mg	E

VI. Estimate of OPPS Transitional Pass-Through Spending for Drugs, Biologicals, Radiopharmaceuticals, and Devices

A. Background

Section 1833(t)(6)(E) of the Act limits the total projected amount of transitional pass-through payments for drugs, biologicals, radiopharmaceuticals, and categories of devices for a given year to an “applicable percentage” of total program payments estimated to be made under section 1833(t) of the Act for all covered services furnished for that year under the hospital OPPS. For a year before CY 2004, the applicable percentage was 2.5 percent; for CY 2004 and subsequent years, we specify the applicable percentage up to 2.0 percent.

If we estimate before the beginning of the calendar year that the total amount of pass-through payments in that year would exceed the applicable percentage, section 1833(t)(6)(E)(iii) of the Act requires a uniform reduction in the amount of each of the transitional pass-through payments made in that year to ensure that the limit is not exceeded. We make an estimate of pass-through spending to determine not only whether payments exceed the applicable percentage, but also to determine the appropriate reduction to the conversion factor for the projected level of pass-through spending in the following year in order to ensure that total estimated pass-through spending for the prospective payment year is budget neutral as required by section 1883(t)(6)(E) of the Act.

For devices, developing an estimate of pass-through spending in CY 2010 entails estimating spending for two groups of items. The first group of items consists of device categories that were recently made eligible for pass-through payment and that would continue to be eligible for pass-through payment in CY

2010. The CY 2008 OPPS/ASC final rule with comment period (72 FR 66778) describes the methodology we have used in previous years to develop the pass-through spending estimate for known device categories continuing into the applicable update year. The second group contains items that we know are newly eligible, or project would be newly eligible, for device pass-through payment in the remaining quarters of CY 2009 or beginning in CY 2010. As discussed in section V.A.4. of this final rule with comment period, we proposed that, beginning in CY 2010, the pass-through evaluation process and pass-through payment for implantable biologicals newly approved for pass-through payment beginning on or after January 1, 2010, that are always surgically inserted or implanted (through a surgical incision or a natural orifice) would be the device pass-through process and payment methodology only. Therefore, we proposed that the estimate of pass-through spending for implantable biologicals newly eligible for pass-through payment beginning in CY 2010 would be included in the pass-through spending estimate for this second group of device categories. The sum of the CY 2010 pass-through estimates for these two groups of device categories equals the total CY 2010 pass-through spending estimate for device categories with pass-through status.

For devices eligible for pass-through payment, section 1833(t)(6)(D)(ii) of the Act establishes the pass-through amount as the amount by which the hospital’s charges for the device, adjusted to cost, exceeds the portion of the otherwise applicable Medicare OPD fee schedule that the Secretary determines is associated with the device. As discussed in section IV.A.2. of this final rule with comment period, we deduct from the pass-through payment for an identified

device category eligible for pass-through payment an amount that reflects the portion of the APC payment amount that we determine is associated with the cost of the device, defined as the device APC offset amount, when we believe that predecessor device costs for the device category newly approved for pass-through payment are already packaged into the existing APC structure. For each device category that becomes newly eligible for device pass-through payment, including an implantable biological under our CY 2010 proposal, we estimate pass-through spending to be the difference between payment for the device category and the device APC offset amount, if applicable, for the procedures that would use the device. If we determine that predecessor device costs for the new device category are not already included in the existing APC structure, the pass-through spending estimate for the device category would be the full payment at charges adjusted to cost.

For drugs and biologicals eligible for pass-through payment, section 1833(t)(6)(D)(i) of the Act establishes the pass-through payment amount as the amount by which the amount authorized under section 1842(o) of the Act (or, if the drug or biological is covered under a competitive acquisition contract under section 1847B of the Act, an amount determined by the Secretary equal to the average price for the drug or biological for all competitive acquisition areas and year established under such section as calculated and adjusted by the Secretary) exceeds the portion of the otherwise applicable fee schedule amount that the Secretary determines is associated with the drug or biological. As we proposed, we are paying for most nonpass-through separately payable drugs and nonimplantable biologicals under the

CY 2010 OPPS at ASP+4 percent, which represents the otherwise applicable fee schedule amount associated with most pass-through drugs and biologicals, and because we are paying for CY 2010 pass-through drugs and nonimplantable biologicals at ASP+6 percent or the Part B drug CAP rate, if applicable, our estimate of drug and nonimplantable biological pass-through payment for CY 2010 is not zero. Furthermore, payment for certain drugs, specifically diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals without pass-through status, is always packaged into payment for the associated procedures because these products would never be separately paid. However, all pass-through diagnostic radiopharmaceuticals, contrast agents, and those implantable biologicals with pass-through status approved prior to CY 2010 are paid at ASP+6 percent or the Part B drug CAP rate, if applicable, like other pass-through drugs and biologicals. Therefore, our estimate of pass-through payment for all diagnostic radiopharmaceuticals and contrast agents and those implantable biologicals with pass-through status approved prior to CY 2010 is also not zero.

In section V.A.6. of this final rule with comment period, we discuss our policy to determine if the cost of certain “policy-packaged” drugs, including diagnostic radiopharmaceuticals and contrast agents, are already packaged into the existing APC structure. If we determine that a “policy-packaged” drug approved for pass-through payment resembles predecessor diagnostic radiopharmaceuticals or contrast agents already included in the costs of the APCs that would be associated with the drug receiving pass-through payment, as we proposed, we are offsetting the amount of pass-through payment for diagnostic radiopharmaceuticals and contrast agents. For these drugs, the APC offset amount is the portion of the APC payment for the specific procedure performed with the pass-through diagnostic radiopharmaceutical or contrast agent that is attributable to diagnostic radiopharmaceuticals or contrast agents, which we refer to as the “policy-packaged” drug APC offset amount. If we determine that an offset is appropriate for a specific diagnostic radiopharmaceutical or contrast agent receiving pass-through payment, we reduce our estimate of pass-through payment for these drugs by this amount. We have not established a policy to offset pass-through payment for implantable biologicals when approved for pass-through payment as a drug or

biological, that is, for CY 2009 and earlier, so we consider full payment at ASP+6 percent for these implantable biologicals receiving biological pass-through payment in our estimate of CY 2010 pass-through spending for drugs and biologicals.

We note that the Part B drug CAP program has been suspended beginning January 1, 2009. We refer readers to the Medicare Learning Network (MLN) Matters Special Edition article SE0833 for more information on this suspension. As of the publication of the CY 2010 OPPS/ASC proposed rule and this final rule with comment period, the Part B drug CAP program has not been reinstated. Therefore, for this final rule with comment period, we are continuing to not have an effective Part B drug CAP rate for pass-through drugs and biologicals. Similar to estimates for devices, the first group of drugs and biologicals requiring a pass-through payment estimate consists of those products that were recently made eligible for pass-through payment and that continue to be eligible for pass-through payment in CY 2010. The second group contains drugs and nonimplantable biologicals that we know are newly eligible, or project would be newly eligible, in the remaining quarters of CY 2009 or beginning in CY 2010. The sum of the CY 2010 pass-through estimates for these two groups of drugs and biologicals equals the total CY 2010 pass-through spending estimate for drugs and biologicals with pass-through status.

B. Estimate of Pass-Through Spending

For CY 2010, we proposed to set the applicable pass-through payment percentage limit at 2.0 percent of the total projected OPPS payments for CY 2010, consistent with our OPPS policy from CY 2004 through 2009 (74 FR 35339).

For the first group of devices for pass-through payment estimate purposes, there were no device categories receiving pass-through payment in CY 2009 that would continue for payment during CY 2010 (74 FR 35339) and, therefore, we proposed a device pass-through payment estimate for the first group of pass-through device categories of \$0.

We also proposed for CY 2010 to use the device pass-through process and payment methodology for implantable biologicals that are always surgically inserted or implanted (through a surgical incision or a natural orifice). We proposed to consider existing implantable biologicals approved for pass-through payment under the drugs

and biologicals pass-through provision prior to CY 2010 as drugs and biologicals for pass-through payment estimate purposes. We proposed to continue to consider these implantable biologicals that have been approved for pass-through status prior to CY 2010 drugs and biologicals until they expire from pass-through status. Therefore, the proposed pass-through spending estimate for the first group of pass-through devices did not include implantable biologicals that were granted pass-through status prior to CY 2010. Finally, we proposed to provide payment for implantable biologicals newly eligible for pass-through payment beginning in CY 2010 based on hospital charges adjusted to cost, rather than the ASP methodology that is applicable to pass-through drugs and biologicals. Therefore, we proposed that, beginning in CY 2010, the estimate of pass-through spending for implantable biologicals first paid as pass-through devices in CY 2010 would be based on the payment methodology for pass-through devices and would be included in the device pass-through spending estimate.

In estimating our proposed CY 2010 pass-through spending for device categories in the second group, that is, device categories that we knew at the time of the development of the proposed rule would be newly eligible for pass-through payment in CY 2010 (of which there were none), additional device categories (including categories that describe implantable biologicals) that we estimated could be approved for pass-through status subsequent to the development of the proposed rule and before January 1, 2010, and contingent projections for new categories (including categories that describe implantable biologicals in the second through fourth quarters of CY 2010), we proposed to use the general methodology described in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66778), while also taking into account recent OPPS experience in approving new pass-through device categories. There were no new device categories (including categories that describe implantable biologicals) for CY 2010 of which we were aware at the time of development of the proposed rule. The estimate of CY 2010 pass-through spending for this second group of device categories was \$10.0 million for the proposed rule (74 FR 35339).

Employing our established methodology that the estimate of pass-through device spending in CY 2010 incorporates CY 2010 estimates of pass-through spending for known device categories continuing in CY 2010, those known or projected to be first effective

January 1, 2010, and those device categories projected to be approved during subsequent quarters of CY 2009 or CY 2010, our proposed CY 2010 estimate of total pass-through spending for device categories was \$10.0 million (74 FR 35339).

To estimate CY 2010 proposed pass-through spending for drugs and biologicals in the first group, specifically those drugs (including radiopharmaceuticals and contrast agents) and biologicals (including implantable biologicals) recently made eligible for pass-through payment and continuing on pass-through status for CY 2010, we proposed to utilize the most recent Medicare physician's office data regarding their utilization, information provided in the respective pass-through applications, historical hospital claims data, pharmaceutical industry information, and clinical information regarding those drugs or biologicals, in order to project the CY 2010 OPPS utilization of the products.

For the known drugs and biologicals (excluding diagnostic radiopharmaceuticals, contrast agents, and implantable biologicals) that would be continuing on pass-through status in CY 2010, we estimated the proposed pass-through payment amount as the difference between ASP+6 percent or the Part B drug CAP rate, as applicable, and ASP+4 percent, aggregated across the projected CY 2010 OPPS utilization of these products. Because payment for a diagnostic radiopharmaceutical or contrast agent would be packaged if the product were not paid separately due to its pass-through status, we included in the pass-through estimate the difference between payment for the drug or biological at ASP+6 percent (or WAC+6 percent, or 95 percent of AWP, if ASP information is not available) and the "policy-packaged" drug APC offset amount, if we determined that the diagnostic radiopharmaceutical or contrast agent approved for pass-through payment resembles predecessor diagnostic radiopharmaceuticals or contrast agents already included in the costs of the APCs that would be associated with the drug receiving pass-through payment. Because payment for an implantable biological continuing on pass-through status in CY 2010 would be packaged if the product were not paid separately due to its pass-through status and because we have not established a pass-through payment offset policy for implantable biologicals when approved for pass-through payment as biologicals, that is, for CY 2009 and earlier, we included in the proposed pass-through spending estimate the full payment for these

implantable biologicals at ASP+6 percent (or WAC+6 percent or 95 percent of AWP, if ASP information is not available). We note that our spending estimate for this first group of drugs and biologicals was stated in the CY 2010 OPPS/ASC proposed rule as \$8.9 million (74 FR 35340), while our estimate for the second group of drugs and biologicals was reported as \$19.1 million. We inadvertently mislabeled these two spending estimates in the CY 2010 OPPS/ASC proposed rule. For this first group of drugs and biologicals, the proposed spending estimate should have been reported as \$19.1 million and the second group should have been reported as \$8.9 million.

To estimate CY 2010 pass-through spending for drugs and nonimplantable biologicals in the second group (that is, drugs and nonimplantable biologicals that we knew at the time of development of the proposed rule would be newly eligible for pass-through payment in CY 2010, additional drugs and nonimplantable biologicals that we estimated could be approved for pass-through status subsequent to the development of the proposed rule and before January 1, 2010, and projections for new drugs and nonimplantable biologicals that could be initially eligible for pass-through payment in the second through fourth quarters of CY 2010), we proposed to use utilization estimates from pass-through applicants, pharmaceutical industry data, clinical information, recent trends in the per unit ASPs of hospital outpatient drugs, and projected annual changes in service volume and intensity as our basis for making the CY 2010 proposed pass-through payment estimate. We also considered the most recent OPPS experience in approving new pass-through drugs and nonimplantable biologicals. As noted earlier, we also proposed to include new implantable biologicals that we expect to be approved for pass-through status as devices beginning in CY 2010 in the second group of items considered for device pass-through estimate purposes. Therefore, we did not include implantable biologicals in the second group of items in the proposed drug and biological pass-through spending estimate.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35314 through 35317), we proposed to revise our pass-through payment policy regarding "new" drugs and biologicals that were not receiving hospital outpatient payment as of December 31, 1996, and that also met the other criteria for receiving pass-through payment. Specifically, we proposed to change the start date of the

pass-through payment eligibility period for a "new" drug or biological from the first date on which pass-through payment is made to the date on which payment is first made for a drug or biological as an outpatient hospital service under Part B, using the date of first sale of the drug or biological in the United States after FDA approval as a proxy, to better reflect the statutory provisions for pass-through payment under section 1833(t)(6) of the Act. We expected that a number of the drugs and biologicals currently receiving pass-through payment in CY 2009 would not be eligible for pass-through payment under the proposed revised definition of the pass-through payment eligibility period. Accordingly, for the CY 2010 OPPS/ASC proposed rule, we reduced our estimate of CY 2010 pass-through spending for new drugs and nonimplantable biologicals in the second group that could be initially eligible for pass-through payment beginning in CY 2010 to take into consideration the potential effect of the proposed CY 2010 pass-through payment eligibility period policy on the future number of drugs and biologicals newly approved for pass-through payment, in comparison with our historical OPPS experience over the past several years.

As noted above, we inadvertently mislabeled the spending estimates for the two groups of drugs and biologicals in the CY 2010 OPPS/ASC proposed rule. Therefore, while we reported that the spending estimate for this second group of drugs and biologicals in the CY 2010 OPPS/ASC proposed rule was \$19.1 million (74 FR 35340), the estimate that should have been reported for this second group of drugs and biologicals was \$8.9 million.

We did not receive any public comments on the proposed methodology to estimate pass-through spending for drugs, biologicals, radiopharmaceuticals, and device categories in CY 2010. However, we did receive public comments on our proposal to use the first date of sale in the United States after FDA approval as a proxy for the first date of payment under Medicare Part B as an outpatient hospital service for determining the pass-through payment eligibility period for pass-through drugs and nonimplantable biologicals under the OPPS for CY 2010, which would have reduced the pass-through payment estimate for drugs and nonimplantable biologicals. These public comments, our responses, and our final policy for CY 2010 are discussed in section V.A.5 of this final rule with comment period. As with our current policy, in CY 2010 the

pass-through payment eligibility period and the period of pass-through payment period will run concurrently. Thus, for our final CY 2010 pass-through spending estimate for new drugs and nonimplantable biologicals that could be initially eligible for pass-through payment beginning in CY 2010, we have no need to reduce our estimate of pass-through spending to take into consideration a policy change in the pass-through payment eligibility period. Therefore, we are finalizing our proposed methodology to estimate annual pass-through spending for devices and drugs and nonimplantable biologicals, with the modification as described above.

As stated in section V.A.4. of this final rule with comment period, as we proposed, beginning in CY 2010, implantable biologicals that are always surgically inserted or implanted (through a surgical incision or a natural orifice) will be evaluated under the device pass-through process and paid according to the device payment methodology. We are continuing to consider implantable biologicals approved for pass-through payment under the drug and biological pass-through provision prior to CY 2010 as drugs and biologicals for pass-through payment estimate purposes. These implantable biologicals that have been approved for pass-through status prior to CY 2010 continue to be considered drugs and biologicals until they expire from pass-through status. Therefore, the final pass-through spending estimate for the first group of pass-through device categories does not include implantable biologicals that have been granted pass-through status prior to CY 2010.

In section V.A.4. of this final rule with comment period, as we proposed, we are providing that payment for implantable biologicals newly eligible for pass-through payment beginning in CY 2010 is based on hospital charges adjusted to cost, rather than the ASP methodology that is applicable to pass-through drugs and biologicals. Therefore, we are providing that, beginning in CY 2010, the estimate of pass-through spending for implantable biologicals first paid as pass-through devices in CY 2010 is based on the payment methodology for pass-through devices, and is included in the final CY 2010 device pass-through spending estimate for the second group of pass-through device categories.

The final CY 2010 pass-through spending estimate for the first group of pass-through device categories is \$0. The estimate of CY 2010 pass-through spending for the second group of pass-through device categories is \$10.0

million for this final rule with comment period, as it was for the proposed rule (74 FR 35339). Our final CY 2010 estimate of total pass-through spending for device categories is \$10.0 million.

The estimate for pass-through spending for the first group of drugs and biologicals is \$28.9 million for CY 2010. The estimate for pass-through spending for the second group of drugs and biologicals is \$6.7 million for CY 2010. As stated above, the final estimates differ, in part, from our proposed rule estimates in order to reflect our final policy regarding the pass-through payment eligibility period. As discussed in section V.A. of this final rule with comment period, radiopharmaceuticals are considered drugs for pass-through purposes. Therefore, we have included radiopharmaceuticals in our CY 2010 pass-through spending estimate for drugs and biologicals. Our final CY 2010 estimate of total pass-through spending for drugs and biologicals is \$35.6 million.

In summary, in accordance with the methodology described above in this section, we estimate that total pass-through spending for the device categories and the drugs and biologicals that are continuing to receive pass-through payment in CY 2010 and those device categories, drugs, and nonimplantable biologicals that first become eligible for pass-through payment during CY 2010 is approximately \$45.5 million, which represents 0.14 percent of total OPSS projected payments for CY 2010. We estimate that pass-through spending in CY 2010 will not amount to 2.0 percent of total projected OPSS CY 2010 program spending.

VII. OPSS Payment for Brachytherapy Sources

A. Background

Section 1833(t)(2)(H) of the Act, as added by section 621(b)(2)(C) of Public Law 108-173 (MMA), mandated the creation of additional groups of covered OPD services that classify devices of brachytherapy consisting of a seed or seeds (or radioactive source) ("brachytherapy sources") separately from other services or groups of services. The additional groups must reflect the number, isotope, and radioactive intensity of the brachytherapy sources furnished and include separate groups for palladium-103 and iodine-125 sources.

Section 1833(t)(16)(C) of the Act, as added by section 621(b)(1) of Public Law 108-173, established payment for brachytherapy sources furnished from January 1, 2004, through December 31,

2006, based on a hospital's charges for each brachytherapy source furnished adjusted to cost. Under section 1833(t)(16)(C) of the Act, charges for the brachytherapy sources may not be used in determining any outlier payments under the OPSS for that period of payment. Consistent with our practice under the OPSS to exclude items paid at cost from budget neutrality consideration, these items were excluded from budget neutrality for that time period as well.

In our CY 2007 annual OPSS rulemaking, we proposed and finalized a policy of prospective payment based on median costs for the 11 brachytherapy sources for which we had claims data. We based the prospective payment rates on median costs for each source from our CY 2005 claims data (71 FR 68102 through 71 FR 68115).

Subsequent to publication of the CY 2007 OPSS/ASC final rule with comment period, section 107 of Public Law 109-432 (MIEA-TRHCA) amended section 1833(t)(16)(C) of the Act by extending the payment period for brachytherapy sources based on a hospital's charges adjusted to cost for 1 additional year, through December 31, 2007. Therefore, we continued to pay for brachytherapy sources based on charges adjusted to cost for CY 2007.

Section 107(b)(1) of Public Law 109-432 amended section 1833(t)(2)(H) of the Act by adding a requirement for the establishment of separate payment groups for "stranded and non-stranded" brachytherapy sources furnished on or after July 1, 2007, in addition to the existing requirements for separate payment groups based on the number, isotope, and radioactive intensity of brachytherapy sources under section 1833(t)(2)(H) of the Act. Section 107(b)(2) of Pub. L. 109-432 authorized the Secretary to implement this requirement by "program instruction or otherwise." We note that public commenters who responded to the CY 2007 OPSS/ASC proposed rule asserted that stranded sources, which they described as embedded into the stranded suture material and separated within the strand by material of an absorbable nature at specified intervals, had greater production costs than non-stranded sources (71 FR 68113 through 68114).

As a result of the statutory requirement to create separate groups for stranded and non-stranded sources as of July 1, 2007, we established several coding changes through a transmittal, effective July 1, 2007 (Transmittal 1259, dated June 1, 2007). Based on public

comments received on the CY 2007 OPPS/ASC proposed rule and industry input, we were aware of three sources available in stranded and non-stranded forms at that time: iodine-125; palladium-103; and cesium-131 (72 FR 42746). We created six new HCPCS codes to differentiate the stranded and non-stranded versions of iodine, palladium, and cesium sources.

In Transmittal 1259, we indicated that if we receive information that any of the other sources now designated as non-stranded are also FDA-approved and marketed as a stranded source, we would create a code for the stranded source. We also established two "Not Otherwise Specified" (NOS) codes for billing stranded and non-stranded sources that are not yet known to us and for which we do not have source-specific codes. We established HCPCS code C2698 (Brachytherapy source, stranded, not otherwise specified, per source) for stranded NOS sources and HCPCS code C2699 (Brachytherapy source, non-stranded, not otherwise specified, per source) for non-stranded NOS sources.

In the CY 2008 OPPS/ASC final rule with comment period (72 FR 66784), we again finalized prospective payment for brachytherapy sources, beginning in CY 2008, with payment rates determined using the CY 2006 claims-based costs per source for each brachytherapy source. Consistent with our policy regarding APC payments made on a prospective basis, we finalized the policy in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66686) to subject the cost of brachytherapy sources to the outlier provision of section 1833(t)(5) of the Act, and to also subject brachytherapy source payment weights to scaling for purposes of budget neutrality. Therefore, brachytherapy sources could receive outlier payments if the costs of furnishing brachytherapy sources met the criteria for outlier payment. In addition, as noted in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66683), implementation of prospective payment for brachytherapy sources would provide opportunities for hospitals to receive additional payments under certain circumstances through the 7.1 percent rural SCH adjustment.

For CY 2008, we also proposed and finalized a policy regarding payment for new brachytherapy sources for which we have no claims data (72 FR 42749 and 72 FR 66786, respectively). We indicated we would assign future new HCPCS codes for new brachytherapy sources to their own APCs, with prospective payment rates set based on our consideration of external data and

other relevant information regarding the expected costs of the sources to hospitals. Finally, we proposed and finalized our policy to discontinue using status indicator "H" (Pass-Through Device Categories. Separate cost based pass-through payment; not subject to co-payment) because we would not be paying charges adjusted to costs after December 31, 2007, and instead adopted a policy of using status indicator "K" (which includes, among others, "Brachytherapy Sources. Paid under OPPS; separate APC payment") for CY 2008 (72 FR 42749 and 72 FR 66785, respectively).

After we finalized these proposals for CY 2008, section 106(a) of Pub. L. 110-173 (MMSEA) extended the charges-adjusted-to-cost payment methodology for brachytherapy sources for an additional 6 months, through June 30, 2008. Because our final CY 2008 policies paid for brachytherapy sources at prospective rates based on median costs, we were unable to implement these policies during this extension.

In the CY 2009 OPPS/ASC proposed rule (73 FR 41502), we again proposed prospective payment rates for brachytherapy sources for CY 2009. We proposed to pay for brachytherapy sources at prospective rates based on their source-specific median costs as calculated from CY 2007 claims data available for CY 2009 ratesetting. Subsequent to issuance of the CY 2009 OPPS/ASC proposed rule, Public Law 110-275 (MIPPA) was enacted on July 15, 2008. Section 142 of Public Law 110-275 amended section 1833(t)(16)(C) of the Act, as amended by section 106(a) of Public Law 110-173 (MMSEA), to further extend the payment period for brachytherapy sources based on a hospital's charges adjusted to cost from July 1, 2008, through December 31, 2009. Therefore, we continued to pay for brachytherapy sources at charges adjusted to cost in CY 2008 from July 1 through December 31, and we maintained the assignment of status indicator "H" to brachytherapy sources for claims processing purposes in CY 2008. For CY 2009, we have continued to pay for all separately payable brachytherapy sources based on a hospital's charges adjusted to cost. Because brachytherapy sources are paid at charges adjusted to cost, we did not subject them to outlier payments under section 1833(t)(5) of the Act, or subject brachytherapy source payment weights to scaling for purposes of budget neutrality. Moreover, during the CY 2009 period of payment at charges adjusted to cost, brachytherapy sources are not eligible for the 7.1 percent rural SCH adjustment (as discussed in detail

in section II.E. of this final rule with comment period).

Furthermore, for CY 2009, we did not adopt the policy we established in the CY 2008 OPPS/ASC final rule with comment period of paying stranded and non-stranded NOS codes for brachytherapy sources, HCPCS codes C2698 and C2699, based on a rate equal to the lowest stranded or non-stranded prospective payment for such sources. Also, for CY 2009, we did not adopt the policy we established in the CY 2008 OPPS/ASC final rule with comment period regarding payment for new brachytherapy sources for which we have no claims data. NOS HCPCS codes C2698 and C2699 and newly established specific source codes are paid at charges adjusted to cost through December 31, 2009, consistent with section 142 of Public Law 110-275.

For CY 2009, we finalized our proposal to create new status indicator "U" (Brachytherapy Sources. Paid under OPPS; separate APC payment) for brachytherapy source payment, instead of using status indicator "K" as proposed and finalized for CY 2008 for prospective payment, or status indicator "H," used during the period of charges adjusted to cost payment. As noted in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68670), assigning a status indicator, such as status indicator "K," to several types of items and services with potentially differing payment policies added unnecessary complexity to our operations. Status indicator "U" is used only for brachytherapy sources, regardless of their specific payment methodology for any period of time.

At the February 2009 meeting, the APC Panel recommended paying for brachytherapy sources in CY 2010 using a prospective payment methodology based on median costs from claims data. The APC Panel reviewed CY 2007 and CY 2008 brachytherapy source median costs from claims data and noted the stability of the data from year to year.

B. OPPS Payment Policy

Under section 142 of Public Law 110-275, payment for brachytherapy sources is mandated at charges adjusted to cost only through CY 2009. In the CY 2010 OPPS/ASC proposed rule (74 FR 35342), we proposed to adopt for CY 2010 the general OPPS prospective payment methodology for brachytherapy sources, consistent with section 1833(t)(2)(C) of the Act.

As we have previously stated (72 FR 66780 and 73 FR 41502), we believe that adopting the general OPPS prospective payment methodology for brachytherapy sources is appropriate for

a number of reasons. The general OPPS payment methodology uses median costs based on claims data to set the relative payment weights for hospital outpatient services. This payment methodology results in more consistent, predictable, and equitable payment amounts per source across hospitals by eliminating some of the extremely high and low payment amounts resulting from payment based on hospitals' charges adjusted to cost. We believe the OPPS prospective payment methodology would also provide hospitals with incentives for efficiency in the provision of brachytherapy services to Medicare beneficiaries. Moreover, this approach is consistent with our payment methodology for the vast majority of items and services paid under the OPPS.

We proposed to use CY 2008 claims data for setting the CY 2010 payment rates for brachytherapy sources, as we proposed for most other items and services that will be paid under the CY 2010 OPPS. For CY 2008, we have a full year of claims data for each of the separately payable sources, including iodine, palladium, and cesium sources that have stranded and non-stranded configurations. As indicated earlier, the APC Panel, at the February 2009 meeting, recommended using the median cost data for CY 2010 rates. Our proposal was consistent with the APC Panel's recommendation.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35342), we proposed to adopt the other payment policies for brachytherapy sources we finalized in previous final rules. We proposed to pay for the stranded and non-stranded NOS codes, HCPCS codes C2698 and C2699, at a rate equal to the lowest stranded or non-stranded prospective payment rate for such sources, respectively, on a per source basis (as opposed, for example, to a per mCi), which is based on the policy we established in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66785). The proposed payment methodology for NOS sources would provide payment to a hospital for new sources, while encouraging interested parties to quickly bring new sources to our attention so that specific coding and payment could be established.

We also proposed to implement the policy we established in the CY 2008 OPPS/ASC final rule with comment period (which was superseded by section 142 of Public Law 110-275) regarding payment for new brachytherapy sources for which we have no claims data, based on the same reasons we discussed in that final rule with comment period (72 FR 66786).

That policy is intended to enable us to assign future new HCPCS codes for new brachytherapy sources to their own APCs, with prospective payment rates set based on our consideration of external data and other relevant information regarding the expected costs of the sources to hospitals.

Consistent with our policy regarding APC payments made on a prospective basis, we proposed to subject brachytherapy sources to outlier payments under section 1833(t)(5) of the Act, and also to subject brachytherapy source payment weights to scaling for purposes of budget neutrality. Therefore, brachytherapy sources could receive outlier payments if the costs of furnishing brachytherapy sources meet the criteria for outlier payment. In addition, as noted in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66683), implementation of prospective payments for brachytherapy sources would provide opportunities for hospitals to receive additional payments in CY 2010 under certain circumstances through the 7.1 percent rural adjustment as described in section II.E. of the proposed rule (74 FR 35295) and this final rule with comment period.

Therefore, in the CY 2010 OPPS/ASC proposed rule, we proposed to pay for brachytherapy sources at prospective payment rates based on their source-specific median costs for CY 2010. The separately payable brachytherapy source HCPCS codes, long descriptors, APCs, status indicators, and approximate median costs that we proposed for CY 2010 were presented in Table 32 of the proposed rule (74 FR 35342).

Comment: A number of commenters recommended that CMS continue to pay for brachytherapy sources separately based on hospitals' charges adjusted to cost due to the commenters' ongoing concerns regarding Medicare hospital claims data for brachytherapy sources; the commenters provided various examples of issues of concern. Some commenters were concerned that characteristics of high dose rate (HDR) iridium-192, which is a renewable source whose life decays over a 90-day period and is used to treat multiple patients, makes establishment of fair and adequate payment difficult on a fixed prospective basis. The commenters also claimed that the CMS brachytherapy source data continue to show huge variations in per unit costs on claims across hospitals. Several commenters stated that one half of the current brachytherapy sources have proposed payment rates based on 50 or fewer hospitals reporting claims for these sources. Some commenters also indicated that "rank order anomalies"

exist in proposed payment rates for brachytherapy sources, citing that HCPCS code C2635 (Brachytherapy source, non-stranded, High Activity, Palladium-103, greater than 2.2 mCi (NIST), per source) always costs more than low activity sources (HCPCS code C2640, Brachytherapy source, stranded, Palladium-103, per source, and HCPCS code C2641, Brachytherapy source, non-stranded, Palladium-103, per source), yet hospital claims data do not reflect this difference. A number of commenters believed that the current charges-adjusted-to-cost methodology is more accurate and has been tested over time. A few commenters argued that the charges-adjusted-to-cost methodology provides overall cost savings to the Medicare program compared to the prospective payment methodology proposed for CY 2010, according to an analysis performed by the brachytherapy source industry. The commenters thus concluded that implementing prospective brachytherapy source payment would increase aggregate Medicare expenditures for brachytherapy sources compared with the charges-adjusted-to-cost payment methodology.

Several commenters supported the CY 2010 proposal to pay for brachytherapy sources prospectively based on median costs from claims data. One commenter asserted that hospital-specific payments based on the charges-adjusted-to-cost payment methodology violate the intent of a prospective payment system, namely to provide incentives to improve efficiency and control costs. The commenter believed that hospital-specific payments could be manipulated because hospitals know the CCR used to determine payments for brachytherapy sources.

Response: As we stated in the CY 2008 final rule with comment period (72 FR 66782), we believe that median costs based on hospital claims data for brachytherapy sources have produced reasonably consistent per-source cost estimates over the past several years, comparable to the patterns we have observed for many other OPPS services whose payments are set based upon relative payment weights from claims data. We believe that our per-source payment methodology specific to each source's radioisotope, radioactive intensity, and stranded or non-stranded configuration, supplemented by payment based on the number of sources used in a specific clinical case, adequately accounts for the major expected sources of variability across treatments.

As we also explained in the CY 2008 OPPS/ASC final rule with comment

period (72 FR 66782), a prospective payment system such as the OPSS relies on the concept of averaging, where the payment may be more or less than the estimated cost of providing a service for a particular patient, but with the exception of outlier cases, it is adequate to ensure access to appropriate care. In the case of brachytherapy sources for which the law requires separate payment groups, without packaging, the costs of these individual items could be expected to show greater variation than some other APCs under the OPSS because higher variability in costs for some component items and services is not balanced with lower variability for others and because relative weights are typically estimated using a smaller set of claims.

Nevertheless, we believe that prospective payment for brachytherapy sources based on median costs from claims calculated according to the standard OPSS methodology is appropriate at this time and would provide hospitals with the greatest incentives for efficiency in furnishing brachytherapy treatment. Under the budget-neutral OPSS, it is the relativity of costs of services, not their absolute costs, that is important, and we believe that brachytherapy sources can now be appropriately paid according to the standard OPSS payment approach. Moreover, OPSS payments for all services are similarly subjected to the same 2-year lag in costs from claims data available for ratesetting. Therefore, we believe the relative costs of OPSS services should generally be appropriate. It is important that the same measure of central tendency (median cost) from claims be used to establish the payment weights for all OPSS services in order to provide appropriate payment for all of these services. The inflation rate of medical services is taken into consideration through the conversion factor, which is updated annually to account for inflation and used to calculate payment rates from the relative payment weights based on median costs.

It is not uncommon for OPSS prospective payment rates to be based on claims from a relatively small number of hospitals that furnished the service in the year of claims data available for the OPSS update year. We are not concerned that some sources may have median costs and proposed payment rates based on 50 or fewer providers, as are some commenters. Fifty hospitals may report hundreds of brachytherapy source claims for many cases and comprise the universe of providers using particular low volume sources, for which we are required to

pay separately by statute. Further, our methodology for estimating median costs for brachytherapy sources utilizes all line-item charges for those sources, which allows us to use all hospital reported charge and estimated cost information to set payment rates for these items. This is in contrast to our limitation of relying on "natural" single and "pseudo" single procedure claims to set APC payment rates for services with packaged costs. We have no reason to believe that prospective payment rates based on claims from those providers furnishing a particular source do not appropriately reflect the cost of that source to hospitals.

As for most other OPSS services, we note that the median costs for brachytherapy sources are based upon the costs of those providers that furnished the sources in CY 2008. Hospitals individually determine their charge for an item or service, and one of Medicare's primary requirements for setting a charge is that it be reasonably and consistently related to the cost of the item or service for that facility (Medicare Provider Reimbursement Manual-I, Section 2203). We then estimate a cost from that charge using the hospital's most recent Medicare hospital cost report data in our standard OPSS ratesetting process. In as much as we paid hospitals at charges adjusted to cost for brachytherapy sources in CY 2008 based on these exact charges, we believe hospital's individual charges to be accurate for their institution.

In the case of high and low activity iodine-125 sources, our claims data showed that the cost of the high activity source is greater than the low activity sources, yet this relationship is reversed for palladium-103 sources, as the commenter pointed out. We have no information about the expected cost differential between high and low activity sources of various isotopes other than what is available in our claims and hospital cost report data. For high activity palladium-103, only 16 hospitals provided this source in CY 2008, compared to 166 and 268 providers for low activity palladium sources described by HCPCS codes C2640 and C2641, respectively. Clearly, fewer providers furnished high activity palladium-103 sources, and we expect that the hospital cost distribution for those hospitals could be different than the cost distribution of the large number of providers reporting the low activity sources. These varied cost distributions clearly contribute to the observed relationship in median costs between the different types of sources, yet we see no reason why our standard ratesetting methodology for brachytherapy sources

that relies on all claims from all hospitals furnishing brachytherapy sources would not yield valid median costs for those hospitals furnishing the different brachytherapy sources upon which CY 2010 prospective payments rates are based.

When the statutory requirement for payment of brachytherapy sources at a hospital's charges adjusted to cost ends on December 31, 2009 (section 1833(t)(16)(C) of the Act), prospective payment for brachytherapy sources based on their median costs would make the source payment an integral part of the OPSS, rather than a separate cost-based payment methodology within the OPSS. We believe that consistent and predictable prospectively established payment rates under the OPSS for brachytherapy sources are appropriate because we do not believe that the hospital resource costs associated with specific brachytherapy sources would vary greatly across hospitals or clinical conditions under treatment, other than through differences in the numbers of sources utilized that would be accounted for in the standard OPSS payment methodology we are finalizing.

We agree that sources such as HDR iridium-192 have a fixed active life and must be replaced every 90 days; as a result, hospitals' per-treatment cost for the source would be dependent on the number of treatments furnished per source. The source cost must be amortized over the life of the source. Therefore, in establishing their charges for HDR iridium, we expect hospitals to project the number of treatments that would be provided over the life of the source and establish their charges for the source accordingly, as we have stated previously (72 FR 66783). For most such OPSS services, our practice is to establish prospective payment rates based on the median costs from hospitals claims data, to provide incentives for efficient and cost-effective delivery of these services.

We do not agree with the commenters that prospective brachytherapy source payment based on median costs would increase aggregate Medicare expenditures compared to the charges-adjusted-to-cost methodology, or that the charges-adjusted-to-cost methodology would provide overall cost savings to the Medicare program compared to the prospective payment methodology. We also do not believe that the beneficiary copayment in the aggregate would increase under the prospective payment methodology. We have traditionally estimated charge inflation for brachytherapy sources as higher than the market basket inflation

update applicable to prospective payment under the OPSS. We estimated charge inflation for brachytherapy sources between the 2 most recent years of hospital claims data by comparing the per-unit charge in CY 2008 claims to the per-unit charge in CY 2007 claims across all sources, and we used this estimate in our budget neutrality calculations. We are currently estimating a charge inflation factor of 17.1 percent for brachytherapy sources between CY 2007 and CY 2008 and, over the past several years, we have consistently estimated brachytherapy source charge inflation factors higher than 8 percent. Inflating payment at hospitals' charges adjusted to cost in the CY 2008 claims to CY 2010 using this most recent charge inflation factor and comparing it to an estimate of prospective payment for the same sources suggest that aggregate brachytherapy source payment for CY 2010 at charges adjusted to cost would be slightly higher than prospective payment for brachytherapy sources in CY 2010. Although the commenters did not include the details of their analysis in their comments, it is possible that the analysis did not include a charge inflation factor to increase payment estimated at charges adjusted to cost from CY 2008 to CY 2010.

Comment: One commenter indicated that the proposed source-specific payments were consistent with its experience with the cost per unit of the sources, except for the proposed payment for HCPCS code C2634 (Brachytherapy source, non-stranded, High Activity, Iodine-125, greater than 1.01 mCi (NIST), per source). The commenter noted that the proposed payment rate for HCPCS code C2634 is \$60, yet its invoices for high activity I-125, that is, HCPCS code C2634, range between \$174 and \$689. The commenter also stated that high activity I-125 sources are ordered based on a range of activity levels. The commenter suggested that there may have been errors in hospital reporting of HCPCS code C2634 in CY 2008 that resulted in the low proposed payment rate. The commenter requested that CMS reevaluate the proposed payment rate for HCPCS code C2634 for CY 2010 using average cost data from manufacturers.

Response: We are pleased that the proposed CY 2010 payment rates for all but one of the brachytherapy sources are consistent with the commenter's experience. The CY 2008 median cost of HCPCS code C2634 for this final rule with comment period is approximately \$59, compared with approximately \$31 in CY 2006 and approximately \$38 in

CY 2007. The CY 2008 median cost is somewhat higher than the previous 2 years, and we acknowledge that the variability in the activity of sources reported with HCPCS code C2634 could explain some of the variability in cost for this source. Furthermore, we note that the CY 2008 median cost for HCPCS code C2634 is based on 18,602 units, over 267 days, from 48 providers. We believe that some variation in relative cost from year to year is to be expected in a prospective payment system, particularly for low volume items.

For all APCs whose payment rates are based upon relative payment weights, we note that the quality and accuracy of reported units and charges significantly influence the final median costs that are the basis for our payments. Beyond our standard OPSS trimming methodology (described in section II.A.2. of this final rule with comment period) that we apply to those claims that have passed various types of claims processing edits, it is not our policy to judge the accuracy of hospital coding and charging for the purpose of ratesetting. Moreover, we do not believe it is necessary to incorporate external cost data from manufacturers of brachytherapy sources or others because, in a relative weight system like the OPSS, it is the relativity of the costs to one another, rather than absolute cost, that is important in setting payment rates. External data lack relativity to the estimated costs derived from the claims and cost report data and generally are not appropriate for determining relative weights that result in payment rates when costs derives from hospital claims and cost report data for services are available.

Comment: One commenter suggested that brachytherapy sources are not reported consistently by all providers using a specific revenue center and recommended that CMS maintain payment at charges adjusted to cost until cost data are improved by refined information resulting from the new cost center for high cost supplies.

Response: In analyzing the reporting of brachytherapy sources in CY 2008 claims, we found that the great majority of brachytherapy sources are reported under revenue code 0278 (Other Implants). Under the policy finalized in the FY 2009 IPPS final rule (73 FR 48462 through 48463), we finalized a definition of a new "Implantable Devices Charged to Patients" cost center to which costs and charges under revenue code 0278 would map in the future. Thus, brachytherapy sources would generally be subject to the "Implantable Devices Charged to Patients" cost center for future cost estimation under the OPSS, potentially

leading to greater accuracy in cost estimation for these devices as noted by the commenter. This new cost center was available for use for cost reporting periods beginning on or after May 1, 2009, and was discussed in Transmittal 20 (dated July 2009) that updated Chapter 36, Hospital and Hospital Health Care Complex Cost Report (Form CMS 2552-96) of the Medicare Provider Reimbursement Manual, Part 2, to provide Line 55.30 to report "Implantable Devices Charged to Patients." The proposed draft cost report Form CMS-2552-10, published in the **Federal Register** for public comment on July 2, 2009 (74 FR 31738), provides new line 69 to report "Implantable Devices Charged to Patients." The proposed cost report can be viewed at: <http://www.cms.hhs.gov/PaperworkReductionActof1995/PRAL/itemdetail.asp?filterType=none&filterByDID=-99&sortByDID=2&sortOrder=descending&itemID=CMS1224069&intNumPerPage=10>.

We have stated previously that we continue to emphasize our preference for long-term cost reporting changes and broad education initiatives to address the accuracy of claims data (73 FR 68524). This recent change to include a new cost center will ultimately influence both the IPPS and OPSS relative weights in the future. Nevertheless, in the meantime, we believe it is fully appropriate to utilize our current cost estimates for brachytherapy sources and all other implantable devices in calculating payment weights under the OPSS because these are our best current estimates of costs as derived from claims and cost report data. When hospital-specific CCRs from the new cost center are available for ratesetting in several years, we will incorporate those into the revenue code-to-cost center crosswalk that we use for OPSS cost estimation. However, at the present time, we believe our current methodology that generally utilizes the available single medical supply CCR leads to appropriate cost estimates for brachytherapy sources, and we see no reason why payment at charges adjusted to cost, which applies an hospital-specific overall ancillary CCR to hospital charges for brachytherapy sources, would lead to a more accurate cost estimate for these items. The hospital-specific overall ancillary CCR is based on costs and charges for a wide range of OPSS services, and we have no reason to believe that hospital markup practices for brachytherapy sources are similar to the relationship between costs

and charges represented in this very general CCR.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to pay for brachytherapy sources prospectively based on CY 2008 median costs from historical hospital claims data. In addition, we will pay the stranded and non-stranded NOS codes, HCPCS codes C2698 and C2699, at a rate equal to the lowest stranded or non-stranded

prospective payment rate for such sources, respectively, on a per source basis. Payment for new brachytherapy sources, which may be established quarterly, will be made through their own APCs, with prospective payment rates set based on our consideration of external data and other relevant information regarding the expected costs of the sources to hospitals because we would have no information from claims data on the costs of these new sources to hospitals. Finally, in CY

2010, brachytherapy sources will be subject to outlier payments, their payment weights subject to scaling for purposes of budget neutrality, and, under some circumstances, their payment subject to the 7.1 percent rural adjustment as discussed in section II.E. of this final rule with comment period.

Table 45 below displays the separately payable brachytherapy source HCPCS codes, long descriptors, APCs, status indicators, and approximate median costs for CY 2010.

TABLE 45—SEPARATELY PAYABLE BRACHYTHERAPY SOURCES FOR CY 2010

CY 2010 HCPCS Code	CY 2010 long descriptor	Final CY 2010 APC	Final CY 2010 SI	Final CY 2010 approximate APC median cost
A9527	Iodine I-125, sodium iodide solution, therapeutic, per millicurie	2632	U	\$38
C1716	Brachytherapy source, non-stranded, Gold-198, per source	1716	U	42
C1717	Brachytherapy source, non-stranded, High Dose Rate Iridium-192, per source ..	1717	U	229
C1719	Brachytherapy source, non-stranded, Non-High Dose Rate Iridium-192, per source.	1719	U	63
C2616	Brachytherapy source, non-stranded, Yttrium-90, per source	2616	U	15,635
C2634	Brachytherapy source, non-stranded, High Activity, Iodine-125, greater than 1.01 mCi (NIST), per source.	2634	U	59
C2635	Brachytherapy source, non-stranded, High Activity, Palladium-103, greater than 2.2 mCi (NIST), per source.	2635	U	28
C2636	Brachytherapy linear source, non-stranded, Palladium-103, per 1MM	2636	U	19
C2638	Brachytherapy source, stranded, Iodine-125, per source	2638	U	42
C2639	Brachytherapy source, non-stranded, Iodine-125, per source	2639	U	36
C2640	Brachytherapy source, stranded, Palladium-103, per source	2640	U	60
C2641	Brachytherapy source, non-stranded, Palladium-103, per source	2641	U	57
C2642	Brachytherapy source, stranded, Cesium-131, per source	2642	U	109
C2643	Brachytherapy source, non-stranded, Cesium-131, per source	2643	U	65
C2698	Brachytherapy source, stranded, not otherwise specified, per source	2698	U	*42
C2699	Brachytherapy source, non-stranded, not otherwise specified, per source	2699	U	*28

* Median cost is that of the lowest cost stranded or non-stranded source upon which CY 2010 payment for the NOS HCPCS code is based.

We continue to invite hospitals and other parties to submit recommendations to us for new HCPCS codes to describe new brachytherapy sources consisting of a radioactive isotope, including a detailed rationale to support recommended new sources. Such recommendations should be directed to the Division of Outpatient Care, Mail Stop C4-05-17, Centers for Medicare and Medicaid Services, 7500 Security Boulevard, Baltimore, MD 21244. We will continue to add new brachytherapy source codes and descriptors to our systems for payment on a quarterly basis.

VIII. OPPTS Payment for Drug Administration Services

A. Background

In CY 2005, in response to the recommendations made by public commenters and the hospital industry, OPPTS transitioned from Level II HCPCS Q-codes to the use of CPT codes for drug administration services. These CPT codes allowed specific reporting of

services regarding the number of hours for an infusion and provided consistency in coding between Medicare and other payers. (For a discussion regarding coding and payment for drug administration services prior to CY 2005, we refer readers to the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66787).)

While hospitals began adopting CPT codes for outpatient drug administration services in CY 2005, physicians paid under the MPFS were using HCPCS G-codes in CY 2005 to report office-based drug administration services. These HCPCS G-codes were developed in anticipation of substantial revisions to the drug administration CPT codes by the CPT Editorial Panel that were expected for CY 2006.

In CY 2006, as anticipated, the CPT Editorial Panel revised its coding structure for drug administration services and incorporated new concepts, such as initial, sequential, and concurrent services, into a structure that previously distinguished services based

on type of administration (chemotherapy/nonchemotherapy), method of administration (injection/infusion/push), and for infusion services, first hour and additional hours. For CY 2006, we implemented the CY 2006 drug administration CPT codes that did not reflect the concepts of initial, sequential, and concurrent services under the OPPTS, and we created HCPCS C-codes that generally paralleled the CY 2005 CPT codes for reporting these other services.

For CY 2007, as a result of public comments on the proposed rule and feedback from the hospital community and the APC Panel, we implemented the full set of CPT codes for drug administration services, including codes incorporating the concepts of initial, sequential, and concurrent services. In addition, the CY 2007 update process offered us the first opportunity to consider data gathered from the use of CY 2005 CPT codes for purposes of ratesetting. For CY 2007, we used CY 2005 claims data to implement a six-

level APC structure for drug administration services. In CY 2008, we continued to use the full set of CPT codes for drug administration services and continued our assignment of drug administration services to this six-level APC structure.

For CY 2009, we continued to allow hospitals to use the full set of CPT codes for drug administration services but moved from a six-level APC structure to a five-level APC structure. We note that, while there were changes in the CPT numerical coding for nonchemotherapy drug administration services in CY 2009, the existing CPT codes were only renumbered, and there were no significant changes to the code descriptors themselves. As we discussed in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68672), the CY 2009 ratesetting process afforded us the first opportunity to examine hospital claims data for the full set of CPT codes that reflected the concepts of initial, sequential, and concurrent services. For CY 2009, we performed our standard annual OPSS review of the clinical and resource characteristics of the drug administration CPT codes assigned to the six-level CY 2008 APC structure based on the CY 2007 claims data available for the CY 2009 OPSS/ASC proposed rule. As a result of our hospital cost analysis and detailed clinical review, we adopted a five-level APC structure for CY 2009 drug administration services to more appropriately reflect their resource utilization in APCs that also group clinically similar services. As we noted in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68671), these APCs generally demonstrated the clinically expected and actually observed comparative relationships between the median costs of different types of drug administration services, including initial and additional services; chemotherapy and other diagnostic, prophylactic, or therapeutic services; injections and infusions; and simple and complex methods of drug administration. In the CY 2009 OPSS/ASC final rule with comment period (73 FR 68673), we indicated our belief that the five-level APC structure was the most appropriate structure based on updated hospital claims data for the full range of CPT codes for drug administration for the CY 2009 OPSS/ASC final rule with comment period because the structure resulted in payment groups with greater clinical and resource homogeneity.

B. Coding and Payment for Drug Administration Services

In the CY 2010 OPSS/ASC proposed rule (74 FR 35343), we proposed for CY 2010 to continue to use the full set of CPT codes for drug administration services. In addition, as a part of our standard annual review, we analyzed the assignments of CPT codes for drug administration into the five-level APC structure and, based on the results of this review, proposed to continue a five-level APC structure for CY 2010. Further, we proposed some minor reconfigurations of the APCs as described below to account for changes in HCPCS code-specific median costs resulting from updated CY 2008 claims data and the most recent cost report data, and the CY 2010 drug payment proposal that is discussed in section V.B.3.b. of the proposed rule (74 FR 35326 through 35333) and this final rule with comment period.

In the CY 2007 OPSS/ASC final rule with comment period (71 FR 68117), we explained that we expected CPT codes for additional hours of infusion to be reported with CPT codes for the initial hour of drug infusion. This would result in a substantial number of claims for drug administration services that were unusable for ratesetting purposes because multiple services would be present on the same bill and result in essentially no correctly coded claims upon which to set the median costs for the CPT codes describing additional hours of infusion. (We refer readers to section II.A.1.b. of the proposed rule (74 FR 35239 through 35241) and this final rule with comment period for a further discussion of multiple bills and our ratesetting methodology.) In order to use these claims for ratesetting purposes for both the initial drug administration codes and the additional hour drug administration codes, we adopted the policy of adding the additional hour drug administration codes to the bypass list in order to create "pseudo" single claims that would be useable for OPSS ratesetting purposes. After the creation of these "pseudo" single claims, we applied the standard OPSS methodology to calculate HCPCS code-specific median costs for these initial and additional hour drug administration codes.

As we explained further in the CY 2007 OPSS/ASC final rule with comment period, bypassing these additional hour drug administration CPT codes and developing additional "per unit" claims provided a methodology for calculating median costs for these previously packaged drug administration services which attributed

all of their line-item cost data to their assigned APCs. However, we noted that this methodology allocates all packaged costs on claims for drug administration services to the associated initial hour of infusion code. Because these additional hours of infusion codes were always reported with other drug administration services, we expected that the packaging related to additional hours of infusion would be appropriately assigned to the initial drug administration service also included on the same claim. While we stated our belief that there are some packaged costs that are clinically related to the second and subsequent hours of infusion, especially for infusions of packaged drugs spanning several hours, we were not able to accurately assign representative portions of packaged costs to multiple different services due to the limitations of our claims data.

We indicated that, while this methodology did not assign any packaged costs to the additional hours of drug administration codes, we believed this methodology took into account all of the packaging on claims for drug administration services and provided a reasonable framework for developing median costs for drug administration services that were often provided in combination with one another.

Since this approach was first adopted for CY 2007, we have updated and expanded the bypass methodology to reflect changing drug administration HCPCS codes that are recognized under the OPSS. We placed all of the add-on CPT codes for drug administration services, including the sequential infusion and intravenous push codes, on the bypass list in CY 2009 (73 FR 68513) and proposed to include them in CY 2010 (74 FR 35242 through 35252) in order to continue this framework for transforming these otherwise unusable multiple bills into "pseudo" single claims that can be used for OPSS ratesetting purposes. Table 33 of the proposed rule (74 FR 35345 through 35349) displayed the proposed configurations of the five drug administration APCs for CY 2010. In proposing to reassign several HCPCS codes for CY 2010, we took into consideration the resource characteristics of the services, as reflected in their HCPCS code-specific median costs and their clinical characteristics. We believed the proposed APC configurations group drug administration services that share sufficiently similar clinical and resource characteristics, taking into consideration updated CY 2008 claims data and the most recent cost report data and

common clinical scenarios that have been described to us.

Comment: Several commenters supported the proposal to include the drug administration add-on codes on the bypass list. The commenters stated that, by including these codes in the bypass methodology, more single bills can be used for ratesetting purposes.

One commenter recommended that CPT code 96368 (Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug): concurrent infusion (List separately in addition to code for primary procedure)) be included on the bypass list in order to ensure consistency with the treatment of other drug administration codes.

Response: As stated in the CY 2007 OPPS/ASC final rule with comment period (71 FR 68117), we expect CPT codes for additional hours of infusion to be reported with CPT codes for the initial hour of drug infusion. This would result in a substantial number of claims for drug administration services that would be unusable for ratesetting purposes because multiple services would be present on the same bill and result in essentially no correctly coded claims upon which to set the median costs for the CPT codes describing additional hours of infusion. In order to use these claims for ratesetting purposes for both the initial drug administration codes and the additional hour drug administration codes, we adopted the policy of adding the additional hour drug administration codes on the bypass list in order to create “pseudo” single claims that would be useable for OPPS ratesetting purposes. We continue to believe that bypassing these additional hour drug administration CPT codes and developing additional “per unit” claims provide a methodology for calculating median costs for these previously packaged additional hour drug administration services, which attributes all of their line-item cost data to their assigned APCs. Although we understand that this methodology does not assign any packaged costs to the additional hours of drug administration codes, we continue to believe this methodology takes into account all of the packaging on claims for drug administration services and provides a reasonable framework for developing median costs for drug administration services that are often provided in combination with one another.

As discussed above, since this approach was first adopted for CY 2007, we have updated and expanded the bypass methodology to reflect changing drug administration HCPCS codes that are recognized under the OPPS. We placed all of the add-on CPT codes for

drug administration services, including the sequential infusion and intravenous push codes, on the bypass list in CY 2009 (73 FR 68513) and proposed to include them in CY 2010 (74 FR 35242 through 35252) in order to continue this framework for transforming these otherwise unusable multiple bills into “pseudo” single claims that can be used for OPPS ratesetting purposes.

We have not added CPT code 96368 (or its predecessor CPT code 90768) on the bypass list because our CY 2010 policy unconditionally packages payment for this service and, therefore, it is not a candidate for the bypass list. The purpose of the bypass list is to develop “pseudo” single claims so that there are more data available to determine the median costs of separately payable services for ratesetting purposes. Including packaged codes would be contrary to the purpose of the bypass list.

We refer readers to section II.A.1.b. of this final rule with comment period for a full discussion of our final bypass policy and list for CY 2010.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to use the full set of CPT codes for drug administration and include all separately paid drug administration add-on HCPCS codes on the CY 2010 bypass list. We will not add CPT code 96368 on the bypass list because it is not a separately paid service and, therefore, it is not a candidate for the bypass list.

Comment: Several commenters expressed support for the proposed five-level APC structure for drug administration services. Some commenters requested that CMS continue to evaluate the five-level structure annually. In addition, several commenters specifically supported the proposed CY 2010 reconfiguration of the HCPCS code assignments to the drug administration APCs.

One commenter stated that the data used to propose reassignments of drug administration codes to drug administration APCs for the proposed rule were inadequate. The commenter explained that changes to CPT codes require hospitals to train staff and implement guidelines for code use and, therefore, accurate claims hospital data for updated CPT codes are not immediately available from the first year of their use. The commenter added that differences in definitions for drug administration codes by Medicare contractors contribute to incomplete and inconsistent data.

Response: In proposing to reassign several HCPCS codes for CY 2010, we

took into consideration the resource characteristics of the services, as reflected in their HCPCS code-specific median costs and their clinical characteristics. We believe the proposed APC configurations group drug administration services that share sufficiently similar clinical and resource characteristics, taking into consideration updated CY 2008 claims data and the most recent cost report data and common clinical scenarios that have been described to us.

We disagree with the commenter who believed that our costs from hospital claims data are inadequate. We believe that the hospital claims data for drug administration HCPCS codes are robust and representative of the costs of the many hospitals performing these services. Multiple drug administration HCPCS codes are reported on several hundred thousand hospital outpatient claims annually and almost all drug administration HCPCS codes are reported on at least several thousand claims. The data that we have reviewed for CY 2010 do not dramatically differ from previous years' data for these high volume services furnished by thousands of hospitals. This is evidenced in the number of hospitals billing for drug administration services, the frequency of specific drug administration services, and the resulting median costs of the drug administration services.

Finally, we note that it is our standard practice to annually review the configuration of all APCs. Therefore, as part of our standard methodology, we expect to continue to review the configuration of drug administration APCs in future years.

Comment: A few commenters requested that HCPCS code C8957 (Intravenous infusion for therapy/diagnosis; initiation of prolonged infusion (more than eight hours), requiring use of portable or implantable pump) not be reassigned to APC 0439 (Level IV Drug Administration) as proposed for CY 2010. Instead, the commenters requested that HCPCS code C8957 continue to be assigned to APC 0440 (Level V Drug Administration) for CY 2010. In addition, one commenter requested that CPT code 96521 (Refilling and maintenance of portable pump) not be reassigned to APC 0439 as proposed for CY 2010. Instead, the commenter requested that CPT code 96521 continue to be assigned to APC 0440. The commenters stated that HCPCS code C8957 and CPT code 96521 represent prolonged infusions that require the use of a pump and a significant amount of time and nursing resources.

Response: As is our standard process, for the CY 2010 proposed rule, we reviewed each APC for clinical cohesiveness and resource homogeneity. As the commenters noted, we proposed to reassign HCPCS code C8957 to APC 0439 because we believed that the proposed HCPCS-specific median cost more closely matched the proposed median cost of APC 0439. Upon further review, we agree with the commenters that the clinical characteristics of the procedure described by HCPCS code C8957 that describes a prolonged intravenous infusion lasting more than 8 hours more closely resemble those of procedures assigned to APC 0440. Further, the HCPCS-specific median cost of HCPCS code C8957 (approximately \$179) is only slightly less than the median cost of APC 0440 (approximately \$218), resulting in our belief that APC 0440 would be the most appropriate assignment of HCPCS code C8957 for CY 2010.

In addition, we proposed to reassign CPT code 96521 to APC 0439 because we believed that the proposed HCPCS-specific median cost more closely matched the median cost of APC 0439. We continue to believe that the HCPCS-specific median cost of CPT code 96521 (approximately \$133) closely resembles the median cost of APC 0439 (approximately \$126). In addition, we note that while HCPCS code C8957 describes the initiation of a prolonged infusion that we would expect to be resource-intensive, CPT code 96521 describes the refilling and maintenance of a portable infusion pump, a drug administration service that we would expect to require less hospital resources. Therefore, while we believe that there is a compelling reason to assign HCPCS code C8957 to the higher level drug administration APC 0440, we do not find a compelling reason to do the same for CPT code 96521.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal for the five-level APC structure for drug administration services, with a modification to not reassign HCPCS code C8957 to APC 0439 as proposed. Instead, we will continue to assign HCPCS code C8957 to APC 0440 for CY 2010, with a final APC median cost of approximately \$218. We are finalizing our proposed CY 2010 assignment of CPT code 96521 to APC 0439, with a final APC median cost of approximately \$126.

Comment: A few commenters requested that CPT codes 96376 (Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); each additional sequential

intravenous push of the same substance/drug provided in a facility) and 96368 (Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); concurrent infusion) be paid separately in CY 2010. Some commenters stated that CPT code 96376 is similar to CPT code 96374 (Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); intravenous push, single or initial substance/drug) and should be assigned to the same APC as CPT code 96374 for CY 2010. In addition, some commenters indicated that CPT code 96368 is similar to CPT code 96375 (Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); each additional sequential intravenous push of a new substance/drug (List separately in additional to code for primary procedure)) and should be assigned to the same APC as CPT code 96375 for CY 2010. Other commenters noted that because CMS now has claims data upon which to set specific payment rate for these services, the OPSS should pay separately for CPT codes 96376 and 96368.

Response: We agree with the commenters that we have cost data for these CPT codes based on historical hospital claims data. However, we also believe that these codes remain appropriate for packaging and, therefore, we include their costs in payment for the independent services with which they are always associated. As we discussed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66787 through 66788) and in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68674), in deciding whether to package a service or pay for it separately, we consider a variety of factors, including whether the service is normally provided separately or in conjunction with other services; how likely it is for the costs of the packaged code to be appropriately mapped to the separately payable codes with which it was performed; and whether the expected cost of the service is relatively low. CPT codes 96376 and 96368, by definition, are always provided in association with other drug administration services, and we continue to believe that they are most appropriately packaged under the OPSS.

Furthermore, we do not agree with the commenters that the services described by CPT code 96376 are similar to those described by CPT code 96374. CPT code 96374 is an initial intravenous push code, and, per CPT instructions, special billing guidelines apply. Commonly, this service requires the initial establishment of intravenous access in a patient, a resource-intensive task

performed by hospital staff using special supplies. In contrast, CPT code 96376 is an add-on code and is reported for each additional sequential intravenous push of the same substance/drug. In the case of this sequential service, the patient already has established intravenous access, so we would expect the service to require fewer hospital resources. In addition, we do not agree with commenters that the services described by CPT code 96368 are similar to those described by CPT code 96375. CPT code 96368 describes a concurrent intravenous infusion while CPT code 96375 describes a sequential intravenous push, and we would expect these services to require different hospital resources.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to unconditionally package payment for CPT codes 96368 and 96376. These CPT codes are, therefore, assigned status indicator "N" in Addendum B to this final rule with comment period.

Comment: Several commenters submitted questions related to coding for drug administration services. Some commenters requested information on how to code for specific clinical scenarios, while other commenters were concerned about documentation requirements for a stop time for an infusion.

Response: Each of these comments and questions is outside of the scope of the proposals in the CY 2010 OPSS/ASC proposed rule. However, we will consider the possibility of addressing these concerns through other available mechanisms, as appropriate.

In summary, after review of the public comments we received, we are finalizing our proposed coding and payment structure for drug administration as follows. We are finalizing, without modification, our proposal to include all separately payable drug administration add-on codes on the bypass list for CY 2010. In addition, we are finalizing our proposed five-level APC structure for payment of drug administration services in the HOPD for CY 2010, with the exception of a modification to continue to assign HCPCS code C8957 to APC 0440 for CY 2010, rather than APC 0439 as we proposed. Finally, we are finalizing our CY 2010 proposal, without modification, to continue to package payment for CPT codes 96376 and 96368 for CY 2010.

Table 46 below displays the final configurations of the five drug administration APCs for CY 2010.

TABLE 46.—CY 2010 DRUG ADMINISTRATION APCs

CY 2010 HCPCS Code	Final CY 2010 APC	Final CY 2010 Approximate APC Median Cost	CY 2010 Long Descriptor
90471	0436	\$25	Immunization administration (includes percutaneous, intradermal, subcutaneous, or intramuscular \injections); one vaccine (single or combination vaccine/toxoid)
90472			Immunization administration (includes percutaneous, intradermal, subcutaneous, or intramuscular injections); each additional vaccine (single or combination vaccine/toxoid) (List separately in addition to code for primary procedure)
90473			Immunization administration by intranasal or oral route; one vaccine (single or combination vaccine/toxoid)
90474			Immunization administration by intranasal or oral route; each additional vaccine (single or combination vaccine/toxoid) (List separately in addition to code for primary procedure)
95115			Professional services for allergen immunotherapy not including provision of allergenic extracts; single injection
95117			Professional services for allergen immunotherapy not including provision of allergenic extracts; 2 or more injections
95165			Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy; single or multiple antigens (specify number of doses)
96361			Intravenous infusion, hydration; each additional hour (List separately in addition to code for primary procedure)
96366			Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); each additional hour (List separately in addition to code for primary procedure)

CY 2010 HCPCS Code	Final CY 2010 APC	Final CY 2010 Approximate APC Median Cost	CY 2010 Long Descriptor
96371			Subcutaneous infusion for therapy or prophylaxis (specify substance or drug); additional pump set-up with establishment of new subcutaneous infusion site(s) (List separately in addition to code for primary procedure)
96372			Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); subcutaneous or intramuscular
96379			Unlisted therapeutic, prophylactic, or diagnostic intravenous or intra-arterial injection or infusion
96549			Unlisted chemotherapy procedure
95144	0437	\$37	Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy, single dose vial(s) (specify number of vials)
95145	0437	\$37	Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy (specify number of doses); single stinging insect venom
95148	0437	\$37	Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy (specify number of doses); 4 single stinging insect venoms
95149	0437	\$37	Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy (specify number of doses); 5 single stinging insect venoms
95170	0437	\$37	Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy; whole body extract of biting insect or other arthropod (specify number of doses)
96367			Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); additional sequential infusion, up to 1 hour (List separately in addition to code for primary procedure)

CY 2010 HCPCS Code	Final CY 2010 APC	Final CY 2010 Approximate APC Median Cost	CY 2010 Long Descriptor
96370			Subcutaneous infusion for therapy or prophylaxis (specify substance or drug); each additional hour (List separately in addition to code for primary procedure)
96373			Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); intra-arterial
96374			Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); intravenous push, single or initial substance/drug
96375			Therapeutic, prophylactic, or diagnostic injection (specify substance or drug); each additional sequential intravenous push of a new substance/drug (List separately in addition to code for primary procedure)
96401			Chemotherapy administration, subcutaneous or intramuscular; non-hormonal anti-neoplastic
96402			Chemotherapy administration, subcutaneous or intramuscular; hormonal anti-neoplastic
96405			Chemotherapy administration; intralesional, up to and including 7 lesions
96415			Chemotherapy administration, intravenous infusion technique; each additional hour (List separately in addition to code for primary procedure)
95146			0438
95147	Professional services for the supervision of preparation and provision of antigens for allergen immunotherapy (specify number of doses); 3 single stinging insect venoms		
96360	Intravenous infusion, hydration; initial, 31 minutes to 1 hour		
96411	Chemotherapy administration; intravenous, push technique, each additional substance/drug (List separately in addition to code for primary procedure)		

CY 2010 HCPCS Code	Final CY 2010 APC	Final CY 2010 Approximate APC Median Cost	CY 2010 Long Descriptor
96417			Chemotherapy administration, intravenous infusion technique; each additional sequential infusion (different substance/drug), up to 1 hour (List separately in addition to code for primary procedure)
96420			Chemotherapy administration, intra-arterial; push technique
96423			Chemotherapy administration, intra-arterial; infusion technique, each additional hour (List separately in addition to code for primary procedure)
96542			Chemotherapy injection, subarachnoid or intraventricular via subcutaneous reservoir, single or multiple agents
95990	0439	\$126	Refilling and maintenance of implantable pump or reservoir for drug delivery, spinal (intrathecal, epidural) or brain (intraventricular);
95991			Refilling and maintenance of implantable pump or reservoir for drug delivery, spinal (intrathecal, epidural) or brain (intraventricular); administered by physician
96365			Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug); initial, up to 1 hour
96369			Subcutaneous infusion for therapy or prophylaxis (specify substance or drug); initial, up to 1 hour, including pump set-up and establishment of subcutaneous infusion site(s)
96406			Chemotherapy administration; intralesional, more than 7 lesions
96409			Chemotherapy administration; intravenous, push technique, single or initial substance/drug
96440			Chemotherapy administration into pleural cavity, requiring and including thoracentesis
96521			Refilling and maintenance of portable pump
96522			Refilling and maintenance of implantable pump or reservoir for drug delivery, systemic (eg, intravenous, intra-arterial)

CY 2010 HCPCS Code	Final CY 2010 APC	Final CY 2010 Approximate APC Median Cost	CY 2010 Long Descriptor
96413	0440	\$218	Chemotherapy administration; intravenous infusion technique; up to 1 hour, single or initial substance/drug
96416			Chemotherapy administration, intravenous infusion technique; initiation of prolonged chemotherapy infusion (more than 8 hours), requiring use of a portable or implantable pump
96422			Chemotherapy administration, intra-arterial; infusion technique, up to 1 hour
96425			Chemotherapy administration, intra-arterial; infusion technique, initiation of prolonged infusion (more than 8 hours), requiring the use of a portable or implantable pump
96445			Chemotherapy administration into peritoneal cavity, requiring and including peritoneocentesis
96450			Chemotherapy administration, into CNS (eg, intrathecal), requiring and including spinal puncture
C8957			Intravenous infusion for therapy/diagnosis; initiation of prolonged infusion (more than eight hours), requiring use of portable or implantable pump

IX. OPSS Payment for Hospital Outpatient Visits

A. Background

Currently, hospitals report visit HCPCS codes to describe three types of OPSS services: Clinic visits, emergency department visits, and critical care services. For OPSS purposes, we recognize clinic visit codes as those codes defined in the CPT codebook to report evaluation and management (E/M) services provided in the physician’s office or in an outpatient or other ambulatory facility. We recognize emergency department visit codes as those codes used to report E/M services provided in the emergency department.

Emergency department visit codes consist of five CPT codes that apply to Type A emergency departments, and five Level II HCPCS codes that apply to Type B emergency departments. For OPSS purposes, we recognize critical care codes as those CPT codes used by hospitals to report critical care services that involve the “direct delivery by a physician(s) of medical care for a critically ill or critically injured patient,” as defined by the CPT codebook. In Transmittal 1139, Change Request 5438, dated December 22, 2006, we stated that, under the OPSS, the time that can be reported as critical care is the time spent by a physician and/or hospital staff engaged in active face-to-

face critical care of a critically ill or critically injured patient. Under the OPSS, we also recognize HCPCS code G0390 (Trauma response team associated with hospital critical care service) for the reporting of a trauma response in association with critical care services.

As we proposed in the CY 2010 OPSS/ASC proposed rule (74 FR 35349 through 35350), we are continuing to recognize these CPT and HCPCS codes describing clinic visits, Type A and B emergency department visits, critical care services, and trauma team activation provided in association with critical care services for CY 2010. These codes are listed below in Table 47.

TABLE 47—HCPCS CODES USED TO REPORT CLINIC AND EMERGENCY DEPARTMENT VISITS AND CRITICAL CARE SERVICES

CY 2010 HCPCS Code	CY 2010 Descriptor
Clinic Visit HCPCS Codes	
99201	Office or other outpatient visit for the evaluation and management of a new patient (Level 1).

TABLE 47—HCPCS CODES USED TO REPORT CLINIC AND EMERGENCY DEPARTMENT VISITS AND CRITICAL CARE SERVICES—Continued

CY 2010 HCPCS Code	CY 2010 Descriptor
99202	Office or other outpatient visit for the evaluation and management of a new patient (Level 2).
99203	Office or other outpatient visit for the evaluation and management of a new patient (Level 3).
99204	Office or other outpatient visit for the evaluation and management of a new patient (Level 4).
99205	Office or other outpatient visit for the evaluation and management of a new patient (Level 5).
99211	Office or other outpatient visit for the evaluation and management of an established patient (Level 1).
99212	Office or other outpatient visit for the evaluation and management of an established patient (Level 2).
99213	Office or other outpatient visit for the evaluation and management of an established patient (Level 3).
99214	Office or other outpatient visit for the evaluation and management of an established patient (Level 4).
99215	Office or other outpatient visit for the evaluation and management of an established patient (Level 5).
Emergency Department Visit HCPCS Codes	
99281	Emergency department visit for the evaluation and management of a patient (Level 1).
99282	Emergency department visit for the evaluation and management of a patient (Level 2).
99283	Emergency department visit for the evaluation and management of a patient (Level 3).
99284	Emergency department visit for the evaluation and management of a patient (Level 4).
99285	Emergency department visit for the evaluation and management of a patient (Level 5).
G0380	Type B emergency department visit (Level 1).
G0381	Type B emergency department visit (Level 2).
G0382	Type B emergency department visit (Level 3).
G0383	Type B emergency department visit (Level 4).
G0384	Type B emergency department visit (Level 5).
Critical Care Services HCPCS Codes	
99291	Critical care, evaluation and management of the critically ill or critically injured patient; first 30–74 minutes.
99292	Critical care, evaluation and management of the critically ill or critically injured patient; each additional 30 minutes.
G0390	Trauma response associated with hospital critical care service.

During the February 2009 APC Panel meeting, the APC Panel recommended that CMS present at the next APC Panel meeting an analysis of the most common diagnoses and services associated with Type A and Type B emergency department visits, including an analysis by hospital-specific characteristics, as well as an analysis of CY 2008 claims data for clinic and emergency department (Types A and B) visits. The APC Panel also recommended that the work of the Visits and Observation Subcommittee continue. We adopted these recommendations in the CY 2010 OPPS/ASC proposed rule (74 FR 35350) and provided frequency and cost data from CY 2008 claims for clinic and emergency department visits at the August 2009 meeting of the APC Panel. We plan to provide the requested analysis of the most common diagnoses and services associated with Type A and Type B emergency department visits to the APC Panel at the winter 2010 meeting of the APC Panel.

At its August 2009 meeting, the APC Panel recommended that CMS present an analysis of CY 2009 claims data for clinic and emergency department (Type A and B) visits at the next meeting of the APC Panel. The APC Panel recommended again that CMS provide

analyses of the most common diagnoses and services associated with Type A and Type B emergency department visits at the next meeting of the APC Panel, including analysis by hospital-specific characteristics. We are accepting all of these recommendations and will present the available requested data at the winter 2010 meeting of the APC Panel.

B. Policies for Hospital Outpatient Visits
 1. Clinic Visits: New and Established Patient Visits

As reflected in Table 47, hospitals use different CPT codes for clinic visits based on whether the patient being treated is a new or an established patient. Beginning in CY 2009, we refined the definitions of new and established patients to reflect whether or not the patient has been registered as an inpatient or outpatient of the hospital within the past 3 years. A patient who has been registered as an inpatient or outpatient of the hospital within the 3 years prior to a visit would be considered to be an established patient for that visit, while a patient who has not been registered as an inpatient or outpatient of the hospital within the 3 years prior to a visit would be considered to be a new patient for that visit. We refer readers to the CY 2009

OPPS/ASC final rule with comment period (73 FR 68677 through 68680) for a full discussion of the refined definitions.

We stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35350) that we continue to believe that defining new or established patient status based on whether the patient has been registered as an inpatient or outpatient of the hospital within the 3 years prior to a visit will reduce hospitals' administrative burden associated with reporting appropriate clinic visit CPT codes. For CY 2010, we proposed to continue recognizing the refined definitions of new and established patients, and our policy of calculating median costs for clinic visits under the OPPS using historical hospital claims data.

Comment: Several commenters recommended that CMS remove the distinction between new and established patient clinic visits, arguing that facilities must expend the same level of resources regardless of whether the patient was registered as an inpatient or an outpatient in the hospital within the past 3 years. According to some commenters, CMS' use of the CPT codes for visits, which differentiate between new and established patients, is contrary to CMS'

past statements that the CPT guidelines and definitions for E/M visit codes are not applicable in the hospital outpatient setting because they fail to reflect hospital services and resource consumption. In addition, some commenters stated there are significant operational issues involved with implementing the 3-year criterion for hospital clinic visit billing purposes, and expressed concerns that hospitals' incorrect compliance with this requirement could be targeted by Recovery Audit Contractor (RAC) audits and other types of audits. Some commenters argued that any differential in costs that is evident in claims data for new versus established patient visits would be the result of hospitals' erroneous reporting of these codes, rather than any real difference in the level of resources expended treating a new versus an established patient. One commenter characterized the median cost differences between new and established patient visit codes as random and suggested that some providers report CPT codes 99201 and 99211 as "default codes" when reporting clinic visits, which, according to the commenter, raises the costs reflected in the claims data for these codes and artificially impacts the overall APC ratesetting process.

Some commenters asserted that CMS' proposal in the CY 2010 MPFS proposed rule to eliminate the use of consultation codes for physician payment purposes provides a precedent for discontinuing the use of the CPT E/M codes by hospitals. According to the commenters, CMS cited findings in the March 2006 OIG report entitled "Consultations in Medicare: Coding and Reimbursement" that physicians frequently misuse CPT codes for consultation services as a basis for no longer recognizing those codes under the MPFS. The commenters stated that hospitals similarly misuse the clinic visit codes, and that CMS should cease to recognize the clinic visit CPT codes under the OPSS for this reason.

Many commenters suggested that, as an alternative to the clinic visit CPT codes for new and established patients, hospitals bill for visits based on the resources expended in the visit at a level determined by the hospitals' internal reporting guidelines, regardless of whether the patient is new or established. Some commenters supported the use of Level II HCPCS G-codes for hospital clinic visits to represent hospital resources expended, without the distinction between new and established patients. According to the commenters, creation of these HCPCS G-codes would streamline

hospital reporting of visits, enable hospitals to correctly code for visits based on established definitions, and facilitate elimination of the new versus established patient visit concept for hospital reporting. The commenters noted that, in the past, providers have resisted implementing hospital-specific HCPCS codes for reporting visits before the implementation of national visit reporting guidelines for hospitals, but suggested that providers may now favor HCPCS G-codes over the existing CPT codes for visits that are tied to CMS' definition of new and established patients for purposes of reporting those codes. Some commenters suggested that CMS discuss the development of clinic visit HCPCS G-codes at the winter 2010 APC Panel meeting and include a proposal for clinic visit HCPCS G-codes in the CY 2011 OPSS/ASC proposed rule.

Response: Because hospital claims data continue to show significant cost differences between new and established patient visits, we continue to believe it is necessary and appropriate to recognize the CPT codes for both new and established patient visits and, in some cases, provide differential payment for new and established patient visits of the same level. For example, the final CY 2010 median cost for the level 3 new patient clinic visit, described by CPT code 99203 and calculated using over 200,000 single claims from CY 2008, is approximately \$96, while the final CY 2010 median cost for the level 3 established patient clinic visit, described by CPT code 99213 and calculated using over 4.5 million single claims from CY 2008, is approximately \$70. We believe this difference in median costs warrants continued assignment of these CPT codes to different APCs for CY 2010.

Given that we have a substantial volume of single claims from a significant number of hospitals upon which to calculate the median costs for all levels of clinic visits, we do not agree with the commenters that the differences in costs for new versus established patient visits are random or the result of erroneous billing. We expect hospitals to report all HCPCS codes in accordance with correct coding principles, CPT code descriptions, and relevant CMS guidance, which, in this case, specifies that the meanings of "new" and "established" patients as included in the clinic visit CPT code descriptors pertain to whether or not the patient has been registered as an inpatient or an outpatient of the hospital within the past 3 years (73 FR 68679). We have no reason to believe that

hospitals are systematically disregarding these principles to the extent that our median costs for clinic visits, which are based on data from millions of single claims, would be artificially skewed.

We also do not agree with the commenters that CMS' proposal in the CY 2010 MPFS proposed rule to eliminate the use of consultation codes for physician payment purposes (74 FR 33553 through 33554) has any direct relevance to the distinction between new and established patient visits under the OPSS. As we stated previously, we have no reason to believe that hospitals are not correctly reporting these services. Furthermore, unlike the MPFS proposal that would require physicians to report the existing CPT codes for new and established patient visits instead of the consultation CPT codes, we could not easily implement a policy to eliminate the use of the clinic visit CPT codes under the OPSS, because there are no other existing codes that hospitals could use to report these services.

We recognize that some commenters believe hospitals would now support the creation of Level II HCPCS G-codes for hospital clinic visits, whereas in the past they generally opposed hospital-specific codes for visits in the absence of national visit reporting guidelines. We welcome any comments hospitals have on alternative coding schemes for reporting hospital clinic visits that would not require hospitals to distinguish between new and established patients, such as the creation of hospital-specific clinic visit HCPCS G-codes or the exclusive use of established patient clinic visit codes. We are particularly interested in commenters' thoughts on how we would develop payment rates for clinic visits under another coding scheme, considering the claims data that we have now for these services demonstrate significant differences in costs between new and established patient clinic visits and could not be easily crosswalked to a structure that does not distinguish between new and established patients. We will consider any ideas that we receive as we prepare for the CY 2011 OPSS/ASC proposed rule.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue to define new or established patient status for the purpose of reporting the clinic visit CPT codes, on the basis of whether or not the patient has been registered as an inpatient or outpatient of the hospital within the past 3 years. We also are finalizing our CY 2010 proposal, without modification, to continue our policy of calculating median costs for

clinic visits under the OPSS using historical hospital claims data. As discussed in detail in section II.A.2.e.(1) of this final rule with comment period and consistent with our CY 2009 policy, when calculating the median costs for the clinic visit APCs (0604 through 0608), we utilized our methodology that excludes those claims for visits that are eligible for payment through the extended assessment and management composite APC 8002 (Level I Extended Assessment and Management Composite). We continue to believe that this approach results in the most accurate cost estimates for APCs 0604 through 0608 for CY 2010.

2. Emergency Department Visits

Since CY 2007, we have recognized two different types of emergency departments for payment purposes under the OPSS—Type A emergency departments and Type B emergency departments. As described in greater detail below, by providing payment for two types of emergency departments, we recognize for OPSS payment purposes both the CPT definition of an emergency department, which requires the facility to be available 24 hours, and the requirements for emergency departments specified in the provisions of the Emergency Medical Treatment and Labor Act (EMTALA) (Pub. L. 99–272), which do not stipulate 24-hour availability but do specify other obligations for hospitals that offer emergency services. For more detailed information on the EMTALA provisions, we refer readers to the CY 2009 OPSS/ASC final rule with comment period (73 FR 68680).

In the CY 2007 OPSS/ASC final rule with comment period (71 FR 68132), we finalized the definition of Type A emergency departments to distinguish them from Type B emergency departments. A Type A emergency department must be available to provide services 24 hours a day, 7 days a week, and meet one or both of the following requirements related to the EMTALA definition of a dedicated emergency department specified at § 489.24(b), specifically: (1) It is licensed by the State in which it is located under the applicable State law as an emergency room or emergency department; or (2) it is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment. For CY 2007 (71 FR 68140), we assigned the five CPT E/M emergency department visit codes for services provided in Type A emergency departments to five

created Emergency Visit APCs, specifically APC 0609 (Level 1 Emergency Visits), APC 0613 (Level 2 Emergency Visits), APC 0614 (Level 3 Emergency Visits), APC 0615 (Level 4 Emergency Visits), and APC 0616 (Level 5 Emergency Visits). We defined a Type B emergency department as any dedicated emergency department that incurred EMTALA obligations, but did not meet the CPT definition of an emergency department. For example, a hospital department that may be characterized as a Type B emergency department would meet the definition of a dedicated emergency department, but may not be available 24 hours a day, 7 days a week. Hospitals with such dedicated emergency departments incur EMTALA obligations with respect to an individual who presents to the department and requests, or has a request made on his or her behalf, examination or treatment for a medical condition.

To determine whether visits to Type B emergency departments have different resource costs than visits to either clinics or Type A emergency departments, in the CY 2007 OPSS/ASC final rule with comment period (71 FR 68132), we finalized a set of five HCPCS G-codes for use by hospitals to report visits to all entities that meet the definition of a dedicated emergency department under the EMTALA regulations but that are not Type A emergency departments. These codes are called “Type B emergency department visit codes.” In the CY 2007 OPSS/ASC final rule with comment period (71 FR 68132), we explained that these new HCPCS G-codes would serve as a vehicle to capture median cost and resource differences among visits provided by Type A emergency departments, Type B emergency departments, and clinics. We stated that the reporting of specific HCPCS G-codes for emergency department visits provided in Type B emergency departments would permit us to specifically collect and analyze the hospital resource costs of visits to these facilities in order to determine if, in the future, a proposal for an alternative payment policy might be warranted. We expected hospitals to adjust their charges appropriately to reflect differences in Type A and Type B emergency department visit costs.

As we noted in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68681), the CY 2007 claims data used for that rulemaking were from the first year of claims data available for analysis that included hospital’s cost data for these new Type B emergency department HCPCS visit codes. Based

on our analysis of the CY 2007 claims data, we confirmed that the median costs of Type B emergency department visits were less than the median costs of Type A emergency department visits for all but the level 5 visit. In other words, the median costs from the CY 2007 hospital claims represented real differences in the hospital resource costs for the same level of visits in a Type A or Type B emergency department. Therefore, for CY 2009, we adopted the August 2008 APC Panel recommendation to assign levels 1 through 4 Type B emergency department visits to their own APCs and to assign the level 5 Type B emergency department visit to the same APC as the level 5 Type A emergency department visit.

We now have CY 2008 cost data for CY 2010 ratesetting for the Type B emergency department HCPCS G-codes, representing a second year of claims data for these Type B emergency department visit HCPCS codes. In the CY 2010 OPSS/ASC proposed rule (74 FR 35351 through 35353), we presented our observation of frequency and patterns of billing based on the CY 2008 claims available at that time. We also repeated some of our analyses of Type B emergency department visits using the available CY 2008 claims and cost report data to confirm that Type B emergency department visit costs are generally lower than Type A emergency department visit costs and to assess whether there are systematic differences in the costs of Type A and Type B emergency department visits by Medicare contractor. The pattern of relative cost differences between Type A and Type B emergency department visits was largely consistent with the distributions we observed in the CY 2007 data, with the exception that, in the CY 2008 claims data available for the proposed rule, we observed a relatively lower HCPCS code-specific median cost associated with level 5 Type B emergency department visits compared to the HCPCS-code specific median cost of level 5 Type A emergency department visits. In contrast, in our CY 2007 claims data, we observed similar resource costs for level 5 Type A and Type B emergency department visits. We also determined that there are no significant differences in how Medicare contractors have interpreted our Type A and Type B emergency department visit reporting policies.

We shared cost and frequency data with the Visits and Observation Subcommittee of the APC Panel during the February 2009 meeting, and in the CY 2010 OPSS/ASC proposed rule (74

FR 35353), we proposed to pay for Type B emergency department visits in CY 2010 consistent with their median costs. Specifically, we proposed to pay for levels 1 through 4 Type B emergency department visits through four levels of APCs: APC 0626 (Level 1 Type B Emergency Visits), APC 0627 (Level 2 Type B Emergency Visits), APC 0628 (Level 3 Type B Emergency Visits), and APC 0629 (Level 4 Type B Emergency Visits). In addition, we proposed to adopt new APC 0630 (Level 5 Type B Emergency Visits) and to pay for level 5 Type B emergency department visits through this new APC. We proposed to assign HCPCS codes G0380, G0381, G0382, G0383, and G0384 (the levels 1, 2, 3, 4, and 5 Type B emergency department visit Level II HCPCS codes) to APCs 0626, 0627, 0628, 0629, and 0630, respectively, for CY 2010. These HCPCS codes would be the only HCPCS codes assigned to these APCs. Furthermore, to distinguish new APC 0630 from the APC for the level 5 Type A emergency visits, we proposed to modify the title of the current level 5 Type A emergency visit APC to incorporate Type A in the title.

Therefore, the revised title of APC 0616 would be "Level 5 Type A Emergency Visits."

We noted in the CY 2010 OPPI/ASC proposed rule (74 FR 35353) that the proposed policy to pay for Type B emergency department visits based on their median costs is consistent with the APC Panel's March 2008 recommendation for payment of Type B emergency department visits. As part of its recommended configuration of APCs for Type B emergency department visits in CY 2009, the APC Panel also stated that, given the limited CY 2007 claims data available for Type B emergency department visits, CMS should reconsider payment adjustments as more claims data become available. In general, the APC Panel's recommended CY 2009 configuration paid appropriately for each level of the Type B emergency department visits, based on the resource costs of the Type B emergency department visits that are reflected in claims data. We stated in the proposed rule that we believe our proposed CY 2010 configuration also would pay appropriately for each level of Type B emergency department visits

based on estimated resource costs from more recent claims data.

For this final rule with comment period, based on updated CY 2008 claims data, we note that 344 hospitals billed at least one Type B emergency department visit code in CY 2008, with a total frequency of visits provided in Type B emergency departments of approximately 220,000. All except 5 of the 344 hospitals reporting Type B emergency department visits in CY 2008 also reported Type A emergency department visits. Overall, many more hospitals (approximately 3,238 total hospitals) reported Type A emergency department visits than Type B emergency department visits. For comparison purposes, the total frequency of visits provided in hospital outpatient clinics and Type A emergency departments is approximately 17.5 million and 11.6 million, respectively. The median costs for the Type B emergency department visit APCs, as compared to the Type A emergency department visit APCs and the clinic visit APCs, are shown in Table 48 below.

TABLE 48—COMPARISON OF MEDIAN COSTS FOR CLINIC VISIT APCs, TYPE B EMERGENCY DEPARTMENT VISIT APCs, AND TYPE A EMERGENCY DEPARTMENT VISIT APCs

Visit level	Final CY 2010 clinic visit approximate APC median cost	Final CY 2010 type B emergency department approximate APC median cost	Final CY 2010 type A emergency visit approximate APC median cost
Level 1	\$57	\$45	\$53
Level 2	69	62	87
Level 3	88	97	139
Level 4	112	141	221
Level 5	166	230	327

As demonstrated in Table 48, the median costs of the lowest level visits, based on the CY 2008 claims and cost report data available for this final rule with comment period, continue to be similar across all settings, including clinic and Type A and B emergency departments. Visit levels 2 and 3 share similar resource costs in the clinic and Type B emergency department settings, while visits provided in Type A emergency departments have higher estimated resource costs at these levels. The level 4 clinic visit APC is less resource-intensive than the level 4 Type B emergency department visit APC, which is similarly less resource-intensive than the level 4 Type A emergency department visit APC. Similarly, the level 5 clinic visit APC is less resource-intensive than the level 5

Type B emergency department visit APC, which is less resource-intensive than the level 5 Type A emergency department visit APC.

This pattern of relative cost differences between Type A and Type B emergency department visits is largely consistent with the distributions we observed in the CY 2007 data, with the exception that, in the updated CY 2008 claims data, we observe a relatively lower HCPCS code-specific median cost associated with level 5 Type B emergency department visits compared to the HCPCS-code specific median cost of level 5 Type A emergency department visits. In contrast, in our CY 2007 claims data, we observed similar resource costs for level 5 Type A and Type B emergency department visits. In the CY 2009 OPPI/ASC final rule with

comment period (73 FR 68683), we hypothesized that, for the highest level of emergency department visits, the resources required would be the same in both emergency department settings. Now that more data on Type B emergency department visits are available and hospitals have more experience billing for Type B services, we observe differences in the resources for the highest level emergency department visits to Type A and Type B emergency departments.

As noted in the CY 2009 OPPI/ASC final rule with comment period (73 FR 68683), we performed data analyses regarding the costs of Type A and Type B emergency department visits in addition to our standard median cost calculations. These analyses included studying the emergency department

visit costs of hospitals that billed Type B emergency department visits only, analyzing the cost data for hospitals that billed both Type A and Type B emergency department visits, and evaluating whether there were differences in the costs of Type A and Type B emergency department visits by Medicare contractor to ascertain whether there were differences in how Medicare contractors have interpreted our Type A and Type B emergency department visit policies. In the CY 2007 data, we observed that hospitals that billed both Type A and Type B emergency department visits had lower costs for Type B emergency department visits than Type A emergency department visits at all levels except for the level 5 Type B emergency department visit. Our analyses of the differences in Type A and Type B emergency department visit median costs by Medicare contractors did not identify concerning differences. Overall, we observed a distribution of visit costs as expected, including generally lower Type B emergency department visit costs in comparison with Type A emergency department visits, and increasing costs for Type B emergency department visits from levels 1 through 5, similar to the cost increases we observed from levels 1 through 5 for Type A emergency department visits. We also observed a few contractors with more unusual cost distributions for Type B emergency department visits, including relatively similar or higher costs across levels 1 through 5 for Type B emergency department visits. For CY 2009, we concluded that we had no reason to believe that the cost differences between Type A and Type B emergency department visits evident in our aggregate OPPS claims data resulted from varying Medicare contractor criteria as to what defines Type A and Type B emergency departments. We also committed to monitoring these distributions in future years.

As we did for the CY 2010 OPPS/ASC proposed rule, for this final rule with comment period, we repeated some of our analyses of Type B emergency department visits using updated CY 2008 claims data to confirm that Type B emergency department visit costs are generally lower than Type A emergency department visit costs and to again assess whether there are systematic differences in the costs of Type A and Type B emergency department visits by Medicare contractor. As noted above, we observed that hospitals that billed both Type A and Type B emergency department visits had lower costs for Type B emergency department visits

than Type A emergency department visits, including level 5 Type B emergency department visits, which is a change from the CY 2007 data. We further evaluated differences in the costs of Type A and Type B emergency department visits by Medicare contractor. Based on our updated analysis of CY 2008 claims, we continue to observe similar patterns in HCPCS code-specific median cost differences between Type A and Type B emergency department visits as observed in the CY 2007 claims. Hospitals in the jurisdictions of most Medicare contractors have generally lower Type B emergency department visit costs in comparison with Type A emergency department visits, as well as increasing costs for Type B emergency department visits from levels 1 through 5, similar to the cost increases we observed from levels 1 through 5 for Type A emergency department visits.

Like last year, we also continue to observe a few Medicare contractors with more unusual cost distributions for Type B emergency department visits, including those with Type B emergency department visit costs that are relatively similar or higher than Type A emergency department visit costs across levels 1 through 5. Some of these Medicare contractors are the same contractors that we noted had more unusual cost distributions for Type B emergency department visits relative to Type A emergency department visit costs in the CY 2007 claims data. In order to confirm that these Medicare contractors are applying our policies consistently, we examined the HCPCS code-specific median costs for Type A and Type B emergency department visits for the hospitals in each Medicare contractor's area. For almost all of these Medicare contractors, we see one or two hospitals with relatively high Type B emergency department visit costs relative to Type B emergency department visit costs nationwide or with Type B emergency department visit costs that are relatively similar to or higher than Type A emergency department visit costs. These one or two hospitals have sufficient visit volumes to influence the calculation of the HCPCS code-specific median costs for their respective Medicare contractors.

Comment: Several commenters supported CMS' proposal to create a new APC for level 5 Type B emergency department visits. One commenter encouraged CMS to adopt the recommendation made by the APC Panel at the August 2009 meeting to provide an analysis of the most common diagnoses and services associated with Type A and Type B emergency

department visits at the next meeting of the APC Panel, including analysis by hospital-specific characteristics, as well as an analysis of CY 2009 claims data for Type A and B emergency department visit APCs.

Response: We appreciate commenters' support of our proposal to create a new APC for level 5 Type B emergency department visits. Our updated analyses of Type B emergency department visits costs for this CY 2010 OPPS/ASC final rule with comment period confirm that the median costs of Type B emergency department visits are less than the median costs of Type A emergency department visits across all levels. Our updated analyses also confirm that there are no significant differences in how Medicare contractors have interpreted our Type A and Type B emergency department visit reporting policies. The median costs from CY 2008 hospital claims represent real differences in the hospital resource costs for the same level of visit in a Type A or Type B emergency department.

Therefore, as we proposed, for the CY 2010 OPPS, we are continuing to pay for Type B emergency department visits in CY 2010 consistent with their median costs. Specifically, we are continuing to pay levels 1 through 4 Type B emergency department visits through four levels of APCs: APC 0626 (Level 1 Type B Emergency Visits), APC 0627 (Level 2 Type B Emergency Visits), APC 0628 (Level 3 Type B Emergency Visits), and APC 0629 (Level 4 Type B Emergency Visits). In addition, we are adopting new APC 0630 (Level 5 Type B Emergency Visits) and will pay for level 5 Type B emergency department visits through this new APC. We are assigning HCPCS codes G0380, G0381, G0382, G0383, and G0384 (the levels 1, 2, 3, 4, and 5 Type B emergency department visit Level II HCPCS codes) to APCs 0626, 0627, 0628, 0629, and 0630, respectively, for CY 2010. These HCPCS codes are the only HCPCS codes assigned to these APCs. Furthermore, to distinguish new APC 0630 from the APC for the level 5 Type A emergency visits, as we proposed, we are modifying the title of the current level 5 Type A emergency visit APC to incorporate Type A in the title. Therefore, the revised title of APC 0616 is "Level 5 Type A Emergency Visits." We believe our CY 2010 configuration pays appropriately for each level of Type B emergency department visits based on estimated resource costs from more recent claims data.

As stated previously, we plan to provide the requested analysis of the most common diagnoses and services associated with Type A and Type B

emergency department visits to the APC Panel at the winter 2010 meeting of the APC Panel, as well as an analysis of CY 2009 claims data for Type A and B emergency department visit APCs available at that time.

Comment: One commenter expressed concerns regarding the 30 minute minimum to bill critical care services, described by CPT code 99291. The commenter argued that the resources expended in less than 30 minutes warrant payment at the highest level of E/M payment, and recommended that CMS change the criteria for payment for critical care services to include instances of 15 minutes of critical care and instances in which the patient expires in less than 30 minutes, despite the critical care services furnished. According to the commenter, the significant resources utilized during these critical care episodes are not appropriately recognized for payment purposes because they cannot be reported with CPT code 99291 under existing guidelines.

Another commenter requested that CMS consider extending payment for trauma team activations, described by HCPCS code G0390, to level 5 emergency department visits, in addition to critical care services when all other trauma activation criteria are met. According to the commenter, an emergency department that is extremely efficient can send a patient in need of a trauma team to surgery before the 30 minute time threshold for reporting critical care services is met. The commenter stated that, because the hospital would bill a level 5 emergency department visit code, rather than a critical care code, the encounter would not qualify for trauma response payment even though a trauma response team was utilized. The commenter argued that hospitals should receive an APC payment for HCPCS code G0390 under these circumstances because equivalent trauma team resources are expended even though the encounter lasted fewer than 30 minutes and cannot be reported with CPT code 99291.

Response: As we have stated in the past (72 FR 66806), the CPT instructions for reporting of critical care services with CPT code 99291 and the CPT code descriptor specify that the code can only be billed if 30 minutes or more of critical care services are provided. Hospitals must continue to provide a minimum of 30 minutes of critical care services in order to bill CPT code 99291, according to the CPT code descriptor and CPT instructions. We note that hospitals can report the appropriate clinic or emergency department visit

code consistent with their internal guidelines if fewer than 30 minutes of critical care is provided. These CPT instructions and our payment policies for covered hospital outpatient services do not apply any differently if the patient dies while undergoing treatment. We do allow hospitals to use the HCPCS-CA modifier to address situations where a procedure on the OPPI inpatient list must be performed to resuscitate or stabilize a patient (whose status is that of an outpatient) with an emergent, life-threatening condition, and the patient dies before being admitted as an inpatient. We refer readers to section II.A.2.d.(7) of this final rule with comment period for more information on how these services are paid under the OPPI.

We do not agree with the commenter that we should modify our policy to recognize HCPCS code G0390 for the reporting of a trauma response in association with critical care services when the hospital provides fewer than 30 minutes of critical care and cannot report CPT code 99291. We believe that it would be extremely unusual for a patient to require trauma team services, be rushed to surgery within 30 minutes of arrival in the emergency department, and not be subsequently admitted to the hospital as an inpatient. Furthermore, hospitals that provide less than 30 minutes of critical care when trauma activation occurs under the circumstances described by the NUBC guidelines that would permit reporting a charge under revenue code series 68x may report a charge under 68x, but they may not report HCPCS code G0390. In these cases, payment for the trauma team activation is packaged into payment for the other services provided to the patient in the encounter, including the associated emergency department visit that is reported.

Comment: One commenter requested clarification regarding “triage only” visits in which a patient is seen by a nurse and triaged in the hospital emergency department but leaves prior to a physician’s examination and treatment. The commenter asked if hospitals can bill visit codes for such cases if the patient is not seen by a physician.

Response: As we have stated in the past (73 FR 68686), under the OPPI, unless indicated otherwise, we do not specify the type of hospital staff (for example, nurses or pharmacists) who may provide services in hospitals because the OPPI only makes payment for services provided incident to physicians’ services. Hospitals providing services incident to

physicians’ services may choose a variety of staffing configurations to provide those services, taking into account other relevant factors, including State and local laws, hospital policies, and other Federal requirements such as EMTALA and the Medicare conditions of participation related to hospital staffing. Billing a visit code in addition to another service merely because the patient interacted with hospital staff or spent time in a room for that service is inappropriate. A hospital may bill a visit code based on the hospital’s own coding guidelines which must reasonably relate the intensity of hospital resources to different levels of HCPCS codes. Services furnished must be medically necessary and documented.

As described previously in this section, we are adopting our proposal, without modification, to continue paying for Type B emergency department visits in CY 2010 consistent with their median costs through 5 levels of Type emergency department visit APCs.

Table 49 below displays the APC median costs for each level of Type B emergency department visit under our CY 2010 configuration. As more cost data become available and hospitals gain additional experience with reporting visits to Type B emergency departments, we will continue to regularly reevaluate patterns of Type A and Type B emergency department visit reporting to ensure that hospitals continue to bill appropriately and differentially for these services. In addition, according to our usual practice, we will examine trends in cost data over time and consider proposing alternative emergency department visit APC configurations in the future if updated data indicate that changes to the payment structure should be considered.

We also note that, as discussed in section II.A.2.e.(1) of this final rule with comment period and consistent with our CY 2009 policy, when calculating the median costs for the emergency department visit and critical care APCs (0609 through 0617 and 0626 through 0630), we utilized our methodology that excludes those claims for visits that are eligible for payment through the extended assessment and management composite APC 8002 (Level I Extended Assessment and Management Composite). We continue to believe that this approach will result in the most accurate cost estimates for APCs 0604 through 0608 for CY 2010.

TABLE 49—CY 2010 TYPE B EMERGENCY DEPARTMENT VISIT APC ASSIGNMENTS AND MEDIAN COSTS

Type B emergency department level	Final CY 2010 APC assignment	Final CY 2010 approximate APC median cost
Level 1	0626	\$45
Level 2	0627	62
Level 3	0628	97
Level 4	0629	141
Level 5	0630	230

3. Visit Reporting Guidelines

Since April 7, 2000, we have instructed hospitals to report facility resources for clinic and emergency department hospital outpatient visits using the CPT E/M codes and to develop internal hospital guidelines for reporting the appropriate visit level. Because a national set of hospital-specific codes and guidelines do not currently exist, we have advised hospitals that each hospital's internal guidelines that determine the levels of clinic and emergency department visits to be reported should follow the intent of the CPT code descriptors, in that the guidelines should be designed to reasonably relate the intensity of hospital resources to the different levels of effort represented by the codes.

As noted in detail in the CY 2008 OPPI/ASC final rule with comment period (72 FR 66802 through 66805), we observed a normal and stable distribution of clinic and emergency department visit levels in hospital claims over the past several years. The data indicated that hospitals, on average, were billing all five levels of visit codes with varying frequency, in a consistent pattern over time. Overall, both the clinic and emergency department visit distributions indicated that hospitals were billing consistently over time and in a manner that distinguished between visit levels, resulting in relatively normal distributions nationally for the OPPI, as well as for specific classes of hospitals. The results of these analyses were generally consistent with our understanding of the clinical and resource characteristics of different levels of hospital outpatient clinic and emergency department visits. In the CY 2008 OPPI/ASC proposed rule (72 FR 42764 through 42765), we specifically invited public comment as to whether a pressing need for national guidelines continued at this point in the maturation of the OPPI, or if the current system where hospitals create and apply their own internal guidelines to report visits was currently more practical and appropriately flexible for hospitals. We

explained that although we have reiterated our goal since CY 2000 of creating national guidelines, this complex undertaking for these important and common hospital services was proving more challenging than we initially thought as we received new and expanded information from the public on current hospital reporting practices that led to appropriate payment for the hospital resources associated with clinic and emergency department visits. We stated our belief that many hospitals had worked diligently and carefully to develop and implement their own internal guidelines that reflected the scope and types of services they provided throughout the hospital outpatient system. Based on public comments, as well as our own knowledge of how clinics operate, it seemed unlikely that one set of straightforward national guidelines could apply to the reporting of visits in all hospitals and specialty clinics. In addition, the stable distribution of clinic and emergency department visits reported under the OPPI over the past several years indicated that hospitals, both nationally in the aggregate and grouped by specific hospital classes, were generally billing in an appropriate and consistent manner as we would expect in a system that accurately distinguished among different levels of service based on the associated hospital resources.

Therefore, we did not propose to implement national visit guidelines for clinic or emergency department visits for CY 2008. Since publication of the CY 2008 OPPI/ASC final rule with comment period, we have again examined the distribution of clinic and Type A emergency department visit levels based upon updated CY 2008 claims data available for the CY 2010 proposed rule and for this final rule with comment period and confirmed that we continue to observe a normal and stable distribution of clinic and emergency department visit levels in hospital claims. We continue to believe that, based on the use of their own internal guidelines, hospitals are

generally billing in an appropriate and consistent manner that distinguishes among different levels of visits based on their required hospital resources. As a result of our updated analyses, we are encouraging hospitals to continue to report visits during CY 2010 according to their own internal hospital guidelines. In the absence of national guidelines, we will continue to regularly reevaluate patterns of hospital outpatient visit reporting at varying levels of disaggregation below the national level to ensure that hospitals continue to bill appropriately and differentially for these services. As originally noted in detail in the CY 2008 OPPI/ASC final rule with comment period (72 FR 66648), we continue to expect that hospitals will not purposely change their visit guidelines or otherwise upcode clinic and emergency department visits for purposes of extended assessment and management composite APC payment.

In addition, we note our continued expectation that hospitals' internal guidelines will comport with the principles listed in the CY 2008 OPPI/ASC final rule with comment period (72 FR 66805). We encourage hospitals with more specific questions related to the creation of internal guidelines to contact their local fiscal intermediary or MAC.

Comment: Several commenters supported CMS' policy of requiring hospitals to use their own internal guidelines to distinguish among different levels of visits based on their required hospital resources and did not favor the implementation of national guidelines at any point in the future. Other commenters expressed appreciation for CMS' approach of studying the challenges associated with national guidelines prior to their implementation. However, many commenters urged CMS to move forward with the implementation of national guidelines for hospitals to report clinic visits because of several problems that they believe continue to exist due to the lack of such guidelines, such as variations in hospitals' internal guidelines that may result in

inconsistent cost data upon which payment rates for visits are based. Some commenters noted that some Medicare contractors, including RACs, use their own auditing methods rather than reviewing each hospital's internal guidelines while conducting medical review.

The commenters urged CMS to adopt national guidelines no later than CY 2011 due to the burden hospitals would face if they had to implement national visit coding guidelines concurrently with the ICD-10-CM and ICD-10-PCS changes required by FY 2013. According to the commenters, the national guidelines should be clear, concise, and specific with little or no room for varying interpretations, and hospitals should have at least 1 year to prepare for the transition. Many commenters indicated that the American Hospital Association (AHA) will reconvene an expert panel to submit a request to the AMA CPT Editorial Panel to create CPT codes for hospital visits and encouraged CMS to be engaged in and supportive of the recommendations of the expert panel.

Several commenters also recommended that, in the absence of national guidelines for hospital visit reporting, CMS provide additional guidance relating to the specific services that should be included or bundled into the visit levels reported by hospitals. One commenter requested that CMS ask the AMA to supplement its CPT Codebook to include a compilation of instructions from CMS regarding appropriate reporting of hospital visits, such as the 11 principles specified in the CY 2008 OPSS/ASC final rule with comment period that hospitals should follow in developing internal guidelines for reporting visits.

Another commenter performed extensive review of a large sample of hospital emergency department visits to determine whether the distributions observed in this sample resembled the distribution described by CMS and printed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66804). The commenter explained that the results are similar to those of CMS at the national level, but that emergency departments have increased the proportion of level 4 and 5 emergency department visits in recent years, and that several outlier providers are billing significantly more high level visits than expected based on their geographic location and hospital type. Therefore, the commenter concluded that national guidelines could help slow rapidly increasing health care costs.

Response: We acknowledge that it would be desirable to many hospitals to

have national guidelines. However, we also understand that it would be disruptive and administratively burdensome to other hospitals that have successfully adopted internal guidelines to implement any new set of national guidelines while we address the problems that would be inevitable in the case of any new set of guidelines that would be applied by thousands of hospitals. As noted in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66806), we encourage fiscal intermediaries and MACs to review a hospital's internal guidelines when an audit occurs. While we also would encourage RACs to review a hospital's internal guidelines when an audit occurs, we note that currently there are no RAC activities involving visit services. RAC audits may involve CMS-approved issues only and must be posted to each RAC's Web site.

We agree with the commenters that national guidelines should be clear, concise, and specific with little or no room for varying interpretations, and that hospitals should have at least 1 year to prepare for the transition. We look forward to reviewing any recommendations that result from the AHA-convened expert panel referenced by the commenters. If the AMA were to create facility-specific CPT codes for reporting visits provided in HOPDs, we would certainly consider such codes for OPSS use. We also appreciate the visit level distribution analysis provided to us by one commenter and note that, in the absence of national guidelines, we will continue to regularly reevaluate patterns of hospital outpatient visit reporting at varying levels of disaggregation below the national level to ensure that hospitals continue to bill appropriately and differentially for these services. We reiterate our expectation that hospitals' internal guidelines fully comply with the principles listed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 68805).

Regarding the public comments requesting clarification of services that should be included or bundled into visit codes, as we have stated in the past (73 FR 68685), hospitals should separately report all HCPCS codes in accordance with correct coding principles, CPT code descriptions, and any additional CMS guidance, when applicable. We refer readers to the July 2008 OPSS quarterly update (Transmittal 1536, Change Request 6094, issued on June 19, 2008) for further clarification about the reporting of CPT codes for hospital outpatient services paid under the OPSS. In that transmittal, we note that, while CPT codes generally are created to

describe and report physician services, they also are used by other providers/suppliers to describe and report services that they provide. Therefore, the CPT code descriptors do not necessarily reflect the facility component of a service furnished by the hospital. Some CPT code descriptors include reference to a physician performing a service. For OPSS purposes, unless indicated otherwise, the usage of the term "physician" does not restrict the reporting of the code or application of related policies to physicians only, but applies to all practitioners, hospitals, providers, or suppliers eligible to bill the relevant CPT codes in accordance with applicable portions of the Act, the Medicare regulations, and other Medicare guidance. In cases where there are separate codes for the technical component, professional component, and/or complete procedure, hospitals should report the code that represents the technical component for their facility services. If there is no separate technical component code for the service, hospitals should report the code that represents the complete procedure. Consistent with past input we have received from many hospitals, hospital associations, the APC Panel, and others, we will continue to utilize CPT codes for reporting services under the OPSS whenever possible to minimize hospitals' reporting burden.

We do not agree with the commenter that we should ask the AMA to supplement its CPT Codebook to include a compilation of instructions from CMS regarding appropriate reporting of hospital visits. Under the OPSS, we develop policies specifically and exclusively for purposes of the Medicare program, while the CPT Codebook provides instructions that are applicable to hospital coding for all payers, unless those payers choose to implement different individual policies. If hospitals believe the inclusion of such information in the CPT Codebook is necessary and appropriate, they may directly request the AMA to do so.

Comment: One commenter requested that CMS recognize CPT codes 99363 (Anticoagulation management for an outpatient taking warfarin, physician review and interpretation of International Normalized Ratio (INR) testing, patient instructions, dosage adjustment (as needed), and ordering of additional tests; initial 90 days of therapy (must include a minimum of 8 INR measurements)) and 99364 (Anticoagulation management for an outpatient taking warfarin, physician review and interpretation of International Normalized Ratio (INR) testing, patient instructions, dosage

adjustment (as needed), and ordering of additional tests; each subsequent 90 days of therapy (must include a minimum of 3 INR measurements)), which are currently assigned status indicator "B" (Codes that are not recognized by OPSS when submitted on an outpatient hospital Part B bill type (12x and 13x)), as payable under the OPSS. The commenter stated that making these CPT codes payable under the OPSS is appropriate because they accurately describe anticoagulation management services. The commenter argued that recognition of these CPT codes would reduce patient liability because they are billed only once every 90 days.

Response: While we appreciate the commenter's concern about patient liability, we cannot assess whether recognition of CPT codes 99363 and 99364 as payable under the OPSS would actually reduce the cumulative amount of copayment a patient may have to pay for all of the different services that may be involved in anticoagulation management, which may be provided at varying time intervals and with very different levels of intensity for individual patients. We expect that a patient undergoing anticoagulation management by hospital staff for a significant medical condition would periodically have hospital visits, and we would package payment for the non-face-to-face management of the patient's therapy between visits into payment for the visits themselves. Our usual policy is to package payment for the hospital resources associated with managing patients' medical conditions between hospital encounters for patients who undergo surgery or receive hospital visits for any medical condition, including diabetes, hypertension, or cardiac arrhythmias, and we do not believe that payment for anticoagulation management services should be made differently than payment for other medical or surgical management services. Therefore, we see no reason to recognize CPT codes 99363 and 99364 for payment under the OPSS.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign status indicator "B" to CPT codes 99363 and 99364 to indicate that these codes are not recognized for payment under the OPSS. We expect that hospitals will continue to consider the hospital resources required to manage patients, including patients requiring anticoagulation management, between hospital encounters when setting their charges for the services furnished in those encounters.

As we have stated in the past (73 FR 68686), we note that billing a visit code in addition to another service merely because the patient interacted with hospital staff or spent time in a room for that service is inappropriate. A hospital may bill a visit code based on the hospital's own coding guidelines, which must reasonably relate the intensity of hospital resources to the different levels of HCPCS codes. Services furnished must be medically necessary and documented. For example, CPT code 85610 (Prothrombin time) is a code that describes performance of the prothrombin time test. If the only service provided is a venipuncture and laboratory test to determine the prothrombin time, this service is the only service that should be billed. If a hospital provides a distinct, separately identifiable service in addition to the test, the hospital is responsible for billing the code that most closely describes the service provided.

We appreciate all of the comments we have received in the past from the public on visit guidelines, and we encourage continued submission of comments throughout the year that would assist us and other stakeholders interested in the development of national guidelines. Until national guidelines are established, hospitals should continue using their own internal guidelines to determine the appropriate reporting of different levels of clinic and emergency department visits. While we understand the interest of some hospitals in having us move quickly to promulgate national guidelines that would ensure standardized reporting of hospital outpatient visit levels, we believe that the issues and concerns identified both by us and others are important and require serious consideration prior to the implementation of national guidelines. Because of our commitment to provide hospitals with 6 to 12 months notice prior to implementation of national guidelines, we would not implement national guidelines prior to CY 2011. Our goal is to ensure that OPSS national or hospital-specific visit guidelines continue to facilitate consistent and accurate reporting of hospital outpatient visits in a manner that is resource-based and supportive of appropriate OPSS payments for the efficient and effective provision of visits in hospital outpatient settings.

X. Payment for Partial Hospitalization Services

A. Background

Partial hospitalization is an intensive outpatient program of psychiatric

services provided to patients as an alternative to inpatient psychiatric care for individuals who have an acute mental illness. Section 1833(t)(1)(B)(i) of the Act provides the Secretary with the authority to designate the HOPD services to be covered under the OPSS. The Medicare regulations at § 419.21 that implement this provision specify that payments under the OPSS will be made for partial hospitalization services furnished by community mental health centers (CMHCs) as well as those services furnished by hospitals to their outpatients. Section 1833(t)(2)(C) of the Act requires the Secretary to establish relative payment weights for covered HOPD services (and any APCs) based on median (or mean, at the election of the Secretary) hospital costs using data on claims from 1996 and data from the most recent available cost reports. Because a day of care is the unit that defines the structure and scheduling of partial hospitalization services, we established a per diem payment methodology for the partial hospitalization program (PHP) APC, effective for services furnished on or after August 1, 2000 (65 FR 18452).

Historically, the median per diem cost for CMHCs greatly exceeded the median per diem cost for hospital-based PHPs and fluctuated significantly from year to year, while the median per diem cost for hospital-based PHPs remained relatively constant (\$200–\$225). We believe that CMHCs may have increased and decreased their charges in response to Medicare payment policies. In developing the CY 2008 update, we began an effort to strengthen the PHP benefit through extensive data analysis and policy and payment changes. We began this effort as a result of the significant fluctuations and declines in the CMHC PHP median per diem costs (we refer readers to the CY 2008 OPSS/ASC final rule with comment period (72 FR 66670 through 66676) for a detailed discussion). The analysis included an examination of revenue-to-cost center mapping, refinements to the per diem methodology, and an in-depth analysis of the number of units of service furnished per day.

For CY 2008, we proposed and finalized two refinements to the methodology for computing the PHP median that we believe resulted in more accurate per diem medians. First, we remapped 10 revenue codes that are common among hospital-based PHP claims to the most appropriate cost centers (72 FR 66671 through 66672). Typically, we map the revenue code to the most specific cost center with a provider-specific CCR. However, if the hospital does not have a CCR for any of

the listed cost centers, we consider the overall hospital CCR as the default. For partial hospitalization services, the revenue center codes billed by hospital-based PHPs are mapped to Primary Cost Center 3550 (Psychiatric/Psychological Services). If that cost center is not available, they are mapped to the Secondary Cost Center 6000 (Clinic). We use the overall facility CCR for CMHCs because PHPs are CMHCs' only Medicare cost, and CMHCs do not have the same cost structure as hospitals. Therefore, for CMHCs, we use the CCR from the outpatient provider-specific file. A closer examination of the revenue-code-to-cost-center crosswalk revealed that 10 of the revenue center codes used by hospital-based PHPs did not map to a Primary Cost Center of 3550 or a Secondary Cost Center of 6000. We believe this occurred because these codes may also be used for services that are not furnished in a PHP or services that are not psychiatric related (for example, occupational therapy). As discussed in the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66671 through 66672), we updated this analysis using more recent PHP claims and CCR data. After remapping codes, we computed an alternate cost for each line item of the hospital-based PHP claims. Remapping those 10 revenue center codes reduced the number of lines that defaulted to the hospitals' overall CCR and thus created a more accurate estimate of PHP per diem costs for a significant number of claims.

Secondly, we refined our methodology for calculating median PHP per diem costs by computing a separate per diem cost for each day rather than for each claim. When there were multiple days of care entered on a claim, a unique cost was computed for each day of care. We only assigned costs for line items on days when a payment was made. All of these costs were then arrayed from lowest to highest, and the middle value of the array was considered the median per diem cost. A complete discussion of the refined method of computing the PHP median per diem cost can be found in the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66672).

After completing extensive data analysis, we continued to observe a clear downward trend in the median per diem cost based on the CY 2006 data used to develop the median per diem cost under the CY 2008 OPPTS/ASC final rule with comment period. The analysis revealed that fewer PHP services were being provided in a given day. We believed, and continue to believe, that the data reflect the level of cost for the

type of services that were being provided and continue to be provided.

Because partial hospitalization is provided in lieu of inpatient care, it should be a highly structured and clinically intensive program, usually lasting most of the day. In order to improve the level of services furnished in a PHP day, in the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66673), we reiterated our expectation that hospitals and CMHCs must provide a comprehensive program consistent with the statutory intent. We also indicated our intent to explore changes to our regulations and claims processing systems in order to deny payment for low intensity days.

For CY 2009, we implemented several regulatory, policy, and payment changes, including a two-tiered payment approach for PHP services under which we pay one amount for days with 3 units of service (APC 0172 (Level I Partial Hospitalization)) and a higher amount for days with 4 or more units of service (APC 0173 (Level II Partial Hospitalization)). We implemented this payment approach to reflect the lower costs of a less intensive day while still paying programs that provide 4 or more units of service an amount that recognizes that they have provided a more intensive day of care. In this way, we pay more appropriately for the level of care provided while still allowing PHPs necessary scheduling flexibility (73 FR 68689). As we stated in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68688), it was never our intention that days with only 3 units of service become the number of services provided in a typical day. Our intention was to provide payment to cover days that consisted of 3 units of service only in certain limited circumstances. For example, we believe 3 units of service a day may be appropriate when a patient is transitioning towards discharge or when a patient is required to leave the PHP early for the day due to an unexpected medical appointment. We refer readers to section X.C.2. of the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68688 through 68695) for a full discussion of this requirement.

For CY 2009, we proposed to calculate the payment rates for PHP APCs 0172 and 0173 using both hospital-based and CMHC PHP data (73 FR 41513). After consideration of the public comments received on our proposal, we decided to base payment rates for the two-tiered approach on hospital-based PHP data only. As we explained in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68689), using the CMHC data for CY

2009 would have significantly reduced the CY 2009 PHP rates and negatively impacted hospital-based PHPs. Because hospital-based PHPs are geographically diverse, whereas CMHCs are located in only a few States, we were concerned that a significant drop in the rate could result in hospital-based PHPs closing and lead to possible beneficiary access to care problems. To calculate the CY 2009 PHP payment rate for APC 0172, we used the median per diem cost for hospital-based PHP days with 3 units of service to derive a PHP payment rate of \$157. For APC 0173, we used the median per diem cost for hospital-based PHP days with 4 or more units of service to derive a CY 2009 PHP payment rate of \$200.

In addition, for CY 2009, we finalized our policy to deny payment for any PHP claims for days when fewer than 3 units of therapeutic services are provided. As noted in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68694), we believe that 3 units of service should be the minimum number of services allowed in a PHP day because a day with 1 or 2 units of service does not meet the statutory intent of a PHP program. Three units of service are a minimum threshold that permits unforeseen circumstances, such as medical appointments, while allowing payment, but maintains the integrity of the PHP benefit.

Further, for CY 2009, we revised the regulations at § 410.43 to codify existing basic PHP patient eligibility criteria and added a reference to current physician certification requirements at § 424.24. We believed these changes would help strengthen the PHP benefit by conforming our regulations to our longstanding policy (73 FR 68694 through 68695). Specifically, we revised § 410.43 to add a reference to existing regulations at § 424.24(e) that require that PHP services be furnished pursuant to a physician certification and plan of care. While the requirements at § 424.24(e) are not new, we included the reference in § 410.43 to provide a more complete description of our expectations for PHP programs in one regulatory section. We also revised § 410.43 to add the following patient eligibility criteria and clarify that PHPs are intended for patients who—(1) require a minimum of 20 hours per week of therapeutic services as evidenced in their plan of care; (2) are likely to benefit from a coordinated program of services and require more than isolated sessions of outpatient treatment; (3) do not require 24-hour care; (4) have an adequate support system while not actively engaged in the program; (5) have a mental health

diagnosis; (6) are not judged to be dangerous to self or others; and (7) have the cognitive and emotional ability to participate in the active treatment process and can tolerate the intensity of the PHP. We refer readers to section X.C.2. of the CY 2009 OPPS/ASC final rule with comment period (73 FR 68694 through 68695) for a full discussion of this requirement.

Lastly, in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68695 through 68697), we revised the partial hospitalization benefit to include several coding updates. We removed three PHP billable codes (CPT codes 90899 (Unlisted psychiatric service or procedure), 90853 (Group psychotherapy other than of a multiple-family group), and 90857 (Interactive group psychotherapy)), and created two new timed HCPCS codes (G0410 (Group psychotherapy other than of a multiple-family group, in a partial hospitalization

setting, approximately 45 to 50 minutes) and G0411 (Interactive group psychotherapy in a partial hospitalization setting, approximately 45 to 50 minutes)). The elimination of CPT code 90899 was a result of our concerns about the type of services that may be billed using an unlisted CPT code when a more appropriate code may be available that better describes the services for which PHP payment may be made. The decision to eliminate the two group therapy CPT codes (90853 and 90857) and replace them with two new parallel timed HCPCS G-codes (G-0410 and G-0411) was based on the need for consistency. As most of the current PHP codes already include time estimates, we wanted to maintain consistency with the existing HCPCS codes used in the PHP by applying a time descriptor to the group therapy codes. In addition to these coding updates, we also decided

to eliminate CPT code 90849 (multi-family group psychotherapy) as a billable PHP code because we believed that CPT code 90849 focuses the service on the needs of the family and not specifically on the needs of the patient, which is not consistent with the intent of the statute that treatment in a PHP be focused on the patient's condition (73 FR 68696).

B. PHP APC Update for CY 2010

For the CY 2010 OPPS/ASC proposed rule (74 FR 35356), we used CY 2008 claims data and computed median per diem costs in the following three categories: (1) All days; (2) days with 3 units of service; and (3) days with 4 or more units of service. These updated median per diem costs were computed separately for CMHCs and hospital-based PHPs and are shown in Table 50 below.

TABLE 50—PHP MEDIAN PER DIEM COSTS FOR CMHCs AND HOSPITAL-BASED PHPs, BY CATEGORY, BASED ON CY 2008 CLAIMS DATA

	CMHCs	Hospital-based PHPs	Combined
All Days	\$140	\$200	\$144
Days with 3 units of service	129	149	131
Days with 4 units or more units of service	173	213	175

Using CY 2008 data and the refined methodology for computing PHP per diem costs that we adopted in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66672), we computed the median per diem cost from all claims of \$144. The data indicate that CMHCs continue to provide far fewer days with 4 or more units of service (33 percent compared to 70 percent for hospital-based PHPs) and that the CMHC median per diem cost for 4 or more units of service (\$173) is substantially lower than the comparable data from hospital-based PHPs (\$213). The median per diem cost for claims containing 4 or more units of service for

all PHP claims, regardless of site of service, is \$175. Medians for claims containing 3 units of service are \$129 for CMHCs, \$149 for hospital-based PHPs, and \$131 for all PHP claims, regardless of site of service. In the CY 2010 OPPS/ASC proposed rule (74 FR 35356), for CY 2010, we proposed to continue to use the two-tiered payment approach for PHP services established in CY 2009. In addition, for CY 2010, we proposed to use only hospital-based PHP data to develop the two PHP APC per diem payment rates for the following reasons. If we used combined CMHC and hospital-based PHP data to develop the

rates, the two per diem payment rates would be reduced by approximately \$26 for APC 0172 and \$25 for APC 0173. We are concerned about further reducing both PHP APC per diem payment rates without knowing the impact of the policy and payment changes we made in CY 2009. Because there is a 2-year delay between data collection and rulemaking, the changes we made in CY 2009 will not be reflected in the claims data until next year when we are developing the update for CY 2011. The two proposed APCs median per diem rates for PHP services were as follows:

TABLE 51—CY 2010 PROPOSED MEDIAN PER DIEM RATES FOR PHP SERVICES

Proposed APC	Group title	Proposed median per diem rate
0172	Level I Partial Hospitalization (3 services)	\$149
0173	Level II Partial Hospitalization (4 or more services)	213

In general, public commenters supported the two-tiered PHP APC per diem payment approach in the proposed rule, the proposed CY 2010 payment rates, and the use of hospital-based PHP

data only for generating the PHP APC payment rates. *Comment:* A majority of the commenters supported the proposed PHP rates for CY 2010, as well as the two-tiered PHP APC payment structure

(with high and low intensity rates). Many of these commenters strongly recommended that CMS use only hospital-based PHP data to determine the final rates. The commenters believed

that hospital-based data are more reliable, predictable, national in scope, consistent, and stable, and that hospital-based data are meeting the intent of the PHP statute and CMS rules.

Several commenters urged CMS not to combine the CMHC and hospital-based PHP data to set the final CY 2010 PHP payment rates. The commenters pointed out that the volatility and significant fluctuation of the CMHC costs and changes in data since 2000 has continued to place the reliability of the CMHC data in doubt. One commenter urged CMS to use PHP hospital-based data only to set the rates because CMS does not know the impact of the comprehensive policy and payment changes made to PHP services during 2009.

The commenters also recommended that, if CMS were to change the methodology to establish the per diem payment rate in CY 2010 and beyond, CMS adopt two additional APCs for separate CMHCs PHP payment rates. The commenter recommended establishing site specific APCs for PHP services where the hospital-based PHP APCs for Level I (3 units of service) and Level II (4 or more units of service) would be established using hospital cost data and CMHC-based PHP APCs for Level I (3 units of service) and Level II (4 or more units of service) would be established using CMHC data. One commenter pointed out that while the aggregate number of PHP service providers has remained relatively stable over time, the number of hospital-based PHPs has dropped by 16 percent, while the number of CMHC PHPs has increased by 53 percent (with the majority of CMHCs located in Florida, Louisiana, and Texas). The commenters reported that 80 percent of the States have two or more hospital-based programs, and only 30 percent of States have more than one CMHC.

Response: We appreciate the commenters' support for our proposals on the two-tiered payment approach and use of hospital-based PHP data only to develop the CY 2010 payment rates. As we continue to evaluate ways to reflect CMHC costs in establishing PHP future rates, we will take the recommendation to establish site-specific PHP APCs into consideration. After consideration of the public comments we received, we have decided to retain the two-tiered payment approach for CY 2010, using hospital-based PHP data only.

Comment: Several commenters urged CMS to propose an APC code or payment rate for PHP claims for days with fewer than 3 units of service or, at a minimum, the commenters want CMS

to suspend for medical review claims with fewer than units of services per day.

Response: We continue to believe that days with 1 or 2 units of service are inconsistent with a benefit designed as a full-day program to substitute for inpatient care. PHP is furnished in lieu of an inpatient psychiatric hospitalization and is intended to be a highly structured and clinically intense program, usually lasting most of the day. We believe that 3 units of service should be the minimum number of services allowed in a PHP day because a day with 1 or 2 units of service does not meet the statutory intent of a PHP program. Our intention was to provide payment to cover days that consisted of 3 units of service only in certain limited circumstances. Therefore, we believe that 3 units of service are a minimum threshold that permits unforeseen circumstances, such as medical appointments, while allowing payment, but still maintains the integrity of a comprehensive program. If there are legitimate instances when 1 or 2 units of service days are justified, the provider has the option of appealing the denial of payment for that day, as specified in the Medicare Claims Processing Manual (Pub. 100-04), Chapter 29 and Chapter 30, Section 30.2.2.

Comment: One commenter stated that the number of rural hospital-based PHPs has declined during the CY 2003–2006 period and indicated that a study conducted last year found that rural areas are being hit with the loss of PHP programs. A few commenters were greatly relieved by the projection of an increase in the reimbursement rate for Level II PHP services. They believed another cut in rates would jeopardize the existence of the PHP benefit, reduce the financial viability of PHPs, and probably lead to closure of many PHPs, thus affecting access to care for this vulnerable population. In addition, because hospital outpatient mental health services paid under the OPSS are capped at the PHP per diem payment rate, many commenters were concerned about overall access to outpatient mental health treatment. The commenters urged CMS to keep mental health services available to all.

Response: We have established the final CY 2010 payment rate based on hospital-based PHP data, yielding an increase in the median for days with 4 or more units of service compared to CY 2009 payment rates. This increase will benefit all PHP programs, including those in rural areas. The CY 2010 payment rates for Level I (3 units of service) shows a decrease in CY 2010

compared to CY 2009. We believe using the CMHC data would significantly reduce the current rate and negatively impact hospital-based PHPs and CMHCs, resulting possibly in reduced access to care. Because hospital-based PHPs are geographically diverse, whereas CMHCs are located in only a few states, we are concerned that a significant drop in the rate could result in hospital-based PHPs closing and leading to possible access problems. For this reason, we are using hospital-based PHP data only to calculate the CY 2010 median per diem payment rates.

Comment: One commenter expressed concern that CMS' data and analysis regarding the median per diem costs for PHP services have been and remain fundamentally flawed. The commenter recommended that CMS conduct further research and, in particular, conduct detailed provider-level research to better understand the costs necessary to deliver PHP services in hospital and CMHC settings. The commenter recommended the payment rate for PHP services (APC 0173) be set at approximately \$325 per patient day and no less than the CY 2007 payment rate of \$234.73.

Response: We base the PHP APC per diem payment rates on providers' charges reported on claims adjusted by the providers' CCRs. This approach is consistent with the method used to compute payment rates for other APCs under the OPSS, except that, for PHPs for which the unit of service is a day, we sum the charges for a given day and then determine the median cost of all days. We expect that a provider's charges will reflect the level of services provided, which has a relationship to the cost of providing those services. In Medicare cost reporting, the total charges are to be reported along with the provider's cost. To the extent that a provider is submitting bills that have charges that do not directly relate to the delivery or provision of services, its CCRs will be unpredictable and would distort the costs of the services provided.

In developing the CY 2010 PHP APC per diem payment rates, we excluded days that have only 1 or 2 units of service. In addition, we did not include days where no payment was made to avoid diluting the cost. To calculate the Level I PHP APC payment rate, we used days with 3 units of service, and to calculate the Level II PHP APC payment rate, we used days with 4 or more units of service. We believe our methodology accurately reflects the median cost of providing these two levels of PHP services. As discussed previously, we made several refinements to our

methodology for computing the per diem costs that more accurately reflect the per diem cost of providing PHP services.

Comment: Many commenters stated that cost report information for CMHCs is not currently included in the Healthcare Cost Report Information System (HCRIS) and recommended that CMS base its calculations only on the cost report information that the agency can verify directly and not on data provided by the fiscal intermediaries or MACs. The commenters believed that CMS should calculate payment rates using only cost data from those cost reports currently in and accessible through the HCRIS.

Response: We understand the commenters concern about making CMHC data available through the HCRIS, and we are taking steps to make the data available in the future. For CY 2010, we will use PHP hospital-based data only to set the PHP APC payment rates. The hospital-based PHP data are based on cost report data currently in and accessible through the HCRIS.

Comment: Several commenters expressed concern that there are additional services furnished by CMHCs that are currently provided to PHP patients for which the providers are not reimbursed. The commenters pointed out that the Substance Abuse and Mental Health Services Administration recognizes these additional services by including payment for treatment of mental illness, not just for substance abuse treatment, and for costs for other services, including locating housing. The commenter included a list of HCPCS H-codes as an example of additional services as specified in Table 52 below.

TABLE 52—ADDITIONAL HCPCS H-CODES RECOMMENDED BY A COMMENTER FOR PAYMENT AS PHP SERVICES

HCPCS H-code	Description
H0001	Alcohol and/or drug assessment.
H0004	Behavioral health counseling and therapy, per 15 minutes.
H0028	Alcohol and/or drug prevention problem identification and referral service (e.g., student assistance and employee assistance programs), does not include assessment.

TABLE 52—ADDITIONAL HCPCS H-CODES RECOMMENDED BY A COMMENTER FOR PAYMENT AS PHP SERVICES—Continued

HCPCS H-code	Description
H0029	Alcohol and/or drug prevention alternatives service (services for populations that exclude alcohol and other drug use, e.g. alcohol free social events).
H0030	Behavioral health hotline service.
H0031	Mental health assessment, by non-physician.
H0032	Mental health service plan development by non-physician.
H0033	Oral medication administration, direct observation.
H0034	Medication training and support, per 15 minutes.
H0047	Alcohol and/or other drug abuse services, not otherwise specified.
H0049	Alcohol and/or drug screening.
H0050	Alcohol and/or drug services, brief intervention, per 15 minutes.
H1011	Family assessment by licensed behavioral health professional for state defined purposes.
H2000	Comprehensive multidisciplinary evaluation.
H2010	Comprehensive medication services, per 15 minutes.
H2011	Crisis intervention service, per 15 minutes.
H2014	Skills training and development, per 15 minutes.
H2027	Psycho educational service, per 15 minutes.

Response: Partial hospitalization services are specifically defined in section 1861(ff) of the Act and are a Medicare benefit category. Because there is no benefit category for substance abuse programs, any such program would have to meet requirements established for PHPs, including the requirements that a physician certify that the patient would otherwise require inpatient psychiatric care in the absence of the partial hospitalization services and that the program provides active treatment (section 1835(a)(2)(F) of the Act and 42 CFR 424.27(e)). PHP services involving direct patient care costs are payable under Medicare. The HCPCS H-codes listed above are not payable by Medicare. However, certain services for substance abuse are payable under a PHP because a PHP provides for patient education, mental health assessment, occupational therapy, and behavioral

health treatment/services among other services. For a complete list of services covered under a PHP under Medicare, we refer readers to the Medicare Claims Processing Manual (Pub. 100-04), Chapter 4, Section 260. Medicare does not pay for services such as 12-step programs. Many of the HCPCS H-codes listed duplicate allowable PHP service codes, for example, patient education and training. However, the PHP service codes generally are not defined in 15-minute increments.

Comment: One commenter suggested that CMS consider alternative arrangements for partial hospitalization and hospital outpatient services for the future. The commenter suggested removing PHP from the APC codes and instead establish a separate payment system (similar to home health) by establishing a reasonable base rate for PHP for Level II PHP services at a slightly higher level (such as \$220–\$225 per day), annually adjusting the base rate by a conservative inflation factor such as the CPI, establishing quality criteria to judge performance, and dropping the payment level for Level I PHP services so that only 4 or more services are recognized for payment.

Response: Currently, the statute does not provide for a separate payment system for partial hospitalization services. Therefore, a statutory change would be required to establish an independent payment system for PHPs. Regarding the commenter's recommendation to establish quality criteria, we agree that establishing benchmarks and indicators would be useful, and we encourage providers to share that information with us. We believe that creating a rate specific to days with three services is consistent with our policy to require CMHCs and hospital-based PHPs to provide a minimum of 3 units of service per day in order to receive payment. Although we do not expect Level I PHP service days to be frequent, we do recognize that there are times when a patient may need a less intensive day of service. Therefore, we continue to recognize the need for a two-tiered payment system: one payment for those less intensive days with three services; and another payment for those more intensive days with four or more services.

Comment: A few commenters urged CMS to find a way to strengthen the integrity of the program by developing and implementing standards of participation. The commenters recommended the implementation of standards of care with emphasis on quality of services and urged CMS to develop conditions of participation that would be a useful regulatory tool for

PHP providers. The commenters suggested that the development process must include all providers and other stakeholders. Several commenters offered to work with CMS to reform and improve the PHP by establishing standards for quality of PHP services provided; adopting CMHC facility-level quality measures and a reporting regime; and assisting with accreditation and cost reporting reforms.

Response: We agree with the commenters that standards of participation is an area that should be addressed, and we are exploring proposing conditions for coverage for CMHCs to establish minimum standards for patient rights, physical environment, staffing, and documentation requirements. We believe that adding conditions for coverage would contribute to more consistency between CMHCs and hospital-based PHPs.

Comment: One commenter suggested that CMS use fiscal intermediaries and MACs to work with hospitals and CMHC providers to establish separate PHP lines on their appropriate Medicare cost reports to arrive at a CCR for PHPs rather than the default psychiatric, clinics, or overall outpatient CCR lines. The commenter believed that, nationally, the CCRs for the PHPs are being understated by applying overall CCRs and/or clinic CCRs and, thus, penalizing the structured intensive partial programs.

Response: We note that most hospitals do not have a cost center for partial hospitalization; therefore, we have used the CCR as specific to PHP as possible. As described earlier in section X.A. of this final rule with comment period, for CY 2008, we proposed and finalized two refinements to the methodology for computing the PHP median per diem cost that we believe resulted in a more accurate median per diem cost. The first of the two CY 2008 refinements was a remapping of 10 revenue codes-to-cost centers for hospital-based PHP claims. We believe that the CY 2008 refinement

to the mapping approach continues to be the best method for assigning the most appropriate cost center for hospital-based PHP claims. For a detailed explanation of the remapping of revenue codes for hospital-based PHP claims, we refer readers to the CY 2008 OPPS/ASC proposed rule (72 FR 42691 through 42692) and the CY 2008 OPPS/ASC final rule with comment period (72 FR 66671 through 66672).

In addition, we note that this remapping refinement applies only to hospital-based PHP claims and not to CMHC claims. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68690), in response to commenters' request that CMS apply the remapping of revenue codes to cost centers to CMHC claims, we stated that we cannot apply the same mapping method to CMHCs because PHP is the CMHCs' only Medicare cost and CMHCs do not have the same cost structure as hospitals.

Comment: One commenter expressed concern that the PHP benefit lacked flexibility. The commenter believed that the rigid guidelines for attendance of 5 plus days a week (20 hours) could create excessive overutilization at times. The commenter stated that it would be more beneficial to restructure PHP to be a more flexible, less costly, outcome-based system of care.

Response: Partial hospitalization is an intensive outpatient program of psychiatric services provided to patients as an alternative to inpatient psychiatric hospitalization or as a step down to shorten an inpatient stay and transition a patient to a less intensive level of care. We understand the commenters' concerns about the 20 hours per week requirement with regard to scheduling flexibility, but we were concerned that if we reduce the minimum number of hours lower than the current guideline, the low end of the range will become the new minimum. Therefore, instead of reducing the number of hours a patient needs in order to be eligible to receive

the benefit, we clarified that the patient eligibility requirement that patients require 20 hours of therapeutic services is evidenced in a patient's plan of care rather than in the actual hours of therapeutic services a patient receives. The intent of this eligibility requirement is that, for most weeks, we expect attendance conforming to the patient's plan of care. We recognize that there may be times at the beginning (or end) of a patient's transition into (or out of) a PHP where the patient may not receive 20 hours of therapeutic services.

Comment: One commenter expressed concern about the proposed cuts for Partial Hospitalization (APC 0173) and the impact the cuts will have on its hospital and community. The commenter reviewed the history of APC payment rates for these services and noted the trend of the payment rates decreasing over the past 7 years. The commenter pointed out that it experienced a significant increase in the staff salary and benefit costs. The commenter expressed concern that the decreasing payment rate and increasing expenses will make it difficult for the hospital to sustain these services and access to care for Medicare beneficiaries could worsen.

Response: Hospital costs per day for PHP services for APC 0173 have remained in the range of \$200—\$225 for CY 2000 through CY 2010. This is the reason we have decided to use hospital-based PHP data only for computing the CY 2010 PHP payment rates, as this approach will lead to payment stability for CY 2010.

In summary, after consideration of the public comments we received, we are adopting as final our CY 2010 proposal to retain the two-tiered payment approach for PHP services and to use only hospital-based PHP data in computing the payment. The two updated PHP APC per diem median costs that we are finalizing for CY 2010 are shown in Table 53 below.

TABLE 53—CY 2010 PHP APC PER DIEM MEDIAN COSTS

APC	Group title	Median per diem costs
0172	Level I Partial Hospitalization (3 services)	\$148
0173	Level II Partial Hospitalization (4 or more services)	209

C. Separate Threshold for Outlier Payments to CMHCs

In the November 7, 2003 final rule with comment period (68 FR 63469), we indicated that, given the difference in PHP charges between hospitals and

CMHCs, we did not believe it was appropriate to make outlier payments to CMHCs using the outlier percentage target amount and threshold established for hospitals. Prior to that time, there was a significant difference in the amount of outlier payments made to

hospitals and CMHCs for PHP services. In addition, further analysis indicated that using the same OPPS outlier threshold for both hospitals and CMHCs did not limit outlier payments to high cost cases and resulted in excessive outlier payments to CMHCs. Therefore,

beginning in CY 2004, we established a separate outlier threshold for CMHCs. The separate outlier threshold for CMHCs has resulted in more commensurate outlier payments.

In CY 2004, the separate outlier threshold for CMHCs resulted in \$1.8 million in outlier payments to CMHCs. In CY 2005, the separate outlier threshold for CMHCs resulted in \$0.5 million in outlier payments to CMHCs. In contrast, in CY 2003, more than \$30 million was paid to CMHCs in outlier payments. We believe this difference in outlier payments indicates that the separate outlier threshold for CMHCs has been successful in keeping outlier payments to CMHCs in line with the percentage of OPPS payments made to CMHCs. Table 54 below includes a listing of the outlier target amounts and the portion of the target amount allocated to CMHCs for PHP outliers for CYs 2004 through 2009.

TABLE 54—OUTLIER TARGET AMOUNT PERCENTAGES AND PORTIONS ALLOCATED TO CMHCs FOR PHP OUTLIERS—CY 2004 THROUGH CY 2007

Year	Outlier target amount percentage	Portion of target amount allocated to CMHCs for PHP outliers (in percent)
CY 2004	2.0	0.5
CY 2005	2.0	0.6
CY 2006	1.0	0.6
CY 2007	1.0	0.15
CY 2008	1.0	0.02
CY 2009	1.0	0.12

As noted in section II.F. of the CY 2010 OPPS/ASC proposed rule (74 FR 35296), for CY 2010, we proposed to continue our policy of identifying 1.0 percent of the aggregate total payments under the OPPS for outlier payments. We proposed that a portion of that 1.0 percent, an amount equal to 0.02 percent of outlier payments (or 0.0002 percent of total OPPS payments), would be allocated to CMHCs for PHP outliers. As discussed in section II.F. of the CY 2010 OPPS/ASC proposed rule (74 FR 35296), we proposed to set a dollar threshold in addition to an APC multiplier threshold for OPPS outlier payments. However, because the PHP APC is the only APC for which CMHCs may receive payment under the OPPS, we would not expect to redirect outlier payments by imposing a dollar threshold. Therefore, we did not propose to set a dollar threshold for CMHC outliers. As noted in section II.F. of the CY 2010 OPPS/ASC proposed

rule (74 FR 35296), we proposed to set the outlier threshold for CMHCs for CY 2010 at 3.40 times the APC payment amount and the CY 2010 outlier payment percentage applicable to costs in excess of the threshold at 50 percent. Specifically, we proposed that if a CMHC's cost for partial hospitalization services, paid under either APC 0172 or APC 0173, exceeds 3.40 times the payment for APC 0173, the outlier payment would be calculated as 50 percent of the amount by which the cost exceeds 3.40 times the APC 0173 payment rate.

Comment: Several commenters suggested that instead of creating separate threshold for outlier payments to CMHCs, it would be beneficial to eliminate the outlier payment and use those funds allocated to outlier payments to bolster payments for services provided by CMHCs.

Response: We note that the Secretary shall provide an outlier payment for each covered OPD service (or group of services) in accordance with the requirements set forth in section 1833(t)(5) of the Act and the applicable regulations. Because CMHCs are a provider of PHP services, outlier payments must be provided for them in accordance with the statute. We note that eliminating outlier payments for CMHCs would not result in an increase in the PHP rate, but rather would provide additional funding for hospital outpatient payments for all HOPD services.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal to set a separate outlier threshold for CMHCs. As discussed in section II.F. of this final rule with comment period, using more recent data for this final rule with comment period, we set the target for hospital outpatient outlier payments at 1.0 percent of total estimated OPPS payments. We allocated a portion of that 1.0 percent, an amount equal to 0.03 percent of outlier payments and 0.0003 percent of total estimated OPPS payments to CMHCs for PHP outliers. For CY 2010, as proposed, we are setting the outlier threshold at 3.40 multiplied by the APC amount and CY 2010 outlier percentage applicable to costs in excess of the threshold at 50 percent.

XI. Procedures That Will Be Paid Only as Inpatient Procedures

A. Background

Section 1833(t)(1)(B)(i) of the Act gives the Secretary broad authority to determine the services to be covered and paid for under the OPPS. Before implementation of the OPPS in August 2000, Medicare paid reasonable costs for

services provided in the HOPD. The claims submitted were subject to medical review by the fiscal intermediaries to determine the appropriateness of providing certain services in the outpatient setting. We did not specify in our regulations those services that were appropriate to provide only in the inpatient setting and that, therefore, should be payable only when provided in that setting.

In the April 7, 2000 final rule with comment period (65 FR 18455), we identified procedures that are typically provided only in an inpatient setting and, therefore, would not be paid by Medicare under the OPPS. These procedures comprise what is referred to as the "inpatient list." The inpatient list specifies those services for which the hospital will be paid only when provided in the inpatient setting because of the nature of the procedure, the underlying physical condition of the patient, or the need for at least 24 hours of postoperative recovery time or monitoring before the patient can be safely discharged. As we discussed in that rule and in the November 30, 2001 final rule with comment period (66 FR 59856), we may use any of a number of criteria we have specified when reviewing procedures to determine whether or not they should be removed from the inpatient list and assigned to an APC group for payment under the OPPS when provided in the hospital outpatient setting. Those criteria include the following:

- Most outpatient departments are equipped to provide the services to the Medicare population.
- The simplest procedure described by the code may be performed in most outpatient departments.
- The procedure is related to codes that we have already removed from the inpatient list.

In the November 1, 2002 final rule with comment period (67 FR 66741), we added the following criteria for use in reviewing procedures to determine whether they should be removed from the inpatient list and assigned to an APC group for payment under the OPPS:

- A determination is made that the procedure is being performed in numerous hospitals on an outpatient basis; or
- A determination is made that the procedure can be appropriately and safely performed in an ASC, and is on the list of approved ASC procedures or has been proposed by us for addition to the ASC list.

The list of codes that we proposed to be paid by Medicare in CY 2010 only as inpatient procedures was included as

Addendum E to the CY 2010 OP/ASC proposed rule.

B. Changes to the Inpatient List

In the CY 2010 OP/ASC proposed rule (74 FR 35358), we proposed to use for the CY 2010 OP/ASC the same methodology as described in the November 15, 2004 final rule with comment period (69 FR 65835) to identify a subset of procedures currently on the inpatient list that are being performed a significant amount of the time on an outpatient basis. Using this methodology, we identified three procedures that met the criteria for potential removal from the inpatient list. We then clinically reviewed these three potential procedures for possible removal from the inpatient list and found them to be appropriate candidates for removal from the inpatient list. During the February 2009 meeting of the APC Panel, we solicited the APC Panel's input on the appropriateness of removing the following three procedures from the CY 2010 inpatient list: CPT codes 21256 (Reconstruction of orbit with osteotomies (extracranial) and with bone grafts (includes obtaining autografts) (eg, micro-ophthalmia)); 27179 (Open treatment of slipped femoral epiphysis; osteoplasty of femoral neck (Heyman type procedure)); and 51060 (Transvesical ureterolithotomy).

In addition to presenting to the APC Panel the three procedures above, we also presented utilization data for the first 9 months of CY 2008 for two other specific procedures, in response to a request by the APC Panel from the March 2008 meeting: CPT code 20660 (Application of cranial tongs, caliper or stereotactic frame, including removal (separate procedure)), a procedure that we removed from the inpatient list for CY 2009; and CPT code 64818 (Sympathectomy, lumbar), a procedure that we maintained on the inpatient list for CY 2009.

Following the discussion at the February 2009 meeting, the APC Panel recommended that CMS remove from the CY 2010 inpatient list CPT codes 21256, 27179, and 51060. The APC Panel also recommended that CPT code 64818 remain on the inpatient list for CY 2010. The APC Panel made no recommendation regarding CPT code 20660.

In the CY 2010 OP/ASC proposed rule (74 FR 35358), we proposed to accept the APC Panel's recommendations to remove the procedures described by CPT codes 21256, 27179, and 51060 from the inpatient list because we agree with the APC Panel that the procedures may be

appropriately provided as hospital outpatient procedures for some Medicare beneficiaries. We also proposed to retain CPT code 64818 on the inpatient list because we agree with the APC Panel that this procedure should be provided to Medicare beneficiaries only in the hospital inpatient setting. The three procedures we proposed to remove from the inpatient list for CY 2010 and their CPT codes, long descriptors, and proposed APC assignments were displayed in Table 37 of the CY 2010 OP/ASC proposed rule (74 FR 35358).

Comment: Several commenters supported the proposal to remove the procedures reported by CPT codes 21256, 27179, and 51060 from the inpatient list, one commenter opposed removing the procedure coded by CPT code 51060, and one commenter expressed concern about removing any of the three procedures proposed for removal from the inpatient list. The commenter that requested that CMS retain CPT code 51060 on the inpatient list reported that the procedure is an open surgical procedure that is much more extensive than a hernia repair. In that commenter's experience, patients who undergo this surgery are not able to go home the same day as surgery because most require parenteral pain medication and ongoing monitoring of the pain, and of possible ileus or hematuria. Another commenter provided no rationale for objecting to the removal of the three procedures as proposed by CMS beyond stating that if CMS does not eliminate the entire inpatient list, then the commenter would have serious concerns about removing the three procedures.

Response: We appreciate the commenters' support of our proposal. In response to the commenters who expressed concern about removing one or more of the three procedures from the inpatient list, we reevaluated the three procedures using more recent utilization data and further clinical review by CMS medical advisors. As a result of that reevaluation, we remain convinced that all three procedures may be safely performed in the HOPD for some Medicare beneficiaries. As we have indicated previously, the removal of a procedure from the inpatient list does not signify a determination by CMS that the procedure should be performed in the HOPD for all beneficiaries. The removal only indicated that CMS is relying on the individual beneficiary's surgeon to advise the most appropriate setting for the procedure based on the beneficiary's medical condition. In fact, as evidenced by the utilization data over the years, most of the newly-removed

procedures from the inpatient list continue to be commonly provided in the inpatient setting after they are removed from the list. The removal of a procedure from the inpatient list simply is recognition that there is evidence that the procedure may be safely performed for some beneficiaries who are outpatients and represents no directive about whether the inpatient setting or the outpatient setting is more appropriate in any particular circumstance.

Comment: Several commenters requested the removal of additional procedures from the inpatient list. Although the commenters requested that CMS remove a total of 20 additional procedures from the inpatient list, 4 of the requested codes were not on the proposed CY 2010 inpatient list. All of the codes requested for removal are displayed in Table 55 below.

As identified by asterisks, 11 of the procedures displayed in the chart below were submitted by one commenter representing a group of hospitals. This commenter stated that each of the procedures requested for removal from the inpatient list was carefully reviewed and could be safely provided to Medicare beneficiaries in the HOPD. The commenter reported that research and investigation indicated that clinical criteria sets such as the *Milliman Care Guidelines* support the safe provision of the 11 procedures in outpatient settings. In addition, the commenter stated that the group's hospitals have physicians providing the procedures safely in the outpatient setting for non-Medicare patients who are in the same age group as the Medicare population.

TABLE 55—ADDITIONAL PROCEDURES REQUESTED BY COMMENTERS FOR REMOVAL FROM THE INPATIENT LIST FOR CY 2010

CY 2009 HCPCS code	CY 2009 short descriptor	Proposed CY 2010 SI
01402	Anesth, knee arthroplasty.	C
22548	Neck spine fusion	C
*22554	Neck spine fusion	C
*22585	Additional spinal fusion.	C
*22851	Apply spine prosth device.	T
27447	Total knee arthroplasty.	C
28805	Amputation thru metatarsal.	C
*32662	Thoracoscopy, surgical.	C
*37182	Insert hepatic shunt (tips).	C
*37215	Transcath stent, cca w/eps.	C

TABLE 55—ADDITIONAL PROCEDURES REQUESTED BY COMMENTERS FOR REMOVAL FROM THE INPATIENT LIST FOR CY 2010—Continued

CY 2009 HCPCS code	CY 2009 short descriptor	Proposed CY 2010 SI
*44950	Appendectomy	C
44955	Appendectomy, add-on.	C
44960	Appendectomy	C
55866	Laparo radical prostatectomy.	C
*60505	Explore parathyroid glands.	C
*63047	Removal spinal lamina.	T
63075	Neck spine disk surgery.	T
*63076	Neck spine disk surgery.	C
*63267	Excise intraspinal lesion.	C
64999	Nervous system surgery.	T

* Submitted by commenter representing a group of hospitals.

Response: In response to the commenters' requests, we reviewed utilization and clinical information for each of the procedures suggested for removal from the inpatient list. Of the 16 procedures reviewed (those with status indicator "C" in the chart above) our medical advisors agreed with the commenters that it would be appropriate to remove 5 of them from the inpatient list. Thus, for CY 2010, we are removing from the inpatient list the procedures reported by CPT codes 28805 (Amputation, foot; transmetatarsal); 37215 (Transcatheter placement of intravascular stent(s), cervical carotid artery, percutaneous; with distal embolic protection); 44950 (Appendectomy); 44955 (Appendectomy; when done for indicated purpose at time of other major procedure (not as separate procedure) (List separately in addition to code for primary procedure)); and 63076 (Discectomy, anterior, with decompression of spinal cord and/or nerve root(s), including osteophytectomy; cervical, each additional interspace (List separately in addition to code for primary procedure)). These procedures and the APCs to which they are assigned for CY 2010 are displayed in Table 56 below.

The clinical and utilization data for the other 11 procedures did not support the appropriateness of providing the procedures to Medicare beneficiaries in the HOPD. We believe that Medicare beneficiaries who undergo any of these 11 procedures should do so only as inpatients.

Comment: Many commenters suggested that CMS eliminate the inpatient list. They believed that each patient's status should be determined by the physician who can establish the most appropriate care plan for the individual. The commenters pointed out the many safety provisions that are met by hospitals participating in the Medicare program as evidence that hospitals would provide care safely and appropriately.

A few of the commenters stated that the inconsistency between the Medicare payment policies for hospitals and physicians of allowing physicians to receive full payment for inpatient procedures that are performed on Medicare beneficiaries in the HOPD who are not inpatients, but denying hospitals payment for those same procedures, gives physicians little incentive to avoid providing inpatient procedures to Medicare beneficiaries who are outpatients. Several commenters suggested that if CMS maintains the inpatient list, the associated payment restrictions be applied consistently to both hospitals and physicians in order to promote a collaborative effort in documentation and clinical care plans.

One commenter stated that the inpatient list, which requires that Medicare beneficiaries be handled differently than the rest of the patient population in some circumstances, creates chaos for the physicians and hospitals who are trying to apply consistent clinical criteria to determine appropriate levels of care for all of their patients.

Many of the commenters argued that there are a variety of circumstances that result in procedures on the inpatient list being performed without an inpatient admission and that hospitals should not be held accountable for those situations. For example, they explained that sometimes during the intraoperative period, due to clinical circumstances, the surgeon performs a procedure that is on the inpatient list in addition to, or rather than, the procedure that was planned.

Finally, the commenters believed that the inpatient list penalizes hospitals unfairly and suggested that if CMS is unwilling to eliminate the inpatient list, it consider developing an appeals process to address circumstances in which payment for a service provided on an outpatient basis is denied due to its presence on the inpatient list. The commenters believed that the appeal would give the hospital the opportunity to submit documentation on the physician's intent, the patient's clinical condition, and the circumstances that

enabled the patient to be sent home safely without an inpatient stay.

Response: While we understand the commenters' reasons for advocating the elimination of the inpatient list, we continue to believe that the inpatient list serves an important purpose in identifying procedures that cannot be safely and effectively provided to Medicare beneficiaries in the HOPD. We are concerned that the elimination of the inpatient list could result in unsafe or prolonged care in hospital outpatient departments for Medicare beneficiaries. Therefore, we are not discontinuing our use of the inpatient list at this time.

Although the commenters suggested that we apply the same payment restrictions to physicians and hospitals when inpatient procedures are performed inappropriately, payment for physicians' services are outside of the scope of OPSS payment policy and of this OPSS/ASC final rule with comment period. We continue to believe that it is very important for hospitals to educate physicians on Medicare services covered under the OPSS to avoid inadvertently providing services in a hospital outpatient setting that only are covered during an inpatient stay.

We also are concerned about the potential results of eliminating the inpatient list on Medicare beneficiary liability. For instance, we are concerned that, without the inpatient list, Medicare beneficiaries could experience longer stays in HOPDs after some procedures. The APC Panel has discussed its concern about these long stays that frequently exceed 24 hours. Moreover, the financial liability for OPSS copayments for complex surgical procedures and long periods of care in the HOPD and coverage of items such as usually self-administered drugs differs significantly from a beneficiary's inpatient cost-sharing responsibilities and coverage, and the beneficiary may incur higher out-of-pocket costs for prolonged outpatient encounters than for an inpatient stay for the same surgical intervention.

We continue to encourage physicians' awareness of the implications for Medicare beneficiaries and hospitals of performing inpatient list procedures in the HOPD on beneficiaries who are not inpatients. We do not plan to adopt a specific appeals process for claims related to inpatient list procedures performed in the HOPD at this time. The existing processes established for a beneficiary or a provider to appeal a specific claim remain in effect.

Comment: One commenter suggested that CMS implement a method to identify scheduled outpatient procedures that become, through

intraoperative circumstances, inpatient procedures. The commenter asserted that, due to hospital billing practices, hospital coding staff do not know until well after the surgery is completed that an unscheduled inpatient procedure was performed on an outpatient who was not admitted as an inpatient. The commenter suggested that CMS implement a HCPCS modifier that the hospital could append to the inpatient procedure on the claim and that payment for the claim could be made at the same rate as those coded with the -CA modifier (APC 0375 (Ancillary Outpatient Services When Patient Expires)).

Response: While we appreciate the commenter's suggestion for addressing circumstances when unplanned inpatient list procedures are performed during operative sessions where outpatient surgical procedures were planned, we do not believe there is a need for a modifier to identify those situations. We continue to believe that the inpatient list procedures are not appropriate for performance on Medicare beneficiaries in the HOPD and, therefore, we expect that when such a procedure is performed, the beneficiary would be admitted as an inpatient.

We established payment for ancillary services reported in association with an inpatient procedure to which the -CA modifier is appended in order to provide payment to hospitals for services furnished in those rare cases in which procedures on the inpatient list are performed to resuscitate or stabilize a patient (whose status is that of an outpatient) with an emergent, life-threatening condition, and the patient dies before being admitted as an inpatient. In these situations, hospitals are unable to admit the patients as inpatients. In the circumstances described by the commenter, we see no insurmountable hospital barriers to admitting the patients as inpatients of the hospital and do not believe it would be appropriate to provide payment (through APC 0375) for a mix of surgical procedures provided to patients who survive at the rate developed specifically to pay for the ancillary services furnished in association with procedures reported with the -CA modifier when a patient dies prior to admission as an inpatient. The calculation of the payment rate for APC 0375 is discussed in detail in section II.A.2.d.(7) of this final rule with comment period.

We understand hospitals' dilemma when the decision is made intraoperatively to perform an unscheduled procedure. However, we

continue to believe that it is important for hospitals to educate physicians on Medicare services paid under the OPDS to avoid inadvertently providing services in a hospital outpatient setting that would be paid only during an inpatient stay because we believe that the HOPD is not an appropriate site of service for inpatient list procedures.

Comment: Another commenter recommended that CMS expand the use of the -CA modifier to allow payments to the hospitals for performance of procedures on the inpatient list that must be performed to resuscitate or stabilize an outpatient with an emergency, life-threatening condition, but the patient is stabilized medically and transferred to another acute care hospital for admission. In other words, the commenter added, the patient is never admitted to the hospital where the inpatient list procedure was performed.

Response: We established the -CA modifier policy to provide payment to hospitals for services provided in the specific and rare situations in which procedures on the inpatient list are performed in the HOPD to resuscitate or stabilize a patient (whose status is that of an outpatient) with an emergent, life-threatening condition, and the patient dies before being admitted as an inpatient, not as a method for hospitals to recoup costs incurred when inpatient procedures are performed inappropriately on a Medicare beneficiary in the HOPD.

We see no rationale for allowing hospitals to report the -CA modifier for any circumstances other than those for which it was created. In the scenario described by the commenter, there is no evidence of the hospital's insurmountable barriers to admitting the patient, a criterion for use of the -CA modifier. In addition, we are not convinced that there is a need for a modifier to describe these rare events. We also do not believe it would be appropriate to provide payment at the rate developed specifically to pay for the ancillary services furnished in association with procedures reported with the -CA modifier when a patient dies prior to admission as an inpatient (APC 0375), for a mix of surgical procedures provided to patients who survive. The calculation of the payment rate for APC 0375 is discussed in detail in section II.A.2.d.(7) of this final rule with comment period.

Comment: One commenter suggested that CMS may lack information on the types of inpatient procedures that are performed for beneficiaries who are outpatients because those claims are returned to the provider by the I/OCE, thereby preventing CMS from capturing

the unpaid services in its claims data. To address this situation, the commenter recommended that CMS modify the I/OCE so that claims for inpatient procedures provided on an outpatient basis could be processed as not payable rather than returned to the provider. The commenter believed that this modification would enable CMS to gather claims data on these procedures and to see how many and what types of procedures physicians believe are appropriate for performance in the HOPD. The claims data also would provide CMS information about the hospital resources expended to care for these Medicare beneficiaries. In addition, the commenter suggested that CMS could share the data with the APC Panel for discussion and review in support of their evaluations of which procedures may appropriately be removed from the inpatient list.

Response: We believe that our outpatient claims data include the claims for inpatient procedures that are performed on Medicare beneficiaries who are outpatients. As would be expected, the volume for these nonpayable procedures is low compared to the number of payable outpatient claims, but we believe that the claims hospitals submit are available to us for examination in our OPDS claims data each year.

The I/OCE logic does not result in claims for inpatient procedures being returned to the provider. Rather, once the inpatient procedure is identified, it is line-item denied and then, with very few exceptions (for example, claims with the -CA modifier and an indication that the beneficiary expired), it assigns line-item edits to result in payment denial for each of the other services on the claim because these services were furnished on the same day as the inpatient procedure. A full description of the I/OCE logic and edits is available on the CMS Web site at: <http://www.cms.hhs.gov/OutpatientCodeEdit/>.

After consideration of the public comments we received, we are finalizing our proposal to remove the procedures reported by CPT codes 21256, 27179, and 51060 from the inpatient list. We also are removing five additional procedures that public commenters requested be removed from the inpatient list. These procedures are reported by CPT codes 28805 (Amputation, foot; transmetatarsal); 37215 (Transcatheter placement of intravascular stent(s), cervical carotid artery, percutaneous; with distal embolic protection); 44950 (Appendectomy); 44955 (Appendectomy; when done for

indicated purpose at time of other major procedure (not as separate procedure) (List separately in addition to code for primary procedure); and 63076 (Discectomy, anterior, with decompression of spinal cord and/or nerve root(s), including osteophyctomy; cervical, each additional interspace (List separately in addition to code for primary procedure)). The final eight procedures we are removing from the inpatient list for CY 2010 and their CPT codes, long descriptors, and final APC assignments are displayed in Table 56 below.

TABLE 56—PROCEDURES REMOVED FROM THE INPATIENT LIST AND THEIR FINAL APC ASSIGNMENTS FOR CY 2010

CY 2010 HCPCS Code	CY 2010 long descriptor	Final CY 2010 APC assignment	Final CY 2010 status indicator
21256	Reconstruction of orbit with osteotomies (extracranial) and with bone grafts (includes obtaining autografts) (eg, micro-ophthalmia).	0256	T
27179	Open treatment of slipped femoral epiphysis; osteoplasty of femoral neck (Heyman type procedure).	0052	T
28805	Amputation, foot; transmetatarsal	0055	T
37215	Transcatheter placement of intravascular stent(s), cervical carotid artery, percutaneous; with distal embolic protection.	0229	T
44950	Appendectomy	0153	T
44955	Appendectomy; when done for indicated purpose at time of other major procedure (not as separate procedure) (List separately in addition to code for primary procedure).	0153	T
51060	Transvesical ureterolithotomy	0163	T
63076	Discectomy, anterior, with decompression of spinal cord and/or nerve root(s), including osteophyctomy; cervical, each additional interspace (List separately in addition to code for primary procedure).	0208	T

XII. OPPTS Nonrecurring Technical and Policy Changes and Clarifications

A. Kidney Disease Education Services

1. Background

Section 152(b) of Public Law 110–275 (MIPPA) amended section 1861(s)(2) of the Act by adding a new subsection (EE) to provide for coverage of kidney disease education (KDE) services as a Medicare Part B benefit for Medicare beneficiaries diagnosed with stage IV chronic kidney disease (CKD) who, according to accepted clinical guidelines identified by the Secretary, will require dialysis or a kidney transplant, effective for services furnished on or after January 1, 2010. Section 152(b) also added a new subsection (ggg) to section 1861 of the Act to define “kidney disease education services” and to specify who may furnish these services as a “qualified person.” Section 1861(ggg)(2)(A)(i) of the Act, as added by section 152(b) of Public Law 110–275, defines a qualified person as a physician (as defined in section 1861(r)(1) of the Act); or a physician assistant, nurse practitioner, or clinical nurse specialist (as defined in section 1861(aa)(5) of the Act) who furnishes services for which payment may be made under the fee schedule established under section 1848 of the Act. Section 1861(ggg)(2)(A)(ii) of the Act also defines a qualified person as a “provider of services located in a rural area (as defined in section 1886(d)(2)(D) [of the Act]).” The definition of a “qualified person” for this benefit includes certain rural providers of

services, such as hospitals, critical access hospitals (CAHs), skilled nursing facilities (SNFs), home health agencies (HHAs), comprehensive outpatient rehabilitation facilities (CORFs), and hospices. Section 1861(ggg)(2)(B) of the Act provides that a qualified person does not include a provider of services (other than a provider of services described in section 1861(ggg)(2)(A)(ii) or a renal dialysis facility).

In the CY 2010 OPPTS/ASC proposed rule (74 FR 35358), we proposed to implement the provisions of section 1861(s)(2)(EE) and 1861(ggg) of the Act, as added by section 152(b) of Pub. L. 110–275, mainly through the June 2009 CY 2010 MPFS proposed rule (CMS–1413–P; Medicare Program; Payment Policies under the Physician Fee Schedule and Other Revisions to Part B for CY 2010), hereinafter referred to as the CY 2010 MPFS proposed rule. Specifically, in section II.G.10. of the CY 2010 MPFS proposed rule (74 FR 33617), we proposed to define the Medicare coverage criteria that would be applicable to KDE services and who may provide these services (that is, a “qualified person”), consistent with the provisions of sections 1861(s)(2)(EE) and 1861(ggg) of the Act. In that proposed rule, we also proposed to define a provider of services in a rural area as defined in section 1886(d)(2)(D) of the Act as a hospital, CAH, SNF, CORF, HHA, or hospice that is physically located in a rural area as defined in § 412.64(b)(ii)(C) of the regulations or a hospital or CAH that is reclassified from urban to rural status

pursuant to section 1886(d)(8)(E) of the Act, as defined in § 412.103 of the regulations. According to the proposal included in the CY 2010 MPFS proposed rule, a hospital, CAH, SNF, CORF, HHA, or hospice would not be considered to be a qualified person if the facility providing KDE services is located outside of a rural area unless the service is furnished in a hospital or CAH that has reclassified from urban to rural status under § 412.103.

In addition, in the CY 2010 MPFS proposed rule (74 FR 33614), consistent with the provisions of section 1861(ggg) of the Act, we proposed a payment amount for KDE services furnished by a “qualified person.” Specifically, we proposed to establish two new Level II HCPCS G-codes to describe KDE services and to specify the associated relative value units (RVUs) under the MPFS for payment for these codes.

We instructed individuals who wished to comment on the proposed coverage criteria for KDE services under section 1861(ggg) of the Act, including the definition of a “qualified person,” the proposed HCPCS G-codes, and the proposed RVUs for KDE services to submit their comments to CMS in response to the CY 2010 MPFS proposed rule that we describe above. Below we discuss our proposed payment for KDE services furnished by providers of services located in a rural area. We instructed individuals who wished to submit public comments relating to payment for KDE services furnished by providers of services located in a rural area to submit those

comments in response to the CY 2010 OPPS/ASC proposed rule.

2. Payment for Services Furnished by Providers of Services Located in a Rural Area

In the CY 2010 OPPS/ASC proposed rule (74 FR 35358), we proposed to pay under the MPFS for KDE services under section 1861(ggg) of the Act when the services are furnished by a qualified person that is a hospital, CAH, SNF, CORF, HHA, or hospice that is located in a rural area as defined in section 1886(d)(2)(D) of the Act or a hospital or CAH that is reclassified from urban to rural status pursuant to section 1886(d)(8)(E) of the Act, as defined in § 412.103 of the regulations. Section 152(b) of Public Law 110-275 amended section 1848(j)(3) of the Act to add section 1861(s)(2)(EE) (kidney disease education services) to the list of subsections of section 1861(s)(2) of the Act, which are included in the definition of physician services in section 1848(j)(3) of the Act. However, the statute does not specify the payment methodology for KDE services furnished by providers of these services located in rural areas.

Given that the statute provides the Secretary with the flexibility to pay all qualified persons under the MPFS and there is precedent for paying both diabetes self-management training and medical nutrition therapy services (which we believe KDE is similar in terms of resource use, specifically staffing and infrastructure) under the MPFS, we proposed to pay all qualified persons for KDE services under the MPFS. This single payment methodology would apply to all qualified persons, including providers of services in a rural area as we proposed to define such providers in the CY 2010 MPFS proposed rule.

The language in section 1861(ggg) of the Act that defines KDE services is similar to the language in section 1861(qq) of the Act that defines "diabetes self-management training services," which is a medical or other health service under section 1861(s)(2)(S) of the Act. In addition, the language in section 1861(ggg) of the Act is similar to the language in section 1861(vv) of the Act that defines "medical nutrition therapy services," which is also a medical or other health service under section 1861(s)(2)(V) of the Act. Finally, both diabetes self-management training and medical nutrition therapy are included in the definition of "physicians' services" for purposes of the MPFS at section 1848(j)(3) of the Act, and our standard policy is to pay for both services under

the MPFS when they are furnished in an HOPD. Given that the statute permits us to pay all qualified persons under the MPFS and the precedent for paying both diabetes self-management training and medical nutrition therapy under the MPFS when these services are provided in the hospital outpatient setting, we believe that payment under the MPFS is the most appropriate methodology for payment to qualified persons who are providers of services located in a rural area or who are hospitals or CAHs that have been reclassified from urban to rural status pursuant to § 412.103 of the regulations for the KDE services they furnish.

The proposed CY 2010 MPFS payments for HCPCS codes GXX26 (Educational services related to the care of chronic kidney disease; individual, per session; face-to-face), now finalized in the CY 2010 MPFS final rule with comment period as G0420 (Educational services related to the care of chronic kidney disease; individual per session, per hour, face-to-face), and GXX27 (Educational services related to the care of chronic kidney disease; group, per session; face-to-face), now finalized in the CY 2010 MPFS final rule with comment period as G0421 (Educational services related to the care of chronic kidney disease; group, per session, per hour, face-to-face), are discussed in the CY 2010 MPFS proposed rule (74 FR 33619). When the qualified person is a rural provider, we proposed to pay the provider the applicable amount under the MPFS and a single payment would be made for each KDE session, limited to no more than six sessions as discussed in the CY 2010 MPFS proposed rule. Subsequently, we would not provide separate payment for both a physician's professional services and the associated facility services if a single session of KDE services was furnished in a rural hospital or CAH. Therefore, because of operational constraints, we proposed that payment would be made to only one qualified person for KDE services on the same day for the same beneficiary. We also note that the MPFS' geographic practice cost index would apply to the calculation of the payment in a particular fee schedule locality because this locality adjustment methodology is applicable to payment for all services paid under the MPFS. We proposed to assign status indicator "A" to HCPCS codes GXX26 and GXX27 in Addendum B to the CY 2010 OPPS/ASC proposed rule to signify that these services, when covered, would be paid under a payment system other than the OPPS, specifically the MPFS in the case of both HCPCS codes.

We instructed individuals who wished to submit public comments on this proposal to pay under the MPFS for covered KDE services furnished by qualified persons who are hospitals, CAHs, SNFs, CORFs, HHAs, or hospices that are located in a rural area or are treated as being rural under § 412.103 to submit those comments in accordance with the instructions for commenting on the CY 2010 OPPS/ASC proposed rule. We instructed individuals who wished to submit public comments on all other aspects of the proposed implementation of sections 1861(s)(2)(EE) and 1861(ggg) of the Act, including, but not limited to, the proposed criteria for coverage of the services, the proposed definition of "session," the proposed HCPCS G-codes, and the proposed content of the program, to submit those comments in response to the CY 2010 MPFS proposed rule.

Comment: A few commenters objected to the proposed payment to rural providers for KDE services. The commenters believed that the proposed rates were too low for appropriate payment for KDE services and recommended that CMS revise its KDE payment rates to reflect the greater resources required for rural provider to furnish KDE services.

Response: As a result of our review of the public comments and further analysis, we are adjusting the final CY 2010 MPFS RVUs for HCPCS codes G0420 and G0421. Specifically, we reviewed the medical nutrition therapy CPT codes, 97802 (Medical nutrition therapy; initial assessment and intervention, individual, face-to-face with the patient, each 15 minutes) and 97804 (Medical nutrition therapy; group (2 or more individual(s)), each 30 minutes), that we are crosswalking to the KDE codes for payment under the MPFS. We have adjusted the final CY 2010 values for HCPCS codes G0420 and G0421 to reflect not only the final specification of the time of one hour for an individual or group KDE session but also to reflect the appropriate supplies and equipment without duplication. We multiplied the physician work RVUs for HCPCS code G0420 by four and the work RVUs for HCPCS code G0421 by two to account for the fact that we are crosswalking a 15 minute MNT code to a 60 minute KDE code for the individual service and a 30 minute MNT code to a 60 minute KDE code for the group service case. We refer readers to the CY 2010 MPFS final rule with comment period for the CY 2010 RVUs for KDE services that determine payment to rural providers for HCPCS codes G0420 and G0421.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to pay under the MPFS for covered KDE services furnished by qualified persons that are hospitals, CAHs, SNFs, CORFs, HHAs, or hospices that are located in a rural area or are treated as being rural under § 412.103. Public comments concerning the definition of a “qualified person,” the proposed HCPCS G-codes, the proposed RVUs for KDE services, the proposed criteria for coverage of the services, the proposed definition of “session,” and the proposed content of the program are discussed in the CY 2010 MPFS final rule with comment period.

B. Pulmonary Rehabilitation, Cardiac Rehabilitation, and Intensive Cardiac Rehabilitation Services

1. Legislative Changes

Section 144(a) of Public Law 110–275 (MIPPA) made a number of changes to the Act to provide Medicare Part B coverage and payment for pulmonary and cardiac rehabilitation services furnished to beneficiaries with chronic obstructive pulmonary disease and certain other conditions, respectively, effective January 1, 2010. Specifically, section 144(a)(1) of the Act amended section 1861(s)(2) of the Act by adding new subparagraphs (CC) and (DD) to specify Medicare Part B coverage of items and services furnished under (1) a cardiac rehabilitation (CR) program (as defined in an added new section 1861(eee)(1) of the Act) or under a pulmonary rehabilitation (PR) program (as defined under an added new section 1861(fff)(1) of the Act; and (2) an intensive cardiac rehabilitation (ICR) program (as defined in an added new section 1861(eee)(4) of the Act). The amendments made by section 144(a) of Public Law 110–275 provide for coverage of CR, PR, and ICR services provided in a physician’s office, in a hospital on an outpatient basis, or in other settings as the Secretary determines appropriate. Section 144(a)(2) of Public Law 110–275 amended section 1848(j)(3) to provide for payment for services furnished in an ICR program under the MPFS and also added a new section 1848(b)(5) to provide specific language governing payment for ICR services. Under that specific section, the Secretary shall substitute the Medicare OPD fee schedule amount established under the prospective payment system for hospital outpatient department services under section 1833(t)(3)(D) of the Act for CR (under HCPCS codes 93797 (Physician

services for outpatient cardiac rehabilitation; without continuous ECG monitoring (per session)) and 93798 (Physician services for outpatient cardiac rehabilitation; with continuous ECG monitoring (per session)) for CY 2007, or any succeeding HCPCS codes established for cardiac rehabilitation). Section 144(a)(2) also defines under the new section 1848(b)(5) a “session” for each of the component CR items and services defined in subparagraphs (A) through (E) of section 1861(eee)(3) of the Act, when furnished for one hour, as a separate session of ICR, and specified that payment may be made for up to 6 sessions per day of the series of 72 one-hour sessions of ICR services. Section 144(a)(1)(B) also requires that a physician must be immediately available and accessible for medical consultations and medical emergencies at all times items and services are being furnished under CR, ICR, and PR programs, except that in the case of such items and services furnished under such a program in a hospital, such availability shall be presumed.

As we discuss in detail in section II.G.8. of the CY 2010 MPFS proposed rule (74 FR 33606), we proposed to use the MPFS and the OPPS rulemaking processes, and may use the national coverage determination (NCD) process as well, to implement the amendments made by section 144(a) of Public Law 110–275. In the CY 2010 MPFS proposed rule, we specified our policy proposals for implementing Medicare Part B coverage and payment for services furnished in a CR, ICR, and PR program under the MPFS. In the CY 2010 OPPS/ASC proposed rule (74 FR 35360), we proposed the CY 2010 OPPS payment for services in a CR, ICR, or PR program furnished to hospital outpatients.

Comment: A number of commenters asked that CMS confirm that the services of physical therapists are not part of the PR, CR, or ICR benefits authorized by section 144(a)(1) of Public Law 110–275 and are always paid under the physical therapy benefit and that, therefore, the therapy services do not require physician supervision when furnished as part of a PR, CR, or ICR program, including in the HOPD. With regard to PR, they stated that CMS has a longstanding history of recognizing the services of a physical therapist as an integral part of a PR program and requiring that these services be reported and paid as physical therapy services. Specifically, the commenters indicated that in the CY 2002 MPFS regulation (66 FR 55246) and in the current Medicare Claims Processing Manual (Pub. 100–04, Chapter 5, Section 20.A), CMS specifies

that when physical therapists treat respiratory conditions, they should report CPT codes for physical therapy in the 97000 series and should not report HCPCS codes G0237 (Therapeutic procedures to increase strength or endurance of respiratory muscles, one on one, face to face, per 15 minutes (includes monitoring)); G0238 (Therapeutic procedures to improve respiratory function or increase strength or endurance of respiratory muscles, one on one, face to face, per 15 minutes (includes monitoring)); or G0239 (Therapeutic procedures to improve respiratory function or increase strength or endurance of respiratory muscles, two or more individuals (includes monitoring)). The commenters added that in the September 25, 2007 Decision Memo for Pulmonary Rehabilitation (CAG–00356N), CMS recognized the importance of physical therapy to patients with pulmonary conditions and stated that these services should be billed and paid under the physical therapy benefit. The commenters argued that a plan of care developed by a physical therapist to improve pulmonary function for a patient with chronic obstructive pulmonary disease (COPD), which meets the medical necessity criteria for physical therapy services, is covered and paid under the physical therapy benefit. They explained that, although it is a covered PR service, the therapy plan of care is separate from the PR benefit authorized by section 144(a)(1) of Public Law 110–275, should continue to be reported under the CPT codes for physical therapy services, and should be paid under the physical therapy benefit. In addition, the commenters requested that CMS confirm that skilled physical therapy services that are rendered in the CR setting by a qualified physical therapist should be conducted, reported, and paid as physical therapy services, and that physician supervision is not necessary in the CR setting when the physical therapist is delivering treatment that clearly meets the criteria for a physical therapy service. The commenters explained that CMS has recognized and codified that physical therapy is a separate benefit and that physical therapists are qualified to perform certain services independent of direct physician supervision.

Response: We expect that most patients participating in PR, CR, or ICR programs authorized by section 144(a)(1) of Public Law 110–275 and covered by Medicare Part B will be debilitated based on their underlying medical condition, age, or other factors. In order to develop a PR, CR, or ICR

treatment plan, some debilitated patients may require evaluations by therapists on the multidisciplinary team, in addition to assessments by other team members. In order to participate successfully in the prescribed exercise component of the PR, CR, or ICR program, we also expect that these patients may receive individualized treatments by therapists on the multidisciplinary team and others to promote the increased functionality that is a principle goal of PR, CR, and ICR programs. As we stated in the CY 2010 MPFS proposed rule, the items and services furnished by a CR or PR program are individualized and set forth in written treatment plans for each beneficiary (74 FR 33607 and 33611). We believe these evaluations and individualized treatments are a part of the PR, CR, or ICR program. As such, we believe they should be conducted by one or more members of the multidisciplinary team of the PR, CR, or ICR program with the appropriate expertise.

While we have not defined PR, CR, or ICR services as always including therapists' services as part of the comprehensive benefit (74 FR 33608 and 33614), we acknowledged that written treatment plans are highly individualized and that there should be flexibility in the type, amount, frequency, and duration of services provided in each session (74 FR 33607). We expect that physical therapists could conduct assessments and individualized treatments as part of the PR, CR, or ICR program because physical therapists have the knowledge and skills to assist in addressing common problems that lead to physicians ordering PR, CR, or ICR services for their patients, including poor aerobic capacity, poor endurance, and shortness of breath in the context of chronic pulmonary or cardiovascular disease. In the context of PR, while we also stated that individuals requiring PR services have a chronic respiratory disease and are in need of supervised aerobic exercise, we acknowledged that patients require assessments to address individualized needs and the provision of a mix of services necessary to address those needs (74 FR 33613).

Patients in PR, CR, or ICR programs must receive the full complement of care as defined under these benefits, in accordance with their individualized treatment plan, including assessments and prescribed exercise. Additionally, the standard HCPCS coding guidance instructs practitioners and providers to report the code for the procedure or service that most accurately describes the service performed. As stated in Section 20.12.1.b. of Chapter 5 of the

Medicare Contractor Beneficiary and Provider Communications Manual, in instances where several component services, which have different CPT/HCPCS codes, may be described in one more comprehensive code, only the single code most accurately describing the procedure performed or service rendered should be reported. Therefore, we would expect that when therapists provide these evaluations and individualized treatment services under a comprehensive PR, CR, or ICR treatment plan, these services would be billed by the hospital as PR, CR, or ICR services under the comprehensive PR, CR, or ICR CPT codes or Level II HCPCS G-codes that apply, and not as physical therapy services. Furthermore, as discussed in section XII.B.4. of this final rule with comment period, for purposes of PR, CR, and ICR services, direct supervision must be provided by a doctor of medicine or osteopathy as defined in section 1861(r)(1) of the Act. This direct supervision rule would also apply to services furnished by therapists on the multidisciplinary team, and these services would be paid to the hospital as PR, CR, or ICR services.

We expect that most patients who meet the diagnosis requirements for coverage of PR, CR, or ICR would receive component services covered under the PR, CR, or ICR benefit as part of a comprehensive PR, CR, or ICR program, subject to the coverage and payment policies that we are finalizing in this CY 2010 OPFS/ASC final rule with comment period and the CY 2010 MPFS final rule with comment period. We understand that some component services of PR, CR, or ICR have previously been furnished to beneficiaries and paid by Medicare under other benefits, such as the outpatient physical therapy benefit. Because section 144(a)(1) of Public Law 100-275 authorized a new comprehensive PR benefit and codified specific benefits for CR and ICR, we believe that hospitals should furnish the full scope of the PR, CR, or ICR benefit, where services will be paid as hospital outpatient services under the OPFS, as comprehensive programs to those patients who qualify for coverage. We would not expect the component services of PR, CR, and ICR programs to be unbundled and billed separately by different providers or practitioners under other benefit categories, such as the physical therapy benefit where services would be paid under the MPFS. Therefore, we expect that it would be uncommon for a patient receiving care under a PR, CR, or ICR treatment plan to also be receiving physical therapy

services under a separate physical therapy plan of care. There may be patients with therapy needs that are outside the treatment plan appropriate for PR, CR, or ICR, and such patients should receive medically necessary physical therapy services specific to those other needs. However, we would not expect this to be the norm. Clearly, a single period of care can only be billed as one type of treatment service, so hospitals could never bill both physical therapy and PR, CR, or ICR services for the same time period for the same patient (for example, an hour session from 10 a.m. to 11 a.m. on a single date of service).

We plan to monitor claims data for PR, CR, and ICR services, as well as any additional claims for therapy services. If we detect patterns of care that are inconsistent with our stated expectations for PR, CR, or ICR services and therapy services, we may encourage Medicare contractors to review cases in which a hospital reports both types of services for the same patient during the same span of time (for example, over a several month period) or we may propose changes to our payment methodologies for these services.

After considering the public comments we received, we are clarifying that we would expect component services that are furnished under a PR, CR, or ICR treatment plan to beneficiaries who qualify for PR, CR, or ICR services to be furnished as PR, CR, or ICR services, regardless of whether they are furnished by a physical therapist or other healthcare practitioner, and that all of the coverage and payment requirements for hospital outpatient services, including, but not limited to, the physician supervision requirements for hospital outpatient therapeutic services, apply to those PR, CR, or ICR services. We refer readers to section XII.D.3. of this final rule with comment period for a discussion of the final CY 2010 policies for the direct supervision of hospital outpatient therapeutic services. We expect that hospitals will furnish the comprehensive set of services that is contemplated in the criteria for PR, CR, or ICR programs to beneficiaries who qualify for the benefit.

2. Payments for Services Furnished to Hospital Outpatients in a Pulmonary Rehabilitation Program

In the CY 2010 OPFS/ASC proposed rule (74 FR 35360), we proposed to create for CY 2010 one new Level II HCPCS code for hospitals to report and bill for the services furnished under a PR program as specified in section 1861(fff) of the Act. Specifically, we

proposed to use HCPCS code GXX30 (Pulmonary rehabilitation, including aerobic exercise (includes monitoring), per session, per day). This proposed new HCPCS G-code would be used by hospitals to report PR services furnished to patients performing physician-prescribed exercises that are targeted to improving the patient's physical functioning and may also include the provision of other aspects of PR, such as education and training. Consistent with our proposal in the CY 2010 MPFS proposed rule, we proposed that hospitals would use HCPCS code GXX30 to report sessions lasting a minimum of 60 minutes each, generally for two to three sessions of PR per week, under the OPSS. We also proposed to allow no more than one session per day because individuals who are furnished services in a PR program have significant respiratory compromise and would not typically be capable of performing more than one session of exercise per day.

We proposed that PR described by HCPCS code GXX30 would be a new comprehensive service. We did not believe there was an existing clinical APC to which this service could be appropriately assigned under the OPSS based on the information currently available to us. We did not believe that any services currently paid under the OPSS were sufficiently similar to PR, based on both clinical and resource characteristics, to justify the initial assignment of HCPCS code GXX30 to the same clinical APC as an existing service. Historically, individual component services that comprise comprehensive PR have been reported separately with existing HCPCS codes that are paid under the OPSS through the individual APC that is most appropriate for each service described by the specific HCPCS code reported.

For payment under the MPFS, we proposed relative value units for new HCPCS code GXX30 for CY 2010 based on the estimated resources and work intensity associated with existing CR and respiratory therapy services. The nonfacility practice expense amount is the component of the MPFS payment that is most comparable to what Medicare pays under the OPSS. Both the MPFS nonfacility practice expense payment and the OPSS payment include payment for the service costs other than the physician professional services that are billed and paid under the MPFS in all service settings. The CY 2010 proposed nonfacility practice expense payment amount under the MPFS was between \$10 and \$20.

For the CY 2010 OPSS, we proposed to assign HCPCS code GXX30 to New

Technology APC 1492 (New Technology—Level IB (\$10-\$20)), the New Technology APC that provides payment for new services with estimated facility costs between \$10 and \$20, because we believed that we lacked appropriate hospital cost data from claims to guide the initial assignment of the new HCPCS code that would describe services furnished under the new PR benefit. The New Technology APC payment of \$15, at the midpoint of the cost band, would be approximately the same as the proposed CY 2010 MPFS nonfacility practice expense amount for PR services described by HCPCS code GXX30. As discussed above, this is the portion of the proposed MPFS payment that is most comparable to what Medicare would pay under the OPSS. We believed that this proposed temporary assignment to a New Technology APC would allow us to pay appropriately for the service under the OPSS at a rate that is similar to the corresponding physician's office payment amount, while we gathered hospital claims data and experience with the new service on which to base a clinically relevant APC assignment in the future.

Comment: Many commenters claimed that 90 percent of PR services are furnished in HOPDs and that CMS is currently paying considerably more for these services than the proposed payment. The commenters indicated the PR services are commonly furnished in the group setting in the HOPD, although they added that patients commonly require significant one-on-one assistance from hospital staff to encourage and facilitate their full participation in supervised exercise. The commenters objected to the proposal to pay \$15 per session under New Technology APC 1492 because they believed that it would reduce payment for PR services to such an extent that hospitals would no longer be able to afford to furnish the services and that access to care would be diminished, rather than expanded as the law intended. Specifically, the commenters were concerned about the proposed payment rate because PR services would be reported under a new HCPCS G-code that would include all component services and assessments furnished in a single session per day with a minimum session duration of 60 minutes and a maximum of 1 session per day. The commenters estimated that the proposed payment for PR services would result in less than 25 percent of the current payment to hospitals for the same component services that are reported under existing HCPCS codes.

The commenters opposed the proposed payment of PR services under a New Technology APC because they believe that CMS has considerable experience with payment for these services under HCPCS codes G0237, G0238, or G0239, which are currently assigned to APC 0077 (Level I Pulmonary Treatment) under the OPSS. They argued that these three existing HCPCS G-codes are now being reported for PR services in HOPDs and that OPSS payment should be made for comprehensive PR services as authorized by section 144(a)(1) of Public Law 100-275 based on the historical payments made for services reported using these codes. They stated that the OPSS currently pays approximately \$27 per unit for HCPCS codes G0237, G0238 and G0239 and that, therefore, the proposed payment for PR services would be much less than hospitals are currently being paid for the same services and would seriously erode the quality and quantity of care currently being furnished. The commenters indicated that PR services require considerable hospital staff and overhead costs that include, but are not limited to, the costs of respiratory therapists and the purchase and maintenance of a wide range of expensive exercise equipment, such as oximeters with printers, one channel ECG monitors, dedicated emergency cart/resuscitation equipment, and portable oxygen equipment. They reasoned that payment at \$15 per session of one hour's duration would be insufficient to permit hospital PR programs to continue to function because the current sessions are at least 1 hour (and typically 2 hours) long and, therefore, OPSS payment for 1 hour of service reported using HCPCS codes G0237 and G0238 is currently about \$100 per hour. Further, they noted that the OPSS pays separately for all assessments and tests under the current policy. They identified the assessment services that are currently paid separately but for which payment would be included in the per session payment for PR services according to the proposal as including, but not limited to, CPT codes 94620 (Pulmonary stress testing; simple (e.g., 6-minute walk test, prolonged exercise test for bronchospasm with pre- and post-spirometry and oximetry)); 94664 (Demonstration and/or evaluation of patient utilization of an aerosol generator, nebulizer, metered dose inhaler or IPPB device); and 94667 (Manipulation chest wall, such as cupping, percussion and vibration to facilitate lung function; initial demonstration and/or evaluation). One

commenter requested that CMS create two new Level II HCPCS codes for PR sessions, one including the assessment services and one excluding the assessment services.

Some commenters claimed that it is not appropriate to compare the costs of CR to PR services because the overhead costs for PR services are higher than for CR services. They also indicated that it is contradictory for CMS to require that lung volume reduction surgery patients receive no less than 2 hours of PR services per day but to limit coverage of PR for moderate to severe COPD patients to 1 session per day.

The commenters suggested three alternatives to the proposed payment for a single PR HCPCS code through a New Technology APC. One alternative would be to continue to permit hospitals to bill HCPCS codes G0237, G0238, and G0239 for PR services and also to bill separately for the assessment services proposed to be included in the single comprehensive PR HCPCS code (with a separate visit payment for the initial physician or hospital staff assessment of the patient that commenters claimed requires 60 to 90 minutes of professional time). Payment for HCPCS codes G0237, G0238, and G0239 would continue to be made through APC 0077 and the assessment services would be paid through their existing clinical APCs as well. Some commenters suggested a variation of this alternative that would establish an APC with a payment rate of \$50 to which HCPCS codes G0237, G0238, and G0239, as currently defined, would be assigned. As a second alternative, the commenters suggested that CMS assign the new comprehensive PR HCPCS code to APC 0078 (Level II Pulmonary Treatment), which had a proposed payment rate of approximately \$96, and not allow separate reporting and payment for CPT codes 94620, 94664, and 94667 and the 60 to 90 minute intake visit. Finally, the commenters stated that CMS could establish a payment rate for the new comprehensive PR HCPCS code by multiplying the OPPS payment rate for APC 0077 by 4 because the cost of the new per hour PR code would be at least 4 times the cost of HCPCS codes G0237 and G0238 that describe 15 minutes of care and are assigned to APC 0077. The commenters believed that any of these alternatives would result in a PR payment to hospitals that is appropriate for the cost of the services being furnished.

Response: We examined our hospital and physician claims data for HCPCS codes G0237, G0238, and G0239, and we agree with the commenters that most services for restoration of pulmonary

function have historically been furnished in hospitals and that there is considerable evidence of hospital outpatient utilization and cost in our claims data. We found that, in CY 2008, Medicare paid approximately \$20 million in aggregate to about 900 hospitals for services reported under HCPCS codes G0237, G0238, and G0239, which describe pulmonary therapy services that commenters believe reflect the costs of the same types of hospital staff, supplies, and overhead that would be used to furnish PR services.

Therefore, for the CY 2010 OPPS, we will pay for PR services in HOPDs under the OPPS through a new clinical APC with a median “per session” cost simulated from historical hospital claims data for similar pulmonary therapy services, rather than assigning the new PR HCPCS G-code to a New Technology APC as we proposed. We have previously used a simulation approach to develop a median cost estimate for a single new CPT or Level II HCPCS code that would previously have been reported under several existing HCPCS codes when furnished in the HOPD, most recently for echocardiography services as discussed in detail in section II.A.2.d.(4) of this final rule with comment period. Specifically, we are assigning the comprehensive Level II HCPCS code G0424 (Pulmonary rehabilitation, including exercise (includes monitoring), per hour, per session) to new clinical APC 0102 (Level II Pulmonary Treatment), with payment based on the aggregate per day simulated “per session” hospital median cost of approximately \$50 as calculated from claims data for the existing pulmonary therapy HCPCS G-codes and associated assessments and tests. No other HCPCS codes are assigned to APC 0102 for CY 2010. (APC 0078 has been renamed Level III Pulmonary Treatment without any change in its configuration for CY 2010.) Our claims data show that, in most cases patients furnished the pulmonary therapy services reported by HCPCS codes G0237, G0238, and G0239 in the HOPD on a single date of service received some individual and some group services (approximately 2.4 units of service per day) and, less often, associated assessments and tests. We found approximately 19,000 days of care for patients who had diagnoses consistent with moderate to very severe COPD, consistent with the coverage requirements for PR in CY 2010, and that included both an individual and group service on the same day. The

claims for these days of care, for which we calculated a “per session” median cost of approximately \$50, were similar to the time duration and services that will be reported under new HCPCS code G0424 in CY 2010.

Specifically, to simulate the “per session” median cost of new HCPCS code G0424 from claims data for existing services, we used only claims that contained at least one unit of HCPCS code G0239, the group code that is without limitation on time duration, and one unit of HCPCS code G0237 or G0238, the individual, face-to-face codes that report 15 minutes of service, on the same date of service. We reasoned that patients in a PR program would typically receive individual and group services in each session of approximately 1 hour in duration. The approach was consistent with the public comments that suggested that PR is often provided in group sessions in the HOPD, although patients commonly require additional one-on-one care in order to fully participate in the program. We note that our use of “per session” claims reporting one unit of HCPCS code G0237 or G0238 and one unit of HCPCS code G0239 in this simulation methodology was also consistent with our overall finding of approximately 2.4 service units of the HCPCS G-codes per day on a single date of service, usually consisting of both individual and group services, for patients receiving pulmonary therapy services in the HOPD based upon CY 2008 claims. We concluded that the typical session of PR would be 1 hour based on public comments that indicated that a session of PR is typically 1 hour and based on our findings that the most commonly reported HCPCS code for pulmonary treatment is HCPCS code G0239, which has no time definition for this group service.

We included all costs of the related tests and assessment services (CPT codes 94620, 94664, and 94667 and all CPT codes for established patient clinic visits) on the same date of service as the HCPCS G-codes in the claims we used to simulate the median cost for HCPCS code G0424. After identifying these “per session” claims, which we believe to represent 1 hour of care, we summed the costs on them and calculated the median cost for the set of selected claims. In light of the cost and clinical similarities of PR and the existing services described by HCPCS codes G0237, G0238, and G0239 and the CPT codes for related assessments and tests, and the significant number of “per session” hospital claims we found, we are confident that the simulated median cost for HCPCS code G0424 is a valid

estimate of the expected hospital cost of a PR session. Since there is no existing clinical APC to which HCPCS code G0424 could be appropriately assigned based on considerations of the clinical and resource characteristics of PR services, we are creating new APC 0102 (Level II Pulmonary Treatment) for CY 2010. Existing APC 0078 (Level II Pulmonary Treatment) has been renamed "Level III Pulmonary Treatment" for CY 2010, with no change in its configuration.

We also are adding the phrase "per hour" to the new HCPCS code G0424 descriptor to conform the descriptor of the code to the basis for the payment being made for one unit of the code and to enable providers to determine when one session of PR ends and the second session begins. Because we are modifying our final policy to cover up to 2 sessions of PR per day in response to public comments that we should permit more than 1 session of PR on the same date, it became necessary to add a time to the definition of HCPCS code G0424 so that providers could determine when they could report a second session. Moreover, when we based the payment for new APC 0102 on the per session costs derived from the OPFS claims data for HCPCS codes G0237, G0238, and G0239, we assumed that a session represented 1 hour of care; and therefore, adding "per hour" to the descriptor for the new HCPCS G-code for PR conforms the HCPCS G-code to the payment we are establishing.

We do not agree with the commenters' suggestion that we permit hospitals to report and be paid for PR services under the existing HCPCS codes G0237, G0238, and G0239 and the CPT codes for assessments and tests because PR is a comprehensive program that consists of multiple component items and services as specified in the statute and discussed in the CY 2010 MPFS final rule with comment period. These three HCPCS G-codes were developed principally to describe services furnished by respiratory therapists in CORFs, and the services consist of therapeutic procedures to increase the strength and endurance of respiratory muscles or to improve respiratory function. When the HCPCS G-codes were established, we indicated that these were developed to provide more specific information about the services being delivered since those services were not previously clearly described by existing CPT codes (66 FR 55311). We also noted that there was no respiratory or pulmonary rehabilitation benefit (67 FR 79999).

The existing HCPCS G-codes do not represent the full scope of services in a

comprehensive PR program as now authorized by section 144(a)(1) of Public Law 100-275. We want to ensure that when hospitals bill and are paid for PR services, they attest to meeting all requirements of the comprehensive PR program by the reporting of a HCPCS G-code specific to a PR session. As discussed above in the context of therapy services, patients in PR, CR, or ICR programs must receive the full complement of care defined under the benefit, including assessments and individualized treatments in accordance with their PR treatment plan. When hospitals furnish these evaluations and individualized treatment services under a PR treatment plan, we would not expect the component services of PR, CR, and ICR programs to be unbundled and billed separately. Instead, the services must be billed by the hospital as PR services under the new Level II HCPCS G-code for PR services, not the existing HCPCS codes G0237, G0238, and G0239 for respiratory treatment services, the CPT codes for the individual assessment services discussed above, or HCPCS codes for physical therapy or other services. Furthermore, we also expect that it would be uncommon for a patient receiving care under a PR treatment plan to also be receiving services under a separate plan of care to improve their respiratory strength and function, such as physical therapy or respiratory treatment. Only those patients whose medical needs for treatment to improve their respiratory strength and function are outside the treatment plan appropriate for PR should receive medically necessary services specific to those other needs. However, we would not expect this to be the norm.

As discussed above, a single period of care can only be billed as one type of treatment service, so hospitals could never bill services reported by HCPCS codes G0237, G0238, or G0239 and PR services for the same time period for the same patient (for example, an hour session from 10:00 a.m. to 11:00 a.m. on a single date of service). We plan to monitor claims data for PR services, as well as any additional claims for these other services. If we detect patterns of care that are inconsistent with our stated expectations for PR services, we may encourage Medicare contractors to review cases in which a hospital reports both types of services for the same patient during the same span of time (for example, over a period of several months) or we may propose changes to our payment methodologies for these services.

In addition, we note that a specific code for PR services allows us to

administratively account for the limit on the number of covered sessions. Furthermore, we are not creating different HCPCS codes for PR sessions provided with and without assessments because we continue to believe PR is a comprehensive program and all sessions should be reported and paid through a single HCPCS G-code. We believe that it is most appropriate to pay for all of the assessments and tests that may be furnished in the program as required for coverage through our single per-session PR payment, and we took the hospital costs of these related services into consideration in establishing the CY 2010 OPFS payment rate for HCPCS code G0424. We also do not agree with the commenters' recommendation to assign the new PR per session HCPCS code to APC 0078 because the median cost of APC 0078 is almost twice the simulated "per session" median cost of HCPCS code G0424 that we calculated based on historical hospital claims data for existing, similar HCPCS codes through our "per session" methodology as described above. Finally, we do not agree with the commenters' suggestion to establish a payment rate for PR HCPCS code G0424 at the rate of 4 times the payment for APC 0077 because that also would pay for PR services at more than twice the simulated median cost for a typical session of PR as reported in our claims data.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, with modification, for OPFS payment for PR services furnished as a part of the comprehensive PR program benefit for CY 2010. We are adopting new HCPCS code G0424 (Pulmonary rehabilitation, including exercise (includes monitoring), per hour, per session) and are assigning the G-code to new APC 0102 (Level II Pulmonary Treatment), with a simulated "per-session" median cost of approximately \$50. As discussed in the CY 2010 MPFS final rule with comment period, PR is covered for up to 36 one-hour sessions, with a maximum of 2 sessions per day, and with contractor discretion to approve up to 72 sessions.

3. Payment for Services Furnished to Hospital Outpatients Under a Cardiac Rehabilitation or an Intensive Cardiac Rehabilitation Program

Currently, CR services furnished by hospitals are reported using CPT codes 93797 and 93798. In the CY 2010 MPFS proposed rule (74 FR 33607), we proposed that each day CR items and services are furnished to a patient, aerobic exercises along with other exercises must be included (that is, a

patient must exercise aerobically every day he or she attends a CR session). In addition, we proposed that each session must be a minimum of 60 minutes and patients must participate in a minimum of two CR sessions a week, with a maximum of two CR sessions a day.

With respect to ICR services, section 1861(eee)(4)(C) of the Act, states that "an intensive cardiac rehabilitation program may be provided in a series of 72 one-hour sessions (as defined in section 1848(b)(5)), up to 6 sessions per day, over a period of up to 18 weeks." In the CY 2010 OPPTS/ASC proposed rule (74 FR 35361), for the CY 2010 OPPTS, we proposed to create two new Level II HCPCS codes to report the services of an ICR program that are furnished to hospital outpatients, consistent with the provisions of section 1861(eee)(4)(C) of the Act: HCPCS code GXX28 (Intensive cardiac rehabilitation; with or without continuous ECG monitoring with exercise, per session) and HCPCS code GXX29 (Intensive cardiac rehabilitation; with or without continuous ECG monitoring; without exercise, per session). These proposed new HCPCS G-codes would be used to report ICR services furnished by hospitals that have an ICR program that has received a designation as a qualified ICR program. Consistent with the proposal in the CY 2010 MPFS proposed rule, we proposed that each session of ICR must be a minimum of 60 minutes and that each day ICR items and services are provided to a patient, aerobic exercises along with other exercises must be included (that is, a patient must exercise aerobically every day he or she attends a ICR session).

For the CY 2010 OPPTS, we proposed to assign HCPCS codes GXX28 and GXX29 to APC 0095 (Cardiac Rehabilitation) with a status indicator of "S." The proposed median cost of APC 0095 for CY 2010 was approximately \$39. This proposed median cost reflected historical hospital cost data for one session of general CR services reported with CPT code 93797 or 93798. Both CR and ICR programs consist of exercise, cardiac risk factor modification, psychosocial assessment, outcomes assessment, and other services, as described in the CY 2010 MPFS proposed rule (74 FR 33607). Although more sessions per day for a beneficiary may be provided in an ICR program than a CR program, in the CY 2010 OPPTS/ASC proposed rule (74 FR 35361) we noted our belief the hospital costs for a single session would be similar, and OPPTS payment for CR and ICR services would be provided on a per session basis. Therefore, because CR and ICR services are similar from both

clinical and resource perspectives, we believed that it would be appropriate to assign the two proposed new Level II HCPCS codes for ICR services to APC 0095 while we collect cost information from hospitals specific to ICR. We proposed to make a single payment through APC 0095 for each session of ICR reported on hospital outpatient claims.

Comment: Some commenters supported the proposal to create two new HCPCS G-codes for ICR services and to pay them at the same rate as CR services through APC 0095. Other commenters objected to setting the payment for ICR services at the same level as CR services on the basis that ICR services should be more costly than CR services because of their intensive nature. Some commenters were concerned about basing payment for CR on historical hospital costs because these costs do not include the new level of physician and clinical staff work required for coverage of CR and ICR, in particular, the psychosocial and outcome assessments that are required by section 144(a)(1) of Public Law 100-275. Under CMS' proposal, these assessments would not be paid separately but, instead, would be paid through payment for the per session CR CPT codes or ICR HCPCS codes.

Some commenters believed the proposed median cost for APC 0095 was too low on the basis that the July 2008 RTI final report for CMS entitled "Refining Cost to Charge Ratios for Calculating APC and DRG Relative Payment Weights" indicated that the current payment calculations for CR are significantly flawed because hospitals misclassify CR costs and that, therefore, the resulting APC 0095 median cost understates the cost of CR. The commenters indicated that the study showed that the cost of CR is approximately \$100 per session when an appropriate CCR, based on the costs and charges for CR and not on the application of a hospital-specific overall ancillary CCR, is applied to the charges for CR services. The commenters were concerned about CMS' stated delay in implementing a specific nonstandard cost center for CR until 2011 because the true cost of CR would not be captured by the OPPTS for ratesetting until CY 2013 or later. The commenters encouraged CMS to correct the underpayment of CR services for CY 2010 in order to ensure the availability of CR programs for Medicare beneficiaries.

Response: We have no reason to believe that ICR services as defined in the coverage criteria are more costly to furnish per session than CR services. We

note that one of the major differences between CR and ICR services is that ICR may be furnished in up to six sessions per day, in comparison with the two sessions per day that are covered for CR. In the case of a Medicare beneficiary who receives six ICR sessions in one day, payment also would be six times the payment for one session and, therefore, the total hospital payment would appropriately pay for the additional costs of more services in a day.

With regard to the comment that our claims data do not reflect the costs of the additional assessments that are now required for CR and ICR coverage, we analyzed the per session cost of CR in our historical hospital claims data and incorporated the costs of hospital outpatient visits that we believe would have been previously reported for assessments furnished on the same day to CR patients into our estimate of the median cost for APC 0095. We found that the APC median cost was essentially unchanged. We believe that psychosocial and outcomes assessments are already a part of high quality CR/ICR programs and that our per session payment for CR and ICR services through APC 0095 will appropriately provide payment for all required components of CR and ICR, including the necessary assessments.

The median cost on which the CY 2010 payment for APC 0095 is based is approximately \$38, which is an increase over the median cost of approximately \$36 on which the CY 2009 payment is based. As we explained in the CY 2009 OPPTS/ASC final rule with comment period (74 FR 68525), we recognize that there are areas of concern with the cost report that are integral to our estimation of hospital costs for OPPTS ratesetting and we are taking steps to address some of them, including adopting a nonstandard cost center for CR. This cost center will be available for use in cost reporting periods beginning on or after February 2010. We refer readers to section II.A.1.c.(2) of this final rule with comment for further discussion of the creation of this new cost center. However, as we have previously explained (74 FR 68522), modifying the cost report data from its submitted form for use in OPPTS ratesetting based upon assumptions about the data typically would be contrary to our principle of using the data as submitted by hospitals. Therefore, we are not making changes or adjustments to our OPPTS cost estimation for CR services for CY 2010 but, instead, will await more accurate information for future ratesetting that may be submitted to us by hospitals when the nonstandard cost center for CR services is available.

We currently have no reason to believe that Medicare beneficiaries have limited access to CR services or that our payment is inappropriate. We have over 2.5 million claims for CR sessions from CY 2008 claims data, and further note that the overall number of services and the number of providers furnishing CR have been stable over the past several years.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to continue to assign the CPT codes for CR, specifically CPT codes 93797 and 93798, to APC 0095. We also are finalizing our CY 2010 proposal to assign the new HCPCS G-codes for ICR, specifically G0422 (Intensive cardiac rehabilitation; with or without continuous ECG monitoring, with exercise, per hour, per session) and G0423 (Intensive cardiac rehabilitation; with or without continuous ECG monitoring, without exercise, per hour, per session) to APC 0095. We have added the phrase “per hour” to the descriptors of these codes because we expect that the OPFS cost data for CR services from the claims submitted for CPT codes 93978 and 93979 generally reflect 1 hour of CR services, in accordance with our reporting instructions for more than one session per day of CR services in Section 200.1 of Chapter 4 of the Medicare Claims Processing Manual. We believe that 1 hour of service is the standard for a session of CR. Section 1861(eee)(4)(C) of the Act provides for up to 72 one-hour sessions of ICR and hence, adding “per hour” to the two new HCPCS code descriptors for ICR services implements the statutory definition of an ICR session as being one hour of service. Moreover, we have established the payment for ICR services on the presumption that one session represents one hour of care. Therefore, we believe that it is appropriate to specify in the descriptors of the HCPCS codes for ICR services that one unit of the code represents one hour of care. The final APC median cost of APC 0095 is approximately \$38. As discussed in the CY 2010 MPFS final rule with comment period, CR is covered for up to 36 one-hour sessions, with a minimum of 1 session per week and a maximum of 2 sessions per day, and Medicare contractors have the authority to approve additional sessions, up to 72 sessions, over an additional period of time. With respect to ICR, section 144(a)(1) of Public Law 100–275 authorizes coverage of ICR programs in a series of 72 one-hour sessions, up to

6 sessions per day, over a period of 18 weeks.

We also note that as discussed in section II.G.9. of the CY 2010 MPFS final rule with comment period, we are requiring that all ICR programs be approved through the NCD process. Once we have approved an ICR program or programs through the NCD process, individual sites wishing to furnish ICR items and services via an approved ICR program may enroll with their local Medicare contractor to become an ICR program supplier as outlined in § 424.510. This enrollment is designed to ensure that the specific sites meet the specific statutory and regulatory requirements to furnish these services and will provide a mechanism to appeal a disapproval of a prospective ICR program site. With regards to billing and payment for CR and ICR services, hospital providers will continue to use their CMS Certification Number (CCN or provider number) and appeals related to the payment of claims will follow those established processes. CMS will provide further instructions for the NCD and individual site enrollment processes.

4. Physician Supervision for Pulmonary Rehabilitation, Cardiac Rehabilitation, and Intensive Cardiac Rehabilitation Services

Section 144 of Public Law 110–275 includes requirements for immediate and ongoing physician availability and accessibility for both medical consultations and medical emergencies at all times items and services are being furnished under CR, ICR, and PR programs. In section II.G.8. of the CY 2010 MPFS proposed rule (74 FR 33606), we proposed that these requirements would be met through existing definitions for direct physician supervision in physicians’ offices and hospital outpatient departments at § 410.26(a)(2) (defined through cross reference to § 410.32(b)(3)(ii) and § 410.27, respectively). We noted that direct supervision, as defined in the regulations, is consistent with the requirements of Public Law 110–275 because the physician must be present and immediately available where the services are being furnished. The physician must also be able to furnish assistance and direction throughout the performance of the services, which would include medical consultations and medical emergencies.

For CR, ICR, and PR services provided to hospital outpatients, direct physician supervision is the standard set forth in the April 7, 2000 OPFS final rule with comment period (68 FR 18524 through 18526) for supervision of hospital outpatient therapeutic services covered

and paid by Medicare in hospitals and provider-based departments of hospitals. We noted in the discussions of CR and PR in the CY 2010 MPFS proposed rule (74 FR 33609 and 33614) that if we were to propose future changes to the physician office or hospital outpatient policies for direct physician supervision, we would provide our assessment of the implications of those proposals for the supervision of CR and PR services at that time.

As discussed in more detail in the CY 2010 OPFS/ASC proposed rule (74 FR 35362), we proposed to refine the definition of the direct supervision of hospital outpatient therapeutic services for those services provided in the hospital and in an on-campus PBD of the hospital. For services, including CR, ICR, and PR services, provided in the hospital and in an on-campus PBD of the hospital, direct supervision would mean that the physician must be present on the same campus, in the hospital or the on-campus PBD of the hospital, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. We also proposed to define “in the hospital” in proposed new paragraph § 410.27(g) to mean areas in the main building(s) of the hospital that are under the ownership, financial, and administrative control of the hospital; are operated as part of the hospital; and for which the hospital bills the services furnished under the hospital’s CCN. We did not propose significant change to the definition or requirements for direct supervision of hospital outpatient therapeutic services provided in off-campus PBDs of a hospital. Thus, with respect to CR, ICR, and PR services furnished in off-campus PBDs of the hospital, direct supervision would continue to mean that the physician must be in the off-campus PBD and immediately available to furnish assistance and direction throughout the performance of the procedure. We believe that direct supervision, as defined in the proposed regulations for hospital outpatient therapeutic services, continues to be consistent with the requirements of Pub. L. 110–275 for CR, ICR, and PR services because the physician must be present and immediately available where the services are being furnished. The physician must also be able to furnish assistance and direction throughout the performance of the services, which would include medical consultations and medical emergencies. For a complete discussion of the current and proposed requirements for the direct

supervision of hospital outpatient therapeutic services, we refer readers to section XII.D. of the CY 2010 OPPS/ASC proposed rule (74 FR 35362 through 35368).

Section 144 of Public Law 110–275 also states that in the case of items and services furnished under such a CR, ICR, or PR program in a hospital, physician availability shall be presumed. As we have stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68702 through 68704), the longstanding presumption of direct physician supervision for hospital outpatient services means that direct physician supervision is the standard for supervision of hospital outpatient therapeutic services covered and paid by Medicare in hospitals and PBDs of hospitals, and we expect that hospitals are providing services in accordance with this standard.

We note that in the CY 2010 OPPS/ASC proposed rule (74 FR 35362), we also proposed that nonphysician practitioners, defined for the purpose of proposed revised § 410.27 of the regulations as clinical psychologists, physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives, may directly supervise all hospital outpatient therapeutic services that they may perform themselves within their State scope of practice and hospital-granted privileges, provided that they meet all additional requirements, including any collaboration or supervision requirements as specified in §§ 410.71, 410.74, 410.75, 410.76, and 410.77. However, in the CY 2010 MPFS proposed rule and in the corresponding proposed regulation text (74 FR 33674 and 33675, respectively), we proposed a different requirement for the direct supervision of CR, ICR, and PR services. We proposed that services provided in CR, ICR, and PR programs must be supervised by a doctor of medicine or osteopathy, as defined in section 1861(r)(1) of the Act. In addition, we proposed specific requirements for the expertise and licensure of physicians supervising CR and ICR services. It would not be in accordance with the proposed regulations for a nonphysician practitioner to supervise services furnished in a CR, ICR, or PR program. The physician supervision and expertise requirements proposed in the coverage policy and regulations for CR, ICR, and PR services must be met for those services to be covered and, therefore, paid by Medicare in hospital outpatient settings.

Comment: One commenter supported the proposed requirements for physician supervision of CR programs and

requested that CMS confirm that the proposed definition of “direct supervision” that would apply to therapeutic services in the HOPD would also apply to CR services. Many commenters objected to the requirement that a physician must be present when PR or CR/ICR services are furnished. They indicated that the law presumes that physician supervision exists in hospitals and that, therefore, the same rules that apply to other hospital outpatient therapeutic services provided in hospitals should apply to PR, CR, and ICR services. The commenters also asked that CMS permit physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse midwives who are functioning within their State licensure and scope of practice and who are permitted to supervise the services under the hospital bylaws to supervise PR, CR, and ICR services. The commenters expressed concern that the CY 2010 proposals in the MPFS proposed rule did not include a justification for why it would be necessary to impose physician-only direct supervision for PR, CR, and ICR services in HOPDs than for other outpatient services. Some commenters explained that rural providers have great difficulty securing physician services and rely heavily on nonphysician practitioners to furnish care in hospitals. They argued that to require physician-only supervision would mean that some PR, CR, and ICR programs in rural areas would have to close for lack of physician supervision and that there would be no access to the PR, CR, and ICR services for beneficiaries in those communities.

Response: We understand the reasoning of the commenters that PR, CR, and ICR services should require direct supervision by physicians and certain nonphysician practitioners, as we proposed for other hospital outpatient therapeutic services, given that PR, CR and ICR services are similar to other hospital outpatient therapeutic services. However, we are unable to revise the regulations to permit nonphysician practitioners to supervise PR, CR, and ICR services. We do not believe that the law provides the flexibility for us to permit anyone other than a physician to supervise hospital outpatient PR, CR, and ICR services because nonphysician practitioners are not physicians as defined in section 1861(r)(1) of the Act. The statutory language of section 144(a)(1) of Public Law 100–275 defines PR, CR and ICR programs as “physician-supervised.” More specifically, it establishes in section 1861(eee)(2)(B) of the Act that

for PR, CR and ICR programs, “a physician is immediately available and accessible for medical consultation and medical emergencies at all times items and services are being furnished under the program, except that, in the case of items and service furnished under such a program in a hospital, such availability shall be presumed * * *.” The text of the statute uses the word “physician” and does not include nonphysician practitioners. Also, as we explained in the CY 2009 OPPS/ASC proposed rule and final rule with comment period (73 FR 41518 through 41519 and 73 FR 68702 through 68704, referencing the April 7, 2000 OPPS final rule (65 FR 18525)), the “presumption” or “assumption” of direct supervision means that direct physician supervision is the standard for all hospital outpatient therapeutic services. We have assumed this requirement is met on hospital premises (meaning we have expected that hospitals are meeting this requirement) because staff physicians would always be nearby in the hospital. In other words, the requirement is not negated by a presumption that the requirement is being met. Hence, unlike the standards for the direct supervision of other hospital outpatient therapeutic services, which we have established through regulation based on section 1861(s) of the Act, in the case of PR, CR, and ICR services, the authorizing provision of the Act at section 1861(eee)(2)(B) explicitly requires direct physician supervision of these services. While we have some flexibility to determine the type of practitioner who may supervise other hospital outpatient therapeutic services, as discussed in XII.D.3. of this final rule with comment period, in the case of PR, CR, and ICR services specifically, the statutory language of section 144(a)(1) of Public Law 100–275 does not provide such flexibility. Instead, the statute imposes strict requirements, describing the direct physician supervision standard for PR, CR, and ICR services, and gives us no flexibility to modify the requirement to allow for other supervisory practitioners.

After consideration of the public comments we received, and in accordance with the final policies set forth in sections II.G.8. and II.G.9. of the CY 2010 MPFS final rule with comment period, we are finalizing our CY 2010 proposal, without modification, to require the direct physician supervision (by a doctor of medicine or doctor of osteopathy) of PR, CR, and ICR services that are furnished to hospital outpatients. We note that we define “direct supervision” with regard to

what it means to be immediately available and accessible for medical consultation and medical emergencies in the same manner for PR, CR, and ICR programs as we do for other therapeutic services furnished in HOPDs. The final CY 2010 definitions of direct supervision for hospital outpatient therapeutic services provided on the campus and in off-campus provider-based departments also apply. These definitions are discussed in detail in section XII.D.3. of this final rule with comment period, including the new and revised regulations.

C. Stem Cell Transplant

Stem cell transplantation is a treatment in which stem cells that are harvested from either a patient's or a donor's bone marrow or peripheral blood are later infused into that patient to treat an illness. Autologous stem cell transplantation is a technique for providing additional stem cells using the patient's own previously harvested stem cells. Allogeneic stem cell transplantation is a procedure in which stem cells from a healthy donor are acquired and prepared to provide a patient with new stem cells.

We recently revised section 90.3.3 of Chapter 3 of the Medicare Claims Processing Manual (Pub. 100-04) and created new section 231.10 of Chapter 4 of the Medicare Claims Processing Manual in order to clarify billing under Medicare for autologous and allogeneic stem cell transplant services. As stated in the cited new and revised manual sections, autologous stem cell transplants performed on Medicare beneficiaries may be provided on an inpatient or an outpatient basis. Hospitals are instructed to bill and show all charges for autologous stem cell harvesting, processing, and transplant procedures based on the status of the patient (that is, inpatient or outpatient) when the individual services are furnished. The CPT codes describing these services may be billed and are separately payable under the OPSS when the services are provided in the hospital outpatient setting.

In contrast, we stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35362) that we believe allogeneic stem cell transplants performed on Medicare beneficiaries are provided on an inpatient basis only, and all services related to acquiring the stem cells from a donor (whether performed on an inpatient or outpatient basis) are billed and are payable under Medicare Part A through the IPPS MS-DRG payment for the stem cell transplant. In addition to payment for the stem cell transplant procedure itself, the MS-DRG payment

for the stem cell transplant includes payment for stem cell acquisition services, which include, but are not limited to, National Marrow Donor Program fees for stem cells from an unrelated donor (if applicable); tissue typing of a donor and a recipient; donor evaluation; physician pre-admission/pre-procedure donor evaluation services; costs associated with the harvesting procedure; post-operative/post-procedure evaluation of a donor; and preparation and processing of stem cells. While certain acquisition services, such as donor harvesting procedures, may be performed in the hospital outpatient setting, hospitals are instructed to include the charges for these services in the recipient's inpatient transplant bill as acquisition services and not to bill them under the OPSS.

In order to be consistent with the revised Section 90.3.3 and the new Section 231.10 of the Medicare Claims Processing Manual cited earlier, which reflect what we believed at the time to be the current clinical practice of performing allogeneic stem cell transplants on Medicare beneficiaries on an inpatient basis only, in the CY 2010 OPSS/ASC proposed rule (74 FR 35362), we proposed to revise the status indicator assignments of certain stem cell transplant-related CPT codes under the OPSS. Specifically, we proposed to change the status indicator for CPT code 38205 (Blood-derived hematopoietic progenitor cell harvesting for transplantation, per collection; allogeneic) from "S" to "E" for the CY 2010 OPSS to reflect that, while an allogeneic stem cell harvesting procedure performed on the donor may take place in the HOPD, payment for the service is made through the IPPS MS-DRG payment for the associated transplant procedure performed on the recipient. We also proposed to change the status indicators for CPT code 38240 (Bone marrow or blood-derived peripheral stem cell transplantation; allogeneic) and CPT code 38242 (Bone marrow or blood-derived peripheral stem cell transplantation; allogeneic donor lymphocyte infusions) from "S" to "C" for the CY 2010 OPSS to reflect that these allogeneic stem cell transplant procedures are payable by Medicare as inpatient procedures only.

At its August 2009 meeting, the APC Panel heard from presenters who stated that reduced intensity conditioning (RIC) regimens have made outpatient allogeneic stem cell transplants feasible for some Medicare beneficiaries. The presenters stated that revising the status indicators for CPT codes 38205, 38240, and 38242 to make them nonpayable on

outpatient claims would impede the current clinical practice of providing allogeneic stem cell transplant and harvesting procedures on an outpatient basis, which, according to public commenters on the proposed rule, would lower costs, provide greater patient comfort, optimize hospital resources, and decrease the risk of nosocomial infections. The APC Panel agreed with the presenters and recommended that CMS maintain the CY 2009 APC assignments and status indicators for CPT codes 38205 and 38242, which are both currently assigned to APC 0111 (Blood Product Exchange), and for CPT code 38240, which is currently assigned to APC 0112 (Apheresis and Stem Cell Procedures) for CY 2009.

Comment: Several commenters on the CY 2010 proposed rule disagreed with CMS' assertion that allogeneic stem cell transplants performed on Medicare beneficiaries are provided on an inpatient basis only. According to the commenters, allogeneic stem cell transplants are currently performed in both the inpatient and outpatient settings and are considered safe and clinically appropriate for many Medicare patients. The commenters argued that CMS' proposal would create strong financial incentives for hospitals to unnecessarily admit patients who could have otherwise been treated in the outpatient setting and requested that CMS maintain the current CY 2009 status indicators and APC assignments of CPT codes 38205, 38240, and 38242 for CY 2010. The commenters urged CMS to continue to pay for CPT code 38205 separately under the OPSS, regardless of where the transplant ultimately occurs, because CMS' policy of bundling payment for stem cell harvesting procedures into the payment for the transplant procedure is overly complicated and burdensome, particularly when the harvesting procedure is performed in a different hospital from the hospital where the transplant is performed. The commenters also requested that CMS revise its manual guidance to reflect that hospitals should report all allogeneic stem cell harvesting, processing, and transplant services and their associated charges based on the status of the patient (that is, inpatient or outpatient) when the individual services are furnished.

Some commenters summarized other billing and payment issues related to allogeneic stem cell transplants for which they are seeking further direction and policy development from CMS, such as how hospitals could be paid for search and procurement costs related to

allogeneic stem cell transplants that do not occur due to a change in the patient's health status and the potential creation of separate MS-DRGs for allogeneic and autologous stem cell transplant services. The commenters recognized that greater analysis of these complex and unique issues would be required before CMS could address them fully.

Response: We appreciate the information on current clinical practice for allogeneic stem cell transplants provided by commenters. Upon further review, we agree with the commenters and the APC Panel that the allogeneic stem cell transplant procedures described by CPT codes 38240 and 38242 can be safely and appropriately performed on some Medicare beneficiaries on an outpatient basis. Therefore, we are not adopting our CY 2010 proposal to change the status indicators for CPT codes 38240 and 38242 from "S" to "C" in order to indicate that Medicare would only pay for these procedures when furnished in the hospital inpatient setting. Rather, we are continuing to assign CPT code 38240 to APC 0112 and CPT code 38242 to APC 0111 for CY 2010 for purposes of hospital outpatient payment, and are maintaining the assignment of status indicator "S" to both CPT codes that describe these procedures. CPT codes 38240 and 38242 are assigned to these same APCs for CY 2009, and we believe the allogeneic stem cell transplant-related procedures they describe share the clinical and resource characteristics of other procedures assigned to those APCs for CY 2010.

We do not agree with the public commenters and the APC Panel that the allogeneic stem cell harvesting procedure described by CPT code 38205 should be separately payable with status indicator "S" under the OPFS for CY 2010 because hospitals may bill and receive payment only for services provided to the Medicare beneficiary who is the recipient of the stem cell transplant and whose illness is being treated with the stem cell transplant. We do not agree with the commenters that we should pay for allogeneic stem cell harvesting services separately because these services are not directly furnished to beneficiaries. Instead, we believe that it continues to be appropriate to pay for these services through payment for the associated stem cell transplant procedure. The hospital should report all allogeneic stem cell acquisition charges, including costs associated with the harvesting procedure, on the recipient's inpatient or outpatient transplant bill under revenue code 0819. Payment for the allogeneic stem cell

harvesting procedure performed on the donor and reported under revenue code 0819 is packaged into the IPPS MS-DRG payment or the OPFS APC payment for the associated transplant procedure performed on the recipient, depending on whether the transplant procedure is performed in the inpatient setting or the outpatient setting. Therefore, we are modifying our CY 2010 proposal to change the status indicator for CPT code 38205 from "S" to "E" for the CY 2010 OPFS. Instead, we are assigning status indicator "B" to CPT code 38205 for CY 2010 to reflect that the code is not recognized by OPFS when submitted on an outpatient hospital Part B bill type. We will update section 90.3.3 of Chapter 3 and section 231.10 of Chapter 4 of the Medicare Claims Processing Manual to comport with these billing and payment changes for allogeneic stem cell transplants and related services as described in this section.

We appreciate the commenters' summaries of other billing and payment issues related to allogeneic stem cell transplants for which they sought further direction and policy development from CMS. While we consider these issues to be outside of the scope of the proposed rule, we will consider the commenters' suggestions as we explore this policy area more broadly in the future. We note that current guidance in Section 90.3.3 of Chapter 3 of the Medicare Claims Processing Manual instructs providers to include on the Medicare cost report any costs associated with acquisition services for allogeneic stem cell acquisition services in cases that do not result in transplant due to death of the intended recipient or other causes.

After consideration of the public comments we received, we are modifying our CY 2010 proposal for allogeneic stem cell transplant procedures. Specifically, for CY 2010, we are accepting the APC Panel's recommendation and continuing to assign CPT code 38240 to APC 0112 and CPT code 38242 to APC 0111, which is consistent with their CY 2009 assignments. The final APC median costs of APC 0112 and APC 0111 are approximately \$2,225 and \$798, respectively, for CY 2010. We are maintaining status indicator "S" for the procedures described by both of these CPT codes. In addition, we are not adopting the APC Panel's recommendation or the public commenters' suggestion to maintain the CY 2009 status indicator and APC assignment for CPT code 38205. Instead, we are assigning status indicator "B" to CPT code 38205 for CY 2010 to reflect that the code is not recognized by OPFS

when submitted on an outpatient hospital Part B bill type because payment is made for allogeneic stem cell harvesting through payment for the recipient's transplant procedure, whether the transplant is provided in the hospital inpatient setting or the outpatient setting.

D. Physician Supervision

1. Background

In the CY 2009 OPFS/ASC proposed rule and final rule with comment period (73 FR 41518 through 41519 and 73 FR 68702 through 68704, respectively), we provided a restatement and clarification of the requirements for physician supervision of hospital outpatient diagnostic and therapeutic services that were set forth in the April 2000 OPFS final rule with comment period (65 FR 18524 through 18526). As we stated in those rules, section 1861(s)(2)(C) of the Act authorizes payment for diagnostic services that are furnished to a hospital outpatient for the purpose of diagnostic study. We have further defined the requirements for diagnostic services furnished to hospital outpatients, including requirements for physician supervision of diagnostic services, in §§ 410.28 and 410.32 of our regulations. Section 410.28(e) states that Medicare Part B makes payment for diagnostic services furnished at provider-based departments (PBDs) of hospitals "only when the diagnostic services are furnished under the appropriate level of physician supervision specified by CMS in accordance with the definitions in §§ 410.32(b)(3)(i), (b)(3)(ii), and (b)(3)(iii)." In addition, in the April 2000 OPFS final rule with comment period (65 FR 18526), we stated that our model for the requirement was the requirement for physician supervision of diagnostic tests payable under the MPFS that was set forth in the CY 1998 MPFS final rule (62 FR 59048). In 2000, we also explained with respect to the supervision requirements for individual diagnostic tests that we intended to instruct hospitals and fiscal intermediaries to use the MPFS as a guide pending issuance of updated requirements. For diagnostic services not listed in the MPFS, we stated that fiscal intermediaries, in consultation with their medical directors, would define appropriate supervision levels in order to determine whether claims for these services are reasonable and necessary. Since 2000, we have continued to follow the supervision requirements for individual diagnostic tests as listed in the MPFS Relative Value File. The file is updated quarterly and is available on the CMS Web site at:

<http://www.cms.hhs.gov/PhysicianFeeSched/>.

In the CY 2009 OPPTS/ASC proposed rule and final rule with comment period (73 FR 41518 through 41519 and 73 FR 68702 through 68704, respectively), we also reiterated that direct physician supervision is the standard for physician supervision as set forth in the April 2000 OPPTS final rule with comment period for supervision of hospital outpatient therapeutic services covered and paid by Medicare in hospitals and PBDs of hospitals. We noted that section 1861(s)(2)(B) of the Act authorizes payment for hospital services “incident to physicians’ services rendered to outpatients.” We have further defined the supervision requirements for hospital outpatient therapeutic services and supplies “incident to” a physician’s service in § 410.27 of our regulations. More specifically, § 410.27(f) states: “Services furnished at a department of a provider, as defined in § 413.65(a)(2) of this subchapter, that has provider-based status in relation to a hospital under § 413.65 of this subchapter, must be under the direct supervision of a physician. ‘Direct supervision’ means the physician must be present and on the premises of the location and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician must be present in the room when the procedure is performed.” This language makes no distinction between on-campus and off-campus PBDs.

In the preamble of the April 2000 OPPTS final rule with comment period (65 FR 18525), we further discussed the requirement for physician supervision and the finalization of the proposed regulation text. In that discussion, we stated that the language of § 410.27(f) “applies to services furnished at an entity that is located off the campus of a hospital that we designate as having provider-based status as a department of a hospital in accordance with § 413.65.” We also stated that, for services furnished in a department of a hospital that is located on the campus of a hospital, “we assume the direct supervision requirement to be met as we explain in Section 3112.4(a) of the Intermediary Manual.” We further stated that “we assume the physician supervision requirement is met on hospital premises because staff physicians would always be nearby within the hospital.”

In the CY 2009 OPPTS/ASC proposed rule and final rule with comment period (73 FR 41518 through 41519 and 73 FR 68702 through 68704, respectively), we

restated the existing physician supervision policy for hospital outpatient therapeutic services because we were concerned that some stakeholders may have misunderstood our use of the term “assume” in the April 2000 OPPTS final rule with comment period, believing that our statement meant that we do not require any supervision in the hospital or in an on-campus PBD for hospital outpatient therapeutic services, or that we only require general supervision for those services. This is not the case. It has been our expectation that hospital outpatient therapeutic services are provided under the direct supervision of physicians in the hospital and in all PBDs of the hospital, specifically, both on-campus and off-campus departments of the hospital. The expectation that a physician would always be nearby predates the OPPTS and is related to the statutory authority for payment of hospital outpatient services—that Medicare makes payment for hospital outpatient services “incident to” the services of physicians in the treatment of patients as described in section 1861(s)(2)(B) of the Act. Section 410.27(a)(1)(ii) of the regulations states that Medicare Part B pays for hospital services and supplies furnished incident to a physician service to outpatients if they are provided “as an integral though incidental part of a physician’s services.” In addition, we have stated in Section 20 of Chapter 6 of the Medicare Benefit Policy Manual (Pub. 100–2) that hospitals provide two distinct types of services to outpatients: Services that are diagnostic in nature, and other services that aid the physician in the treatment of the patient. We further defined these therapeutic services and supplies in Section 20.5.1 of the Medicare Benefit Policy Manual, stating “therapeutic services and supplies which hospitals provide on an outpatient basis are those services and supplies (including the use of hospital facilities) which are incident to the services of physicians in the treatment of patients.” We also provide in Section 20.5.1 that services and supplies must be furnished on a physician’s order and delivered under physician supervision. However, the manual indicates further that each occasion of a service by a nonphysician does not need to also be the occasion of the actual rendition of a personal professional service by the physician responsible for the care of the patient. Nevertheless, as stipulated in that same section of the manual “during any course of treatment rendered by auxiliary personnel, the physician must personally see the patient periodically

and sufficiently often enough to assess the course of treatment and the patient’s progress and, where necessary, to change the treatment regimen.”

The expectation that a physician would always be nearby within the hospital also dates back to a time when hospital inpatient services provided in a single hospital building represented the majority of hospital payments by Medicare. Since that time, advances in medical technology, changes in the patterns of health care delivery, and changes in the organizational structure of hospitals have led to the development of extensive hospital campuses, sometimes spanning several city blocks, as well as off-campus and satellite provider-based campuses at different locations. In the April 2000 OPPTS final rule with comment period (65 FR 18525), we described the focus of the direct physician supervision requirement for off-campus PBDs. In the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68703), we stated that we do expect direct physician supervision of all hospital outpatient therapeutic services, regardless of their on-campus or off-campus location, but that we would continue to emphasize the physician supervision requirement for off-campus PBDs. However, we also noted that if there were problems with outpatient care in a hospital or in an on-campus PBD where direct supervision was not in place (that is, the expectation of direct physician supervision was not met), we would consider that to be a quality concern. We noted that we want to ensure that payment is made for high quality hospital outpatient services provided to beneficiaries in a safe and effective manner and consistent with Medicare requirements.

Finally, we noted that the definition of direct supervision in § 410.27(f) for PBDs requires that the physician must be present and on the premises of the location and immediately available to furnish assistance and direction throughout the performance of the procedure. In the April 2000 OPPTS final rule with comment period (65 FR 18525), we further distinguished “on the premises of the location” by stating “* * * a physician must be present on the premises of the entity accorded status as a department of the hospital and therefore, immediately available to furnish assistance and direction for as long as patients are being treated at the site.” We also stated that this characterization does not mean that the physician must be physically in the room where a procedure or service is furnished. We noted in the CY 2009 OPPTS/ASC final rule with comment

period (73 FR 68703) that although we have not further defined the term “immediately available” for this specific context, the lack of timely physician response to a problem in the HOPD would represent a quality concern from our perspective that hospitals should consider in structuring their provision of services in ways that meet the direct physician supervision requirement for HOPD services.

In response to a comment requesting clarification, we also discussed in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68703 through 68704) that a nonphysician practitioner may not provide the physician supervision in a PBD, even if a nurse practitioner’s or a physician assistant’s professional service was being billed as a nurse practitioner or a physician assistant service and not a physician service. We noted that section 1861(r) of the Act defines a physician as follows: “[t]he term ‘physician’, when used in connection with the performance of any function or action, means (1) a doctor of medicine or osteopathy legally authorized to practice medicine and surgery by the State in which he performs such function or action * * *; (2) a doctor of dental surgery or of dental medicine [legally authorized to practice in the State and acting within the scope of his license]; (3) a doctor of podiatric medicine [for certain purposes and to the extent authorized by the State]; (4) a doctor of optometry [for certain purposes and to the extent legally authorized by the State]; or (5) a chiropractor [for certain purposes and to the extent legally authorized by the State and consistent with the Secretary’s standards].” In addition, we pointed out that the conditions of participation for hospitals under § 482.12(c)(1)(i) through (c)(1)(vi) of our regulations require that every Medicare hospital patient is under the care of a doctor of medicine or osteopathy, a doctor of dental surgery or dental medicine, a doctor of podiatric medicine, a doctor of optometry, a chiropractor, or a clinical psychologist; each practicing within the extent of the Act, the Federal regulations, and State law. Further, § 482.12(c)(4) of our regulations requires that a doctor of medicine or osteopathy must be responsible for the care of each Medicare patient with respect to any medical or psychiatric condition that is present on admission or develops during hospitalization and is not specifically within the scope of practice of one of the other practitioners listed in § 482.12(c)(1)(ii) through (c)(1)(vi).

Moreover, section 1861(s)(2)(B) of the Act authorizes payment for hospital services “incident to physicians’

services rendered to outpatients.” We have further defined the requirements for hospital outpatient therapeutic services and supplies “incident to” a physician’s service in § 410.27 of our regulations. Section 410.27(a)(1)(ii) describes payment for hospital outpatient services when they are “an integral though incidental part of a physician’s services.” Also, § 410.27(f) requires that hospital outpatient services provided in PBDs must be under the direct supervision of a physician. We stated that the language of the statute and regulations does not include nonphysician practitioners other than clinical psychologists. Therefore, it would not be in accordance with the law and regulations for a nonphysician practitioner other than a clinical psychologist to be providing the physician supervision in a PBD, even if a nurse practitioner’s or a physician assistant’s professional service was being billed as a nurse practitioner or a physician assistant service and not a physician service.

2. Issues Regarding the Physician Supervision of Hospital Outpatient Services Raised by Hospitals and Other Stakeholders

Although we received a few public comments on the discussion of physician supervision in the CY 2009 OPSS/ASC proposed rule, since publication of the CY 2009 OPSS/ASC final rule with comment period on November 18, 2008, we have received many questions and concerns about the current policies from hospitals and other stakeholders, as we discussed in the CY 2010 OPSS/ASC proposed rule (74 FR 35364). Some stakeholders expressed appreciation for the CMS clarification, stating that the supervision policies were clear and represented needed safeguards for beneficiaries. On the other hand, we have received numerous questions about the application of the policies to hospital outpatient therapeutic services furnished in areas of the hospital that some stakeholders believe have not clearly been discussed, such as the application of direct supervision to hospital outpatient therapeutic services furnished within the main buildings of the hospital that may not be PBDs of the hospital. Some hospitals expressed difficulty in determining whether certain areas of their hospitals were considered provider-based. Other stakeholders cited the direct supervision policy as first articulated in 2000 as problematic because they believe that CMS failed to consider hospitals’ current organizational structures. Some hospitals and other stakeholders

inquired about a physician’s qualifications for providing supervision or questioned whether physician supervision must be provided by a physician in a particular medical specialty. A number of stakeholders challenged the current policy that nonphysician practitioners cannot provide direct supervision for those hospital outpatient therapeutic services they may personally perform or that they may order to be provided by other hospital staff incident to the nonphysician practitioner’s services. In addition, numerous stakeholders, especially rural hospitals, raised budgetary and patient access concerns related to ensuring adequate physician staffing, especially because nonphysician practitioners may not directly supervise the delivery of hospital outpatient therapeutic services. Furthermore, rural hospitals and CAHs raised concerns regarding the inconsistency of the direct supervision requirements for CAHs with other CAH policies, pointing out that the Medicare conditions of participation for CAHs allow nurse practitioners and physician assistants to be responsible for the care of Medicare patients in CAHs. Some stakeholders specifically questioned whether § 410.27(f) applied to CAHs because the term “CAH” is not in the heading of § 410.27, which currently reads “Outpatient hospital services and supplies incident to a physician service: Conditions.” Other stakeholders complained about the significant burden on hospitals to provide direct physician supervision because they believe there is no clear clinical need for such supervision, particularly a uniform level of supervision for all types of hospital outpatient therapeutic services. Some stakeholders challenged the applicability of the direct supervision requirements to specific types of hospital outpatient services, such as partial hospitalization or chemotherapy administration services.

Similar to the issues related to direct supervision of hospital outpatient therapeutic services raised by hospitals and other stakeholders, we have received questions since publication of the CY 2009 OPSS/ASC final rule with comment period regarding the application of physician supervision policies for hospital outpatient diagnostic services, especially with respect to services provided within the main buildings of the hospital that are not PBDs. In addition, some stakeholders have pointed out that there is no site-of-service requirement for hospital outpatient diagnostic services, and that, therefore, hospitals may send

patients to independent diagnostic testing facilities (IDTFs) or other entities to receive diagnostic services under arrangement. They added that although these facilities are not PBDs, the hospital would bill for these services as hospital outpatient services in accordance with the hospital bundling rules. Some of these stakeholders have asked what type of physician supervision is required for diagnostic services provided under arrangement.

A number of stakeholders urged CMS to withdraw or delay the physician supervision policies discussed in the CY 2009 OPPS/ASC final rule with comment period, arguing that this rule included policy changes rather than clarification and, therefore, sufficient opportunity for public notice and comment was not provided. Some further argued that CMS should suspend enforcement of these policies while CMS gathers additional public input and considered alternatives. These stakeholders suggested a variety of additional approaches to soliciting full feedback from the hospital and physician communities on the supervision policies and their impact, including holding an open door forum or town hall meeting and reopening the discussion during the CY 2010 OPPS rulemaking process.

As discussed in the CY 2010 OPPS/ASC proposed rule (74 FR 35365), we provided a restatement and clarification of existing policy in the CY 2009 OPPS/ASC proposed rule (73 FR 41518 through 41519), citing numerous existing statutory, regulatory, manual, and prior rule preamble statements in section XII.A. of that rule specifically titled, "Physician Supervision of HOPD Services." The CY 2009 OPPS/ASC proposed rule provided for a 60-day comment period. We stated that we continue to believe that the CY 2009 restatement and clarification made no change to longstanding hospital outpatient physician supervision policies as incorporated in prior statements of policy, including the codified Federal regulations. In addition, we provided for public notice and comment regarding these physician supervision policies through the CY 2009 OPPS/ASC proposed rule in which, as noted above, we discussed physician supervision in a distinct section of the proposed rule. However, we received only a few public comments on that section. We also noted that the physician supervision policies for hospital outpatient diagnostic and therapeutic services as described in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68702 through 68704) continue to be in

effect for CY 2009. In the CY 2010 OPPS/ASC proposed rule (74 FR 35365), we stated that we have not instructed contractors to delay initiation of enforcement actions or to discontinue pursuing pending enforcement actions regarding the physician supervision of hospital outpatient services.

However, while we did not propose to withdraw the longstanding physician supervision policies for hospital outpatient services in CY 2009, we stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35365) that we had extensively considered the many questions and concerns on this topic raised to us by stakeholders in the course of developing the proposed rule in order to assess whether proposed changes to the existing policies should be considered. We appreciated the many detailed comments and suggestions interested stakeholders have raised in the first few months since publication of the CY 2009 OPPS/ASC final rule with comment period. We considered a wide variety of potential modifications to our physician supervision policies in response to this information about current health care delivery practices and challenges. The dialogue with interested stakeholders provided us with sufficient information to develop proposals for certain changes to the supervision policies for hospital outpatient services for CY 2010 in order to take into full consideration current clinical practice and patterns of care, the need to ensure patient access, the associated hospital and physician responsibilities, consistency among requirements for different sites of services, and other important factors. We indicated our belief that the proposals we were making for CY 2010 would address many of the concerns and questions regarding our existing policies that had been raised to us by stakeholders over the first 6 months of CY 2009. In the CY 2010 OPPS/ASC proposed rule (74 FR 35365), we welcomed robust public comments regarding our CY 2010 proposals for physician supervision in order to inform our decisions regarding final policies for CY 2010.

In considering the questions and concerns that had been raised over the first 6 months of CY 2009, we identified three areas within our existing hospital outpatient physician supervision policies for which we stated our belief that proposals of policy changes were appropriate for CY 2010, two related to the supervision of therapeutic services and one related to the supervision of diagnostic services. Our specific CY 2010 proposals, including the proposed changes to our regulations to conform to

these proposals, the public comments on the proposals and our responses, and our final CY 2010 supervision policies for hospital outpatient therapeutic and diagnostic services are discussed below.

3. Policies for Direct Supervision of Hospital and CAH Outpatient Therapeutic Services

In the CY 2010 OPPS/ASC proposed rule (74 FR 35365), we proposed that nonphysician practitioners, specifically physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives, may directly supervise all hospital outpatient therapeutic services that they may perform themselves in accordance with their State law and scope of practice and hospital-granted privileges, provided that they continue to meet all additional requirements, including any collaboration or supervision requirements as specified in the regulations at §§ 410.74 through 410.77. Clinical psychologists may already provide direct supervision, as we mentioned in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68703 through 68704) because they, along with physicians (as defined in section 1861(r)(1) of the Act), may be responsible for the care of a hospital patient, as discussed in the Medicare conditions of participation for hospitals in § 482.12(c) of our regulations. We stated our belief that allowing certain nonphysician practitioners (nurse practitioners, physician assistants, clinical nurse specialists, and certified nurse-midwives) to provide direct supervision of certain hospital outpatient therapeutic services is appropriate because, even though these practitioners are not physicians, they are recognized in statute and regulation as providing services that are analogous to physicians' services. Medicare Part B covers the professional services of clinical psychologists, nurse practitioners, physician assistants, clinical nurse specialists, and certified nurse-midwives when the services would be physicians' services if furnished by a physician (a doctor of medicine or osteopathy, as set forth in section 1861(r)(1) of the Act). Their services are described in §§ 410.71(a), 410.74(a), 410.75(a) and (c), 410.76(a) and (c), and 410.77(a), respectively, of our regulations. Medicare also makes payment for services provided incident to the services of these nonphysician practitioners as specified in §§ 410.71(a)(2)(iii), 410.74(b), 410.75(d), 410.76(d), and 410.77(c), respectively.

We also noted that section 1861(r) of the Act does not include clinical psychologists, nurse practitioners,

physician assistants, clinical nurse specialists, or certified nurse-midwives in the definition of a physician. However, as previously mentioned, the conditions of participation for hospitals at § 482.12(c)(1)(vi) of our regulations do include clinical psychologists as practitioners who may be responsible for the care of Medicare patients. The conditions of participation at §§ 482.12(c)(1)(i) through (c)(1)(vi) require that every Medicare hospital patient be under the care of a doctor of medicine or osteopathy, a doctor of dental surgery or dental medicine, a doctor of podiatric medicine, a doctor of optometry, a chiropractor, or a clinical psychologist; each practicing in accordance with the Act, Federal regulations, and State law. Further, § 482.12(c)(4) of our regulations requires that a doctor of medicine or osteopathy must be responsible for the care of each Medicare patient with respect to any medical or psychiatric condition that is present on admission or develops during hospitalization and is not specifically within the scope of practice of one of the other practitioners listed in §§ 482.12(c)(1)(ii) through (c)(1)(vi). Also, as permitted by State law, certain nonphysician practitioners may admit individuals to a hospital or CAH and order and provide therapeutic services to them. Since 1998, we have allowed payment for the professional services of these nonphysician practitioners in addition to payment for physicians' services when the nonphysician practitioner's professional services are furnished in an HOPD. We also have made outpatient facility payments to the hospital for those facility services provided incident to the professional services of these nonphysician practitioners (63 FR 58873). In addition, the conditions of participation for CAHs at § 485.631 require that a doctor of medicine or osteopathy, a nurse practitioner, a physician assistant, or a clinical nurse specialist be available to furnish patient care services at all times the CAH operates. A doctor of medicine or osteopathy must be present for sufficient periods of time to provide medical direction, medical care services, consultation and supervision as described in the conditions of participation and must be available through radio or telephone contact for assistance with medical emergencies or patient referral.

Taking into consideration the totality of these existing conditions and requirements, we proposed to revise § 410.27 of the regulations to make clear that Medicare Part B payment may be made for hospital outpatient services

and supplies furnished incident to the services of a physician, clinical psychologist, nurse practitioner, physician assistant, clinical nurse specialist, or certified nurse-midwife service; and to add that, effective January 1, 2010, clinical psychologists, nurse practitioners, physician assistants, clinical nurse specialists, or certified nurse-midwives may provide direct supervision for hospital outpatient therapeutic services that they may perform themselves under State law and within their scope of practice and hospital-granted privileges in the context of the existing requirements in §§ 410.71, 410.74, 410.75, 410.76, and 410.77. However, we note that, in the CY 2010 OPPTS/ASC proposed rule (74 FR 35366), we proposed that the direct supervision of CR, ICR, and PR services must be furnished by a doctor of medicine or osteopathy, as specified in the proposed coverage policy and regulations for CR, ICR, and PR services. We also noted that Medicare does not make a payment to a physician under the MPFS when the physician solely provides the direct physician supervision of hospital outpatient therapeutic services but furnishes no direct professional services to a patient. This also would apply to the supervision of hospital outpatient therapeutic services provided by nonphysician practitioners.

We also indicated that we did not propose to modify requirements relating to physician supervision or collaboration for these nonphysician practitioners. In regard to the supervision of physician assistants, § 410.74(a)(iv) requires that physician assistants perform services under the general supervision of a physician. We have further defined this general supervision in section 190(c) of Chapter 15 of the Medicare Benefit Policy Manual. Section 190(c) states that "the PA's physician supervisor (or a physician designated by the supervising physician or employer as provided under State law or regulations) is primarily responsible for the overall direction and management of the physician assistant's professional activities and for assuring that the services provided are medically appropriate for the patient. The physician supervisor (or physician designee) need not be physically present with the physician assistant when a service is being furnished to a patient and may be contacted by telephone, if necessary, unless State law or regulations require otherwise."

The requirements for collaboration of nurse practitioners are defined in § 410.75(c)(3) of the regulations and

Section 200(D) of Chapter 15 of the Medicare Benefit Policy Manual. The requirements for clinical nurse specialists are located in § 410.76(c)(3) of the regulations and Section 210(D) of Chapter 15 of the Medicare Benefit Policy Manual. These sections define collaboration as a process in which the nurse practitioner or the clinical nurse specialist works with one or more physicians (doctors of medicine or osteopathy) to deliver health care services within the scope of the practitioner's expertise, with medical direction and appropriate supervision as required by the law of the State in which the services are being furnished. In the absence of more stringent State law requirements governing collaboration, collaboration is to be evidenced by the nurse practitioner or the clinical nurse specialist documenting his or her scope of practice and indicating the relationships that he or she has with physicians to deal with issues outside their scope of practice. The collaborating physician does not need to be present with the nurse practitioner or clinical nurse specialist when the services are furnished or to make an independent evaluation of each patient who is seen by the nurse practitioner or clinical nurse specialist.

In addition, for CY 2010, we proposed to refine the definition of direct supervision of hospital outpatient therapeutic services for those services furnished in a hospital and in on-campus PBDs of a hospital. For services furnished on a hospital's main campus, we proposed that direct supervision means that the supervisory physician or nonphysician practitioner must be present on the same campus, in the hospital or the on-campus PBD of the hospital as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. We proposed to add a new paragraph (a)(1)(iv)(A) to § 410.27 to reflect this requirement. We also proposed to define "in the hospital" in new paragraph § 410.27(g) as meaning areas in the main building(s) of a hospital that are under the ownership, financial, and administrative control of the hospital; that are operated as part of the hospital; and for which the hospital bills the services furnished under the hospital's CCN. Therefore, to be present in the hospital or the on-campus PBD of the hospital and immediately available requires that the physician or nonphysician practitioner must be physically present in areas on the campus of the hospital that are part of

the hospital, including on-campus PBDs, that are operated by the hospital, and where services furnished in those areas are billed under the hospital's CCN. Under the CY 2010 proposal, the supervisory physician or nonphysician practitioner of the hospital's outpatient therapeutic services may not be located in any other entity, such as a physician's office, IDTF, co-located hospital, or hospital-operated provider or supplier such as a skilled nursing facility (SNF), end-stage renal disease (ESRD) facility, or home health agency (HHA), or any other nonhospital space that may be co-located on the hospital's campus, as "hospital campus" is defined in § 413.65(a)(2) of the regulations.

We stated that while we have not previously specified in policy guidance a definition for the term "immediately available" with respect to services provided in areas of the hospital on its main campus that are not PBDs, we believe that the existing definitions of direct supervision in §§ 410.27(f) and 410.32(b)(3)(ii) that apply to PBDs and physician office settings indicate that the physician must be physically present in order to provide direct supervision of services. With regard to services provided in PBDs of hospitals or physicians' offices, these regulations specify that the physician must be present in the PBD or in the office suite, respectively. Thus, we have previously established that direct supervision requires immediate physical presence. While we also have not specifically defined the word "immediate" for direct supervision in terms of time or distance, we noted that the general definition of the word means "without interval of time." Therefore, the supervisory physician or nonphysician practitioner could not be immediately available while, for example, performing another procedure or service that he or she could not interrupt. In addition, we stated that we understand that advances in medical technology, changes in the patterns of health care delivery, and changes in the organizational structure of hospitals have led to the development of extensive hospital campuses, sometimes spanning several city blocks. However, in the context of direct physician or nonphysician practitioner supervision, we believed that it would be neither appropriate nor "immediate" for the supervisory physician or nonphysician practitioner to be so physically far away on the main campus from the location where hospital outpatient services are being furnished that he or she could not intervene right away. As we stated in the CY 2009

OPPS/ASC final rule with comment period (73 FR 68703), if there were problems with outpatient care in a hospital or in an on-campus PBD where the requirement for direct supervision was not met, we would consider that to be a quality concern. Appropriate supervision is a key aspect of the delivery of safe and high quality hospital outpatient services that are paid under Medicare.

In addition, the definition of direct supervision in existing § 410.27(f) has included and would continue to specify under our CY 2010 proposal that the physician or nonphysician practitioner must be available to furnish assistance and direction throughout the performance of the procedure. This means that the physician or nonphysician practitioner must be prepared to step in and perform the service, not just to respond to an emergency. This includes the ability to take over performance of a procedure and, as appropriate to both the supervisory physician or nonphysician practitioner and the patient, to change a procedure or the course of treatment being provided to a particular patient. We originally stated in the April 2000 OPPS final rule (65 FR 18525) that the physician does not "necessarily need to be of the same specialty as the procedure or service that is being performed." We also have stated in manual guidance that hospital medical staff that supervises the services "need not be in the same department as the ordering physician" (Section 20.5.1 of Chapter 6 of the Medicare Benefits Policy Manual). However, in order to furnish appropriate assistance and direction for any given service or procedure, we believed the supervisory physician or nonphysician practitioner must have, within his or her State scope of practice and hospital-granted privileges, the ability to perform the service or procedure.

We did not propose significant changes to the definition or requirements for direct supervision in off-campus PBDs of the hospital other than to allow nonphysician practitioners to provide direct supervision in these PBDs for the services that these practitioners may perform. With respect to off-campus PBDs of hospitals, direct supervision would continue to mean that the physician or nonphysician practitioner must be in the off-campus PBD and immediately available to furnish assistance and direction throughout the performance of the procedure. We proposed to revise existing § 410.27(f) and include the revised language as § 410.27(a)(1)(iv) and make a technical

change in § 410.27(a)(i)(iv)(B) to clarify the current language by removing "present and on the premises of the location" and replacing it with "present in the off-campus provider-based department." While the meaning of this provision is the same, we believed that this proposed modification to the language defining direct supervision is more consistent with the language of the other proposed changes to § 410.27. As we clarified in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68704), the supervisory physician for hospital outpatient therapeutic services must be in each PBD of a particular off-campus outpatient location, but that does not mean that the physician must be in the room when the procedure is performed. In the April 2000 OPPS final rule (65 FR 18525), we responded to public commenters who asserted that requiring a physician to be onsite at a PBD throughout the performance of all "incident to" (therapeutic) services would be burdensome and costly for hospitals where there are a limited number of physicians available to provide coverage, particularly in rural settings. We disagreed then that the supervision requirement was unnecessary and burdensome because hospitals, prior to 2000, were already required to "meet a direct supervision of 'incident to' services requirement that is unrelated to the provider-based rules. That is, we require that hospital services and supplies furnished to outpatients that are incident to physician services be furnished on a physician's order by hospital personnel and under a physician's supervision" (Section 3112.4 of the Medicare Intermediary Manual). In addition, when we discussed the "assumption" or expectation that the physician supervision requirement is met on the hospital's main campus in the April 2000 OPPS final rule (65 FR 18525), we specifically did not extend that assumption to off-campus departments of the hospital. We stated that we continue to believe that it would be inappropriate to allow one physician or nonphysician practitioner to supervise all services being provided in all PBDs at a particular off-campus outpatient location. Since first allowing off-campus sites to be considered PBDs of hospitals, we have placed particular emphasis on ensuring the quality and safety of the services provided in these locations, which can be many miles from the main hospital campus, through both additional provider-based requirements in § 413.65(e) and our emphasis on direct physician supervision under existing § 410.27(f). In addition, because

the physician or nonphysician practitioner must be immediately available and have, within his or her State scope of practice and hospital-granted privileges, the ability to perform the services being supervised, we believed it would be highly unlikely that one physician or nonphysician practitioner would be both immediately available at all times that therapeutic services are being provided and would have the knowledge and ability to adequately supervise all services being performed at once in multiple off-campus PBDs at a single location.

To reflect these proposed changes for the provision of direct supervision of therapeutic services provided to hospital outpatients in our regulations, we proposed to revise the language of the existing § 410.27(f) and include the revised language in a new paragraph (a)(1)(iv) of § 410.27 to specify that direct physician or nonphysician practitioner supervision of hospital outpatient therapeutic services is required for Medicare Part B payment. We proposed to add a new paragraph (a)(1)(iv)(A) to § 410.27 to state that, for services provided on the hospital's main campus, direct supervision means that the physician or nonphysician practitioner must be present on the same campus, in the hospital or on-campus PBD of the hospital, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician or nonphysician practitioner must be in the room when the procedure is performed. We also proposed to add new paragraph (a)(1)(iv)(B) to § 410.27 to reflect that, for off-campus PBDs of hospitals, the physician or nonphysician practitioner must be present in the off-campus PBD, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician or nonphysician practitioner must be in the room when the procedure is performed. As we stated previously, the proposed language of paragraph (a)(1)(iv)(B) is similar to existing § 410.27(f) that we proposed to revise. Furthermore, we proposed to make a technical change to clarify the language in this paragraph to remove "present and on the premises of the location" and replace it with "present in the off-campus provider-based department." Also, in the CY 2010 OPFS/ASC proposed rule (74 FR 35368) and as proposed in the CY 2010 MPFS proposed rule (74 FR 33674), we proposed that the direct supervision of

CR, ICR, and PR services must be furnished by a doctor of medicine or osteopathy, as specified in proposed §§ 410.47 and 410.49, respectively. We proposed to include this exception in proposed paragraphs (a)(1)(iv)(A) and (a)(1)(iv)(B) in § 410.27. In addition, we proposed to add a new paragraph (f) to § 410.27 to define a nonphysician practitioner for purposes of § 410.27 as a clinical psychologist, a physician assistant, a nurse practitioner, a clinical nurse specialist, or a certified nurse-midwife. Proposed new § 410.27(a)(1)(iv) would provide that these nonphysician practitioners may directly supervise services that they could furnish themselves in accordance with State law and within their scope of practice and hospital-granted privileges, as long as all requirements for coverage, including the physician supervision or collaboration for these nonphysician practitioners, are met in accordance with §§ 410.71, 410.74, 410.75, 410.76, and 410.77, respectively. We also proposed to define "in the hospital" in new paragraph § 410.27(g) to mean areas in the main building(s) of the hospital that are under the ownership, financial, and administrative control of the hospital; that are operated as part of the hospital; and for which the hospital bills the services furnished under the hospital's CCN. Finally, we proposed to make a technical correction to the title of § 410.27 to read, "Outpatient hospital or CAH services and supplies incident to a physician service: Conditions" to clarify in the title that the requirements for payment of hospital outpatient therapeutic services incident to a physician or nonphysician practitioner service in that section apply to both hospitals and CAHs. Similarly, we proposed to include the phrase "hospital or CAH" throughout the text of § 410.27 wherever the text currently refers just to "hospital." The omission of the term "CAH" from § 410.27 was a drafting oversight. However, we have applied the requirements of § 410.27, including "incident to" requirements such as the site-of-service requirement and physician supervision, as well as other hospital policies, such as the bundling rules, to CAHs, just as we have in 42 CFR Part 409 (Subparts A through D and F through H) and § 410.28 and § 413.65 of the regulations where CAHs are explicitly mentioned.

Comment: All commenters who addressed the proposed direct supervision by nonphysician practitioners (specifically physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives) supported the proposal.

Some commenters also requested that CMS include licensed clinical social workers in the list of nonphysician practitioners who are permitted to supervise outpatient psychiatric services and partial hospitalization program (PHP) services. Other commenters requested that pharmacists be permitted to supervise medication therapy management services and that specialty certified registered nurses, such as wound care nurses, also be able to provide the direct supervision of hospital outpatient therapeutic services. One commenter stated that because physicians do not furnish nursing services or the services of other ancillary health professionals, they should not be expected to supervise those services and it would be inappropriate to expect physicians to accept responsibility for care that they have not personally furnished.

Response: We appreciate the commenters' support for our proposal to revise § 410.27 of the regulations to allow certain nonphysician practitioners to directly supervise services that they may perform themselves under their State scope of practice and hospital-granted privileges in the context of the existing requirements in §§ 410.71, 410.74, 410.75, 410.76 and 410.77. We agree with the commenters that we should add licensed clinical social workers to the group of nonphysician practitioners permitted to provide direct supervision for hospital outpatient therapeutic services. We believe this is appropriate because licensed clinical social workers are recognized under the Medicare program as independent practitioners who may furnish services for the diagnosis and treatment of mental illness that they are legally authorized to perform under State law of the State in which the services are performed. We further agree with the commenters that the inclusion of licensed clinical social workers would help to ensure continued access to mental health services, especially in rural hospitals and CAHs where other types of practitioners may be less available. We emphasize though, that licensed clinical social workers, like other nonphysician practitioners, may only supervise those therapeutic services within their own scope of practice and hospital-granted privileges. We are not expanding the regulations further to allow supervision by pharmacists, registered nurses, or other medical professionals. These professionals are not recognized in the Social Security Act as providing services that would be physicians' services if performed by a physician and

they may not enroll in Medicare as independent practitioners and receive payment directly for their professional services.

With regard to the comments about physician scope of practice and supervision, we remind hospitals that the only statutory basis for payment of hospital outpatient therapeutic services is incident to the services of a physician, meaning the services are ordered by the physician or qualified practitioner and furnished as an integral though incidental part of his or her services. It follows, then, that a qualified physician or nonphysician practitioner would supervise the provision of those services to ensure the service or procedure is being furnished appropriately.

Comment: One commenter supported the proposed requirements for the direct physician supervision of CR programs and requested that CMS confirm that the proposed definition of “direct supervision” that would apply to therapeutic services in the HOPD would also apply to CR services. However, many commenters, especially CAHs and rural hospitals, asked that CMS permit physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives who are functioning within their State licensure and scope of practice and who are permitted to supervise the services under the hospital or CAH bylaws to supervise CR, ICR, and PR services.

Response: As discussed in detail in sections II.G.8. and II.G.9. of the CY 2010 MPFS final rule with comment period and in section XII.B.4. of this final rule with comment period, section 144(a)(1) of Public Law 100–275 imposes strict requirements for the direct physician supervision of PR, CR, and ICR services and gives us no flexibility to modify the requirement. Therefore, we are finalizing our CY 2010 proposal, without modification, to require the direct physician supervision (by a doctor of medicine or a doctor of osteopathy) of PR, CR, and ICR services that are furnished to hospital outpatients. We note that we define “direct supervision” with regard to what it means to be immediately available and accessible for medical consultation and medical emergencies in the same manner for PR, CR, and ICR services as we do for other therapeutic services furnished in HOPDs, as discussed below. Also, the final definitions of direct supervision in § 410.27 for therapeutic services provided on campus and in off-campus PBDs also apply. These definitions and the final regulation text of

§§ 410.27(a)(1)(iv)(A) and (B) are discussed in detail below.

Comment: With respect to the proposed definition of direct supervision of hospital outpatient therapeutic services, some commenters fully supported the proposals as appropriate and necessary for ensuring that Medicare beneficiaries receive safe and high quality care. Many commenters acknowledged that the proposal to broaden the location of the supervising physician from in the department on campus to “in the hospital” would give hospitals significantly more flexibility. However, the commenters stated that, while the proposal would be more flexible, it would still limit access to care and would cost hospitals and the Medicare program. Other commenters questioned why CMS has a supervision requirement at all, stating that there is no such specified requirement for hospital inpatient services. Many commenters believed that the proposals would not help CAHs and rural hospitals, where physicians often maintain offices off-campus and qualified nonphysician practitioners may not be readily available to provide services in the hospital. The commenters claimed that the proposed definition of direct supervision would mean that CAHs and rural hospitals would be required to hire staff solely to supervise services and that this extra cost would force these hospitals and CAHs to eliminate services. These commenters requested that CMS not apply the “incident to” requirements of § 410.27 to CAHs.

Response: We appreciate the commenters’ support and the acknowledgement that we are striving to provide more flexibility for hospitals, while maintaining appropriate supervision of hospital outpatient therapeutic services that helps to ensure safe, high quality care and providing Medicare payment that is consistent with the statutory requirements for coverage. We have received numerous comments and questions since publication of the CY 2009 OPFS/ASC final rule with comment period, met with interested stakeholders to hear their questions and concerns throughout the year, and reviewed many thoughtful, wide-ranging comments on the CY 2010 OPFS/ASC proposed rule. In considering all of this information, we have taken into full consideration current clinical practice and patterns of care, the need to ensure patient access, the associated hospital and physician responsibilities, consistency among requirements for different sites of services, and other important factors. We believe that our final policies

address many of the concerns and questions regarding our proposals. Through the expansion of supervisory authority to nonphysician practitioners and a modification to the requirement for direct supervision of on-campus therapeutic services, discussed in more detail below, we believe our final policies allow more flexibility for hospitals and CAHs and help ensure access to care for Medicare beneficiaries while maintaining our standards for safe, high quality care and consistent interpretation of longstanding regulations.

We disagree with the commenters who suggested that there should be no supervision requirements for hospital outpatient services because there are not similar codified supervision requirements for hospital inpatient services. Hospitals provide a wide variety of complex services to their outpatients who may or may not have an established relationship with the supervising physician or nonphysician practitioner and hospital staff on the day the hospital outpatient services are furnished. A treating physician or nonpractitioner in the community may not even be aware that ordered outpatient services are being furnished by the hospital on a given day. Therefore, we believe it is appropriate for CMS to set requirements for the safe and effective diagnosis and treatment of Medicare beneficiaries, including standards for the appropriate supervision of hospital outpatient services by a physician or nonphysician practitioner, in accordance with the statutory basis for payment of hospital outpatient services in section 1861(s)(2)(B) of the Act, which is “incident to physicians’ services rendered to outpatients.”

We set the standard for direct supervision of hospital outpatient therapeutic services that we have held since the implementation of the OPFS in 2000, and we have assumed that this standard was being met because we assumed that a physician would always be nearby in the hospital. Given that hospital inpatients generally have medically complex conditions requiring a high level of acute care, we have not established explicit supervision requirements in regulations because we believe hospitals would have physicians or other qualified practitioners available at all times that complex hospital inpatient services are being furnished. If this is not the case, we may consider addressing the supervision of hospital inpatient services in the future.

In regard to hospital and CAH concerns about hiring physician and nonphysician staff solely for

supervisory services, we reiterate that the supervisor need only be available when outpatient therapeutic services and procedures are being furnished, meaning that many services or departments would not require 24 hour per day, 7 days per week direct supervision, as some commenters believed. We also believe that allowing the supervisory physician or nonphysician practitioner to be located anywhere on the hospital campus, as discussed more fully below, should alleviate this concern for many hospitals and CAHs. In addition, we remind CAHs that the conditions of participation for CAHs at § 485.631 require that a doctor of medicine or osteopathy, a nurse practitioner, a physician assistant, or a clinical nurse specialist is available “to furnish patient care services at all times the CAH operates” (emphasis added). It follows then that this physician or nonphysician practitioner who is available at all times the CAH operates would be directly furnishing services that are within his or her State scope of practice and CAH-granted privileges and that the CAH would not be furnishing services that are not within this practitioner’s scope.

Comment: Many commenters stated that it is overly restrictive and arbitrary to not allow direct supervision by practitioners located in other entities on-campus and to require the immediate physical presence of the physician. Several commenters pointed out that, because of the varying ways that hospitals have structured their services and campuses, a physician’s office may be next door or closer to the HOPD in which services that he or she would be supervising are furnished, than a practitioner located in another HOPD. Other commenters stated that the proposal would preclude physicians from taking a lunch break, patronizing the retail establishments in the hospital, or going to other areas such as parking lots.

The commenters were particularly concerned about specialized services such as chemotherapy, blood transfusions, and radiation therapy services. Several hospital associations and other commenters requested that CMS remove the phrase “immediately available to furnish assistance and direction throughout the performance of the procedure.” They instead recommended that CMS redefine direct supervision for therapeutic services to mean that the physician may be on or in close proximity to the campus and able to respond in a timely manner according to hospital’s policies and bylaws. The commenters also believed that “immediate availability” does not

and should not mean immediate physical presence and that requiring physical presence in every instance is impractical. Instead, the commenters believed the supervising practitioner should be able to directly supervise services and procedures remotely by telephone, radio, robotic device, or electronic media.

Response: We appreciate the many public comments that we received on the proposed definition of direct supervision for hospital outpatient therapeutic services provided on the campus of the hospital. We acknowledge the comments related to hospital building and campus structures and the physical proximity of physicians’ offices and other entities to the hospital department where a particular hospital outpatient service is furnished. We agree with the commenters that allowing the supervising physician to be in nonhospital space on the campus could make it easier for a supervising physician or nonphysician practitioner to respond immediately. Therefore, we believe it would be appropriate to allow the supervising physician or nonphysician practitioner to be located anywhere on the same campus of the hospital, as long as he or she was immediately available to furnish assistance and direction throughout the performance of the procedure.

This is consistent with our longstanding definition of “direct supervision” as it has been applied across settings in terms of the immediate physical presence of the physician and what it means to “furnish assistance and direction throughout the performance of the procedure.” However, we continue to believe that the supervisory physician or nonphysician practitioner could not be immediately available while, for example, performing another procedure or service that he or she could not interrupt. It also would be neither appropriate nor “immediate” for the supervisory physician or nonphysician practitioner to be so physically far away on the main campus from the location where hospital outpatient services are being furnished that he or she could not intervene right away.

As we stated in the CY 2010 OPPI/ASC proposed rule (74 FR 35367), we believe that the existing definitions of direct supervision in §§ 410.27(f) and 410.32(b)(3)(ii) that apply to PBDs and physicians’ office settings indicate that the physician must be physically present in order to provide direct supervision of services. With regard to services provided in PBDs of hospitals or physicians’ offices, for at least 9 years

these regulations have specified that the physician must be present in the PBD or in the office suite, respectively. Thus, we have previously established that direct supervision requires immediate physical presence. Medicare only covers hospital outpatient therapeutic services furnished incident to physicians’ services, yet a hospital service ordered by a physician may be furnished on a day when the beneficiary does not receive services directly from a physician. Therefore, we believe it is important to retain direct supervision as the standard.

With regard to the commenters’ recommendations that CMS redefine direct supervision for therapeutic services to mean that the physician may be “on or in close proximity to the campus” and “able to respond in a timely manner” according to hospital’s policies and bylaws, we believe that the suggested new definition of direct supervision for on-campus hospital outpatient therapeutic services would be wholly inconsistent with the definition of the term as previously codified in at least two Medicare sections of the Code of Federal Regulations. We disagree with the commenters that describing immediate availability as without lapse of time would be read so narrowly as to require that the supervising physician or nonphysician practitioner must be standing in the room next to the nonphysician personnel furnishing the service. A similar argument could be made that the phrase “able to respond in a timely manner” is so vague that a supervising physician or nonphysician practitioner could interpret it to mean that arriving within an hour or hours would be reasonable. In addition, we note that the definition proposed by commenters is virtually identical in interpretation to the current existing definition of general supervision. Section 410.32(b)(3)(i) of the regulations defines general supervision to mean that “the procedure is furnished under the physician’s overall direction and control, but the physician’s presence is not required during the performance of the procedure.” We have historically interpreted this to mean that the physician may be available by telephone or other electronic device to supervise and direct the nonphysician personnel furnishing the service. This lower standard of general supervision would not ensure the immediate presence of a qualified practitioner in the hospital to furnish assistance and direction to hospital personnel providing a wide array of complex therapeutic services to hospital outpatients. Several of the

commenters requesting this relaxed definition of direct supervision also asked that certain services be designated as requiring only general supervision. We are unclear about the commenters' understanding of the definition of general supervision if their suggested definition of direct supervision requires no physical presence and only specifies availability in a reasonable period of time by telephone, radio, robotics, or telemedicine.

We have set direct supervision as a minimum standard for supervision of all Medicare hospital outpatient therapeutic services. We do not believe that this standard or the definition, as codified, precludes a hospital from developing bylaws, credentialing procedures, and policies that it believes are appropriate to ensure that all hospital patients receive high quality services in a safe and effective manner. We believe that hospitals take quality of care and patient safety very seriously, and we understand that hospitals are subject to accreditation requirements. Considering that hospitals provide a wide array of very complex services and procedures, including surgeries, we would expect that hospitals already have the leadership, credentialing procedures, bylaws, and other policies in place to ensure that services furnished to Medicare beneficiaries are being provided only by qualified practitioners in accordance with all applicable laws, regulations, and coding guidance. For services not furnished directly by a physician, we would expect that these bylaws and policies would ensure that the services are being supervised in a manner commensurate with the complexity of the service, including personal supervision where appropriate.

Comment: Many commenters disagreed with statements in the proposed rule that the supervising physician should have, within his or her scope of practice and hospital-granted privileges, the knowledge, ability, and hospital privileges to perform the services being supervised. Some commenters stated that the supervisor should be required to provide only medical consultation and attend to medical emergencies, but should not be expected to intervene or change the course of treatment because this usurps the responsibility of treating physician. Other commenters stated that the supervisor should be "clinically appropriate" to supervise the outpatient therapeutic services.

Response: As we explained in the CY 2010 OPPS/ASC proposed rule (74 FR 35367), we believe the supervising physician or nonphysician practitioner

must be prepared to step in and perform the service, not just to respond to an emergency. This includes the ability to take over performance of a procedure and, as appropriate to both the supervisory physician or nonphysician practitioner and the patient, to change a procedure or the course of treatment being provided to a particular patient. We originally stated in the April 2000 OPPS final rule (65 FR 18525) that the physician does not "necessarily need to be of the same specialty as the procedure or service that is being performed." We also have stated in manual guidance that hospital medical staff that supervises the services "need not be in the same department as the ordering physician" (Section 20.5.1 of Chapter 6 of the Medicare Benefits Policy Manual). We understand hospitals' concerns, and note that we would not expect that a supervising physician would operate in a vacuum, making all decisions without informing or consulting the patient's treating physician or nonphysician practitioner. This would be illogical and inappropriate for good medical practice. However, in order to furnish appropriate assistance and direction for any given service or procedure, we continue to believe the supervisory physician or nonphysician practitioner must have, within his or her State scope of practice and hospital-granted privileges, the ability to perform the service or procedure. We believe that our interpretation of the requirement means that the supervisor must be a person who is "clinically appropriate" to supervise the service or procedure. We believe it is inappropriate for a supervisory physician or nonphysician practitioner to be responsible for patients, hospital staff, and services that are outside the scope of their knowledge, skills, licensure, or hospital-granted privileges.

This interpretation of the previously codified language is consistent with our longstanding application of direct supervision across settings in terms of the physical presence of the physician and what it means to "furnish assistance and direction throughout the performance of the procedure." We do not believe that allowing a supervisor to be responsible for emergencies only would satisfy the standard to "furnish assistance and direction throughout the performance of the procedure" as the language has historically been interpreted for physicians' offices and PBDs. We disagree with commenters who stated that the historical intent of direct supervision has been for a supervising physician to provide

guidance and direction without expecting that professional to be able to perform the service or procedure and that performance of the procedure applies only to personal supervision. It would be unreasonable to think that a physician or nonphysician practitioner could competently assist and direct a procedure for which they do not have sufficient knowledge and skills to perform or redirect the procedure or service.

Comment: Several commenters requested additional guidance as to what CMS considers hospital outpatient "incident to" services. One commenter requested clarification on the applicability of "incident to" to emergency department services. The commenter believed that the "incident to" provision for hospital outpatient services may not be applicable to emergency department services because the emergency department physician would be immediately available in the area to care for patients, but would not have previously seen and established a relationship with the patient, as this commenter believed is required. Other commenters believed that the requirements for supervision of hospital outpatient therapeutic services should be specific to each clinical service and should be designated as either general or direct, as for diagnostics. Some commenters asked for clarification of how to report services in a Condition Code 44 situation when the patient received care as an inpatient, with no direct supervision, and the hospital then changed the patient's status to outpatient.

Response: As previously stated in this discussion, § 410.27(a)(1)(ii) of the regulations states that Medicare Part B pays for hospital services and supplies furnished incident to a physician service to outpatients if they are provided "as an integral though incidental part of a physician's services." In addition, we have stated in Section 20 of Chapter 6 of the Medicare Benefit Policy Manual that hospitals provide two distinct types of services to outpatients: services that are diagnostic in nature and other services that aid the physician in the treatment of the patient. We further defined these therapeutic services and supplies in Section 20.5.1 of Chapter 6 of the Medicare Benefit Policy Manual, stating "therapeutic services and supplies which hospitals provide on an outpatient basis are those services and supplies (including the use of hospital facilities) which are incident to the services of physicians in the treatment of patients." In essence, all hospital outpatient services that are not

diagnostic are services that aid the physician in the treatment of the patient, and are called therapeutic services. These are the services for which Medicare makes a hospital facility payment only when they are provided "incident to" the services of a physician.

We also provide in Section 20.5.1 that services and supplies must be furnished on a physician's order and delivered under physician supervision. However, the manual indicates further that each occasion of a service by a nonphysician does not need to also be the occasion of the actual rendition of a personal professional service by the physician responsible for the care of the patient. Nevertheless, as stipulated in that same section of the manual "during any course of treatment rendered by auxiliary personnel, the physician must personally see the patient periodically and sufficiently often enough to assess the course of treatment and the patient's progress and, where necessary, to change the treatment regimen." Section 20.5.1 also explicitly includes clinic and emergency room services as examples of hospital outpatient services that are provided incident to the services of a physician. We note in this section that the policies for hospital services incident to physicians' services rendered to outpatients differ in some respects from policies that pertain to "incident to" services furnished in office and physician-directed clinic settings. Those requirements are discussed in Section 60 of Chapter 15 of the Medicare Benefit Policy Manual. The commenter incorrectly believed that for hospital services to be "incident to" a physician's services, the physician must have previously seen and established a relationship with the patient. When a patient presents to an emergency department or a hospital clinic and a physician examines the patient and orders services to be provided, the provision of those services is incident to the services of that physician.

We recognize the potential benefit to hospitals of specifically designating supervision requirements for individual therapeutic services based on clinical complexity, especially for less complex services that we might deem to require general supervision. We note that the commenters requesting individual designations mentioned only defining services as requiring general or direct supervision. However, as we discussed earlier in this section, direct supervision represents a minimum standard currently applicable to all outpatient therapeutic services. If we were to designate individual supervision levels

for hospital outpatient therapeutic services just as we do for diagnostic services, it would be most consistent and appropriate for CMS to make specific determinations for services that we believe may only be safely provided under the personal supervision of a physician or that must be performed only by a physician. We stated above that we expect that hospitals already have credentialing procedures, bylaws, and other policies established to ensure that services furnished to Medicare beneficiaries are being provided only by qualified practitioners in accordance with all applicable laws, regulations and coding guidance. For services not furnished directly by a physician, we would expect that these bylaws and policies would ensure that the services are being supervised in a manner commensurate with the complexity of the service, including personal supervision where appropriate.

The use of Condition Code 44 pertains to the entire patient encounter, the patient's status, and the hospital bill type submitted. Reporting of individual HCPCS codes on an outpatient claim must be consistent with all instructions and CMS guidance, including the requirements of § 410.27 of the regulations, which specify that direct supervision is required for hospital outpatient therapeutic services.

Comment: One commenter requested clarification of the meaning of the phrase "performance of the procedure" in the definition of direct supervision. The commenter was unclear whether this phrase specifically applied only to surgical procedures or whether it was a general term for services and procedures. The commenter also asked if the direct supervision requirement would apply to recovery room services following a surgical procedure performed by the physician.

Response: The use of the term "procedure" is intended to encompass all services and procedures and includes all components of a service or procedure furnished by a hospital to an outpatient, including recovery room services, and covered and paid by Medicare, regardless of whether separate payment is made for those component services. This is how we have applied the term within the codified definitions of the levels of supervision (general, direct, and personal) in §§ 410.27 and 410.32. While each supervision definition uses the phrase "performance of the procedure," the term "service" may be substituted for the word "procedure" each time "procedure" appears in the regulations.

Comment: Several commenters raised questions about the proposed definition of the phrase "in the hospital." Some requested clarification of the meaning of "on the same campus." Several commenters suggested that the word "ownership" in the definition seems to unintentionally exclude areas that are operated and controlled by the hospital under a lease agreement or a written operations agreement. These commenters suggested either removing this term from the definition of "in the hospital" or clarifying in this final rule with comment period where CMS is referring to the business operation aspect of ownership rather than the physical building.

Response: "On the same campus" was used to denote that the supervising physician or nonphysician practitioner must be physically located on the same campus as the services being furnished by the hospital because it is possible for some hospitals to have more than one main campus, as well as off-campus PBDs.

We appreciate the public comments that raised the questions about the term "ownership." We agree with the commenters that a narrow interpretation of the word "ownership" would exclude spaces that the hospital leases or operates under another type of operations agreement. This was not our intention. The commenters correctly pointed out that the word "ownership" in this context applies to the actual business operation, not solely a physical building. However, we also believe that the rest of the definition includes these aspects. If the definition is read as a whole, instead of parsing individual clauses out of context, then the spirit of the regulation is understood. Because the phrase "in the hospital or CAH" applies broadly to "incident to" requirements such as the site of service requirement for therapeutic services provided by the hospital directly and under arrangement, we are finalizing the proposed definition of "in the hospital" in new paragraph (g) of § 410.27 as meaning areas in the main building(s) of a hospital that are under the ownership, financial, and administrative control of the hospital; that are operated as part of the hospital; and for which the hospital bills the services furnished under the hospital's CCN.

Comment: One commenter requested that CMS reiterate that the "incident to" and physician supervision requirements of § 410.27 do not apply to physical therapy, occupational therapy, and speech-language pathology services furnished under a therapy plan of care.

Response: Section 1833(t) of the Act excludes physical therapy, occupational

therapy, and speech-language pathology from hospital outpatient services paid under the OPSS. In addition, in the April 2000 OPSS final rule (65 FR 18525), we stated in response to a comment about physical therapy services that the coverage provision in section 1861(s)(2)(D) of the Act does not require that physical therapy services be provided incident to the services of a physician. Finally, in Section 20 (Hospital Outpatient Services) of Chapter 6 of the Medicare Benefit Policy Manual, we state, “[t]he following rules pertaining to the coverage of outpatient hospital services are not applicable to physical therapy, speech-language pathology, occupational therapy, or end stage renal disease (ESRD) services furnished by hospitals to outpatients.” This section instructs readers to consult sections 220 and 230 of Chapter 15 of the Medicare Benefit Policy Manual for rules on the coverage of outpatient physical therapy, occupational therapy, and speech-language pathology furnished by a hospital.

Comment: Several commenters from CMHCs and hospital-based PHPs questioned whether the direct physician supervision clarifications and the “incident to” requirements of § 410.27 apply to those programs. Several hospital-based PHP stakeholders noted the discrepancy in the physician supervision requirements for PHP services furnished by CMHCs and hospitals.

Response: Medicare makes payment for hospital outpatient therapeutic services only when provided “incident to” the services of a physician. Outpatient psychiatric and hospital-based PHP services are outpatient hospital services paid under the OPSS and, therefore, must meet all conditions of § 410.27, including the requirement for direct supervision by a physician or qualified nonphysician practitioner.

Because CMHCs are paid under the OPSS for PHP services, the “incident to” requirements of § 410.27 also apply to CMHCs, with the exception of direct supervision for outpatient PHP services. Currently, CMHCs have a different physician supervision standard to meet. On February 11, 1994, CMS issued the Partial Hospitalization Services in Community Mental Health Centers (CMHCs) interim final rule with comment period (59 FR 6570), developing and implementing the coverage criteria and payment methodology for CMHCs under Medicare. At the time when the CY 1994 CMHC rule was published, the decision was made to permit general supervision in CMHCs. As implemented in § 410.110(a), services provided in a

CMHC PHP must be prescribed by a physician and furnished under the general supervision of a physician. General supervision means that a physician must be at least available by telephone, but is not required to be on the premises of the CMHC at all times (59 FR 6573). We recognized that a direct supervision requirement could cause hardship to CMHCs because some of these entities were unable to employ physicians on a full-time basis due to the expense involved. In the CY 1994 interim final rule with comment, we explained that because we believed that less than direct supervision by a full-time physician in a CMHC would not jeopardize a patient’s health or treatment program, in combination with our belief that there would be a number of professionals involved in the care of the patient who are authorized to furnish services that would otherwise be furnished by a physician, we required general physician supervision.

On August 1, 2000 (65 FR 18434), the OPSS was implemented and provided payment for PHP services provided in two settings: hospital outpatient departments to their outpatients and CMHCs. Although PHP is one benefit and both provider types receive the same payment for services rendered, CMHCs have been permitted to furnish PHP services under general supervision; and the OPSS has, since 2000, held a standard of direct supervision for hospital outpatient therapeutic services, including PHP services (42 CFR 410.110 (a)). While the policy was clearly codified for PBDs of hospitals in the CY 2009 OPSS/ASC proposed rule and final rule with comment period, CMS restated and clarified that the policy also applies to hospital outpatient services not provided in PBDs due to questions referencing the assumption of physician supervision in the April 2000 OPSS final rule (73 FR 68702 through 68704 referencing 65 FR 18525). Even though there are different physician supervision standards for CMHCs and hospital-based PHPs, the certification, recertification, and plan of treatment requirements at § 424.24(e) and section 1835(2)(F) of the Act continue to apply to both provider types. The physician would certify and recertify (where services are furnished over a period of time) that the individual would require inpatient psychiatric care in the absence of such services; an individualized, written plan for furnishing such services has been established by a physician and is reviewed periodically by a physician; and such services are or were furnished while the individual is or was under the care of a physician. The physician

recertification must be signed by a physician who is treating the patient and has knowledge of the patient’s response to treatment.

In order to unify the benefit and create more equity and consistency, we are exploring the possibility of extending the same physician supervision requirements to both provider types. Specifically, we are concerned whether the current policy of requiring only general physician supervision in CMHCs continues to be appropriate, taking into account the differences among provider settings. We also plan to analyze PHP claims data from the past several years and assess whether our current policies and payment structures are working. Therefore, we will evaluate the policies and the possibility of extending the same physician supervision requirement to PHP services furnished in both CMHCs and HOPDs in future rulemaking.

With that in mind, we are requesting comments on this final rule with comment period that address: (1) What types of practitioners currently provide the supervision of PHP services in CMHCs; (2) what is the expertise of supervising practitioners in CMHCs and what is the expectation for their availability; (3) based on the final CY 2010 supervision requirements for hospital outpatient therapeutic services (including PHP services furnished in HOPDs), under what circumstances would direct supervision of PHP services furnished in CMHCs not be occurring, according to the applicable definitions for direct supervision of on-campus and off-campus therapeutic services; and (4) what would be the rationale for maintaining different supervision requirements for PHP services furnished in CMHCs and HOPDs, given that all PHP services are paid under the OPSS.

Comment: A few commenters stated that the provisions for off-campus PBDs should not require the supervising practitioner to be in the department because the existing policy is burdensome and costly, especially for rural providers. The commenters requested that a supervising physician or nonphysician practitioner be able to supervise services being furnished in more than one off-campus PBD at a time. Some commenters stated that they know of off-campus PBDs in current operation that operate with a supervisor nearby, for example, in a physician’s office, but not in the PBD.

Response: We first note that the requirement for direct supervision of hospital outpatient therapeutic services furnished in PBDs of the hospital was

codified in § 410.27(f) of the regulations in the April 2000 OPPTS final rule. In that rule, we explicitly stated that “on the premises of the location” means that the supervisor must be on the premises of the PBD (65 FR 18525). We also responded to public commenters who asserted that requiring a physician to be onsite at a PBD throughout the performance of all “incident to” therapeutic services would be burdensome and costly for hospitals where there are a limited number of physicians available to provide coverage, particularly in rural settings. We disagreed then that the supervision requirement was unnecessary and burdensome because hospitals, prior to 2000, were already required to “meet a direct supervision of ‘incident to’ services requirement that is unrelated to the provider-based rules.” We continue to believe that it would be inappropriate to allow one physician or nonphysician practitioner to supervise all services being provided in all PBDs at a particular off-campus outpatient location. Since first allowing off-campus sites to be considered PBDs of hospitals, we have placed particular emphasis on ensuring the quality and safety of the services provided in these locations, which can be many miles from the main hospital campus, through both additional provider-based requirements in § 413.65(e) and our emphasis on direct physician supervision under § 410.27(f). In addition, because the physician or nonphysician practitioner must be immediately available and have, within his or her State scope of practice and hospital-granted privileges, the ability to perform the services being supervised, we believe it would be highly unlikely that one physician or nonphysician practitioner would be both immediately available at all times that therapeutic services are being provided and would have the knowledge and ability to adequately supervise all services being performed at once in multiple off-campus PBDs.

Comment: Many commenters maintained that the CY 2009 OPPTS/ASC final rule with comment period statements on supervision were not a “restatement or clarification” but a significant change in policy. They argued that any enforcement should be prospective beginning in CY 2010 only, with no enforcement regarding prior years’ hospital outpatient therapeutic services. The commenters believed that the policy prior to CY 2009 at best was unclear, and that the regulations were not comprehensive, and therefore, they concluded that hospitals should not be held accountable to the policy stated in

the CY 2009 OPPTS/ASC final rule with comment period for hospital outpatient therapeutic services furnished in 2000 through CY 2008. The commenters stated that CMS should also not enforce the policy as clarified in the CY 2009 OPPTS/ASC final rule with comment period for CY 2009 hospital outpatient therapeutic services because sufficient opportunity for public notice and comment was not provided. Many hospitals were unaware that the policy had been discussed in the CY 2000 OPPTS rulemaking process, and the commenters argued that hospitals may not have had enough time to meet these requirements for CY 2009. Furthermore, hospitals expressed concern about their potential liability due to *qui tam* litigation.

Response: We provided a restatement and clarification of existing policy in the CY 2009 OPPTS/ASC proposed rule (73 FR 41518 through 41519), citing numerous existing statutory, regulatory, manual, and prior rule preamble statements in section XII.A. of that rule, specifically titled, “Physician Supervision of HOPD Services.” The CY 2009 OPPTS/ASC proposed rule provided for a 60-day comment period. We continue to believe that the CY 2009 restatement and clarification made no change to longstanding hospital outpatient physician direct supervision policies as incorporated in prior statements of policy, including the regulations. In addition, we provided for public notice and comment regarding these physician supervision policies through the CY 2009 OPPTS/ASC proposed rule in which, as noted above, we discussed physician supervision explicitly in a distinct section of the proposed rule, and we responded in the final rule to the few public comments we received on the supervision discussion. Therefore, we believe that the usual enforcement practices of Medicare contractors are appropriate for services furnished in CY 2009. Likewise, the final supervision policies described in this CY 2010 final rule with comment period for hospital outpatient therapeutic services are effective and, therefore, subject to enforcement beginning January 1, 2010.

In regard to hospital outpatient therapeutic services provided in CY 2000 through CY 2008, in CY 2009 we recognized the need for clarification of the direct supervision policy. CMS was relatively silent on this topic between 2000 and 2008. Furthermore, the existing regulations at § 410.27(f) only specify the supervision requirements for hospital outpatient therapeutic services furnished in PBDs but do not address services furnished in areas of the

hospital that may not be PBDs. However, we note that the details of the direct supervision policy for hospital outpatient therapeutic services furnished in off-campus PBDs were clearly and consistently stated in the April 2000 OPPTS final rule discussion and the regulations, including the requirement that the supervising practitioner be physically present in an off-campus PBD when such services were being furnished. As stated earlier in this section, we have placed and will continue to place particular emphasis on ensuring the quality and safety of the services provided in these locations and will continue to do so through our enforcement and other efforts. However, in the case of hospital outpatient therapeutic services furnished on the hospital’s campus in 2000 through 2008, we plan to exercise our discretion and decline to enforce in situations involving claims where the hospital’s noncompliance with the direct physician supervision policy resulted from error or mistake.

After consideration of the public comments we received, we are finalizing our proposal to allow, in addition to clinical psychologists, certain other nonphysician practitioners to directly supervise services that they may perform themselves under their State license and scope of practice and hospital-granted or CAH-granted privileges, with one modification. In addition to physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives, in this final rule with comment period, we are allowing licensed clinical social workers to provide direct supervision. Specifically, we are modifying the final text of revised § 410.27(f) to include licensed clinical social workers among the listing of nonphysician practitioners who may directly supervise the provision of hospital outpatient therapeutic services. These nonphysician practitioners may directly supervise services that they may personally furnish in accordance with State law and all additional requirements, including those specified in §§ 410.71, 410.73, 410.74, 410.75, 410.76, and 410.77, respectively. Accordingly, we also are adding the cross-reference to § 410.73 (Clinical social worker services) in the final revision of paragraph (a)(1)(iv) of § 410.27 (which indicates that services must be furnished under the direct supervision of a physician or a nonphysician practitioner as specified in paragraph (f)).

We are finalizing the proposed direct physician supervision requirements for PR, CR, and ICR services furnished in

the HOPD that would require the direct supervision to be provided by a doctor of medicine or osteopathy. Accordingly, we are finalizing the relevant language in proposed §§ 410.27(a)(1)(iv)(A) and (B) which indicates that, for PR, CR, and ICR services, direct supervision must be furnished by a doctor of medicine or osteopathy, as specified in §§ 410.47 and 410.49, respectively.

For services furnished on a hospital's main campus, we are finalizing a modification of our proposed definition of "direct supervision" in new paragraph (a)(1)(iv)(A) of § 410.27 that allows for the supervisory physician or nonphysician practitioner to be anywhere on the hospital campus, including a physician's office, an on-campus SNF, RHC, or other nonhospital space. Therefore, direct supervision means that the supervisory physician or nonphysician practitioner must be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure.

Because the term "in the hospital or CAH" applies broadly to "incident to" requirements such as the site of service requirement for therapeutic services provided by the hospital directly and under arrangement, we also are finalizing the definition of "in the hospital" in new paragraph § 410.27(g) as meaning areas in the main building(s) of a hospital or CAH that are under the ownership, financial, and administrative control of the hospital or CAH; that are operated as part of the hospital; and for which the hospital bills the services furnished under the hospital's or CAH's CCN.

We are finalizing, without modification, the addition of new paragraph (a)(1)(iv)(B) to § 410.27 to reflect that, for off-campus PBDs of hospitals, the physician or nonphysician practitioner must be present in the off-campus PBD, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician or nonphysician practitioner must be in the room when the procedure is performed. As we stated previously, the language of paragraph (a)(1)(iv)(B) is similar to existing § 410.27(f) that we are revising and relocating. Furthermore, we are finalizing the proposed technical change to clarify the language in this paragraph by removing the phrase "present and on the premises of the location" and replacing it with the phrase "present in the off-campus provider-based department."

Additionally, we are finalizing the proposal to make a technical correction to the title of § 410.27 to read, "Outpatient hospital or CAH services and supplies incident to a physician service: Conditions" to clarify in the title that the requirements for payment of hospital outpatient therapeutic services incident to a physician or nonphysician practitioner service in that section apply to both hospitals and CAHs. Similarly, we are including the phrase "hospital or CAH" throughout the text of § 410.27 wherever the text currently refers just to "hospital."

4. Policies for Direct Supervision of Hospital and CAH Outpatient Diagnostic Services

As we discussed in detail in the CY 2010 OPPTS/ASC proposed rule (74 FR 35368), with respect to the physician supervision requirements for individual diagnostic tests, we have continued since the April 2000 OPPTS final rule discussion (65 FR 18526) to instruct hospitals that, for diagnostic services furnished in PBDs of hospitals, hospitals should follow the supervision requirements for individual diagnostic tests as listed in the MPFS Relative Value File. For diagnostic services not listed in the MPFS file, Medicare contractors, in consultation with their medical directors, define appropriate supervision levels in order to determine whether claims for these services are reasonable and necessary. To further specify the supervision policy across service settings and to provide consistency for all hospital outpatient diagnostic services, for CY 2010 we proposed to require that all hospital outpatient diagnostic services that are provided directly or under arrangement, whether provided in the main buildings of the hospital, in a PBD of a hospital, or at a nonhospital location, follow the physician supervision requirements for individual tests as listed in the MPFS Relative Value File. We also proposed that the definitions of general, direct, and personal supervision as defined in §§ 410.32(b)(3)(i) through (b)(3)(iii) would also apply. In the case of direct supervision of diagnostic services furnished directly by the hospital or under arrangement in the main hospital buildings or on-campus in a PBD, we proposed that the definition of direct supervision would be the same as the definition we proposed for therapeutic services provided on-campus as discussed in the CY 2010 OPPTS/ASC proposed rule (74 FR 35369), meaning that the physician would be present on the same campus, in the hospital or the on-campus PBD of the hospital, as defined in § 413.65, and immediately

available to furnish assistance and direction throughout the performance of the procedure. In addition, the definition of "in the hospital or CAH" as defined in proposed § 410.27(g), discussed above, would apply. In the CY 2010 OPPTS/ASC proposed rule, we explained that this means that the supervisory physician may not be located in any entity such as a physician's office, co-located hospital, IDTF, or hospital-operated provider or supplier such as a SNF, ESRD facility, or HHA, or any other nonhospital space that may be co-located on the hospital's campus, as campus is defined in § 413.65(a)(2).

Similarly, in the case of direct physician supervision of diagnostic services furnished directly or under arrangement in an off-campus PBD, we proposed that the definition of direct supervision would be the same as the current definition for therapeutic services provided in an off-campus PBD as discussed in the CY 2010 OPPTS/ASC proposed rule (74 FR 35369), meaning the physician must be present in the off-campus PBD, as defined in § 413.65 and immediately available to furnish assistance and direction throughout the performance of the procedure. As we discussed in the April 2000 OPPTS final rule (65 FR 18524 through 18525) and the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68702 through 68704), we have long made the analogy of the PBD to the physician's office suite, as described in the definition of direct supervision in § 410.32(b)(3)(ii).

In addition to providing diagnostic services directly or under arrangement in the hospital, including PBDs of the hospital, a hospital may also send its outpatients to another entity, such as an IDTF, to furnish these services under arrangement for the hospital. For example, in the April 2000 OPPTS final rule (65 FR 185440 through 185441), in a discussion of the hospital bundling rules, we discussed that an entity, like an IDTF, may be located in the main buildings of a hospital or on the hospital campus but operated independently of the hospital. In addition, these suppliers, providers, or other entities may be located elsewhere, not on a hospital's main campus or other hospital property. These entities, like IDTFs and physicians' offices, may provide services to their own patients (not hospital outpatients) and to hospital outpatients under arrangements with the hospital. They follow the physician supervision requirements of the MPFS and § 410.32 when providing services to Medicare beneficiaries who are not hospital outpatients. For consistency, we proposed for CY 2010

that all diagnostic services provided to hospital outpatients under arrangement in nonhospital entities, whether those entities are located on the main campus of the hospital or elsewhere, would also follow the requirements as described in § 410.32(b)(3)(i) through (iii). When hospitals contract with other entities to provide services under arrangement, the hospital must exercise professional responsibility over the arrangement for services, in accordance with the guidance provided in Section 10.3 (Under Arrangements) of Chapter 5 of the Medicare General Information, Eligibility and Entitlement Manual. This means that for the hospital to receive payment, it is responsible for ensuring that all applicable requirements in §§ 410.28 and 410.32 are met. In the case of hospital outpatient diagnostic services provided under arrangement at nonhospital locations, such as IDTFs, we believe that the term “office suite” used in § 410.32(b)(3)(ii) is directly applicable because these facilities usually also provide diagnostic services to their own patients and, therefore, would be able to apply the direct supervision requirement in § 410.32(b)(3)(ii) without further definition.

We stated in the CY 2010 OPPTS/ASC proposed rule (74 FR 35369) that physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives who operate within the scope of practice under State law may order and perform diagnostic tests, as discussed under § 410.32(a)(3) and in the corresponding manual guidance in section 80 (Requirements for Diagnostic X-Ray, Diagnostic Laboratory, and Other Diagnostic Tests) of Chapter 15 of the Medicare Benefit Policy Manual. However, this manual guidance and the regulations at § 410.32(b)(1) also state that diagnostic x-ray and other diagnostic tests must be furnished under the appropriate level of supervision by a physician as defined in section 1861(r) of the Act. Thus, physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives may not function as supervisory physicians for the purposes of diagnostic tests. In accordance with these existing requirements, we did not propose to allow physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives to provide the supervision of diagnostic tests provided to hospital outpatients. Clinical psychologists may supervise only diagnostic psychological and neuropsychological testing services as described in an exception to the basic rule at § 410.32(b)(2)(iii) for diagnostic

psychological and neuropsychological testing services, when these services are personally furnished by a clinical psychologist or an independently practicing psychologist or when they are furnished under the general supervision of a physician or clinical psychologist.

To reflect these proposed changes for the provision of direct supervision of diagnostic services provided to hospital outpatients in the regulations, we proposed to revise existing § 410.28(e). First, we proposed to specify that the provisions of proposed revised paragraph (e) apply to diagnostic services furnished by the hospital, directly or under arrangement, consistent with our proposal to apply the existing diagnostic services supervision requirement for PBDs to diagnostic services provided directly by the hospital or under arrangement. We would continue to specify that the definitions of general and personal physician supervision included in § 410.32(b)(3)(i) and (b)(3)(iii) apply to these levels of supervision of hospital outpatient diagnostic services. Furthermore, we proposed to add new paragraph (e)(1) to § 410.28 to indicate that, for services furnished directly or under arrangement, in the hospital or in an on-campus department of a provider, as defined in § 413.65, direct supervision means that the physician must be present on the same campus, in the hospital or PBD of the hospital as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. We also would continue to provide that direct supervision does not mean that the physician must be in the room when the procedure is performed. As discussed above, we would apply the definition of “in the hospital” as proposed in § 410.27(g) of the regulations. In addition, we proposed to add new paragraph (e)(2) to § 410.28 to reflect that, for the direct physician supervision of diagnostic services furnished directly or under arrangement in off-campus PBDs of hospitals, the physician must present in the off-campus PBD, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. We would continue to provide that direct supervision does not mean that the physician must be in the room when the procedure is performed. Finally, we proposed to add new paragraph (e)(3) to specify that for the direct supervision of hospital outpatient services provided under arrangement in physicians’ offices and other nonhospital locations, the definition of

direct supervision in § 410.32(b)(3)(ii) applies.

Comment: Some commenters fully supported the proposal for hospital outpatient diagnostic services, agreeing with CMS that it is appropriate to apply the requirements for physician supervision consistently across all sites of service. Other commenters, including major hospitals associations and commenters representing rural hospitals, believed the proposal is unnecessary, unrealistic, costly, and would reduce access to diagnostic services. Several commenters stated that CMS’ interpretation of “furnish assistance and direction throughout the performance of the procedure” would mean that the physician would have to be able to operate sophisticated equipment, replacing the technicians who are trained to operate the equipment. Some commenters indicated that rural hospitals would not be able to meet the proposed requirements. Other commenters requested that CMS add a column to Addendum B specifying the required level of supervision for diagnostic tests to make it easier for hospitals to comply with the requirements.

Response: We disagree with the commenters that requiring hospitals to follow the MPFS levels of supervision for individual diagnostic tests would create additional hospital burden for many services because CMS has already applied the lowest level of supervision (general supervision) to numerous diagnostic services. Additionally, in the April 2000 OPPTS final rule (65 FR 18536), we codified § 410.28(e) of the regulations to apply this requirement to all on and off campus PBDs of hospitals. It is appropriate to apply the same requirements to diagnostic services that are provided “in the hospital” as those that have been long established for on-campus and off-campus PBDs of the hospital, including rural hospitals. We also believe that, in the interest of safe and high quality care for beneficiaries, it is appropriate for hospitals to follow the same requirements for supervision as physicians and IDTFs when furnishing more complex diagnostic tests that we have specifically identified as requiring direct or personal supervision, or should be performed only by the physician. In addition, since hospitals may contract with other entities to have diagnostic services provided under arrangement, it is also appropriate to ensure that those entities are consistently following the supervision levels that we have identified for both their own patients as well as hospital outpatients. We recognize that specially trained

ancillary staff and technicians are the primary operators of some specialized diagnostic testing equipment. However, we also believe it is reasonable for the physician that supervises the provision of the services to be knowledgeable about those tests.

Therefore, we are finalizing our CY 2010 proposal that all hospital outpatient diagnostic services that are provided directly or under arrangement, whether provided in the main buildings of the hospital, in a PBD of the hospital, or at a nonhospital location, follow the physician supervision requirements for individual tests as listed in the MPFS Relative Value File. The definitions of general, direct, and personal supervision as defined in §§ 410.32(b)(3)(i) through (b)(3)(iii) also apply. In the case of direct supervision of diagnostic services furnished directly by the hospital or under arrangement in the main hospital buildings or on-campus in a PBD, the definition of direct supervision is the same as the modified definition that we are finalizing for therapeutic services provided on-campus, as discussed in section XII.D.3. of this final rule with comment period, meaning that the physician would be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. We would continue to specify that direct supervision does not mean that the physician must be in the room when the procedure is performed. As discussed above, we are applying the definition of “in the hospital” as proposed and finalized in § 410.27(g) of the regulations. While the definition of “in the hospital” is no longer a component of the definition of direct supervision, it remains applicable to describe areas operated as part of the hospital that are not PBDs for other purposes, such as services provided under arrangement. It is also appropriate to apply the definition of that term consistently for both diagnostic and therapeutic hospital outpatient services. In addition, we are finalizing our proposal to add new paragraph (e)(2) to § 410.28 to reflect that, for the direct physician supervision of diagnostic services furnished directly or under arrangement in off-campus PBDs of hospitals, the physician must be present in the off-campus PBD of the hospital, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. We would continue to specify that direct supervision does not mean that the physician must be in the

room when the procedure is performed. Also, we are finalizing our proposal to add new paragraph (e)(3) to specify that for the direct supervision of hospital outpatient services provided under arrangement in physicians’ offices and other nonhospital locations, the definition of direct supervision in § 410.32(b)(3)(ii) applies.

We acknowledge the commenters’ request to publish the diagnostic supervision levels in Addendum B. Addendum B currently specifies information related directly to the payment for services described by HCPCS codes, including relative weights, payment rates, and copayments. We do not believe it is necessary to include the supervision levels for diagnostic services in Addendum B that are not directly relevant to the payment rates for those services. These supervision levels are readily available on the CMS Web site in the MPFS RVU File, and, because we make both the MPFS RVU File and Addendum B available in spreadsheet format, an interested hospital can easily modify Addendum B to add whatever code-specific information the hospital believes would be most useful to incorporate in a single electronic file for reference purposes.

Comment: Several commenters asserted that nonphysician practitioners should be able to supervise diagnostic tests because they may order and perform diagnostic tests that are within their scope of practice under State law.

Response: We acknowledged in the CY 2010 OPPS/ASC proposed rule (74 FR 35369) that physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives who operate within the scope of practice under State law may order and perform diagnostic tests, as discussed in § 410.32(a)(3) and corresponding manual guidance in Section 80 of Chapter 15 of the Medicare Benefit Policy Manual. However, we noted that this manual guidance and the long established regulation at § 410.32(b)(1) also state that diagnostic x-ray and other diagnostic tests must be furnished under the appropriate level of supervision by a physician as defined in section 1861(r) of the Act. Thus, CMS historically has not permitted physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives to function as supervisory “physicians” for the purposes of diagnostic tests. In accordance with these existing requirements, we did not propose to allow physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives to provide the supervision of

diagnostic tests provided to hospital outpatients. Because we establish the physician supervision levels in the MPFS Relative Value File based on the policy that only a physician may provide the supervision, we believe it continues to be most appropriate to allow only physicians to provide the supervision of hospital outpatient diagnostic services.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to require that all hospital outpatient diagnostic services that are provided directly or under arrangement, whether provided in the main buildings of the hospital, in a PBD of a hospital, or at a nonhospital location, follow the physician supervision requirements for individual tests as listed in the MPFS RVU File. The definitions of general, direct, and personal supervision as defined in §§ 410.32(b)(3)(i) through (b)(3)(iii) also apply. In the case of direct supervision of diagnostic services furnished directly by the hospital or under arrangement in the main hospital buildings or on-campus in a PBD of a hospital, the definition of direct supervision is the same as the modified definition that we are finalizing for therapeutic services provided on-campus as discussed in section XII.D.3. of this final rule with comment period, meaning that the physician must be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. We continue to provide that direct supervision does not mean that the physician must be in the room when the procedure is performed. As discussed above, we are applying the definition of “in the hospital” as specified in new § 410.27(g) of the regulations. While the definition of in the hospital is no longer a component of the definition of direct supervision, it remains applicable to describe areas operated as part of the hospital that are not PBDs for other purposes, such as services provided under arrangement. It is also appropriate to apply the definition of that term consistently for both diagnostic and therapeutic hospital outpatient services. In addition, we are finalizing our CY 2010 proposal, without modification, to add new paragraph (e)(2) to § 410.28 to reflect that, for the direct physician supervision of diagnostic services furnished directly or under arrangement in off-campus PBDs of hospitals, the physician must be present in the off-campus PBD, as defined in § 413.65, and immediately available to furnish

assistance and direction throughout the performance of the procedure. We continue to provide that direct supervision does not mean that the physician must be in the room when the procedure is performed. Also, we are finalizing the CY 2010 proposal, without modification, to add new paragraph (e)(3) to specify that, for the direct supervision of hospital outpatient services provided under arrangement in physicians' offices and other nonhospital locations, the definition of direct supervision in § 410.32(b)(3)(ii) applies. We did not propose to allow physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives to provide the supervision of diagnostic tests provided to hospital outpatients and we are finalizing this policy.

5. Summary of CY 2010 Physician Supervision Final Policy

In summary, for CY 2010, nonphysician practitioners who are specified under § 410.27 of the final regulations as clinical psychologists, licensed clinical social workers, physician assistants, nurse practitioners, clinical nurse specialists, and certified nurse-midwives, may directly supervise all hospital outpatient therapeutic services that they may perform themselves within their State scope of practice and hospital-granted privileges, provided that they meet all additional requirements, including any collaboration or supervision requirements as specified in §§ 410.71, 410.73, 410.74, 410.75, 410.76, and 410.77. We are finalizing the proposed direct physician supervision requirements for PR, CR, and ICR services furnished in the HOPD that would require the supervision to be provided by a doctor of medicine or osteopathy. Accordingly, we are finalizing proposed §§ 410.27(a)(1)(iv)(A) and (B) which indicate that, for PR, CR, and ICR services, direct supervision must be furnished by a doctor of medicine or osteopathy, as specified in §§ 410.47 and 410.49, respectively.

We also are refining the definition of the direct supervision of hospital outpatient therapeutic services for those services provided in the hospital or on-campus PBD of the hospital. For services provided in the hospital or on-campus PBD of the hospital, direct supervision would mean that the physician or nonphysician practitioner must be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. In addition, we are finalizing the definition

of "in the hospital" in new paragraph § 410.27(g) to mean areas in the main building(s) of a hospital or CAH that are under the ownership, financial, and administrative control of the hospital or CAH; that are operated as part of the hospital or CAH; and for which the hospital or CAH bills the services furnished under the hospital's or CAH's CCN.

We are not making any significant changes to the definition or requirements for direct supervision of hospital outpatient therapeutic services provided in off-campus PBDs of a hospital or CAH, other than to allow nonphysician practitioners to provide direct supervision for the services that they may perform themselves in those locations. Therefore, we are finalizing, without modification, the addition of new paragraph (a)(1)(iv)(B) to § 410.27 to reflect that, for off-campus PBDs of hospitals or CAHs, the physician or nonphysician practitioner must be present in the off-campus PBD, as defined in § 413.65, and immediately available to furnish assistance and direction throughout the performance of the procedure. We state that this requirement does not mean that the physician or nonphysician practitioner must be in the room when the procedure is performed.

Additionally, we are finalizing the proposal to make a technical correction to the title of § 410.27 and the text of § 410.27 to clarify throughout that the requirements for payment of hospital outpatient therapeutic services incident to a physician or nonphysician practitioner service in that section apply to both hospitals and CAHs.

For CY 2010, we are finalizing the proposal to require that all hospital outpatient diagnostic services provided directly or under arrangement, whether provided in the hospital, in a PBD of a hospital, or at a nonhospital location, follow the physician supervision requirements for individual tests as listed in the MPFS Relative Value File. The existing definitions of general and personal supervision as defined in §§ 410.32(b)(3)(i) and (b)(3)(iii) also apply. For services furnished directly or under arrangement in the hospital or on-campus PBD, direct supervision means that the physician must be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. For the purposes of § 410.28, as for the general purposes of § 410.27, the definition of "in the hospital," as defined in § 410.27(g), applies. For diagnostic services furnished directly or under arrangement off-campus in a PBD of the hospital,

direct supervision continues to mean that the physician must be present in the off-campus PBD and immediately available to furnish assistance and direction throughout the performance of the procedures. For all hospital outpatient diagnostic services provided under arrangement in nonhospital locations, such as IDTFs and physicians' offices, the existing definition of direct supervision under § 410.32(b)(3)(ii) applies. We are revising § 410.28 of the regulations to reflect these changes.

E. Direct Referral for Observation Services

Since CY 2003, hospitals have reported a Level II HCPCS code for Medicare billing purposes for a "direct admission" to a hospital for outpatient observation services. In section 290 of Chapter 4 of the Medicare Claims Processing Manual (Pub. 100-4), we define a "direct admission" as the direct referral of a patient by a community physician to a hospital for observation services without an associated emergency room visit, hospital outpatient clinic visit, critical care service, or hospital outpatient surgical procedure (that is, a status indicator "T" procedure) on the day of the initiation of observation services. Since CY 2006, we have instructed hospitals to report a "direct admission" referred for observation services using HCPCS code G0379 (Direct admission of patient for hospital observation care) (70 FR 68688 through 68691).

Observation care is a hospital outpatient service that is reported using HCPCS code G0378 (Hospital observation services, per hour). Hospitals report outpatient observation services, which are commonly provided in association with a hospital clinic visit, emergency department visit, or other major service, on hospital outpatient claims, just like other outpatient services. Physicians order observation care, defined as clinically appropriate services, including ongoing short-term treatment, assessment, and reassessment furnished in order for the physician to determine whether the beneficiary will require further treatment as an inpatient or whether the beneficiary may be safely discharged from the hospital.

We have become aware that, because the word "admission" is generally used in reference to inpatient hospital care, our historical use of the phrase "direct admission" in the code descriptor for HCPCS code G0379 and the use of the phrase "observation status" in the Medicare Claims Processing Manual (Chapter 4, Section 290) and the Medicare Benefit Policy Manual

(Chapter 6, Section 20) may be contributing to confusion for hospitals and beneficiaries related to a beneficiary's status as an inpatient or an outpatient when he or she is receiving observation services. For Medicare payment purposes, there is no patient status termed "observation status." Hospitals may only bill for items and services furnished to inpatients, outpatients, or nonpatients. We believe that using terminology such as "observation status" or "admission to observation" may be confusing for physicians, hospitals, and beneficiaries. Therefore, in the CY 2010 OPPS/ASC proposed rule (74 FR 35370 through 35371), we proposed to modify the code descriptor for HCPCS code G0379 to remove the reference to the word "admission" and to replace it with "referral." The proposed long code descriptor for HCPCS code G0379 would be "Direct referral for hospital observation care." We proposed this change to more accurately reflect that the physician in the community has referred the beneficiary to the hospital for observation services as a hospital outpatient. In addition to the proposed CY 2010 change to the code descriptor for HCPCS code G0379, we modified the Medicare Claims Processing Manual and the Medicare Benefit Policy Manual to remove most references related to "admission" for observation services or "observation status." We refer readers to Transmittal 1760 dated June 23, 2009 (which rescinded and replaced Transmittal 1745, dated May 22, 2009) and Transmittal 107 dated May 22, 2009 (both issued under Change Request 6492), for more information regarding

the specific changes incorporated in the manuals.

We did not propose to change the status indicator or payment methodology for HCPCS code G0379 for CY 2010. Instead, we proposed to continue the payment policy that was finalized for the CY 2009 OPPS (73 FR 68554). In the CY 2010 OPPS/ASC proposed rule (74 FR 35370 through 35371), we proposed to continue to assign HCPCS code G0379 status indicator "Q3," indicating that it would be eligible for payment through APC 8002 (Level I Extended Assessment & Management Composite) when certain criteria are met or through APC 0604 (Level I Hospital Clinic Visits) when other criteria are met; otherwise, its payment would be packaged into payment for other separately payable services in the same encounter. The established criteria for payment of HCPCS code G0379 under either composite APC 8002, as part of the extended assessment and management composite service, or APC 0604, as a separately payable individual service that we would continue for CY 2010 are: (1) both HCPCS codes G0378 and G0379 are reported with the same date of service; and (2) no service with a status indicator of "T" or "V" or Critical Care (APC 0617) is provided on the same date of service as HCPCS code G0379. If either of the above criteria is not met, HCPCS code G0379 is assigned status indicator "N" and its payment is packaged into the payment for other separately payable services provided in the same encounter.

We did not receive any public comments related to our CY 2010 proposal to revise the code descriptor

for HCPCS code G0379 to read "Direct referral for hospital observation care," or on our proposal to continue the CY 2009 status indicator assignment and payment methodology for HCPCS code G0379 for CY 2010. Therefore, we are finalizing these CY 2010 proposals, without modification.

XIII. OPSS Payment Status and Comment Indicators

A. OPSS Payment Status Indicator Definitions

Payment status indicators (SIs) that we assign to HCPCS codes and APCs play an important role in determining payment for services under the OPSS. They indicate whether a service represented by a HCPCS code is payable under the OPSS or another payment system and also whether particular OPSS policies apply to the code. The final CY 2010 status indicator assignments for APCs and HCPCS codes are shown in Addendum A and Addendum B, respectively, to this final rule with comment period.

As we proposed in the CY 2010 OPSS/ASC proposed rule (74 FR 35371), in this final rule with comment period, we are changing the definitions of status indicators "H" and "K." We did not propose to make any changes to the other status indicators that were listed in Addendum D1 of the CY 2009 OPSS/ASC final rule with comment period. The final status indicators are listed in the tables under sections XIII.A.1., 2., 3., and 4. of this final rule with comment period.

1. Payment Status Indicators to Designate Services That Are Paid Under the OPSS

Indicator	Item/code/service	OPSS payment status
G	Pass-Through Drugs and Biologicals	Paid under OPSS; separate APC payment.
H	Pass-Through Device Categories	Separate cost-based pass-through payment; not subject to co-payment.
K	Nonpass-Through Drugs and Nonimplantable Biologicals, including Therapeutic Radiopharmaceuticals.	Paid under OPSS; separate APC payment.
N	Items and Services Packaged into APC Rates	Paid under OPSS; payment is packaged into payment for other services. Therefore, there is no separate APC payment.
P	Partial Hospitalization	Paid under OPSS; per diem APC payment.
Q1	STVX-Packaged Codes	Paid under OPSS; Addendum B displays APC assignments when services are separately payable. (1) Packaged APC payment if billed on the same date of service as a HCPCS code assigned status indicator "S," "T," "V," or "X." (2) In all other circumstances, payment is made through a separate APC payment.
Q2	T-Packaged Codes	Paid under OPSS; Addendum B displays APC assignments when services are separately payable. (1) Packaged APC payment if billed on the same date of service as a HCPCS code assigned status indicator "T." (2) In all other circumstances, payment is made through a separate APC payment.

Indicator	Item/code/service	OPPS payment status
Q3	Codes that may be paid through a composite APC	Paid under OPPS; Addendum B displays APC assignments when services are separately payable. Addendum M displays composite APC assignments when codes are paid through a composite APC. (1) Composite APC payment based on OPPS composite-specific payment criteria. Payment is packaged into a single payment for specific combinations of service. (2) In all other circumstances, payment is made through a separate APC payment or packaged into payment for other services.
R	Blood and Blood Products	Paid under OPPS; separate APC payment.
S	Significant Procedure, Not Discounted When Multiple	Paid under OPPS; separate APC payment.
T	Significant Procedure, Multiple Reduction Applies	Paid under OPPS; separate APC payment.
U	Brachytherapy Sources	Paid under OPPS; separate APC payment.
V	Clinic or Emergency Department Visit	Paid under OPPS; separate APC payment.
X	Ancillary Services	Paid under OPPS; separate APC payment.

Section 142 of Public Law 110-275 (MIPPA) required CMS to pay for therapeutic radiopharmaceuticals for the period of July 1, 2008, through December 31, 2009, at hospitals' charges adjusted to the costs. The status indicator "H" was assigned to therapeutic radiopharmaceuticals to indicate that an item was paid at charges adjusted to cost during CY 2009. In the CY 2010 OPPS/ASC proposed rule (74 FR 35373), we proposed to pay prospectively and separately for therapeutic radiopharmaceuticals with average per day costs greater than the proposed CY 2010 drug packaging threshold of \$65 under the OPPS. Therefore, we proposed to change the status indicator for HCPCS codes used to report separately payable therapeutic radiopharmaceuticals from "H" to "K," which indicates that an item is separately paid under the OPPS at the APC payment rate established for the item. We refer readers to section V.B.5. of the proposed rule (74 FR 35333 through 36336) and this final rule with comment period for discussion of the proposed and final CY 2010 changes to our payment policy for therapeutic radiopharmaceuticals.

We received many comments on our proposal to establish prospective payment rates for therapeutic radiopharmaceuticals. We respond to these comments and discuss our final

policy in section V. B. 5. of this final rule with comment period. However, we did not receive any public comments related to our proposal to change the definitions of status indicators "H" and "K," to reflect this change in policy. Therefore, we are changing the definitions of status indicators "H" and "K" as proposed, without modification, to reflect our final therapeutic radiopharmaceutical payment policy, and we are finalizing assignment of status indicator "K" to therapeutic radiopharmaceuticals with average per day costs greater than the final CY 2010 drug packaging threshold of \$65 under the OPPS.

As we discussed in detail in section V.A.4. of the CY 2010 OPPS/ASC proposed rule (74 FR 35311 through 35314), we proposed to consider implantable biologicals that are not on pass-through status as a biological before January 1, 2010, as devices for pass-through evaluation and payment beginning in CY 2010. Therefore, as devices, pass-through implantable biologicals would be assigned a status indicator of "H," while nonpass-through implantable biologicals would be assigned a status indicator of "N" beginning in CY 2010. Those implantable biologicals that have been granted pass-through status under the drug and biological criteria prior to January 1, 2010, would continue to be

assigned a status indicator of "G" until they are proposed for expiration from pass-through status during our annual rulemaking cycle. In the proposed rule (74 FR 35373), we proposed to assign status indicator "K" to nonimplantable biologicals and to adjust the definition of status indicator "K" accordingly.

We received numerous comments on our proposal to treat implantable biologicals as devices and we respond to them in section V.A.4. of this final rule with comment period. We did not receive any public comments with regard to the proposed changes to status indicator "K" to reflect the implantable biological pass-through payment policy. Therefore, we are finalizing our proposal, without modification, to assign status indicator "K" to nonimplantable biologicals and to adjust the definition of status indicator "K" accordingly.

The final CY 2010 status indicators are displayed in the table above, as well as in Addendum D1 to this final rule with comment period.

2. Payment Status Indicators to Designate Services That Are Paid Under a Payment System Other Than the OPPS

We did not propose any changes to the status indicators listed below for the CY 2010 OPPS.

Indicator	Item/code/service	OPPS payment status
A	Services furnished to a hospital outpatient that are paid under a fee schedule or payment system other than OPPS, for example: <ul style="list-style-type: none"> • Ambulance Services • Clinical Diagnostic Laboratory Services • Non-Implantable Prosthetic and Orthotic Devices • EPO for ESRD Patients • Physical, Occupational, and Speech Therapy • Routine Dialysis Services for ESRD Patients Provided in a Certified Dialysis Unit of a Hospital. • Diagnostic Mammography • Screening Mammography 	Not paid under OPPS. Paid by fiscal intermediaries/MACs under a fee schedule or payment system other than OPPS. Not subject to deductible or coinsurance.
C	Inpatient Procedures	Not subject to deductible. Not paid under OPPS. Admit patient. Bill as inpatient.

Indicator	Item/code/service	OPPS payment status
F	Corneal Tissue Acquisition; Certain CRNA Services; and Hepatitis B Vaccines.	Not paid under OPPS. Paid at reasonable cost.
L	Influenza Vaccine; Pneumococcal Pneumonia Vaccine	Not paid under OPPS. Paid at reasonable cost; not subject to deductible or coinsurance.
M	Items and Services Not Billable to the Fiscal Intermediary/MAC	Not paid under OPPS.
Y	Non-Implantable Durable Medical Equipment	Not paid under OPPS. All institutional providers other than home health agencies bill to DMERC.

We did not receive any public comments regarding the status indicators that designate services that are paid under a payment system other than the OPPS. Therefore, we are finalizing our CY 2010 proposal, without modification. The final CY 2010 status indicators are displayed in the

table above, as well as in Addendum D1 to this final rule with comment period.

3. Payment Status Indicators To Designate Services That are Not Recognized Under the OPPS but That May Be Recognized by Other Institutional Providers

We did not propose any changes to the status indicators listed below for the CY 2010 OPPS.

Indicator	Item/code/service	OPPS payment status
B	Codes that are not recognized by OPPS when submitted on an outpatient hospital Part B bill type (12x and 13x).	Not paid under OPPS. <ul style="list-style-type: none"> • May be paid by fiscal intermediaries/MACs when submitted on a different bill type, for example, 75x (CORF), but not paid under OPPS. • An alternate code that is recognized by OPPS when submitted on an outpatient hospital Part B bill type (12x and 13x) may be available.

We did not receive any public comments regarding the status indicators that designate services that are not recognized under the OPPS but that may be recognized for payment to other institutional providers. Therefore,

we are finalizing our CY 2010 proposal, without modification. The final status indicators are displayed in the table above, as well as in Addendum D1 to this final rule with comment period.

4. Payment Status Indicators To Designate Services That Are Not Payable by Medicare on Outpatient Claims

We did not propose any changes to the payment status indicators listed below for the CY 2010 OPPS.

Indicator	Item/code/service	OPPS payment status
D	Discontinued Codes	Not paid under OPPS or any other Medicare payment system.
E	Items, Codes, and Services: <ul style="list-style-type: none"> • That are not covered by any Medicare outpatient benefit based on statutory exclusion. • That are not covered by any Medicare outpatient benefit for reasons other than statutory exclusion. • That are not recognized by Medicare for outpatient claims; alternate code for the same item or service may be available. • For which separate payment is not provided on outpatient claims. 	Not paid by Medicare when submitted on outpatient claims (any outpatient bill type).

We did not receive any public comments related to payment status indicators that designate services that are not payable by Medicare on outpatient claims and, therefore, we are finalizing our CY 2010 proposal, without modification. The final status indicators are displayed in the table above, as well as in Addendum D1 to this final rule with comment period.

Addendum B, with a complete listing of HCPCS codes that includes their payment status indicators and APC assignments for CY 2010 is available electronically on the CMS Web site under supporting documentation for this final rule with comment period at:

<http://www.cms.hhs.gov/HospitalOutpatientPPS/HORD/list.asp#TopOfPage>.

B. Comment Indicator Definitions

In the CY 2010 OPPS/ASC proposed rule (74 FR35374), we proposed to use the same two comment indicators that are in effect for the CY 2009 OPPS.

- “CH”—Active HCPCS codes in current and next calendar year; status indicator and/or APC assignment have changed or active HCPCS code that will be discontinued at the end of the current calendar year.

- “NI”—New code for the next calendar year or existing code with

substantial revision to its code descriptor in the next calendar year as compared to current calendar year, interim APC assignment; comments will be accepted on the interim APC assignment for the new code.

We proposed to use the “CH” comment indicator in this CY 2010 OPPS/ASC final rule with comment period to indicate HCPCS codes for which the status indicator or APC assignment, or both, would change in CY 2010 compared to their assignment as of December 31, 2009.

We believe that using the “CH” indicator in this CY 2010 OPPS/ASC final rule with comment period will

help facilitate the public's review of the changes that we are finalizing for CY 2010. The use of the comment indicator "CH" in association with a composite APC indicates that the configuration of the composite APC is changed in this CY 2010 OPPTS/ASC final rule with comment period.

In the CY 2010 OPPTS/ASC proposed rule (74 FR 35374 through 35375), we did not propose any changes to the definitions of the OPPTS comment indicators for CY 2010 and we did not receive any public comments on the comment indicators. However, we want to clarify our policy regarding the use of comment indicator "NI" in this CY 2010 OPPTS/ASC final rule with comment period to describe a new code. There are numerous instances in which the descriptor of an existing Category I CPT code is substantially revised for CY 2010 so that it describes a new service or procedure that could have been assigned a new code number by the CPT Editorial Panel and that new code number would then have been assigned the "NI" comment indicator. Because, for CY 2010, not all new services or procedures will be assigned a new CPT code number, but instead will be described by an existing CPT code number with a substantially revised code descriptor, we are assigning the comment indicator "NI" to these codes in order to allow for comment on these substantially revised codes. Therefore, for this final rule with comment period, we have expanded the definition of comment indicator "NI" to include an existing code with a substantial revision to its code descriptor in the next calendar year as compared to the current calendar year to indicate that the code's CY 2010 OPPTS treatment is open to public comment on this final rule with comment period. Like all codes labeled with comment indicator "NI," we will respond to public comments and finalize their OPPTS treatment in the CY 2011 OPPTS/ASC final rule with comment period. In accordance with our usual practice, CPT and Level II HCPCS code numbers that are new for CY 2010 are also labeled with comment indicator "NI" in Addendum B to this final rule with comment period.

Only HCPCS codes with comment indicator "NI" in this CY 2010 OPPTS/ASC final rule with comment period are subject to comment. HCPCS codes that do not appear with comment indicator "NI" in this CY 2010 OPPTS/ASC final rule with comment period are not open to public comment, unless we specifically have requested additional comments elsewhere in this final rule with comment period. The CY 2010

treatment of HCPCS codes that appears in this CY 2010 OPPTS/ASC final rule with comment period to which comment indicator "NI" is not appended was open to public comment during the comment period for the CY 2010 OPPTS/ASC proposed rule, and we are responding to those comments in this final rule with comment period.

We did not receive any public comments regarding comment indicators. Therefore, we are finalizing our proposal, without modification, and we are continuing to use the two comment indicators, "CH" and "NI," for CY 2010. Their definitions are listed in Addendum D2 to this final rule with comment period.

XIV. OPPTS Policy and Payment Recommendations

A. MedPAC Recommendations

MedPAC was established under section 1805 of the Act to advise the U.S. Congress on issues affecting the Medicare program. As required under the statute, MedPAC submits reports to Congress not later than March and June of each year that present its Medicare payment policy recommendations. The following section describes recent recommendations relevant to the OPPTS that have been made by MedPAC.

The March 2009 MedPAC "Report to Congress: Medicare Payment Policy" included the following recommendation relating specifically to the Medicare hospital OPPTS:

Recommendation 2A-1: The Congress should increase payment rates for the acute inpatient and outpatient prospective payment systems in 2010 by the projected rate of increase in the hospital market basket index, concurrent with implementation of a quality incentive payment program.

CMS Response: As proposed in the CY 2010 OPPTS/ASC proposed rule (74 FR 35375), in this final rule with comment period, we are increasing payment rates for the CY 2010 OPPTS by the projected rate of increase in the hospital market basket through adjustment of the full CY 2010 conversion factor. Simultaneously, for CY 2010, we are continuing to reduce the annual update factor by 2.0 percentage points for hospitals that are defined under section 1886(d)(1)(B) of the Act and that do not meet the hospital outpatient quality data reporting required by section 1833(t)(17) of the Act. Specifically, we have calculated two conversion factors: A full conversion factor based on the full hospital market basket increase and a reduced conversion factor that reflects the 2.0 percentage point reduction to the

hospital market basket. We discuss our update of the conversion factor and our adoption and implementation of the reduced conversion factor that will apply to hospitals that fail their quality reporting requirements for the full CY 2010 OPPTS update in section XVI of this final rule with comment period.

The full March 2009 MedPAC report can be downloaded from MedPAC's Web site at: http://www.medpac.gov/documents/Mar09_EntireReport.pdf.

We note that MedPAC also submitted comments on the CY 2010 OPPTS/ASC proposed rule. The specific issues that were the subject of MedPAC's comments and the sections of this final rule with comment period where they are addressed are listed below:

- Pharmacy overhead costs and setting payments for separately paid drugs: Section V.B.3.
- Payment rates for brachytherapy sources and therapeutic radiopharmaceuticals: Sections VII. and V.B.5.
- Collection of quality data through clinical registries and electronic health records (EHRs): Section XVII.I.
- Collection of quality data from ASCs: Section XVI.H.
- Collection of cost data from ASCs: Section XV.G.
- Payment policy for healthcare-associated conditions: Section XVII.

B. APC Panel Recommendations

Recommendations made by the APC Panel at its February 2009 and August 2009 meetings are discussed in the sections of this final rule with comment period that correspond to topics addressed by the APC Panel. The report and recommendations from the APC Panel's February 18-19, 2009 and August 5-6, 2009 meetings are available on the CMS Web site at: http://www.cms.hhs.gov/FACA/05_AdvisoryPanelonAmbulatoryPaymentClassificationGroups.asp.

C. OIG Recommendations

The mission of the Office of the Inspector General (OIG), as mandated by Public Law 95-452, as amended, is to protect the integrity of the U.S. Department of Health and Human Services (HHS) programs as well as the health and welfare of beneficiaries served by those programs. This statutory mission is carried out through a nationwide network of audits, investigations, and inspections. In June 2007, the OIG released a report, entitled "Impact of Not Retroactively Adjusting Outpatient Outlier Payments," that described the OIG's research into sources of errors in CMHC outlier payments. The OIG report included the

following two recommendations relating specifically to the hospital OPSS under which payment is made for outpatient services provided by CMHCs.

Recommendation 1: The OIG recommended that CMS require adjustments of outpatient outlier payments at final cost report settlement, retroactive to the beginning of the cost report period.

Recommendation 2: The OIG recommended that CMS require retroactive adjustments of outpatient outlier payments when an error caused by the fiscal intermediary or provider is identified after a cost report is settled.

We addressed both of these recommendations in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68594). We noted in that final rule that the OIG's findings were based largely on information from the OPSS' early implementation period, between CY 2000 and CY 2003, and that we believed we had taken several steps since that time in order to improve the accuracy and frequency of the Medicare contractors' CCR calculations, including updating our instructions for calculating CCRs, increasing the frequency of CCR calculation, and conducting an annual review of CMHC CCRs.

However, taking into account these OIG recommendations, we proposed and finalized a policy to provide for reconciliation of outlier payments under the OPSS at final cost report settlement as recommended by the OIG, beginning in CY 2009. We discuss our rationale for this policy in detail in section II.F.4. of the CY 2009 OPSS/ASC final rule with comment period (73 FR 68594 through 68599).

Other than the June 2007 recommendations, there have been no other recent OIG recommendations pertaining to the OPSS.

XV. Updates to the Ambulatory Surgical Center (ASC) Payment System

A. Background

1. Legislative Authority for the ASC Payment System

Section 1832(a)(2)(F)(i) of the Act provides that benefits under Medicare Part B include payment for facility services furnished in connection with surgical procedures specified by the Secretary that are performed in an ASC. To participate in the Medicare program as an ASC, a facility must meet the standards specified in section 1832(a)(2)(F)(i) of the Act, which are set forth in 42 CFR part 416, subpart B and Subpart C of our regulations. The regulations at 42 CFR part 416, subpart B describe the general conditions and requirements for ASCs, and the

regulations at subpart C explain the specific conditions for coverage for ASCs.

Section 141(b) of the Social Security Act Amendments of 1994, Public Law 103-432, required establishment of a process for reviewing the appropriateness of the payment amount provided under section 1833(i)(2)(A)(iii) of the Act for intraocular lenses (IOLs) that belong to a class of new technology intraocular lenses (NTIOLs). That process was the subject of a final rule entitled "Adjustment in Payment Amounts for New Technology Intraocular Lenses Furnished by Ambulatory Surgical Centers," published on June 16, 1999, in the **Federal Register** (64 FR 32198).

Section 626(b) of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (MMA), Public Law 108-173, added subparagraph (D) to section 1833(i)(2) of the Act, which required the Secretary to implement a revised ASC payment system to be effective not later than January 1, 2008. Section 626(c) of the MMA amended section 1833(a)(1) of the Act by adding new subparagraph (G), which requires that, beginning with implementation of the revised ASC payment system, payment for surgical procedures furnished in ASCs shall be 80 percent of the lesser of the actual charge for the services or the amount determined by the Secretary under the revised payment system.

Section 5103 of the Deficit Reduction Act of 2005 (DRA), Public Law 109-171, amended section 1833(i)(2) of the Act by adding a new subparagraph (E) to place a limitation on payment amounts for surgical procedures furnished in ASCs on or after January 1, 2007, but before the effective date of the revised ASC payment system (that is, January 1, 2008). Section 1833(i)(2)(E) of the Act provides that if the standard overhead amount under section 1833(i)(2)(A) of the Act for an ASC facility service for such surgical procedures, without application of any geographic adjustment, exceeds the Medicare payment amount under the hospital OPSS for the service for that year, without application of any geographic adjustment, the Secretary shall substitute the OPSS payment amount for the ASC standard overhead amount.

Section 109(b) of the Medicare Improvements and Extension Act of 2006 of the Tax Relief and Health Care Act of 2006 (MIEA-TRHCA), Public Law 109-432, amended section 1833(i) of the Act, in part, by redesignating clause (iv) as clause (v) and adding a new clause (iv) to paragraph (2)(D) and adding paragraph (7)(A), which provide

the Secretary the authority to require ASCs to submit data on quality measures and to reduce the annual update by 2 percentage points for an ASC that fails to submit data as required by the Secretary on selected quality measures. Section 109(b) of the MIEA-TRHCA also amended section 1833(i) of the Act by adding new paragraph (7)(B), which requires that, to the extent the Secretary establishes such an ASC quality reporting program, certain quality of care reporting requirements mandated for hospitals paid under the OPSS, under section 109(a) of the MIEA-TRHCA, be applied in a similar manner to ASCs unless otherwise specified by the Secretary.

For a detailed discussion of the legislative history related to ASCs, we refer readers to the June 12, 1998 proposed rule (63 FR 32291 through 32292).

2. Prior Rulemaking

On August 2, 2007, we published in the **Federal Register** (72 FR 42470) the final rule for the revised ASC payment system, effective January 1, 2008 (the "August 2, 2007 final rule"). We revised our criteria for identifying surgical procedures that are eligible for Medicare payment when furnished in ASCs and adopted the method we would use to set payment rates for ASC covered surgical procedures and covered ancillary services furnished in association with those covered surgical procedures beginning in CY 2008. In that final rule, we also established a policy for updating on an annual calendar year basis the ASC conversion factor, the relative payment weights, the ASC payment rates, and the list of procedures for which Medicare would not make an ASC payment. We also established a policy for treating new and revised HCPCS and CPT codes under the ASC payment system. This policy is consistent with the OPSS to the extent possible (72 FR 42533).

In the CY 2008 OPSS/ASC final rule with comment period (72 FR 66827), we updated and finalized the CY 2008 ASC rates and lists of covered surgical procedures and covered ancillary services. We also made regulatory changes to 42 CFR parts 411, 414, and 416 related to our final policies to provide payments to physicians who perform noncovered ASC procedures in ASCs based on the facility practice expense (PE) relative value units (RVUs), to exclude covered ancillary radiology services and covered ancillary drugs and biologicals from the categories of designated health services (DHS) that are subject to the physician self-referral prohibition, and to reduce

ASC payments for surgical procedures when the ASC receives full or partial credit toward the cost of the implantable device. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68722), we updated and finalized the CY 2009 ASC rates and lists of covered surgical procedures and covered ancillary services.

3. Policies Governing Changes to the Lists of Codes and Payment Rates for ASC Covered Surgical Procedures and Covered Ancillary Services

The August 2, 2007 final rule established our policies for determining which procedures are ASC covered surgical procedures and covered ancillary services. Under §§ 416.2 and 416.166 of the regulations, subject to certain exclusions, covered surgical procedures are surgical procedures that are separately paid under the OPPS, that would not be expected to pose a significant risk to beneficiary safety when performed in an ASC, and that would not be expected to require active medical monitoring and care at midnight following the procedure ("overnight stay"). We adopted this standard for defining which surgical procedures are covered surgical procedures under the ASC payment system as an indicator of the complexity of the procedure and its appropriateness for Medicare payment in ASCs. We use this standard only for purposes of evaluating procedures to determine whether or not they are appropriate for Medicare beneficiaries in ASCs. We define surgical procedures as those described by Category I CPT codes in the surgical range from 10000 through 69999, as well as those Category III CPT codes and Level II HCPCS codes that crosswalk or are clinically similar to ASC covered surgical procedures (72 FR 42478). We note that we added over 800 surgical procedures to the list of covered surgical procedures for ASC payment in CY 2008, the first year of the revised ASC payment system, based on the criteria for payment that we adopted in the August 2, 2007 final rule as described above in this section. Patient safety and health outcomes continue to be important to us as more health care moves to the ambulatory care setting. Therefore, as we gain additional experience with the ASC payment system, we are interested in any information the public may have regarding the comparative patient outcomes of surgical care provided in ambulatory settings, including HOPDs, ASCs, and physicians' offices, particularly with regard to the Medicare population.

In the August 2, 2007 final rule, we also established our policy to make separate ASC payments for the following ancillary items and services when they are provided integral to ASC covered surgical procedures: Brachytherapy sources; certain implantable items that have pass-through status under the OPPS; certain items and services that we designate as contractor-priced, including, but not limited to, procurement of corneal tissue; certain drugs and biologicals for which separate payment is allowed under the OPPS; and certain radiology services for which separate payment is allowed under the OPPS. These covered ancillary services are specified in § 416.164(b) and, as stated previously, are eligible for separate ASC payment (72 FR 42495). Payment for ancillary items and services that are not paid separately under the ASC payment system is packaged into the ASC payment for the covered surgical procedure.

The full CY 2009 lists of ASC covered surgical procedures and covered ancillary services are included in Addenda AA and BB, respectively, to the CY 2009 OPPS/ASC final rule with comment period (73 FR 68840 through 68933 and 69270 through 69308).

We update the lists of, and payment rates for, covered surgical procedures and covered ancillary services, in conjunction with the annual proposed and final rulemaking process to update the OPPS and ASC payment systems (§ 416.173; 72 FR 42535). In addition, because we base ASC payment policies for covered surgical procedures, drugs, biologicals, and certain other covered ancillary services on the OPPS payment policies, we also provide quarterly updates for ASC services throughout the year (January, April, July, and October), just as we do for the OPPS. The updates are to implement newly created Level II HCPCS codes and Category III CPT codes for ASC payment and to update the payment rates for separately paid drugs and biologicals based on the most recently submitted ASP data. New Category I CPT codes, except vaccine codes, are released only once a year and, therefore, are implemented through the January quarterly update. New Category I CPT vaccine codes are released twice a year and thus are implemented through the January and July quarterly updates.

In our annual updates to the ASC list of, and payment rates for, covered surgical procedures and covered ancillary services, we undertake a review of excluded surgical procedures (including all procedures newly proposed for removal from the OPPS

inpatient list), new procedures, and procedures for which there is revised coding, to identify any that we believe meet the criteria for designation as ASC covered surgical procedures or covered ancillary services. Updating the lists of covered surgical procedures and covered ancillary services, as well as their payment rates, in association with the annual OPPS rulemaking cycle is particularly important because the OPPS relative payment weights and, in some cases, payment rates, are used as the basis for the payment of covered surgical procedures and covered ancillary services under the revised ASC payment system. This joint update process ensures that the ASC updates occur in a regular, predictable, and timely manner.

Comment: Several commenters provided a number of general suggestions related to the ASC list of covered surgical procedures. They contended that CMS should not restrict which procedures are payable in ASCs any more than CMS restricts which procedures are payable in HOPDs. The commenters added that if the policy to exclude procedures from the list is maintained, CMS should at least provide the exclusionary criteria for all of the payable OPPS procedures that are excluded from the ASC list so that the public can provide meaningful comments about CMS' decisions. They suggested that CMS publish an addendum to the proposed and final OPPS/ASC rules that would identify which of the criteria at § 416.166(c) triggered CMS' decision to exclude each procedure.

Some commenters urged CMS to eliminate unlisted codes from the exclusionary criteria at § 416.166(c), and other commenters requested that ASCs be allowed to use unlisted codes to bill for procedures that are from anatomic sites that could not possibly pose a potential risk to beneficiary safety. The commenters reported that unlisted codes enable surgeons to utilize innovative techniques or new technologies and are paid under the OPPS and by commercial insurers. They suggested that ASCs could provide documentation to the contractor that explains and justifies the procedure reported by an unlisted code; thus ensuring that Medicare does not make payment for a service that would otherwise be excluded from payment.

Response: We appreciate the commenters' suggestions regarding the consistency of CMS' decisions about which procedures are excluded from the ASC list. However, as we explained in the August 2, 2007 final rule (72 FR 42479), we do not believe that all

procedures that are appropriate for performance in HOPDs are appropriate in ASCs. HOPDs are able to provide much higher acuity care than ASCs. ASCs have neither patient safety standards consistent with those in place for hospitals, nor are they required to have the trained staff and equipment needed to provide the breadth and intensity of care that hospitals are required to maintain. Therefore, we are not modifying our policy and will continue to exclude from the ASC list of covered surgical procedures certain procedures for which payment is made in HOPDs.

We do not agree with the commenters' request that we provide specific reasons for our decisions to exclude each procedure from the ASC list other than that we believe a procedure is expected to pose a significant risk to beneficiary safety or to require an overnight stay. We believe that these reasons are sufficiently specific to enable the public to provide meaningful comments on our decisions to exclude procedures from the list of covered surgical procedures. Our decisions to exclude procedures from the ASC list are based on a number of the criteria listed at § 416.166 of the regulations, and we believe that it would be unnecessary and overly burdensome to list each and every reason for those decisions.

We also do not agree with the commenters' recommendation that we include certain unlisted codes on the list of covered procedures. Even though it may be highly unlikely that any procedures that would be expected to pose a risk to beneficiary safety or to require an overnight stay would be reported by an unlisted code from certain anatomic sites, we cannot know what surgical procedure is being reported by an unlisted code. Therefore, because we cannot evaluate any such procedure, we believe that we must exclude unlisted codes as a group from the list of covered surgical procedures.

We do not believe it is reasonable, or within the scope of our contractors' work, to accept the commenters' suggestion that ASCs could provide documentation to our Medicare contractors, upon request, in order for the contractors to make a retrospective determination about whether or not a procedure that was billed using an unlisted code represented a significant risk to beneficiary safety or would be expected to require an overnight stay.

Comment: One commenter noted that, although CMS specified in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68714) that patients may remain in ASCs up to 24 hours in order to allow adequate time for recovery

following some surgical procedures, CMS did not specify the requirements for physician supervision during the recovery period. The commenter argued that CMS also failed to specify the time period during which the required post-anesthesia assessment is to be performed by requiring only that it be performed prior to discharge from the ASC. The commenter's concern was that patients in ASCs may have no physician supervision for extended periods, a policy in contrast to CMS' policy regarding the direct physician supervision required for hospital outpatient services. The commenter requested that CMS clarify why the same supervision requirements are not applied equally to hospitals and ASCs.

Response: Historically, Medicare has covered surgical procedures performed in ASCs that have relatively short recovery periods and, therefore, we have believed that physicians were always immediately available to furnish assistance and direction in the ASC while ASC services were being furnished, including during the postoperative recovery period. However, as the commenter points out, not only have we recently revised the Conditions for Coverage to allow longer stays in ASCs, we have greatly expanded the list of covered surgical procedures under the revised ASC payment system, including covering some surgical procedures that may require a prolonged recovery period. Given these two revisions, both of which enable ASCs to provide more clinically complex surgical procedures, and taking into consideration patient safety and quality of care, we believe it could be appropriate to consider establishing requirements for physician or nonphysician practitioner supervision in ASCs, similar to the requirements for the direct supervision of hospital outpatient therapeutic services that we are finalizing for HOPDs in this CY 2010 OPPTS/ASC final rule with comment period. We note that, for therapeutic services furnished incident to a physician's professional service in an office setting, there also is a requirement for direct physician supervision, meaning that the physician must be in the office suite and immediately available to furnish assistance and direction throughout the procedure (§ 410.26(a)(2) and (b)(5)). In addition, we note that payment for covered ancillary services may be made to ASCs, including payment for some of the diagnostic tests that would be subject to the physician supervision requirements for hospital outpatient diagnostic services if provided in the HOPD. The

final CY 2010 physician supervision requirements for hospital outpatient diagnostic and therapeutic services are discussed in detail in section XII.E. of this final rule with comment period. For diagnostic services furnished in physicians' offices, IDTFs and other Part B settings, the requirements of § 410.32 of the regulations apply, including supervision of diagnostic services.

We did not propose to adopt supervision requirements for therapeutic and diagnostic services furnished in ASCs similar to the requirements for HOPDs for CY 2010. However, given the overlap in surgical procedures that may be performed in HOPDs and ASCs and the increased breadth and complexity of ASC covered procedures, we are requesting comments on this final rule with comment period that address: (1) What types of practitioners currently provide the diagnostic and therapeutic services in ASCs, particularly during the extended postoperative recovery period; (2) what types of practitioners currently provide the supervision for the diagnostic and therapeutic services in ASCs, particularly during the extended postoperative period; (3) what is the expertise of supervising practitioners in ASCs and what is the expectation for their availability; (4) based on the final CY 2010 supervision requirements for hospital outpatient therapeutic services, under what circumstances would direct supervision of ASC services (including during the postoperative recovery period) not be occurring, according to the applicable definitions for direct supervision for HOPD services; and (5) what would be the rationale for not establishing supervision requirements for ASC services that parallel the supervision requirements in other settings, including HOPDs and physicians' offices.

After consideration of the public comments we received, we are not accepting the commenters' recommendations to not exclude all procedures reported by unlisted codes or all procedures for which Medicare payment is made to HOPDs. We will continue to exclude all procedures that we determine would be expected to pose a significant risk to beneficiary safety or require an overnight stay. Further, we are not accepting the commenters' recommendation that CMS provide more specific reasons for its decisions regarding exclusion of specific procedures from the ASC list of covered surgical procedures. In this final rule with comment period, we are soliciting public comments on the issue of physician supervision of ASC services, especially as related to extended

postoperative stays. In summary, we are making no changes to the final criteria for determining which procedures are excluded from the ASC list of covered surgical procedures.

B. Treatment of New Codes

1. Treatment of New Category I and Category III CPT Codes and Level II HCPCS Codes

We finalized a policy in the August 2, 2007 final rule to evaluate each year all new Category I and Category III CPT codes and Level II HCPCS codes that describe surgical procedures, and to make preliminary determinations in the annual OPPS/ASC final rule with comment period regarding whether or not they meet the criteria for payment in the ASC setting and, if so, whether they are office-based procedures (72 FR 42533). In addition, we identify new codes as ASC covered ancillary services based upon the final payment policies of the revised ASC payment system. New HCPCS codes that are released in the summer through the fall of each year, to be effective January 1, are included in the final rule with comment period updating the ASC payment system for the following calendar year. These new codes are flagged with comment indicator “NI” in Addenda AA and BB to the OPPS/ASC final rule with comment period to indicate that we are assigning a payment indicator to the codes on an interim basis. The interim payment indicators assigned to the new codes under the revised ASC payment system are subject to public comment in that final rule with comment period. These interim determinations must be made in the OPPS/ASC final rule with comment

period because, in general, the new HCPCS codes and their descriptors for the upcoming calendar year are not available at the time of development of the OPPS/ASC proposed rule. We will respond to those comments in the OPPS/ASC final rule with comment period for the following calendar year. In the CY 2010 OPPS/ASC proposed rule (74 FR 35377), we proposed to continue this identification and recognition process for CY 2010.

We did not receive any public comments regarding this proposal. For CY 2010, we are continuing our established policy for recognizing new Category I and Category III CPT codes and Level II HCPCS codes.

In addition, we proposed to continue our policy of implementing through the ASC quarterly update process new mid-year CPT codes, generally Category III CPT codes, that the AMA releases in January to become effective the following July, and released in July to become effective the following January. We proposed to include in Addenda AA or BB, as appropriate, to this CY 2010 OPPS/ASC final rule with comment period the new Category III CPT codes released in January 2009 for implementation on July 1, 2009 (through the ASC quarterly update process), that we identify as ASC covered services. Similarly, we proposed to include in Addenda AA and BB to this final rule with comment period any new Category III CPT codes that the AMA released in July 2009 to be effective on January 1, 2010, that we identify as ASC covered services. However, only those new Category III CPT codes implemented effective January 1, 2010, are designated by

comment indicator “NI” in the Addenda to this CY 2010 OPPS/ASC final rule with comment period to indicate that we have assigned them an interim payment indicator that is subject to public comment. The two Category III CPT codes implemented in July 2009 for ASC payment, which appeared in Table 38 of the CY 2010 OPPS/ASC proposed rule (74 FR 35378), were subject to comment through that proposed rule, and we proposed to finalize their payment indicators in this CY 2010 OPPS/ASC final rule with comment period.

We proposed to assign payment indicator “G2” (Non office-based surgical procedure added in CY 2008 or later; payment based on OPPS relative payment weight) to both of these two new codes. Because new Category III CPT codes that become effective for July are not available to CMS in time for incorporation into the Addenda to the OPPS/ASC proposed rule, our policy is to include the codes, their proposed payment indicators, and proposed payment rates in the preamble to the proposed rule but not in the Addenda to the proposed rule. These codes and their final payment indicators and rates are included in the Addenda to this CY 2010 OPPS/ASC final rule with comment period.

We did not receive any public comments regarding this proposal. For CY 2010, we are continuing our established policy for recognizing new mid-year CPT codes, and the new mid-year codes implemented in July 2009 are displayed in Table 57 below, as well as in Addendum AA to this final rule with comment period.

TABLE 57—NEW CATEGORY III CPT CODES IMPLEMENTED IN JULY 2009 FOR ASC PAYMENT

CY 2010 HCPCS code	CY 2010 long descriptor	Final CY 2010 ASC payment indicator
0200T	Percutaneous sacral augmentation (sacroplasty), unilateral injection(s), including the use of a balloon or mechanical device (if utilized), one or more needles.	G2
0201T	Percutaneous sacral augmentation (sacroplasty), bilateral injections, including the use of a balloon or mechanical device (if utilized), two or more needles.	G2

For CY 2010, there are numerous instances in which the descriptor of an existing Category I CPT code is substantially revised so that it describes a new service or procedure. In each such instance, revision of the code’s descriptor created a more specific description of some of the services or procedures that were reported by the existing CPT code and required that at least one other code be created to describe the other services that were

described by the existing code descriptor. Thus, the services or procedures that were described by the existing CPT code descriptor will be described by two new codes for CY 2010: one newly created code number and descriptor and one code with the same code number for which the code descriptor has been substantially revised. For example, CPT code 21556 (Excision tumor, soft tissue of neck or thorax; deep, subfascial, intramuscular)

was revised to create two new procedures, one of which will be reported by the same CPT code number with a different description. Thus, for CY 2010, two new procedures are reported by revised CPT code 21556 (Excision, tumor, soft tissue of neck or anterior thorax, subfascial (e.g. intramuscular); less than 5 cm) and new CPT code 21554 (Excision, tumor, soft tissue of neck or anterior thorax, subfascial (e.g. intramuscular); 5 cm or

greater). In the past, the more common practice has been to delete an existing code number which was used to report a general description of a procedure and assign a new code number to each new, more specifically-described procedure.

Due to the practice of maintaining an existing CPT code number for a substantially revised descriptor, we have had to make changes to the payment indicators for existing code numbers that now describe different procedures. Specifically, thirty-one of the existing CPT code numbers that are used to represent new procedures in CY 2010 are currently used to report procedures that were on the ASC list of covered surgical procedures in CY 2007 and, therefore, in CY 2009 are assigned payment indicator "A2" (Surgical procedure on ASC list in CY 2007; payment based on OPPS relative payment weight based on their old descriptions). All of the newly created procedures, including the 31 procedures that will be reported by an existing CPT code number, were evaluated for appropriateness for inclusion on the ASC list of covered surgical procedures. For the procedures that we included on the ASC list, we also made an interim determination regarding whether the procedure should be designated as office-based for CY 2010. Therefore, in Addendum AA to this final rule with comment period, the same CPT code number that was assigned payment indicator "A2" for CY 2009 may be assigned payment indicators "G2" (Non office-based surgical procedure added in CY 2008 or later; payment based on OPPS relative payment weight); "P2" (Office-based surgical procedure added to ASC list in CY 2008 or later with MPFS nonfacility PE RVUs; payment based on OPPS relative payment weight); "P3" (Office-based surgical procedure added to ASC list in CY 2008 or later with MPFS nonfacility PE RVUs; payment based on MPFS nonfacility PE RVUs); or "R2" (Office-based surgical procedure added to ASC list in CY 2008 or later without MPFS nonfacility PE RVUs; payment based on OPPS relative payment weight) due to the change in the procedure assigned to the numeric code.

Any existing CPT codes with substantial revisions to their code descriptors for CY 2010 such that we consider them to describe new procedures and for which their ASC payment indicator may change are labeled with comment indicator "NI" in Addendum AA to this final rule with comment period, to indicate that we have assigned them an interim final payment status which is subject to public comment. Like all codes labeled

with comment indicator "NI," we will respond to public comments and finalize their ASC treatment in the CY 2011 OPPS/ASC final rule with comment period. In addition to assigning the "NI" indicator to these new CPT codes, in accordance with our standard practice, all new CPT and Level II HCPCS code numbers for CY 2010 are labeled with comment indicator "NI" in Addendum AA and BB to this final rule with comment period.

2. Treatment of New Level II HCPCS Codes Implemented in April and July 2009

New Level II HCPCS codes may describe covered surgical procedures or covered ancillary services. All new Level II HCPCS codes implemented in April and July 2009 for ASCs describe covered ancillary services. During the second quarter of CY 2009, we added to the list of covered ancillary services two new Level II HCPCS codes because they are drugs or biologicals for which separate payment was newly allowed under the OPPS in the same calendar quarter. The two Level II HCPCS codes added, effective April 1, 2009, were HCPCS code C9247 (Iobenguane, I-123, diagnostic, per study dose, up to 10 millicuries) and HCPCS code C9249 (Injection, certolizumab pegol, 1 mg). Although HCPCS code C9247 was created for use beginning on January 1, 2009, initially it was not paid separately under the hospital OPPS, and therefore its payment also was packaged under the ASC payment system, until April 1, 2009.

After publication of the CY 2010 OPPS/ASC proposed rule, the CMS HCPCS Workgroup created permanent HCPCS codes to replace these two HCPCS C-codes that were implemented in April 2009. We will be recognizing these HCPCS codes for payment of these drugs and biologicals under the CY 2010 ASC payment system, consistent with our general policy to use permanent HCPCS codes, if appropriate, for the reporting of drugs and biologicals. Table 58 shows the new permanent HCPCS codes that replace the HCPCS C-codes that will be deleted effective December 31, 2009.

Specifically, HCPCS code C9247 was replaced with HCPCS code A9582 (Iodine I-123 iobenguane, diagnostic, per study dose, up to 15 millicuries) and HCPCS code C9249 was replaced with HCPCS code J0718 (Injection, certolizumab pegol, 1 mg). The new HCPCS codes, effective January 1, 2010, describe the same drugs. Although HCPCS code A9582 indicates "per study dose, up to 15 millicuries" and the

descriptor of its predecessor C-code designates "per study dose, up to 10 millicuries," we believe that the reporting of one study dose would be the same in most cases under either the new permanent code or the predecessor code. The recommended dose of I-123 iobenguane is 10 millicuries for adult patients, so we expect that hospitals would report 1 unit of new HCPCS code A9582 for the typical dose in CY 2010, just as they would have reported one unit of HCPCS code C9247 previously for the typical dose and, therefore, there would be no effect on the payment indicator.

For the third quarter of CY 2009, we added 11 new Level II drug and biological HCPCS codes to the list of ASC covered ancillary services because they were newly eligible for separate payment under the OPPS effective July 1, 2009. These HCPCS codes are: C9250 (Human plasma fibrin sealant, vapor-heated, solvent-detergent (Artiss), 2 ml); C9251 (Injection, C1 esterase inhibitor (human) 10 units); C9252 (Injection, plerixafor, 1 mg); C9253 (Injection, temozolomide, 1 mg); C9360 (Dermal substitute, native, non-denatured collagen, neonatal bovine origin (SurgiMend Collagen Matrix), per 0.5 square centimeters); C9361 (Collagen matrix nerve wrap (NeuroMend Collagen Nerve Wrap), per 0.5 centimeter length); C9362 (Porous purified collagen matrix bone void filler (Integra Mozaik Osteoconductive Scaffold Strip), per 0.5 cc); C9363 (Skin substitute, Integra Meshed Bilayer Wound Matrix, per square centimeter); C9364 (Porcine implant, Permacol, per square centimeter); Q2023 (Injection, factor viii (antihemophilic factor, recombinant) (Xyntha), per i.u.); and Q4116 (Skin substitute, Alloderm, per square centimeter).

We assigned payment indicator "K2" (Drugs and biologicals paid separately when provided integral to a surgical procedure on ASC list; payment based on OPPS rate) to all of these new Level II HCPCS codes and added the codes to the list of covered ancillary services through either the April 2009 update (Transmittal 1698, Change Request 6424, dated March 13, 2009) or the July 2009 update (Transmittal 1740, Change Request 6496, dated May 22, 2009) to the CY 2009 ASC payment system. Initially, we assigned payment indicator "K2" to new HCPCS code Q4115 (Skin substitute, Alloskin, per square centimeter) for July 2009, but then changed that assignment retroactive to July 2009 to signify that this HCPCS code was not a covered ancillary service because it was not recognized for payment under the OPPS during that

same time period. Subsequently, in the October 2009 quarterly update CR (Transmittal 1806, Change Request 6629, dated August 28, 2009), HCPCS code Q4115 was added as payable (assigned payment indicator “K2”), effective October 1, 2009.

In the CY 2010 OPPS/ASC proposed rule (74 FR 35378), we solicited public comment on the proposed CY 2010 ASC payment indicators and payment rates for the drugs and biologicals, as listed in Tables 39 and 40 of the proposed rule (74 FR 35378 through 35379). Those HCPCS codes became payable in ASCs beginning in April 2009 or July 2009, respectively, based on the ASC rates posted for the appropriate calendar quarter on the CMS Web site at: <http://www.cms.hhs.gov/ASCPayment/>.

The HCPCS codes that were listed in Table 39 of the proposed rule became effective in April 2009 and were included in Addendum BB to the proposed rule. However, because HCPCS codes that become effective for July are not available to CMS in time for incorporation into the Addenda to the OPPS/ASC proposed rule, our policy is to include these HCPCS codes and their proposed payment indicators and payment rates in the preamble to the proposed rule but not in the Addenda

to the proposed rule. These codes and their final payment indicators and rates are included in the appropriate Addendum to this CY 2010 OPPS/ASC final rule with comment period. Thus, the codes implemented by the July 2009 ASC quarterly update and their proposed CY 2010 payment rates (based on July 2009 ASP data) that were displayed in Table 40 of the CY 2010 OPPS/ASC proposed rule were not included in Addendum BB to the CY 2010 OPPS/ASC proposed rule. We proposed to include the services reported using the new HCPCS codes that were displayed in Tables 39 and 40 as covered ancillary services and to incorporate all of them into Addendum BB to this CY 2010 OPPS/ASC final rule with comment period, consistent with our annual update policy.

After publication of the CY 2010 OPPS/ASC proposed rule, the HCPCS Workgroup created permanent HCPCS J-codes for 4 of the 11 separately payable covered ancillary services that were displayed in Table 40 of the proposed rule. Consistent with our general policy of using permanent HCPCS codes, if appropriate, rather than HCPCS C-codes in order to streamline coding, effective for CY 2010, we are adopting the 4

permanent HCPCS J-codes to replace the HCPCS C-codes. As displayed in Table 59 below, HCPCS code C9251 is replaced with J0598 (Injection, C1 esterase inhibitor (human), 10 units); C9252 with J2562 (Injection, plerixafor, 1 mg); C9253 with J9328 (Injection, temozolomide, 1 mg); and Q2023 with J7185 (Injection, factor viii (antihemophilic factor, recombinant) (Xyntha), per i.u.). The HCPCS J-codes, effective January 1, 2010, describe the same drugs and the same dosages as the HCPCS C-codes that will be deleted December 31, 2009. Therefore, there is no effect on the services’ payment indicators.

We did not receive any public comments regarding our proposals. We are adopting as final the ASC payment indicators for the new Level II HCPCS codes implemented in April and July 2009 as shown in Tables 58 and 59, respectively. Moreover, we are adopting as final the replacement HCPCS codes, specifically, A9582 and J0718, as shown in Table 58 and HCPCS codes J0598, J2562, J7185, and J9328 as displayed in Table 59 below. All of the new HCPCS codes and payment indicators also are included in Addendum BB to this final rule with comment period.

TABLE 58—NEW LEVEL II HCPCS CODES FOR COVERED ANCILLARY SERVICES IMPLEMENTED IN APRIL 2009

CY 2010 HCPCS Code	CY 2009 HCPCS Code	CY 2010 Long descriptor	Final CY 2010 payment indicator
A9582	C9247	Iodine I-123 iobenguane, diagnostic, per study dose, up to 15 millicuries	K2
J0718	C9249	Injection, certolizumab pegol, 1 mg	K2

TABLE 59—NEW LEVEL II HCPCS CODES FOR COVERED ANCILLARY SERVICES IMPLEMENTED IN JULY 2009

CY 2010 HCPCS Code	CY 2009 HCPCS Code	CY 2010 Long descriptor	Final CY 2010 ASC payment indicator
C9250	C9250	Human plasma fibrin sealant, vapor-heated, solvent-detergent (Artiss), 2 ml	K2
J0598	C9251	Injection, C1 esterase inhibitor (human), 10 units	K2
J2562	C9252	Injection, plerixafor, 1 mg	K2
J9328	C9253	Injection, temozolomide, 1 mg	K2
C9360	C9360	Dermal substitute, native, non-denatured collagen, neonatal bovine origin (SurgiMend Collagen Matrix), per 0.5 square centimeters.	K2
C9361	C9361	Collagen matrix nerve wrap (NeuroMend Collagen Nerve Wrap), per 0.5 centimeter length.	K2
C9362	C9362	Porous purified collagen matrix bone void filler (Integra Mozaik Osteoconductive Scaffold Strip), per 0.5 cc.	K2
C9363	C9363	Skin substitute, Integra Meshed Bilayer Wound Matrix, per square centimeter	K2
C9364	C9364	Porcine implant, Permacol, per square centimeter	K2
J7185	Q2023	Injection, factor viii (antihemophilic factor, recombinant) (Xyntha), per i.u.	K2
Q4116	Q4116	Skin substitute, Alloderm, per square centimeter	K2

C. Update to the Lists of ASC Covered Surgical Procedures and Covered Ancillary Services

1. Covered Surgical Procedures

a. Additions to the List of ASC Covered Surgical Procedures

In the CY 2010 OPPS/ASC proposed rule (74 FR 35379), we proposed to update the ASC list of covered surgical procedures by adding 28 procedures to the list. Twenty-six of these procedures were among those excluded from the ASC list for CY 2009 because we believed they did not meet the definition of a covered surgical procedure based on our expectation that they would pose a significant safety risk to Medicare beneficiaries or would require an overnight stay if performed in ASCs. The other two procedures, specifically those described by CPT code 0200T (Percutaneous sacral augmentation (sacroplasty), unilateral injection(s), including the use of a balloon or mechanical device (if utilized), one or more needles) and CPT code 0201T (Percutaneous sacral augmentation (sacroplasty), bilateral injections, including the use of a balloon or mechanical device (if utilized), two or more needles), are new Category III CPT codes that became effective July 1, 2009, and were implemented in the July 2009 ASC update (Table 57 above). As a result of our clinical evaluation of the procedures described by the new Category III codes, we determined that these two new procedures may be appropriately provided to Medicare beneficiaries in ASCs.

In response to comments on the CY 2009 proposed rule, we stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68724) that, as we developed the CY 2010 proposed rule, we would perform a comprehensive review of the APCs in order to identify potentially inconsistent ASC treatment of procedures assigned to a single APC under the OPPS. Thus, we examined surgical procedures that are excluded from the current ASC list of covered surgical procedures and the APCs to which they are assigned under the OPPS. We identified for review 223 excluded surgical procedures that were assigned to the same APCs in CY 2009 as one or more ASC covered surgical procedures. Based upon our clinical review of those procedures, we determined that 26 surgical procedures may be appropriate for performance in ASCs and proposed to add them to the CY 2010 ASC list of covered surgical procedures and to assign payment indicator "G2" (Non office-based

surgical procedure added in CY 2008 or later; payment based on OPPS relative payment weight) to each of them. We found that the remaining 197 excluded procedures would pose significant safety risks to beneficiaries or would be expected to require an overnight stay if provided in ASCs. Therefore, we did not propose to add those 197 procedures to the CY 2010 ASC list of covered surgical procedures.

The 28 procedures that we proposed to add to the ASC list of covered surgical procedures, including their HCPCS code short descriptors and proposed CY 2010 payment indicators, were displayed in Table 41 in the CY 2010 OPPS/ASC proposed rule (74 FR 35379 through 35380).

Among the procedures we identified as meeting the criteria for designation as a covered surgical procedure was CPT code 35475 (Transluminal balloon angioplasty, percutaneous; brachiocephalic trunk or branches, each vessel). The volume and utilization data for this procedure indicate that it is most frequently performed in outpatient settings. After review, our CMS medical advisors found that it would be appropriate to propose designation of CPT code 35475 as an ASC covered surgical procedure for CY 2010. Related to our proposal to add CPT code 35475 to the list of covered surgical procedures is our concurrent proposal to delete two Level II HCPCS codes we created effective for CY 2007, HCPCS codes G0392 (Transluminal balloon angioplasty, percutaneous; for maintenance of hemodialysis access, arteriovenous fistula or graft; arterial) and G0393 (Transluminal balloon angioplasty, percutaneous; for maintenance of hemodialysis access, arteriovenous fistula or graft; venous) to enable ASCs to receive Medicare payment for providing the angioplasty services required to maintain the arteriovenous fistulae that are important to individuals who undergo routine dialysis. We proposed to delete HCPCS codes G0392 and G0393 concurrently with the designation of CPT code 35475 as a covered surgical procedure because there no longer would be a need for the two Level II HCPCS G-codes. ASCs would be able to use CPT 35475 and CPT code 35476 (Transluminal balloon angioplasty, percutaneous; venous), which was included on the list of ASC covered surgical procedures beginning in CY 2008, to report the same procedures currently reported by HCPCS codes G0392 and G0393.

Thus, we proposed to add the 28 surgical procedures listed in Table 41 of the OPPS/ASC proposed rule to the list of covered ASC surgical procedures and

to delete the HCPCS codes displayed in Table 42 of the proposed rule (74 FR 35380).

Comment: One commenter requested that CMS not finalize several of its proposed additions to the ASC list for CY 2010. The commenter believed that the procedure described by CPT code 26037 (Decompressive fasciotomy, hand) is inappropriate for the ASC setting. The commenter stated that the typical patient that requires this procedure has had a severe crush injury to the hand and/or an infection and would require at least 23 hours of medical monitoring following the surgery. The commenter also objected to adding procedures described by CPT codes 42225 (Palatoplasty for cleft palate; attachment pharyngeal flap) and 42227 (Lengthening of palate, with island flap) because those procedures are overwhelmingly performed on infants and very young children who would require at least an overnight stay to observe for postoperative swelling, airway compromise, and bleeding. The commenter believed that CMS' inclusion of these procedures on the ASC list would be inappropriate because Medicare claims data are inadequate for CMS' use in making its determination and that inclusion of the procedures on the Medicare ASC list leads to interpretation by commercial insurers that the ASC setting is appropriate for all patient populations and would result in very young patients not being able to receive care in more appropriate settings.

The commenter also reiterated a previous request (73 FR 68729) that CMS remove other cleft lip and palate reconstruction procedures from the ASC list of covered surgical procedures. Those procedures and their CPT codes are: 21215 (Graft, bone; mandible (includes obtaining graft)); 40700 (Plastic repair of cleft lip/nasal deformity; primary, partial or complete, unilateral); 40701 (Plastic repair of cleft lip/nasal deformity, primary bilateral, one stage procedure); 42200 (Palatoplasty for cleft palate, soft and/or hard palate only); 42205 (Palatoplasty for cleft palate, with closure of alveolar ridge; soft tissue only); 42210 (Palatoplasty for cleft palate, with closure of alveolar ridge; with bone graft to alveolar ridge includes obtaining graft); 42215 (Palatoplasty for cleft palate; major revision); and 42220 (Palatoplasty for cleft palate; secondary lengthening procedure). The commenter stated that all of these procedures require general anesthesia and close postoperative monitoring and are often performed in the inpatient setting.

Response: Our medical advisors reviewed the three procedures described by CPT codes 26037, 42225, and 42227 that we proposed to add to the ASC list for CY 2010. As a result of that review, we continue to believe that all three of the procedures may be appropriately provided to a Medicare beneficiary in an ASC. We do not see a basis for removing the eight procedures from the ASC list as requested by the commenter. All of these procedures were on the list of covered surgical procedures even before CY 2007 and, to our knowledge, have been safely performed in ASCs for many years. We continue to believe that these 11 procedures would not pose a significant safety risk to Medicare beneficiaries and would not require an overnight stay if performed in ASCs.

As established at § 416.166(b), decisions regarding whether a surgical procedure should be excluded from the Medicare ASC list of covered surgical procedures are based on assessments of the needs of Medicare beneficiaries and not all patient populations. We include on the ASC list all procedures we believe are appropriate for some Medicare beneficiaries in order to provide physicians and patients with the greatest possible choice for sites-of-service. We expect that physicians will consider for each individual patient which site-of-service is most appropriate. We understand that the procedures on the ASC list are sometimes more appropriately performed on an inpatient basis due to the individual's age or other clinical considerations.

Comment: Several commenters supported the proposal to add 28 procedures to the list of covered surgical procedures and requested that CMS add 24 additional surgical procedures. A few commenters on the CY 2010 OPPTS/ASC proposed rule and on the CY 2009 OPPTS/ASC final rule with comment period requested that a total of 18 specific unlisted codes be added to the ASC list. Some commenters provided specific reasons for their requests for addition of particular procedures, but for most of the requested additions, no specific information was submitted.

The commenters who requested that the procedure reported by CPT code 50593 (Ablation, renal tumor(s) unilateral, percutaneous, cryotherapy) be added to the ASC list stated that the procedure is similar to the procedure reported by CPT code 50592 (Ablation, 1 or more renal tumor(s), percutaneous, unilateral, radiofrequency), which is already on the ASC list, and that CPT code 50593 is compatible with CMS' safety criteria. The commenters also reported that a significant number of the

laminectomy procedures listed in Table 60 below already are being performed in ASCs for commercially insured patients. They stated that the most common of these laminectomy procedures are performed on the cervical and lumbar regions of the spine, take only 60 to 90 minutes to perform, and typically require only about 4 hours of recovery time. They also stated that patients are carefully screened before the ASC is selected as the appropriate site for their surgical procedures, a practice they would expect to see applied to the Medicare population as well.

The commenters who requested the addition of the procedure reported by CPT code 52649 (Laser enucleation of the prostate with morcellation, including control of postoperative bleeding, complete (vasectomy, meatotomy, cystourethroscopy, urethral calibration and/or dilation, internal urethrotomy and transurethral resection of prostate are included if performed)) requested that the procedure be added to the ASC list because it does not require an overnight stay and it is similar to benign prostatic hypertrophy treatment procedures, which are already included on the ASC list. The commenters who requested addition of the procedure described by CPT code 57310 (Closure of urethrovaginal fistula) reported that it should be added to the list because a substantially similar and more complex procedure, described by CPT code 57320 (Closure of vesicovaginal fistula; vaginal approach), is already on the ASC list.

The commenters who requested the addition of the unlisted procedure described by CPT code 19499 (Unlisted procedure, breast) stated that it should be added to the list because it is used for setting breast ductoscopy prior to some surgeries in lieu of a ductogram and because it may be used to report services described by CPT codes 0046T (Catheter lavage of a mammary duct(s) for collection of cytology specimen(s), in high risk individuals (GAIL risk scoring or prior personal history of breast cancer), each breast; single duct) and 0047T (Catheter lavage of a mammary duct(s) for collection of cytology specimen(s), in high risk individuals (GAIL risk scoring or prior personal history of breast cancer), each breast; each additional duct) that were payable in ASCs in CY 2008 but were deleted effective for CY 2009. Some commenters requested the addition of unlisted CPT codes 55899 (Unlisted procedure, male genital system); 58999 (Unlisted procedure, female genital system (nonobstetrical)); and 64999 (Unlisted procedure, nervous system) because these codes may be used to

report the procedures that were described by CPT codes 0027T (Endoscopic lysis of epidural adhesions with direct visualization using mechanical means (eg, spinal endoscopic catheter system) or solution injection (eg, normal saline) including radiologic localization and epidurography); 0031T (Speculoscopy); 0032T (Speculoscopy; with directed sampling); and 53853 (Transurethral destruction of prostate tissue; by water-induced thermotherapy) that were payable in ASCs in CY 2008 but were deleted effective for CY 2009.

Finally, the commenter who requested the addition of intravascular stent placement procedures, CPT codes 37205 (Transcatheter placement of an intravascular stent(s)(except coronary, carotid, and vertebral vessel), percutaneous; initial vessel) and 37206 (Transcatheter placement of an intravascular stent(s) (except coronary, carotid, and vertebral vessel), percutaneous; each additional vessel), claimed that the addition of these procedures to the ASC list of covered surgical procedures would improve access to care for patients with vascular access dysfunction, decrease costs to Medicare, and result in a higher quality of care for these patients. The commenters requested that if CMS is reluctant to add these procedures to the ASC list because CPT codes 37205 and 37206 are not restricted to the treatment of hemodialysis vascular access sites, CMS could allow reporting of the codes for ASC payment with a new and distinct modifier that would apply to hemodialysis vascular access procedures.

All of the procedures requested by commenters for addition to the ASC list of covered surgical procedures are displayed in Tables 60 and 61 below.

TABLE 60—SURGICAL PROCEDURES REQUESTED FOR ADDITION TO THE CY 2010 ASC LIST OF COVERED SURGICAL PROCEDURES

CY 2010 CPT code	CY 2010 short descriptor
27485	Surgery to stop leg growth.
29867	Allgrft implnt, knee w/scope.
29868	Meniscal trnsp, knee w/scope.
35470	Repair arterial blockage.
35474	Repair arterial blockage.
35493	Atherectomy, percutaneous.
35495	Atherectomy, percutaneous.
37205	Transcath iv stent, percut.
37206	Transcath iv stent/perc addl.
50593	Perc cryo ablate renal tum.
52649	Prostate laser enucleation.
57310	Repair urethrovaginal lesion.
60210	Partial thyroid excision.
60220	Partial removal of thyroid.

TABLE 60—SURGICAL PROCEDURES REQUESTED FOR ADDITION TO THE CY 2010 ASC LIST OF COVERED SURGICAL PROCEDURES—Continued

CY 2010 CPT code	CY 2010 short descriptor
63001	Removal spinal lamina.
63005	Removal spinal lamina.
63020	Neck spine disk surgery.
63030	Low back disk surgery.
63035	Spinal disk surgery add-on.
63040	Laminotomy, single cervical.
63042	Laminotomy, single lumbar.
63045	Removal of spinal lamina.
63047	Removal of spinal lamina.
63048	Remove spinal lamina add-on.

TABLE 61—SPECIFIC CPT UNLISTED CODES REQUESTED FOR ADDITION TO ASC LIST OF COVERED SURGICAL PROCEDURES

CY 2010 CPT code	CY 2010 short descriptor
17999	Skin tissue procedure.
19499	Breast surgery procedure.
23929	Shoulder surgery procedure.
27599	Leg surgery procedure.
27899	Leg/ankle surgery procedure.
28899	Foot/toes surgery procedure.
29999	Arthroscopy of joint.
31299	Sinus surgery procedure.
55899	Genital surgery procedure.
58999	Genital surgery procedure.
64999	Nervous system surgery.
66999	Eye surgery procedure.
67299	Eye surgery procedure.
67399	Eye muscle surgery procedure.
67999	Revision of eyelid.
68399	Eyelid lining surgery.
68899	Tear duct system surgery.
92499	Eye service or procedure.

Response: We reviewed all of the surgical procedures that commenters requested be added to the ASC list of covered surgical procedures. We did not review any of the procedures that may be reported by the CPT unlisted codes listed in Table 61 because those codes are not eligible for addition to the ASC list, consistent with our final policy which is discussed in detail in the August 2, 2007 final rule (72 FR 42484 through 42486). We do not agree that any of the procedures recommended by the commenters are appropriate for provision to Medicare beneficiaries in ASCs. Although the commenters asserted that some of the procedures they were requesting for addition to the list are less complex than procedures already on the list and that all of the requested procedures are as safe as procedures on the list, our review did not support those assertions.

We exclude from ASC payment any procedure for which standard medical practice dictates that the beneficiary who undergoes the procedure would typically be expected to require active medical monitoring and care at midnight following the procedure (overnight stay) as well as all surgical procedures that our medical advisors determine may be expected to pose a significant safety risk to Medicare beneficiaries. The criteria used under the revised ASC payment system to identify procedures that would be expected to pose a significant safety risk when performed in an ASC include, but are not limited to, those procedures that: generally result in extensive blood loss; require major or prolonged invasion of body cavities; directly involve major blood vessels; are emergent or life-threatening in nature; or commonly require systemic thrombolytic therapy (see § 416.166).

In our review of the procedures listed in Table 60, we found that all of the procedures either may be expected to pose a threat to beneficiary safety or require active medical monitoring at midnight following the procedure. Specifically, we found that prevailing medical practice called for inpatient hospital stays for beneficiaries undergoing many of the procedures and that some of the procedures directly involve major blood vessels and/or may result in extensive blood loss.

After consideration of the public comments we received, we are not adding any of the procedures requested by the commenters to the list of ASC covered surgical procedures. We also are not removing any of the procedures from the list as requested by commenters. We are finalizing, without modification, our proposal to add 28 procedures to the CY 2010 ASC list and to delete two Level II HCPCS codes. The procedures, their short descriptors, and payment indicators are displayed in Tables 62 and 63 below.

TABLE 62—NEW ASC COVERED SURGICAL PROCEDURES FOR CY 2010

CY 2010 CPT Code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator
26037	Decompress fingers/hand.	G2
27475	Surgery to stop leg growth.	G2
27479	Surgery to stop leg growth.	G2
27720	Repair of tibia	G2
35460	Repair venous blockage.	G2

TABLE 62—NEW ASC COVERED SURGICAL PROCEDURES FOR CY 2010—Continued

CY 2010 CPT Code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator
35475	Repair arterial blockage.	G2
41512	Tongue suspension.	G2
42225	Reconstruct cleft palate.	G2
42227	Lengthening of palate.	G2
43130	Removal of esophagus pouch.	G2
43752	Nasal/ orogastric w/ stent.	G2
45541	Correct rectal prolapsed.	G2
49435	Insert subq exten to ip cath.	G2
49436	Embedded ip cath exit-site.	G2
49442	Place cecostomy tube perc.	G2
50080	Removal of kidney stone.	G2
50081	Removal of kidney stone.	G2
50727	Revise ureter ..	G2
51535	Repair of ureter lesion.	G2
57295	Revise vag graft via vagina.	G2
60210	Partial thyroid excision.	G2
60212	Partial thyroid excision.	G2
60220	Partial removal of thyroid.	G2
60225	Partial removal of thyroid.	G2
61770	Incise skull for treatment.	G2
0193T	Rf bladder neck micro-remodel.	G2
0200T*	Perq sacral augmt unilat inj.	G2
0201T*	Perq sacral augmt bilat inj.	G2

* Indicates codes are new, effective July 2009.

TABLE 63—HCPCS CODES DELETED EFFECTIVE CY 2010

CY 2009 HCPCS Code	CY 2009 short descriptor	CY 2009 ASC payment indicator
G0392	AV fistula or graft arterial.	A2

TABLE 63—HCPCS CODES DELETED
EFFECTIVE CY 2010—Continued

CY 2009 HCPCS Code	CY 2009 short descriptor	CY 2009 ASC payment indicator
G0393	AV fistula or graft venous.	A2

b. Covered Surgical Procedures Designated as Office-Based

(1) Background

In the August 2, 2007 final rule, we finalized our policy to designate as “office-based” those procedures that are added to the ASC list of covered surgical procedures in CY 2008 or later years that we determine are performed predominantly (more than 50 percent of the time) in physicians’ offices based on consideration of the most recent available volume and utilization data for each individual procedure code and/or, if appropriate, the clinical characteristics, utilization, and volume of related codes. In that rule, we also finalized our policy to exempt all procedures on the CY 2007 ASC list from application of the office-based classification (72 FR 42512). The procedures that were added to the ASC list of covered surgical procedures beginning in CY 2008 that we determined were office-based were identified in Addendum AA to that rule by payment indicator “P2” (Office-based surgical procedure added to ASC list in CY 2008 or later with MPFS nonfacility PE RVUs; payment based on OPFS relative payment weight); “P3” (Office-based surgical procedure added to ASC list in CY 2008 or later with MPFS nonfacility PE RVUs; payment based on MPFS nonfacility PE RVUs); or “R2” (Office-based surgical procedure added to ASC list in CY 2008 or later without MPFS nonfacility PE RVUs; payment based on OPFS relative payment weight), depending on whether we estimated it would be paid according to the standard ASC payment methodology based on its OPFS relative payment weight or at the MPFS nonfacility PE RVU amount.

Consistent with our final policy to annually review and update the list of surgical procedures eligible for payment in ASCs, each year we identify surgical procedures as either temporarily or permanently office-based after taking into account updated volume and utilization data.

(2) Changes to Covered Surgical Procedures Designated as Office-Based for CY 2010

In developing the CY 2010 OPFS/ASC proposed rule (74 FR 35381), we followed our policy to annually review and update the surgical procedures for which ASC payment is made and to identify new procedures that may be appropriate for ASC payment, including their potential designation as office-based. We reviewed CY 2008 volume and utilization data and the clinical characteristics for all surgical procedures that are assigned payment indicator “G2” in CY 2009, as well as for those procedures assigned one of the temporary office-based payment indicators, specifically “P2*,” “P3*,” or “R2*” in the CY 2009 OPFS/ASC final rule with comment period (73 FR 68730 through 68733). As a result of that review, we proposed to newly designate 6 procedures as office-based for CY 2010 (74 FR 35381). We also proposed to make permanent the office-based designations of 4 surgical procedures that have temporary office-based designations in CY 2009.

The 6 procedures we proposed to permanently designate as office-based are: CPT code 15852 (Dressing change (for other than burns) under anesthesia (other than local)); CPT code 19105 (Ablation, cryosurgical, of fibroadenoma, including ultrasound guidance, each fibroadenoma); CPT code 20555 (Placement of needles or catheters into muscle and/or soft tissue for subsequent interstitial radioelement application (at the time of or subsequent to the procedure)); CPT code 36420 (Venipuncture, cutdown; younger than age 1 year); CPT code 50386 (Removal (via snare/capture) of internally dwelling ureteral stent via transurethral approach, without use of cystoscopy, including radiological supervision and interpretation); and CPT code 57022 (Incision and drainage of vaginal hematoma; obstetrical/postpartum). These procedures and their HCPCS code short descriptors and proposed CY 2010 payment indicators were displayed in Table 43 of the CY 2010 OPFS/ASC proposed rule (74 FR 35381).

Comment: Many commenters expressed their continued disagreement with the policy to make payment at the lower of the ASC rate or MPFS nonfacility PE RVU payment amount for procedures we identify as office-based and requested that CMS not finalize any of the proposed office-based designations. They believed that, due to the payment limits required by CMS’ payment policy for providing these procedures in ASCs, beneficiaries who

require the level of care provided in ASCs instead have to receive treatment in the more costly HOPD setting and that the policy makes the ASC payment rates subject to the fluctuations of the MPFS nonfacility PE RVUs and other problems of the MPFS.

The commenters recommended that CMS revise its policies in at least three ways. First, they recommended that CMS establish a minimum volume threshold before designating a procedure office-based. They asserted that it is unnecessary and inappropriate to designate as office-based procedures with extremely low volume because the utilization data for those procedures are variable and, thus, not reliable. Second, they recommended that CMS raise the utilization threshold above 50 percent for designating a procedure office-based and use multiple years of data for making the designation because of the variability in the utilization data across years. Third, the commenters recommended that CMS recognize the OPFS median costs for procedures as the best proxy for relative ASC costs and limit the reduction in payment to ASCs for the office-based procedures. The commenters reasoned that, because the ASC payment system was to be based on the relative payment weights under the OPFS, which are updated annually based on hospital claims and cost reports, OPFS median costs are a better source of relative payment weights than the MPFS, which is not based on facility costs estimated from claims and cost reports like the OPFS. In addition, they expressed concern that as more procedures are designated as office-based for ASC payment, the linkage between the OPFS and ASC ratesetting methodologies would be eroded and the ASC payment system would be increasingly affected by the unpredictable inflation updates under the MPFS.

Response: We continue to believe that our policy of identifying low complexity procedures that are usually provided in physicians’ offices and limiting their payment in ASCs to the physician’s office payment amount is necessary and valid. We believe this is the most appropriate approach to preventing the creation of payment incentives for services to move from physicians’ offices to ASCs for the many newly-covered low complexity procedures on the ASC list. Moreover, we are confident that the CY 2008 claims data, the most recent full year of volume and utilization data, are an appropriate source to inform our decisions regarding the site-of-service for procedures. In our review process, when we believe that the available data are inadequate bases

upon which to make a determination that a procedure should be office-based, we either make no change to the procedure's payment status or make the change temporary and reevaluate our decision using data that become available for our next evaluation. We believe that it is appropriate to continue using our judgment regarding whether the volume of cases and the proportion of cases that are provided in the physicians' office setting indicate that the procedure is an office-based procedure in addition to our medical advisors' clinical judgments, utilization data for procedures that are closely related to the procedures being evaluated, and any other information that is available to us. Thus, we will not alter our review and decision processes with respect to establishing or changing volume or utilization thresholds as recommended by the commenters.

We continue to believe that it is appropriate that ASCs be paid no more for performing office-based procedures than those procedures would be paid when performed in physicians' offices, in order to deter inappropriate migration of these surgical procedures to ASCs based on financial

considerations rather than clinical needs. Although our policy to pay for some services at the MPFS non-facility PE RVU amount does introduce payment for a number of procedures at rates not based on the ASC relative payment weights and, as such, may be viewed as an erosion of the linkage between the OPFS and ASC payment system, we do not believe that the alternative of making payments at the higher ASC rate is preferable. None of the office-based procedures was eligible for ASC payment prior to implementation of the revised payment system and we see no inherent unfairness in limiting ASC payment to the rate for the lower-intensity site of service (physician's office) that our data indicate is the care setting for most Medicare cases. In fact, the lower than expected CY 2008 ASC utilization for office-based procedures reported by commenters (discussed in section XV.I.2. of this final rule with comment period) may be an indication that our policy does not encourage inappropriate migration of these services to ASCs, as was our intention. While we acknowledge the potential volatility of the office-based payments under the

MPFS, we continue to believe it is appropriate to treat office-based procedures similarly in the office and ASC settings for purposes of Medicare payment. Therefore, we also will not adopt the commenters' recommendation that ASC payment rates should be based only on OPFS median costs and will continue to update the office-based list of ASC covered surgical procedures annually, to account for changes in medical practice and new surgical procedures that may result in additional surgical procedures that are predominantly performed in physicians' offices.

The utilization data for the procedures listed in Table 43 of the proposed rule and restated in Table 64, below, did not change between the proposed rule and this final rule with comment period. We did not receive any public comments that specifically addressed our proposals to designate the 6 procedures listed in Table 64 as office-based for CY 2010. Therefore, we are finalizing our CY 2010 proposals, without modification, to designate the procedures displayed in Table 64 below as office-based for CY 2010.

TABLE 64—CY 2010 FINAL DESIGNATIONS OF ASC COVERED SURGICAL PROCEDURES NEWLY DESIGNATED AS OFFICE-BASED

CY 2010 HCPCS code	CY 2010 short descriptor	CY 2009 ASC payment indicator	Proposed CY 2010 ASC payment indicator	Final CY 2010 ASC payment indicator*
15852	Dressing change not for burn	G2	R2	R2
19105	Cryosurg ablate fa, each	G2	P3	P2
20555	Place ndl musc/tis for rt	G2	R2	R2
36420	Vein access cutdown < 1 yr	G2	R2	R2
50386	Remove stent via transureth	G2	P2	P2
57022	I & d vaginal hematoma, pp	G2	R2	R2

* Payment indicators are based on a comparison of the rates according to the ASC standard ratesetting methodology and the MPFS final CY 2010 rates. Under current law, the MPFS payment rates have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule with comment period.

In the CY 2010 OPFS/ASC proposed rule (74 FR 35381), we also reviewed CY 2008 volume and utilization data and other information for the 10 procedures with temporary office-based designations for CY 2009. Among these 10 procedures, there were no claims data for the 3 procedures with CPT codes that were new in CY 2009. Those 3 new procedure codes are: CPT code 46930 (Destruction of internal hemorrhoid(s) by thermal energy (eg, infrared coagulation, cautery, radiofrequency)); CPT code 64455 (Injection(s), anesthetic agent and/or steroid, plantar common digital nerve(s) (eg, Morton's neuroma)); and CPT code 64632 (Destruction by neurolytic agent;

plantar common digital nerve). Consequently, in the CY 2010 OPFS/ASC proposed rule (74 FR 35381), we proposed to maintain their temporary office-based designations for CY 2010.

As a result of our review of the remaining 7 procedures that have temporary office-based designations for CY 2009, we proposed to make permanent the office-based designations for 4 procedures for CY 2010. The 4 surgical procedure codes are: CPT code 0084T (Insertion of a temporary prostatic urethral stent); CPT code 21073 (Manipulation of temporomandibular joint(s) (TMJ), therapeutic, requiring an anesthesia service (ie, general or monitored

anesthesia care)); CPT code 55876 (Placement of interstitial device(s) for radiation therapy guidance (eg, fiducial markers, dosimeter), prostate (via needle, any approach), single or multiple); and HCPCS code C9728 (Placement of interstitial device(s) for radiation therapy/surgery guidance (eg, fiducial markers, dosimeter), other than prostate (any approach), single or multiple). Although we have no Medicare volume and utilization data in physicians' offices for HCPCS code C9728 because this code is not recognized for payment under the MPFS, we noted in the CY 2009 OPFS/ASC proposed rule (73 FR 41528) that because HCPCS code C9728 is

analogous to CPT code 55876, we believe they should be paid according to the same ASC payment methodology under the ASC payment system. In the CY 2010 OPPI/ASC proposed rule, we indicated our belief that the volume and utilization data for CPT codes 0084T, 21073, and 55876 are sufficient to support our determination that these procedures are most commonly provided in physicians' offices. Therefore, we proposed to make permanent the office-based designations for the four procedures (including HCPCS code C9728) for CY 2010.

We did not propose to make permanent the office-based designations for the 3 other procedures for which the CY 2009 office-based designations are temporary because we did not believe that the currently available volume and utilization data provided an adequate basis for proposing permanent office-based designations. Rather, available

data supported our determination that maintaining the temporary office-based designation was appropriate for CY 2010 for CPT code 0099T (Implantation of intrastromal corneal ring segments); CPT code 0124T (Conjunctival incision with posterior extrac scleral placement of pharmacological agent (does not include supply of medication)); and CPT code 67229 (Treatment of extensive or progressive retinopathy, 1 or more sessions; preterm infant (less than 37 weeks gestation at birth), performed from birth up to 1 year of age (eg, retinopathy of prematurity), photocoagulation or cryotherapy). Thus, we proposed to maintain the temporary office-based designation for those procedures for CY 2010.

The procedures that we proposed to permanently designate as office-based for CY 2010 that were temporarily designated as office-based procedures in CY 2009 were displayed in Table 44 in

the CY 2010 OPPI/ASC proposed rule (74 FR 35382). The procedures that we proposed to continue to temporarily designate as office-based for CY 2010 were displayed in Table 45 in the CY 2010 OPPI/ASC proposed rule (74 FR 35382). The procedures for which the proposed office-based designation for CY 2010 is temporary also were indicated by an asterisk in Addendum AA to the proposed rule.

We did not receive any public comments that specifically addressed our proposals to designate the 4 procedures listed in Table 44 of the proposed rule and restated in Table 65, below, as permanently office-based for CY 2010. Therefore, we are adopting as final for CY 2010 permanent office-based payment indicators as proposed for the procedures reported by HCPCS codes 0084T, 21073, 55876, and C9728 that were designated temporarily office-based for CY 2009.

TABLE 65—CY 2009 TEMPORARILY DESIGNATED OFFICE-BASED ASC COVERED SURGICAL PROCEDURES THAT ARE DESIGNATED AS PERMANENTLY OFFICE-BASED FOR CY 2010

CY 2009 HCPCS code	CY 2010 HCPCS code	CY 2010 short descriptor	CY 2009 ASC payment indicator	Final CY 2010 ASC payment indicator**
0084T	53855	Temp prostate urethral stent	R2*	P2
21073	21073	Mnpj of tmj w/anesth	P3*	P3
55876	55876	Place rt device/marker, pros	P3*	P3
C9728	C9728	Place device/marker, non pro	R2*	R2

* If designation is temporary.

** Payment indicators are based on a comparison of the rates according to the ASC standard ratesetting methodology and the MPFS final CY 2010 rates. Under current law, the MPFS payment rates have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule with comment period.

Comment: One commenter supported the proposal to maintain CPT codes 64455 and 64632 as temporarily office-based pending the availability of actual claims data.

Response: We appreciate the commenter's support.

After consideration of the public comment we received, we are finalizing our CY 2010 proposal, without modification, to maintain the temporary office-based payment indicators as displayed in Table 66 for the 6 procedures reported by CPT codes 0099T, 0124T, 46930, 64455, 64632, and 67229 that were designated temporarily office-based for CY 2009.

TABLE 66—CY 2009 TEMPORARILY OFFICE-BASED PROCEDURES THAT ARE DESIGNATED AS TEMPORARILY OFFICE-BASED FOR CY 2010 *

CY 2010 HCPCS Code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator**
0099T	Implant corneal ring.	R2*
0124T	Conjunctival drug placement.	R2*
46930	Destroy internal hemorrhoids.	P3*
64455	N block inj, plan-tar digit.	P3*
64632	N block inj, com-mon digit.	P3*
67229	Tr retinal les preterm inf.	R2*

* If designation is temporary.

** Payment indicators are based on a comparison of the rates according to the ASC standard ratesetting methodology and the MPFS final CY 2010 rates. Under current law, the MPFS payment rates have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule with comment period.

Displayed in Table 67 below are new (or substantially revised) CY 2010 HCPCS codes to which we have assigned temporary office-based payment indicators. As explained in section XV.B.1. of this final rule with comment period, we reviewed all of the newly created HCPCS codes that became available after the issuance of the CY 2009 OPPI/ASC proposed rule that will be used to report surgical procedures in CY 2010 to evaluate their appropriateness for the ASC list of covered surgical procedures. Of the procedures reported by new or substantially revised CY 2010 HCPCS codes that we determined should not be excluded from the ASC list based on our clinical review, including assessment of

available utilization and volume data for any closely related procedures and consideration of other available information, we determined that 16 of the procedures would predominantly be performed in physicians' offices. However, because we had no utilization data for the procedures specifically described by these new HCPCS codes, we made the office-based designations temporary rather than permanent and will reevaluate the procedures when data become available. The temporary payment indicators for the 16 office-based procedures displayed in Table 67 are interim designations and are open to public comment during the 60-day comment period for this final rule with comment period. HCPCS codes that are new (or substantially revised) for CY 2010 are designated with an "NI" comment indicator in Addendum AA. We will respond to public comments on the interim designations in the CY 2011 OPPS/ASC final rule with comment period.

TABLE 67—FINAL CY 2010 PAYMENT INDICATORS FOR NEW CY 2010 HCPCS CODES FOR ASC COVERED SURGICAL PROCEDURES DESIGNATED AS TEMPORARILY OFFICE-BASED ON AN INTERIM BASIS

CY 2010 HCPCS Code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator**
21015	Resection of facial tumor.	R2 *
21555	Remove lesion, neck/chest.	P3 *
21930	Remove lesion, back or flank.	P3 *
23075	Removal of shoulder lesion.	P3 *
24075	Remove arm/elbow lesion.	P3 *
25075	Removal forearm lesion subcu.	P3 *
26115	Removal hand lesion, subcut.	P3 *
27047	Remove hip/pelvis lesion.	P3 *
27327	Removal of thigh lesion.	P3 *
27618	Remove lower leg lesion.	P3 *
28039	Exc foot/toe tum sc > 1.5 cm.	P3 *
28041	Exc foot/toe tum deep > 1.5 cm.	R2 *
28043	Excision of foot lesion.	P3 *
28045	Excision of foot lesion.	P3 *
28046	Resection of tumor, foot.	R2 *

TABLE 67—FINAL CY 2010 PAYMENT INDICATORS FOR NEW CY 2010 HCPCS CODES FOR ASC COVERED SURGICAL PROCEDURES DESIGNATED AS TEMPORARILY OFFICE-BASED ON AN INTERIM BASIS—Continued

CY 2010 HCPCS Code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator**
37761	Ligate leg veins open.	R2 *

* If designation is temporary.

** Payment indicators are based on a comparison of the rates according to the ASC standard ratesetting methodology and the MPFS final CY 2010 rates. Under current law, the MPFS payment rates have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule with comment period.

c. ASC Covered Surgical Procedures Designated as Device-Intensive

(1) Background

As discussed in the August 2, 2007 final rule (72 FR 42503 through 42508), we adopted a modified payment methodology for calculating the ASC payment rates for covered surgical procedures that are assigned to the subset of OPPS device-dependent APCs with a device offset percentage greater than 50 percent of the APC cost under the OPPS, in order to ensure that payment for the procedure is adequate to provide packaged payment for the high-cost implantable devices used in those procedures. We assigned payment indicators "H8" (Device-intensive procedure on ASC list in CY 2007; paid at adjusted rate) and "J8" (Device-intensive procedure added to ASC list in CY 2008 or later; paid at adjusted rate) to identify the procedures that were eligible for ASC payment calculated according to the modified methodology, depending on whether the procedure was included on the ASC list of covered surgical procedures prior to CY 2008 and, therefore, subject to transitional payment as discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68739 through 68742). The 52 device-intensive procedures for which the modified rate calculation methodology applies in CY 2009 were displayed in Table 47 and in Addendum AA to the CY 2009 OPPS/ASC final rule with comment period (73 FR 68736 through 68738 and 68840 through 68933).

(2) Changes to List of Covered Surgical Procedures Designated as Device-Intensive for CY 2010

In the CY 2010 OPPS/ASC proposed rule (74 FR 35382), we proposed to update the ASC list of covered surgical procedures that are eligible for payment according to the device-intensive procedure payment methodology for CY 2010, consistent with the proposed OPPS device-dependent APC update, reflecting the proposed APC assignments of procedures, designation of APCs as device-dependent, and APC device offset percentages based on the CY 2008 OPPS claims and cost report data available for the proposed rule. The ASC covered surgical procedures that we proposed to designate as device-intensive and that would be subject to the device-intensive procedure payment methodology for CY 2010 were listed in Table 46 in the CY 2010 OPPS/ASC proposed rule (74 FR 35383 through 35384).

Comment: Some commenters expressed general concerns regarding the sufficiency of ASC payment for device-related services and recommended modifications to the ASC device-intensive payment methodology. First, the commenters argued that CMS should not apply the ASC conversion factor to the device-related portion of the payment for all procedures for which CMS can establish a median device cost, regardless of whether they are designated as device-intensive under the established methodology. The commenters stated that, unlike ASCs' general abilities to achieve greater operational efficiencies than HOPDs, ASCs are unable to extract greater discounts on devices and expensive operative supplies than their hospital counterparts. Second, the commenters argued that CMS should not adjust the device portion of the ASC payment for device-intensive procedures by the wage index. According to the commenters, the acquisition of devices occurs on a national market, and ASCs in rural areas pay approximately the same for medical devices and equipment as facilities in more expensive labor markets. The commenters stated that CMS is underpaying device costs in markets where the wage index is low, and overpaying in markets where the wage index is high.

One commenter specifically remarked that the procedures described by CPT code 19296 (Placement of radiotherapy afterloading balloon catheter into the breast for interstitial radioelement application following partial mastectomy, includes imaging guidance; on date separate from partial

mastectomy) and CPT code 19297 (Placement of radiotherapy afterloading balloon catheter into the breast for interstitial radioelement application following partial mastectomy, includes imaging guidance; concurrent with partial mastectomy), which map to OPPS device-dependent APC 0648 (Level IV Breast Surgery), require the use of a device that has a list price that exceeds 50 percent of the median costs calculated for those CPT codes and, therefore, concluded that these procedures should be added to the ASC list of device-intensive procedures. Another commenter requested that CMS add the procedure described by CPT code 66180 (Aqueous shunt to extraocular reservoir (eg, Molteno, Schocket, Denver-Krupin)) to the ASC list of device-intensive procedures, arguing that the list prices of devices involved in performing this procedure are greater than 50 percent of the proposed ASC payment rate for this procedure for CY 2010.

Response: In the August 2, 2007 final rule (72 FR 42508), we established that the modified payment methodology for calculating ASC payment rates for device-intensive procedures shall apply to ASC covered surgical procedures that are assigned to device-dependent APCs under the OPPS for the same calendar year, where those APCs have a device cost of greater than 50 percent of the APC cost (that is, the device offset percentage is greater than 50). We continue to believe these criteria ensure that ASC payment rates are adequate to provide packaged payment for high cost implantable devices and ensure Medicare beneficiaries have access to these procedures in all appropriate settings of care. We do not agree that we should change our criteria and treat as device-intensive ASC services that are assigned to APCs for which the device offset percentage is less than 50 percent (such as the procedures described by CPT codes 19296 and 19297) or ASC services that are not assigned to OPPS device-dependent APCs (such as the procedure described by CPT code 66180). Under the modified payment methodology for ASC covered surgical procedures designated as device-intensive, we separately determine both the device payment and service payment portions of the ASC payment rate, and apply the ASC conversion factor only to the specifically calculated OPPS relative payment weight for the service portion, while providing the same packaged payment for the device portion as would be made under the OPPS. The 50-percent device offset threshold is established to ensure that

the ASC conversion factor is not applied to the costs of high cost implantable devices, which likely do not vary between ASCs and HOPDs in the same manner service costs have been shown to vary. As we have stated in the past (73 FR 68734), we believe that when device costs comprise less than 50 percent of total procedure costs, those costs are less likely to be as predictable across sites-of-service. Accordingly, we believe that it is possible for ASCs to achieve efficiencies relative to HOPDs when providing those procedures, and that the application of the ASC conversion factor to the entire ASC payment weight is appropriate.

We also do not believe it is appropriate to vary the percentage of the national payment that is wage adjusted for different services. Under the revised ASC payment system, we utilize 50 percent as the labor-related share to adjust national ASC payment rates for geographic wage differences. We apply to ASC payments the IPPS pre-floor, pre-reclassification wage index values associated with the June 2003 OMB geographic localities, as recognized under the IPPS and OPPS, in order to adjust the labor-related portion of the national ASC payment rates for geographic wage differences. Consistent with the OPPS, we apply the ASC geographic wage adjustment to the entire ASC payment rate for device-intensive procedures. As we have noted in the past (73 FR 68735), MedPAC has indicated its intent to evaluate CMS' method for adjusting payments for variations in labor costs in light of differences in labor-related costs for device-implantation services. We look forward to reviewing the results of its evaluation, as well as any recommendations it may provide, regarding the OPPS or ASC wage adjustment policy.

Comment: Some commenters argued that CMS should not subject procedures that were on the ASC list of covered surgical procedures in CY 2007 but were rarely performed in ASCs prior to CY 2008 to the transitional adjustment. One commenter provided a data analysis demonstrating that CPT code 55873 (Cryosurgical ablation of the prostate (includes ultrasonic guidance and monitoring)) was present on three ASC claims in CY 2007, on one claim in CY 2006, and was not billed at all by ASCs in CY 2005. According to the commenter, the transitional payment for CPT code 55873 is inadequate to cover ASCs' costs of providing the procedure and will prevent Medicare beneficiaries from accessing this procedure in the ASC setting.

Response: As established in regulation at § 416.171(c), the transitional adjustment applies to all services on the CY 2007 ASC list of covered services. We cannot make an exception for procedures, such as the one described by CPT code 55873, that were on the CY 2007 list of covered services but were rarely performed in ASCs according to the commenter. Furthermore, as we stated in the August 2, 2007 final rule (72 FR 42520), the transition to the fully implemented revised ASC system payment system should not be asymmetrical, meaning that procedures with decreasing payments under the revised payment system should not be transitioned differently from those with increasing payments.

Comment: One commenter requested that CMS adjust the OPPS device offset percentages for ASC device-intensive payment purposes to account for the effects of charge compression. The commenter suggested that CMS "decompress" the supply median costs to minimize any artificial reductions that charge compression causes in the estimate of the OPPS device offset percentages.

Response: Charge compression is the practice of applying a lower charge markup to higher-cost services and a higher charge markup to lower-cost services. As a result of charge compression, the cost-based OPPS weights incorporate aggregation bias, undervaluing high cost items and overvaluing low cost items when an estimate of average markup, embodied in a single CCR, is applied to items of widely varying costs in the same cost center. As discussed in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68524), we did not adopt any short-term statistical regression-based adjustments under the OPPS that would serve to "decompress" the median costs for procedures involving devices, or for any other procedures. Rather, we chose to focus on long-term changes to Medicare cost reporting to address the effects of charge compression, including the creation of two new cost centers, "Medical Supplies Charged to Patients" and "Implantable Devices Charged to Patients," as discussed in more detail in section II.A.1.c.(2) of this final rule with comment period. We believe that this change to how hospitals report costs for devices and supplies will improve our future estimates of costs related to high cost implantable devices, including the device offset percentages upon which we base the device portions of ASC payment rates for device-intensive procedures.

Comment: Several commenters remarked on the adequacy of the proposed payment rates calculated according to the ASC device-intensive payment methodology for procedures involving cochlear implants, described by CPT code 69930 (Cochlear device implantation, with or without mastoidectomy). According to the commenters, the proposed increase to the ASC payment rate for CPT code 69930 is a necessary improvement to ensure beneficiary access to cochlear implants. Several commenters also supported the proposed payment rate increases for procedures involving auditory osseointegrated devices, described by CPT codes 69714 (Implantation, osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; without mastoidectomy); 69715 (Implantation, osseointegrated implant, temporal bone, with percutaneous attachment to

external speech processor/cochlear stimulator; with mastoidectomy); 69717 (Replacement (including removal of existing device), osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; without mastoidectomy); and 69718 (Replacement (including removal of existing device), osseointegrated implant, temporal bone, with percutaneous attachment to external speech processor/cochlear stimulator; with mastoidectomy). Other commenters encouraged CMS to finalize a somewhat higher payment rate for these procedures.

Response: We appreciate the commenters' support of the proposed payment rates for procedures involving cochlear implants and auditory osseointegrated devices. We believe that the final CY 2010 ASC payment rates for these procedures, calculated according to the ASC device-intensive ratesetting

methodology, are appropriate and adequate to ensure beneficiaries have access to these procedures in the ASC setting.

After consideration of the public comments we received, we are designating the ASC covered surgical procedures displayed in Table 68 below as device-intensive for CY 2010. The HCPCS code, the HCPCS code short descriptor, the CY 2010 ASC payment indicator, the CY 2010 OPSS APC assignment, the OPSS APC Title, and the CY 2010 OPSS APC device offset percentage are listed in Table 68. Each device-intensive procedure is assigned payment indicator "H8" or "J8," depending on whether it is subject to transitional payment. All of these procedures are included in Addendum AA to this final rule with comment period. The OPSS device-dependent APCs are discussed further in section II.A.2.d.(1) of this final rule with comment period.

TABLE 68.—ASC COVERED SURGICAL PROCEDURES DESIGNATED AS DEVICE-INTENSIVE FOR CY 2010

CY 2010 CPT code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator	Final CY 2010 OPSS APC	OPSS APC title	Final CY 2010 device-dependent APC offset percentage
24361	Reconstruct elbow joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
24363	Replace elbow joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
24366	Reconstruct head of radius	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
25441	Reconstruct wrist joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
25442	Reconstruct wrist joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
25446	Wrist replacement	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
27446	Revision of knee joint	J8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
33206	Insertion of heart pace-maker.	J8	0089	Insertion/Replacement of Permanent Pacemaker and Electrodes.	72
33207	Insertion of heart pace-maker.	J8	0089	Insertion/Replacement of Permanent Pacemaker and Electrodes.	72
33208	Insertion of heart pace-maker.	J8	0655	Insertion/Replacement/Conversion of a permanent dual chamber pacemaker.	75
33212	Insertion of pulse generator	H8	0090	Insertion/Replacement of Pacemaker Pulse Generator	74
33213	Insertion of pulse generator	H8	0654	Insertion/Replacement of a permanent dual chamber pacemaker.	75
33214	Upgrade of pacemaker system.	J8	0655	Insertion/Replacement/Conversion of a permanent dual chamber pacemaker.	75
33224	Insert pacing lead & connect.	J8	0418	Insertion of Left Ventricular Pacing Elect.	81
33225	Lventric pacing lead add-on.	J8	0418	Insertion of Left Ventricular Pacing Elect.	81
33240	Insert pulse generator	J8	0107	Insertion of Cardioverter-Defibrillator	89
33249	Eltrd/insert pace-defib	J8	0108	Insertion/Replacement/Repair of Cardioverter-Defibrillator Leads.	88
33282	Implant pat-active ht record	J8	0680	Insertion of Patient Activated Event Recorders	73
53440	Male sling procedure	H8	0385	Level I Prosthetic Urological Procedures	59
53444	Insert tandem cuff	H8	0385	Level I Prosthetic Urological Procedures	59
53445	Insert uro/ves nck sphincter	H8	0386	Level II Prosthetic Urological Procedures	71
53447	Remove/replace ur sphincter.	H8	0386	Level II Prosthetic Urological Procedures	71
54400	Insert semi-rigid prosthesis	H8	0385	Level I Prosthetic Urological Procedures	59
54401	Insert self-contd prosthesis	H8	0386	Level II Prosthetic Urological Procedures	71
54405	Insert multi-comp penis pros.	H8	0386	Level II Prosthetic Urological Procedures	71
54410	Remove/replace penis prosth.	H8	0386	Level II Prosthetic Urological Procedures	71
54416	Remv/repl penis contain pros.	H8	0386	Level II Prosthetic Urological Procedures	71

TABLE 68.—ASC COVERED SURGICAL PROCEDURES DESIGNATED AS DEVICE-INTENSIVE FOR CY 2010—Continued

CY 2010 CPT code	CY 2010 short descriptor	Final CY 2010 ASC payment indicator	Final CY 2010 OPSS APC	OPSS APC title	Final CY 2010 device-dependent APC offset percentage
55873	Cryoablate prostate	H8	0674	Prostate Cryoablation	56
61885	Insrt/redo neurostim 1 array.	H8	0039	Level I Implantation of Neurostimulator Generator	85
61886	Implant neurostim arrays ...	H8	0315	Level II Implantation of Neurostimulator Generator	88
62361	Implant spine infusion pump.	H8	0227	Implantation of Drug Infusion Device	83
62362	Implant spine infusion pump.	H8	0227	Implantation of Drug Infusion Device	83
63650	Implant neuroelectrodes ...	H8	0040	Percutaneous Implantation of Neurostimulator Electrodes.	58
63655	Implant neuroelectrodes ...	J8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr.	64
63685	Insrt/redo spine n generator.	H8	0039	Level I Implantation of Neurostimulator Generator	85
64553	Implant neuroelectrodes ...	H8	0040	Percutaneous Implantation of Neurostimulator Electrodes.	58
64555	Implant neuroelectrodes ...	J8	0040	Percutaneous Implantation of Neurostimulator Electrodes.	58
64560	Implant neuroelectrodes ...	J8	0040	Percutaneous Implantation of Neurostimulator Electrodes.	58
64561	Implant neuroelectrodes ...	H8	0040	Percutaneous Implantation of Neurostimulator Electrodes.	58
64565	Implant neuroelectrodes ...	J8	0040	Percutaneous Implantation of Neurostimulator Electrodes.	58
64573	Implant neuroelectrodes ...	H8	0225	Implantation of Neurostimulator Electrodes, Cranial Nerve.	73
64575	Implant neuroelectrodes ...	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr.	64
64577	Implant neuroelectrodes ...	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr.	64
64580	Implant neuroelectrodes ...	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr.	64
64581	Implant neuroelectrodes ...	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr.	64
64590	Insrt/redo pn/gastr stimulat ...	H8	0039	Level I Implantation of Neurostimulator Generator	85
65770	Revise cornea with implant	H8	0293	Level V Anterior Segment Eye Procedures	62
69714	Implant temple bone w/ stimulat.	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
69715	Temple bne implnt w/ stimulat.	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
69717	Temple bone implant revision.	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
69718	Revise temple bone implant.	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58
69930	Implant cochlear device	H8	0259	Level VII ENT Procedures	85

d. ASC Treatment of Surgical Procedures Removed From the OPSS Inpatient List for CY 2010

As we discussed in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68724), we adopted a policy to include in our annual evaluation procedures proposed for removal from the OPSS inpatient list for possible inclusion on the ASC list of covered surgical procedures. The final list of procedures removed from the inpatient list for CY 2009 may be found in section XI.B. of that final rule with comment period.

We evaluated each of the 3 procedures we proposed to remove from

the OPSS inpatient list for CY 2010 according to the criteria for exclusion from the list of covered ASC surgical procedures. As we stated in the CY 2010 OPSS/ASC proposed rule (74 FR 35384), we believe that all of these procedures should continue to be excluded from the ASC list of covered surgical procedures for CY 2010 because they would be expected to pose a significant risk to beneficiary safety or to require an overnight stay in ASCs. A full discussion about the APC Panel's recommendations regarding the procedures we proposed to remove from the OPSS inpatient list for CY 2010 may be found in section XI.B. of this final

rule with comment period. The HCPCS codes for these 3 procedures and their long descriptors were listed in Table 47 in the CY 2010 OPSS/ASC proposed rule (74 FR 35384).

We did not receive any public comments specifically about our proposal to continue to exclude from the ASC list the 3 procedures reported by CPT codes 21256 (Reconstruction of orbit with osteotomies (extracranial) and with bone grafts (includes obtaining autografts) (eg, micro-ophthalmia); 27179 (Open treatment of slipped femoral epiphysis; osteoplasty of femoral neck (Heyman type procedure); and 51060 (Transvesical

ureterolithotomy). However, we did receive public comments requesting that we remove additional procedures from the OPPS inpatient list. In response to those comments, we removed 5 additional procedures from the OPPS inpatient list for CY 2010. The comments requesting that we remove additional procedures from the

inpatient list and our responses may be found in section XI.B. of this final rule with comment period. Our review of the 5 procedures removed from the OPPS inpatient list in response to comments convinced us that none of them was appropriate for performance in the ASC setting. Our medical advisors determined that the procedures were

expected to pose significant risks to beneficiary safety or to require an overnight stay when provided in ASCs. The final list of procedures that have been removed from the CY 2010 OPPS inpatient list but that continue to be excluded from the ASC list of covered surgical procedures is displayed in Table 69 below.

TABLE 69—PROCEDURES EXCLUDED FROM THE ASC LIST OF COVERED SURGICAL PROCEDURES FOR CY 2010 THAT WERE REMOVED FROM THE CY 2010 OPPS INPATIENT LIST

CY 2010 HCPCS code	CY 2010 long descriptor
21256	Reconstruction of orbit with osteotomies (extracranial) and with bone grafts (includes obtaining autografts) (eg, micro-ophthalmia).
27179	Open treatment of slipped femoral epiphysis; osteoplasty of femoral neck (Heyman type procedure).
28805	Amputation, foot; transmetatarsal.
37215	Transcatheter placement of intravascular stent(s), cervical carotid artery, percutaneous; with distal embolic protection.
44950	Appendectomy.
44955	Appendectomy; when done for indicated purpose at time of other major procedure (not as separate procedure) (List separately in addition to code for primary procedure).
51060	Transvesical ureterolithotomy.
63076	Discectomy, anterior, with decompression of spinal cord and/or nerve root(s), including osteophyctectomy; cervical, each additional interspace (List separately in addition to code for primary procedure).

2. Covered Ancillary Services

In the CY 2010 OPPS/ASC proposed rule (74 FR 35384), consistent with the established ASC payment system policy, we proposed to update the ASC list of covered ancillary services to reflect the proposed payment status for the services under the CY 2010 OPPS. Maintaining consistency with the OPPS resulted in proposed changes to ASC payment indicators for some covered ancillary items and services because of changes that were proposed under the OPPS for CY 2010. Comment indicator “CH,” discussed in section XV.F. of the CY 2010 OPPS/ASC proposed rule (74 FR 35390), was used in Addendum BB to that proposed rule to indicate covered ancillary services for which we proposed a change in the ASC payment indicator to maintain consistency with a proposed change in the OPPS treatment of the service for CY 2010.

Except for the Level II HCPCS codes listed in Table 40 of the proposed rule (74 FR 35379), all ASC covered ancillary services and their proposed payment indicators for CY 2010 were included in Addendum BB to the proposed rule.

We did not receive any public comments on our proposal to update the ASC list of covered ancillary services to reflect the payment status for the services under the OPPS. Therefore, we are finalizing, without modification, our proposed updates to the ASC list of covered ancillary services as proposed. All CY 2010 ASC covered ancillary services and their final payment indicators are included in Addendum

BB to this final rule with comment period.

D. ASC Payment for Covered Surgical Procedures and Covered Ancillary Services

1. Payment for Covered Surgical Procedures

a. Background

Our ASC payment policies for covered surgical procedures under the revised ASC payment system are fully described in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66828 through 66831). Under our established policy for the revised ASC payment system, the ASC standard ratesetting methodology of multiplying the ASC relative payment weight for the procedure by the ASC conversion factor for that same year is used to calculate the national unadjusted payment rates for procedures with payment indicator “G2.” For procedures assigned payment indicator “A2,” our final policy established blended rates to be used during the transitional period and, beginning in CY 2011, ASC rates calculated according to the ASC standard ratesetting methodology. The rate calculation established for device-intensive procedures (payment indicators “H8” and “J8”) is structured so that the packaged device payment amount is the same as under the OPPS, and only the service portion of the rate is subject to the ASC standard ratesetting methodology. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68722 through 68759), we

updated the CY 2008 ASC payment rates for ASC covered surgical procedures with payment indicators of “A2,” “G2,” “H8,” and “J8” using CY 2007 data, consistent with the CY 2009 OPPS update. Payment rates for device-intensive procedures also were updated to incorporate the CY 2009 OPPS device offset percentages.

Payment rates for office-based procedures (payment indicators “P2,” “P3,” and “R2”) are the lower of the MPFS nonfacility PE RVU amount (we refer readers to the CY 2010 MPFS final rule with comment period) or the amount calculated using the ASC standard ratesetting methodology for the procedure. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68722 through 68759), we updated the payment amounts for office-based procedures (payment indicators “P2,” “P3,” and “R2”) using the most recent available MPFS and OPPS data. We compared the estimated CY 2009 rate for each of the office-based procedures, calculated according to the ASC standard ratesetting methodology, to the MPFS nonfacility PE RVU amount to determine which was lower and, therefore, would be the CY 2009 payment rate for the procedure according to the final policy of the revised ASC payment system (see § 416.171(d)).

b. Update to ASC Covered Surgical Procedure Payment Rates for CY 2010

In the CY 2010 OPPS/ASC proposed rule (74 FR 35385), we proposed to update ASC payment rates for CY 2010

using the established rate calculation methodologies under § 416.171. Thus, we proposed to calculate CY 2010 payments for procedures subject to the transitional payment methodology (payment indicators “A2” and “H8”) using a blend of 75 percent of the proposed CY 2010 ASC rate calculated according to the ASC standard ratesetting methodology and 25 percent of the CY 2007 ASC payment rate, incorporating the device-intensive procedure methodology, as appropriate, for procedures assigned ASC payment indicator “H8.” We proposed to use the amount calculated under the ASC standard ratesetting methodology for procedures assigned payment indicator “G2” because these procedures are not subject to the transitional payment methodology.

We proposed payment rates for office-based procedures (payment indicators “P2,” “P3,” and “R2”) and device-intensive procedures not subject to transitional payment (payment indicator “J8”) calculated according to our established policies. Thus, we proposed to update the payment amounts for device-intensive procedures based on the CY 2010 OPPS proposal that reflects updated OPPS device offset percentages, and to make payment for office-based procedures at the lesser of the CY 2010 proposed MPFS nonfacility PE RVU amount or the proposed CY 2010 ASC payment amount calculated according to the ASC standard ratesetting methodology.

Comment: Many commenters requested that CMS modify its packaging policy to provide separate payment for procedures that were eligible for separate ASC payment prior to becoming packaged into separately payable services under the OPPS that are not reported by any of the codes within the CPT surgical code range. The commenters stated that these HCPCS codes into which minor procedure payments are packaged are not on the list of ASC covered surgical procedures. The commenters believed that, as an unintended result of CMS’ OPPS packaging policies, procedural services that meet the criteria for performance in ASCs are being excluded from coverage. They recommended that CMS adopt a policy under which packaging policy changes under the OPPS would not result in surgical procedures that were on the ASC list of covered surgical procedures in CY 2007 or CY 2008 becoming nonpayable.

Response: We did not propose to make any change in our policy to adopt the packaging decisions made under the OPPS for the ASC payment system. Further, we do not know which

procedures the commenters were referring to in their comments and, therefore, are unable to fully address their other concerns.

Comment: One commenter requested that CMS correct the ASC payment rates for the procedures reported by CPT codes 64626 (Destruction by neurolytic agent, paravertebral facet joint nerve; cervical or thoracic, single level); 64627 (Destruction by neurolytic agent, paravertebral facet joint nerve; cervical or thoracic, each additional level); and 64680 (Destruction by neurolytic agent, with or without radiologic monitoring; celiac plexus). The commenter stated that the rates in Addendum AA to the proposed rule for these procedures were incorrectly listed as \$299.12, \$158.13, and \$312.90, respectively.

Response: We reviewed the proposed payment rates for these three procedures and found that they are all correct. We believe that the commenter failed to notice that we proposed to assign two of the procedures, CPT codes 64626 and 64680, to different APCs under the OPPS for CY 2010. The proposed changes in their OPPS APC assignments resulted in lower OPPS relative payment weights and, therefore, lower proposed ASC payment rates for CY 2010. The proposed payment rate for the third procedure, CPT code 64627, is correct as displayed in Addendum AA to the CY 2010 OPPS/ASC proposed rule. There was no proposed change to the APC assignment for that procedure under the OPPS for CY 2010. Therefore, the proposed ASC payment rate change for CY 2010 is due to the recalibration of the OPPS APC relative payment weight, which was subsequently incorporated into the ASC payment system, and also due to the progression to the third year of the transition to ASC rates calculated entirely based on the standard ratesetting methodology.

After consideration of the public comments we received, we are finalizing our CY 2010 proposal, without modification, to calculate the CY 2010 final ASC payment rates according to our established methodologies.

c. Adjustment to ASC Payments for No Cost/Full Credit and Partial Credit Devices

Our ASC policy with regard to payment for costly devices implanted in ASCs at no cost or with full or partial credit as set forth in § 416.179 is consistent with the OPPS policy. The CY 2010 OPPS APCs and devices subject to the adjustment policy are discussed in section IV.B.2. of this final rule with comment period. The established ASC policy includes

adoption of the OPPS policy for reduced payment to providers when a specified device is furnished without cost or with full or partial credit for the cost of the device for those ASC covered surgical procedures that are assigned to APCs under the OPPS to which this policy applies. We refer readers to the CY 2009 OPPS/ASC final rule with comment period for a full discussion of the ASC payment adjustment policy for no cost/full credit and partial credit devices (73 FR 68742 through 68745).

In the CY 2010 OPPS/ASC proposed rule (74 FR 35385), consistent with the OPPS, we proposed to update the list of ASC covered device-intensive procedures and devices that would be subject to the no cost/full credit and partial credit device adjustment policy for CY 2010. Table 48 in the CY 2010 OPPS/ASC proposed rule (74 FR 35386 through 35387) displayed the ASC covered device-intensive procedures that we proposed would be subject to the no cost/full credit and partial credit device adjustment policy for CY 2010. Specifically, when a procedure that is listed in Table 48 is performed to implant a device that is listed in Table 49 of the proposed rule (74 FR 35387), where that device is furnished at no cost or with full credit from the manufacturer, the ASC would append the HCPCS “FB” modifier on the line with the procedure to implant the device. The contractor would reduce payment to the ASC by the device offset amount that we estimate represents the cost of the device when the necessary device is furnished without cost to the ASC or with full credit. We would provide the same amount of payment reduction based on the device offset amount in ASCs that would apply under the OPPS under the same circumstances. We stated in the CY 2010 OPPS/ASC proposed rule (74 FR 35385) that we continue to believe that the reduction of ASC payment in these circumstances is necessary to pay appropriately for the covered surgical procedure being furnished by the ASC.

We also proposed to reduce the payment for implantation procedures listed in Table 48 of the proposed rule by one-half of the device offset amount that would be applied if a device was provided at no cost or with full credit, if the credit to the ASC is 50 percent or more of the cost of the new device. The ASC would append the HCPCS “FC” modifier to the HCPCS code for a surgical procedure listed in Table 48 in the proposed rule when the facility receives a partial credit of 50 percent or more of the cost of a device listed in Table 49. In order to report that they received a partial credit of 50 percent or

more of the cost of a new device, ASCs would have the option of either: (1) Submitting the claim for the device replacement procedure to their Medicare contractor after the procedure's performance but prior to manufacturer acknowledgment of credit for the device, and subsequently contacting the contractor regarding a claim adjustment once the credit determination is made; or (2) holding the claim for the device implantation procedure until a determination is made by the manufacturer on the partial credit and submitting the claim with the "FC" modifier appended to the implantation procedure HCPCS code if the partial credit is 50 percent or more of the cost of the replacement device. Beneficiary

coinsurance would continue to be based on the reduced payment amount. We did not receive any comments on our CY 2010 proposal to continue the no cost/full credit and partial credit device adjustment policy for ASCs. For CY 2010, as we proposed, we will reduce the payment for the device implantation procedures listed in Table 70, below, by the full device offset amount for no cost/full credit cases. ASCs must append the modifier "FB" to the HCPCS procedure code when the device furnished without cost or with full credit is listed in Table 71, below, and the associated implantation procedure code is listed in Table 70. In addition, for CY 2010, we will reduce the payment for implantation procedures listed in Table

70 by one half of the device offset amount that would be applied if a device were provided at no cost or with full credit, if the credit to the ASC is 50 percent or more of the device cost. If the ASC receives a partial credit of 50 percent or more of the cost of a device listed in Table 71, the ASC must append the modifier "FC" to the associated implantation procedure code if the procedure is listed in Table 70. We are adding device HCPCS code L8680 (Implantable neurostimulator electrode, each) to the list of devices in Table 71 because we are recognizing this code as packaged under the OPPS for CY 2010, as described in section IV.B.2. of this final rule with comment period.

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TABLE 70.—CY 2010 PROCEDURES TO WHICH THE NO COST/FULL CREDIT AND PARTIAL CREDIT DEVICE ADJUSTMENT POLICY APPLIES

CY 2010 CPT Code	CY 2010 Short Descriptor	Final CY 2010 ASC Payment Indicator	Final CY 2010 OPSS APC	OPSS APC Title	Final CY 2010 OPSS Full APC Offset Percentage	Final CY 2010 OPSS Partial APC Offset Percentage
24361	Reconstruct elbow joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
24363	Replace elbow joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
24366	Reconstruct head of radius	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
25441	Reconstruct wrist joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
25442	Reconstruct wrist joint	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
25446	Wrist replacement	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
27446	Revision of knee joint	J8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
33206	Insertion of heart pacemaker	J8	0089	Insertion/Replacement of Permanent Pacemaker and Electrodes	72%	36%
33207	Insertion of heart pacemaker	J8	0089	Insertion/Replacement of Permanent Pacemaker and Electrodes	72%	36%
33208	Insertion of heart pacemaker	J8	0655	Insertion/Replacement/Conversion of a permanent dual chamber pacemaker	75%	37%
33212	Insertion of pulse generator	H8	0090	Insertion/Replacement of Pacemaker Pulse Generator	74%	37%
33213	Insertion of pulse generator	H8	0654	Insertion/Replacement of a permanent dual chamber pacemaker	75%	37%
33214	Upgrade of pacemaker system	J8	0655	Insertion/Replacement/Conversion of a permanent dual chamber pacemaker	75%	37%
33224	Insert pacing lead & connect	J8	0418	Insertion of Left Ventricular Pacing Elect.	81%	41%
33225	Lventric pacing lead add-on	J8	0418	Insertion of Left Ventricular Pacing Elect.	81%	41%
33240	Insert pulse generator	J8	0107	Insertion of Cardioverter-Defibrillator	89%	44%
33249	Eltrd/insert pace-defib	J8	0108	Insertion/Replacement/Repair of Cardioverter-Defibrillator Leads	88%	44%
33282	Implant pat-active ht record	J8	0680	Insertion of Patient Activated Event Recorders	73%	36%

CY 2010 CPT Code	CY 2010 Short Descriptor	Final CY 2010 ASC Payment Indicator	Final CY 2010 OPSS APC	OPSS APC Title	Final CY 2010 OPSS Full APC Offset Percentage	Final CY 2010 OPSS Partial APC Offset Percentage
53440	Male sling procedure	H8	0385	Level I Prosthetic Urological Procedures	59%	30%
53444	Insert tandem cuff	H8	0385	Level I Prosthetic Urological Procedures	59%	30%
53445	Insert uro/ves nck sphincter	H8	0386	Level II Prosthetic Urological Procedures	71%	35%
53447	Remove/replace ur sphincter	H8	0386	Level II Prosthetic Urological Procedures	71%	35%
54400	Insert semi-rigid prosthesis	H8	0385	Level I Prosthetic Urological Procedures	59%	30%
54401	Insert self-contd prosthesis	H8	0386	Level II Prosthetic Urological Procedures	71%	35%
54405	Insert multi-comp penis pros	H8	0386	Level II Prosthetic Urological Procedures	71%	35%
54410	Remove/replace penis prosth	H8	0386	Level II Prosthetic Urological Procedures	71%	35%
54416	Remv/repl penis contain pros	H8	0386	Level II Prosthetic Urological Procedures	71%	35%
61885	Insrt/redo neurostim 1 array	H8	0039	Level I Implantation of Neurostimulator Generator	85%	43%
61886	Implant neurostim arrays	H8	0315	Level II Implantation of Neurostimulator Generator	88%	44%
62361	Implant spine infusion pump	H8	0227	Implantation of Drug Infusion Device	83%	41%
62362	Implant spine infusion pump	H8	0227	Implantation of Drug Infusion Device	83%	41%
63650	Implant neuroelectrodes	H8	0040	Percutaneous Implantation of Neurostimulator Electrodes	58%	29%
63655	Implant neuroelectrodes	J8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr	64%	32%
63685	Insrt/redo spine n generator	H8	0039	Level I Implantation of Neurostimulator Generator	85%	43%
64553	Implant neuroelectrodes	H8	0040	Percutaneous Implantation of Neurostimulator Electrodes	58%	29%
64555	Implant neuroelectrodes	J8	0040	Percutaneous Implantation of Neurostimulator Electrodes	58%	29%
64560	Implant neuroelectrodes	J8	0040	Percutaneous Implantation of Neurostimulator Electrodes	58%	29%

CY 2010 CPT Code	CY 2010 Short Descriptor	Final CY 2010 ASC Payment Indicator	Final CY 2010 OPSS APC	OPSS APC Title	Final CY 2010 OPSS Full APC Offset Percentage	Final CY 2010 OPSS Partial APC Offset Percentage
64561	Implant neuroelectrodes	H8	0040	Percutaneous Implantation of Neurostimulator Electrodes	58%	29%
64565	Implant neuroelectrodes	J8	0040	Percutaneous Implantation of Neurostimulator Electrodes	58%	29%
64573	Implant neuroelectrodes	H8	0225	Implantation of Neurostimulator Electrodes, Cranial Nerve	73%	37%
64575	Implant neuroelectrodes	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr	64%	32%
64577	Implant neuroelectrodes	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr	64%	32%
64580	Implant neuroelectrodes	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr	64%	32%
64581	Implant neuroelectrodes	H8	0061	Laminectomy, Laparoscopy, or Incision for Implantation of Neurostimulator Electr	64%	32%
64590	Insrt/redo pn/gastr stimul	H8	0039	Level I Implantation of Neurostimulator Generator	85%	43%
69714	Implant temple bone w/stimul	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
69715	Temple bne implnt w/stimulat	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
69717	Temple bone implant revision	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
69718	Revise temple bone implant	H8	0425	Level II Arthroplasty or Implantation with Prosthesis	58%	29%
69930	Implant cochlear device	H8	0259	Level VII ENT Procedures	85%	42%

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TABLE 71—DEVICES FOR WHICH THE “FB” OR “FC” MODIFIER MUST BE REPORTED WITH THE PROCEDURE CODE IN CY 2010 WHEN FURNISHED AT NO COST OR WITH FULL OR PARTIAL CREDIT

CY 2010 device HCPCS code	CY 2010 short descriptor
C1721	AICD, dual chamber.
C1722	AICD, single chamber.
C1764	Event recorder, cardiac.
C1767	Generator, neurostim, imp.
C1771	Rep dev, urinary, w/sling.
C1772	Infusion pump, programmable.
C1776	Joint device (implantable).
C1778	Lead, neurostimulator.

TABLE 71—DEVICES FOR WHICH THE “FB” OR “FC” MODIFIER MUST BE REPORTED WITH THE PROCEDURE CODE IN CY 2010 WHEN FURNISHED AT NO COST OR WITH FULL OR PARTIAL CREDIT—Continued

CY 2010 device HCPCS code	CY 2010 short descriptor
C1779	Lead, pmkr, transvenous VDD.
C1785	Pmkr, dual, rate-resp.
C1786	Pmkr, single, rate-resp.
C1813	Prosthesis, penile, inflatab.
C1815	Pros, urinary sph, imp.
C1820	Generator, neuro rechgbat sys.
C1881	Dialysis access system.
C1882	AICD, other than sing/dual.

TABLE 71—DEVICES FOR WHICH THE “FB” OR “FC” MODIFIER MUST BE REPORTED WITH THE PROCEDURE CODE IN CY 2010 WHEN FURNISHED AT NO COST OR WITH FULL OR PARTIAL CREDIT—Continued

CY 2010 device HCPCS code	CY 2010 short descriptor
C1891	Infusion pump, non-prog, perm.
C1897	Lead, neurostim, test kit.
C1898	Lead, pmkr, other than trans.
C1900	Lead coronary venous.
C2619	Pmkr, dual, non rate-resp.
C2620	Pmkr, single, non rate-resp.
C2621	Pmkr, other than sing/dual.

TABLE 71—DEVICES FOR WHICH THE “FB” OR “FC” MODIFIER MUST BE REPORTED WITH THE PROCEDURE CODE IN CY 2010 WHEN FURNISHED AT NO COST OR WITH FULL OR PARTIAL CREDIT—Continued

CY 2010 device HCPCS code	CY 2010 short descriptor
C2622	Prosthesis, penile, non-inf.
C2626	Infusion pump, non-prog, temp.
C2631	Rep dev, urinary, w/o sling.
L8614	Cochlear device/system.
L8680	Implt neurostim elctr each.
L8685	Implt nrostm pls gen sng rec.
L8686	Implt nrostm pls gen sng non.
L8687	Implt nrostm pls gen dua rec.
L8688	Implt nrostm pls gen dua non.
L8690	Aud osseo dev, int/ext comp.

2. Payment for Covered Ancillary Services

a. Background

Our final payment policies under the revised ASC payment system for covered ancillary services vary according to the particular type of service and its payment policy under the OPPS. Our overall policy provides separate ASC payment for certain ancillary items and services integrally related to the provision of ASC covered surgical procedures that are paid separately under the OPPS and provides packaged ASC payment for other ancillary items and services that are packaged under the OPPS. Thus, we established a final policy to align ASC payment bundles with those under the OPPS (72 FR 42495).

Our ASC payment policies provide separate payment for drugs and biologicals that are separately paid under the OPPS at the OPPS rates, while we pay for separately payable radiology services at the lower of the MPFS nonfacility PE RVU (or technical component) amount or the rate calculated according to the ASC standard ratesetting methodology (72 FR 42497). In all cases, ancillary items and services must be provided integral to the performance of ASC covered surgical procedures for which the ASC bills Medicare, in order for those ancillary services also to be paid.

ASC payment policy for brachytherapy sources generally mirrors the payment policy under the OPPS. We

finalized our policy in the CY 2008 OPPS/ASC final rule with comment period (72 FR 42499) to pay for brachytherapy sources applied in ASCs at the same prospective rates that were adopted under the OPPS or, if OPPS rates were unavailable, at contractor-priced rates. Subsequent to publication of that rule, section 106 of the Medicare, Medicaid, and SCHIP Extension Act of 2007 (Pub. L. 110–173) mandated that, for the period January 1, 2008 through June 30, 2008, brachytherapy sources be paid under the OPPS at charges adjusted to cost. Therefore, consistent with our final overall ASC payment policy, we paid ASCs at contractor-priced rates for brachytherapy sources provided in ASCs during that period of time. Beginning July 1, 2008, brachytherapy sources applied in ASCs were to be paid at the same prospectively set rates that were finalized in the CY 2008 OPPS/ASC final rule with comment period (72 FR 67165 through 67188). Immediately prior to the publication of the CY 2009 OPPS/ASC proposed rule, section 142 of the Medicare Improvements for Patients and Providers Act of 2008 (Pub. L. 110–275) amended section 1833(t)(16)(C) of the Act (as amended by section 106 of the Medicare, Medicaid, and SCHIP Extension Act of 2007) to extend the requirement that brachytherapy sources be paid under the OPPS at charges adjusted to cost through December 31, 2009. Therefore, consistent with final ASC payment policy, ASCs continued to be paid at contractor-priced rates for brachytherapy sources provided integral to ASC covered surgical procedures during that period of time.

Other separately paid covered ancillary services in ASCs, specifically corneal tissue acquisition and device categories with OPPS pass-through status, do not have prospectively established ASC payment rates according to the final policies of the revised ASC payment system (72 FR 42502 and 42509). Under the revised ASC payment system, corneal tissue acquisition is paid based on the invoiced costs for acquiring the corneal tissue for transplantation. As discussed in section IV.A.1. of this final rule with comment period, new pass-through device categories may be established on a quarterly basis, but currently there are no OPPS device pass-through categories that would continue for OPPS pass-through payment (and, correspondingly, separate ASC payment) in CY 2010.

b. Payment for Covered Ancillary Services for CY 2010

In the CY 2010 OPPS/ASC proposed rule (74 FR 35388), we proposed to update the ASC payment rates and make

changes to ASC payment indicators as necessary to maintain consistency between the OPPS and ASC payment system regarding the packaged or separately payable status of services and the proposed CY 2010 OPPS and ASC payment rates. The proposed CY 2010 OPPS payment methodologies for separately payable drugs and biologicals and brachytherapy sources were discussed in sections V. and VII. of the CY 2010 OPPS/ASC proposed rule (74 FR 35324 through 35333 and 74 FR 35340 through 35343), respectively, and we proposed to set the CY 2010 ASC payment rates for those services equal to the proposed CY 2010 OPPS rates.

Consistent with established ASC payment policy (72 FR 42497), the proposed CY 2010 payment for separately payable covered radiology services was based on a comparison of the CY 2010 proposed MPFS nonfacility PE RVU amounts (74 FR 33687 through 33800) and the proposed CY 2010 ASC payment rates calculated according to the ASC standard ratesetting methodology and then set at the lower of the two amounts. Alternatively, payment for a radiology service may be packaged into the payment for the ASC covered surgical procedure if the radiology service is packaged under the OPPS. The payment indicators in Addendum BB to the CY 2010 OPPS/ASC proposed rule indicated whether the proposed payment rates for radiology services are based on the MPFS nonfacility PE RVU amount or the ASC standard ratesetting methodology, or whether payment for a radiology service is packaged into the payment for the covered surgical procedure (payment indicator “N1”). Radiology services that we proposed to pay based on the ASC standard ratesetting methodology are assigned payment indicator “Z2” (Radiology service paid separately when provided integral to a surgical procedure on ASC list; payment based on OPPS relative payment weight) and those for which the proposed payment is based on the MPFS nonfacility PE RVU amount are assigned payment indicator “Z3” (Radiology service paid separately when provided integral to a surgical procedure on ASC list; payment based on MPFS nonfacility PE RVUs).

All covered ancillary services and their proposed payment indicators were listed in Addendum BB to the CY 2010 OPPS/ASC proposed rule.

Comment: One commenter expressed continued disagreement with the ASC packaging policy related to discography services. According to the policy, the injection procedures reported by CPT codes 62290 (Injection procedure for

discography, each level; lumbar) and 62291 (Injection procedure for discography, each level; cervical or thoracic) are packaged into the services reported by CPT codes 72285 (Discography, cervical or thoracic, radiological supervision and interpretation) and 72295 (Discography, lumbar, radiological supervision and interpretation) and, therefore, separate payment is made to an ASC only when the radiology service is provided integral to a covered surgical procedure. The commenter asserted that the injection procedures reported by CPT codes 62290 and 62291 are the major procedures of the discography because they require more time and resources than the radiological services and, as such, should not be packaged into the lesser radiological services.

The commenter believed that discography has many similarities to vertebroplasty, for which separate payment is made under the ASC payment system. The commenter stated that both procedures require sedation, insertion of a needle into the spine (one into the disc and the other into the bone), and image guidance, and that material (contrast agent or bone cement, respectively) is injected into the spine in both procedures. Based on discography's similarities to the separately payable vertebroplasty procedures, the commenter requested that CMS implement separate payments for discography and radiological supervision and interpretation, recognizing that the injection procedures are the major procedures in discography.

Response: As we explained fully in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68747), we continue to believe that our packaging policy for discography services is appropriate and we do not agree that packaging policies under the ASC payment system should vary from those under the OPPS. Also, we continue to believe that discography is a radiology service, even though a component of it may be defined as surgical, and that radiology services are not appropriate for performance and separate payment in ASCs unless they are integral to covered surgical procedures.

Comment: Several commenters requested that CMS pay for low dose rate (LDR) prostate brachytherapy services under the ASC payment system based on the composite APC methodology used under the OPPS rather than making two separate payments for the services reported by CPT codes 55875 (Transperineal placement of needles or catheters into prostate for interstitial radioelement

application, with or without cystoscopy) and 77778 (Interstitial radiation source application; complex). The composite APCs were developed for procedures like LDR prostate brachytherapy in which two procedures are frequently performed in a single hospital visit. The commenters asserted that basing ASC payments for the services on the composite APC methodology in which one payment is made for the combination of the two services, would result in a more accurate payment than is currently being made to ASCs because ASC payment is based on the median costs from single-service claims that CMS has acknowledged are mostly incorrectly coded claims.

Response: Although we have tried to align the ASC and OPSS packaging policies to the fullest extent, in the case of the LDR prostate brachytherapy composite APC and other composite APCs, the differences in the payment policies across the two payment systems pose some obstacles to making payment to ASCs using the composite packages of services. In the case of the two services included in the LDR brachytherapy composite APC, the surgical procedure was on the ASC list in CY 2007 and, therefore, is subject to the transitional payment methodology in CY 2010. The other service in the LDR brachytherapy composite APC is a covered ancillary service for which the ASC payment is made at the lesser of the ASC rate calculated according to the ASC standard ratesetting methodology or the MPFS nonfacility PE RVU amount for that year. We do not see a method by which to calculate an ASC rate for the package of the two procedures that is consistent with the established ASC payment policies. Further, we did not propose to implement composite payment policies under the ASC payment system.

After consideration of the public comments we received, we are providing CY 2010 payment for covered ancillary services in accordance with the final policies of the revised ASC payment system as described in the CY 2008 OPPS/ASC final rule with comment period (72 FR 42493 through 42501). Covered ancillary services and their final CY 2010 payment indicators are listed in Addendum BB to this final rule with comment period.

E. New Technology Intraocular Lenses (NTIOLs)

1. Background

In the CY 2007 OPPS/ASC final rule with comment period (71 FR 68176), we finalized our current process for reviewing applications to establish new

active classes of new technology intraocular lenses (NTIOLs) and for recognizing new candidate intraocular lenses (IOLs) inserted during or subsequent to cataract extraction as belonging to a NTIOL class that is qualified for a payment adjustment. Specifically, we established the following process:

- We announce annually in the **Federal Register** a document that proposes the update of ASC payment rates for the following calendar year, a list of all requests to establish new NTIOL classes accepted for review during the calendar year in which the proposal is published and the deadline for submission of public comments regarding those requests. Pursuant to Section 141(b)(3) of Public Law 103-432 and our regulations at § 416.185(b), the deadline for receipt of public comments is 30 days following publication of the list of requests.

- In the **Federal Register** document that finalizes the update of ASC payment rates for the following calendar year, we—

- Provide a list of determinations made as a result of our review of all new class requests and public comments; and

- Announce the deadline for submitting requests for review of an application for a new NTIOL class for the following calendar year.

In determining whether a lens belongs to a new class of NTIOLs and whether the ASC payment amount for insertion of that lens in conjunction with cataract surgery is appropriate, we expect that the insertion of the candidate IOL would result in significantly improved clinical outcomes compared to currently available IOLs. In addition, to establish a new NTIOL class, the candidate lens must be distinguishable from lenses already approved as members of active or expired classes of NTIOLs that share a predominant characteristic associated with improved clinical outcomes that was identified for each class. Furthermore, in the CY 2007 OPPS/ASC final rule with comment period (71 FR 68227), we finalized our proposal to base our determinations on consideration of the following factors set out at § 416.195:

- The IOL must have been approved by the FDA and claims of specific clinical benefits and/or lens characteristics with established clinical relevance in comparison with currently available IOLs must have been approved by the FDA for use in labeling and advertising;

- The IOL is not described by an active or expired NTIOL class; that is, it does not share the predominant, class-

defining characteristic associated with improved clinical outcomes with designated members of an active or expired NTIOL class; and

- Evidence demonstrates that use of the IOL results in measurable, clinically meaningful, improved outcomes in comparison with use of currently available IOLs. According to the statute, and consistent with previous examples provided by CMS, superior outcomes that we consider include the following:

- Reduced risk of intraoperative or postoperative complication or trauma;
- Accelerated postoperative recovery;
- Reduced induced astigmatism;
- Improved postoperative visual acuity;
- More stable postoperative vision; and/or
- Other comparable clinical advantages, such as—

- Reduced dependence on other eyewear (for example, spectacles, contact lenses, and reading glasses);

- Decreased rate of subsequent diagnostic or therapeutic interventions, such as the need for YAG laser treatment;

- Decreased incidence of subsequent IOL exchange; and

- Decreased blurred vision, glare, other quantifiable symptom or vision deficiency.

For a request to be considered complete, we require submission of the information that is found in the guidance document entitled “Application Process and Information Requirements for Requests for a New Class of New Technology Intraocular Lens (NTIOL)” posted on the CMS Web site at: http://www.cms.hhs.gov/ASCPayment/08_NTIOls.asp #TopOfPage.

As we stated in the CY 2007 OPPTS/ASC final rule with comment period (71 FR 68180), there are three possible outcomes from our review of a request for establishment of a new NTIOL class. As appropriate, for each completed request for consideration of a candidate IOL into a new class that is received by the established deadline, one of the following determinations is announced

annually in the final rule updating the ASC payment rates for the next calendar year:

- The request for a payment adjustment is approved for the candidate IOL for 5 full years as a member of a new NTIOL class described by a new HCPCS code;

- The request for a payment adjustment is approved for the candidate IOL for the balance of time remaining as a member of an active NTIOL class; or

- The request for a payment adjustment is not approved.

We also discussed our plan to summarize briefly in the final rule with comment period the evidence that we reviewed, the public comments, and the basis for our determinations in consideration of applications for establishment of a new NTIOL class. We established that when a new NTIOL class is created, we identify the predominant characteristic of NTIOls in that class that sets them apart from other IOLs (including those previously approved as members of other expired or active NTIOL classes) and that is associated with improved clinical outcomes. The date of implementation of a payment adjustment in the case of approval of an IOL as a member of a new NTIOL class would be set prospectively as of 30 days after publication of the ASC payment update final rule, consistent with the statutory requirement.

2. NTIOL Application Process for Payment Adjustment

In CY 2007, we posted an updated guidance document to the CMS Web site to provide process and information requirements for applications requesting a review of the appropriateness of the payment amount for insertion of an IOL to ensure that the ASC payment for covered surgical procedures includes payment that is reasonable and related to the cost of acquiring a lens that is approved as belonging to a new class of NTIOls. This guidance document can be accessed on the CMS Web site at:

<http://www.cms.hhs.gov/ASCPayment/downloads/NTIOLprocess.pdf>.

We note that we have also issued a guidance document entitled “Revised Process for Recognizing Intraocular Lenses Furnished by Ambulatory Surgery Centers (ASCs) as Belonging to an Active Subset of New Technology Intraocular Lenses (NTIOls).” This guidance document can be accessed on the CMS Web site at: http://www.cms.hhs.gov/ASCPayment/Downloads/Request_for_inclusion_in_current_NTIOl_subset.pdf.

This second guidance document provides specific details regarding requests for recognition of IOLs as belonging to an existing, active NTIOL class, the review process, and information required for a request to review. Currently, there is one active NTIOL class whose defining characteristic is the reduction of spherical aberration. CMS accepts requests throughout the year to review the appropriateness of recognizing an IOL as a member of an active class of NTIOls. That is, review of candidate lenses for membership in an existing, active NTIOL class is ongoing and not limited to the annual review process that applies to the establishment of new NTIOL classes. We ordinarily complete the review of such a request within 90 days of receipt of all information that we consider pertinent to our review, and upon completion of our review, we notify the requestor of our determination and post on the CMS Web site notification of a lens newly approved for a payment adjustment as an NTIOL belonging to an active NTIOL class when furnished in an ASC.

3. Classes of NTIOls Approved and New Requests for Payment Adjustment

a. Background

Since implementation of the process for adjustment of payment amounts for NTIOls that was established in the June 16, 1999 **Federal Register**, we have approved three classes of NTIOls, as shown in the following table, with the associated qualifying IOLs to date:

NTIOL class	HCPCS code	\$50 approved for services furnished on or after	NTIOL characteristic	IOLs eligible for adjustment
1	Q1001	May 18, 2000, through May 18, 2005.	Multifocal	Allergan AMO Array Multifocal lens, model SA40N.
2	Q1002	May 18, 2000, through May 18, 2005.	Reduction in Preexisting Astigmatism.	STAAR Surgical Elastic Ultraviolet-Absorbing Silicone Posterior Chamber IOL with Toric Optic, models AA4203T, AA4203TF, and AA4203TL.

NTIOL class	HCPCS code	\$50 approved for services furnished on or after	NTIOL characteristic	IOLs eligible for adjustment
3	Q1003	February 27, 2006, through February 26, 2011.	Reduced Spherical Aberration	Advanced Medical Optics (AMO) Tecnis® IOL models Z9000, Z9001, Z9002, ZA9003, and AR40xEM and Tecnis® 1-Piece model ZCB00; Alcon Acrysof® IQ Model SN60WF, Acrysert Delivery System model SN60WS and Acrysof® IQ Toric model SN6ATT; Bausch & Lomb Sofport AO models LI61AO and LI61AOV and Akreos AO models AO60 and MI60; STAAR Affinity Collamer model CQ2015A and CC4204A and Elastimide model AQ2015A; Hoya model FY-60AD, FC-60AD, PY-60AD, and PC-60AD.

b. Request To Establish New NTIOL Class for CY 2010 and Deadline for Public Comment

As explained in the guidance document on the CMS Web site, the deadline for each year's requests for review of the appropriateness of the ASC payment amount for insertion of a candidate IOL as a member of a new class of NTIOLs is announced in the final rule updating the ASC and OPSS payment rates for that calendar year. Therefore, a request for review for a new class of NTIOLs for CY 2010 must have been submitted to CMS by March 2, 2009, the due date published in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68752). We did not receive any requests for review to establish a new NTIOL class for CY 2010 by the March 2, 2009 due date.

4. Payment Adjustment

The current payment adjustment for a 5-year period from the implementation date of a new NTIOL class is \$50. In the CY 2007 OPSS/ASC final rule with comment period, we revised § 416.200(a) through (c) to clarify how the IOL payment adjustment is made and how an NTIOL is paid after expiration of the payment adjustment, and made minor editorial changes to § 416.200(d). For CY 2008 and CY 2009, we did not revise the payment adjustment amount, and, in the CY 2010 OPSS/ASC proposed rule (74 FR 35390), we did not propose to revise the payment adjustment amount for CY 2010 in light of our limited experience with the revised ASC payment system, implemented initially on January 1, 2008. Therefore, the final ASC payment adjustment amount for NTIOLs in CY 2010 is \$50.

5. ASC Payment for Insertion of IOLs

In accordance with the final policies of the revised ASC payment system, for CY 2010, payment for IOL insertion procedures is established according to the standard payment methodology of

the revised payment system, which multiplies the ASC conversion factor by the ASC payment weight for the surgical procedure to implant the IOL. CY 2010 ASC payment for the cost of a conventional lens is packaged into the payment for the associated covered surgical procedures performed by the ASC. The HCPCS codes for IOL insertion procedures were included in Table 50 in the CY 2010 OPSS/ASC proposed rule (74 FR 35390), and their proposed CY 2010 payment rates were included in Addendum AA to that proposed rule.

We did not receive any public comments concerning the proposed CY 2010 payment rates for the insertion of IOL procedures. Therefore, we are finalizing the payment rates for the insertion of IOL procedures, calculated according to the standard methodology of the revised ASC payment system. The HCPCS codes for IOL insertion procedures are displayed in Table 72 below, and their final CY 2010 payment rates may be found in Addendum AA to this final rule with comment period.

TABLE 72—INSERTION OF IOL PROCEDURES

CY 2009 HCPCS code	CY 2009 long descriptor
66983	Intracapsular cataract extraction with insertion of intraocular lens prosthesis (one stage procedure).
66984	Extracapsular cataract removal with insertion of intraocular lens prosthesis (one stage procedure), manual or mechanical technique (eg, irrigation and aspiration or phacoemulsification).
66985	Insertion of intraocular lens prosthesis (secondary implant), not associated with concurrent cataract removal.
66986	Exchange of intraocular lens.

6. Announcement of CY 2010 Deadline for Submitting Requests for CMS Review of Appropriateness of ASC Payment for Insertion of an NTIOL Following Cataract Surgery

In accordance with § 416.185(a) of our regulations as revised by the CY 2007 OPSS/ASC final rule with comment period, CMS announces that in order to be considered for payment effective January 1, 2011, requests for review of applications for a new class of new technology IOLs must be received at CMS by 5 p.m. EST, on March 8, 2010. Send requests to ASC/NTIOL, Division of Outpatient Care, Mailstop C4-05-17, Centers for Medicare and Medicaid, 7500 Security Boulevard, Baltimore, MD 21244-1850.

To be considered, requests for NTIOL reviews must include the information on the CMS Web site at: <http://www.cms.hhs.gov/ASCPayment/downloads/NTIOLprocess.pdf>.

F. ASC Payment and Comment Indicators

1. Background

In addition to the payment indicators that we introduced in the August 2, 2007 final rule, we also created final comment indicators for the ASC payment system in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66855). We created Addendum DD1 to define ASC payment indicators that we use in Addenda AA and BB to provide payment information regarding covered surgical procedures and covered ancillary services, respectively, under the revised ASC payment system. The ASC payment indicators in Addendum DD1 are intended to capture policy-relevant characteristics of HCPCS codes that may receive packaged or separate payment in ASCs, including: their ASC payment status prior to CY 2008; their designation as device-intensive or office-based and the corresponding ASC payment methodology; and their classification as separately payable radiology services,

brachytherapy sources, OPSS pass-through devices, corneal tissue acquisition services, drugs or biologicals, or NTIOLs.

We also created Addendum DD2 that lists the ASC comment indicators. The ASC comment indicators used in Addenda AA and BB to the proposed rules and final rules with comment period serve to identify, for the revised ASC payment system, the status of a specific HCPCS code and its payment indicator with respect to the timeframe when comments will be accepted. The comment indicator "NI" is used in the OPSS/ASC final rule with comment period to indicate new HCPCS codes for the next calendar year or existing codes with substantial revisions to their descriptors such that we consider them to be describing new services or procedures for which their ASC payment indicators may change. All HCPCS codes to which the interim payment indicator is assigned are subject to comment.

The "CH" comment indicator was used in Addenda AA and BB to the CY 2010 OPSS/ASC proposed rule to indicate that a new payment indicator (in comparison with the indicator for the CY 2009 ASC April quarterly update) was proposed for assignment to an active HCPCS code for the next calendar year; an active HCPCS code was proposed for addition to the list of procedures or services payable in ASCs; or an active HCPCS code was proposed for deletion at the end of the current calendar year. The "CH" comment indicators that are published in the final rule with comment period are provided to alert readers that a change has been made from one calendar year to the next, but do not indicate that the change is subject to comment. The full definitions of the payment indicators and comment indicators are provided in Addenda DD1 and DD2 to this final rule with comment period.

2. ASC Payment and Comment Indicators

In the CY 2010 OPSS/ASC proposed rule (74 FR 35390 through 35391), we did not propose any changes to the definitions of the ASC payment and comment indicators for CY 2010 and we did not receive any public comments on the payment and comment indicators. Therefore, we are finalizing our proposed CY 2010 payment and comment indicators in Addenda DD1 and DD2 to this final rule with comment period, with modification to the meaning of comment indicator "NI" as follows. We want to clarify our policy regarding the use of comment indicator "NI" in this CY 2010 OPSS/ASC final

rule with comment period to describe a new code. There are numerous instances in which the descriptor of an existing Category I CPT code is substantially revised for CY 2010 so that it describes a new service or procedure that could have been assigned a new code number by the CPT Editorial Panel and that new code number would then have been assigned the "NI" comment indicator. Because, for CY 2010, not all new services or procedures will be assigned a new CPT code number, but instead will be described by an existing CPT code number with a substantially revised code descriptor, we are assigning the comment indicator "NI" to these codes in order to allow for comment on these substantially revised codes. Like all codes labeled with comment indicator "NI," we will respond to public comments and finalize their ASC treatment in the CY 2011 OPSS/ASC final rule with comment period. In accordance with our usual practice, CPT and Level II HCPCS code numbers that are new for CY 2010 and are ASC covered surgical procedures or covered ancillary items and services are also labeled with comment indicator "NI" in Addenda AA and BB to this final rule with comment period.

G. ASC Policy and Payment Recommendations

MedPAC was established under section 1805 of the Act to advise the U.S. Congress on issues affecting the Medicare program. Sections 1805(b)(1)(B) and 1805(b)(1)(C) of the Act require MedPAC to submit reports to Congress not later than March 1 and June 15 of each year that present its Medicare payment policy reviews and recommendations. The following section describes a recent MedPAC recommendation that is relevant to the ASC payment system.

The March 2009 MedPAC "Report to the Congress: Medicare Payment Policy" included the following recommendation relating specifically to the ASC payment system for CY 2010:

Recommendation 2B-4: The Congress should increase payments for ambulatory surgery center (ASC) services in calendar year 2010 by 0.6 percent. In addition, the Congress should require ASCs to submit to the Secretary cost data and quality data that will allow for an effective evaluation of the adequacy of ASC payment rates.

CMS Response: As we proposed in the CY 2010 proposed rule (74 FR 35391), in this final rule with comment period we are increasing the payment amounts for the ASC payment system according to our established policy as stated in the

August 2, 2007 final rule (72 FR 42518 through 42519). Section 1833(i)(2)(C) of the Act requires that, if the Secretary has not updated the ASC payment amounts in a calendar year, the payment amounts shall be increased by the percentage increase in the Consumer Price Index for All Urban Consumers (CPI-U) as estimated by the Secretary for the 12-month period ending with the midpoint of the year involved. We indicated that we planned to implement the annual updates through an adjustment to the conversion factor under the ASC payment system beginning in CY 2010 when the statutory requirement for a zero update no longer applies. Further, we noted that we were proposing to update the conversion factor for the CY 2010 ASC payment system by the percentage increase in the CPI-U, consistent with our policy as codified under § 416.171(a)(2).

We also did not propose to require ASCs to submit cost data to the Secretary for CY 2010 and, therefore, will not require cost reporting in this final rule with comment period. We explained that the 2006 GAO report, "Medicare: Payment for Ambulatory Surgical Centers Should Be Based on the Hospital Outpatient Payment System" (GAO-07-86), concluded that the APC groups in the OPSS reflect the relative costs of surgical procedures performed in ASCs in the same way they reflect the relative costs of the same procedures when they are performed in HOPDs. Consistent with the GAO findings, CMS is using the OPSS as the basis for the ASC payment system, which provides for an annual revision of the ASC payment rates under the budget neutral ASC payment system. In addition, we noted that under the methodology of the revised ASC payment system, we do not utilize ASC cost information to set and revise the payment rates for ASCs but, instead, rely on the relativity of hospital outpatient costs developed for the OPSS, consistent with the recommendation of the GAO. Furthermore, we explained that we have never required ASCs to routinely submit cost data and expressed our concern that a new Medicare requirement for ASCs to do so could be administratively burdensome for ASCs. However, in light of the MedPAC recommendation, in the CY 2010 OPSS/ASC proposed rule (74 FR 35391), we solicited public comment on the feasibility of ASCs submitting cost information to CMS, including whether costs should be collected from a sample or the universe of ASCs, the administrative burden associated with

such an activity, the form that such a submission could take considering existing Medicare requirements for other types of facilities and the scope of ASC services, the expected accuracy of such cost information, and any other issues or concerns of interest to the public on this topic.

Finally, we noted that section 109(b) of the MIEA-TRHCA (Pub. L. 109-432) gives the Secretary the authority to implement ASC quality measure reporting and to reduce the payment update for ASCs that fail to report those required measures. We restated our belief that promoting high quality care in the ASC setting through quality reporting is highly desirable and fully in line with our efforts under other payment systems. For the reasons discussed in section XVI.G. of this final rule with comment period, we did not require ASC quality data reporting for CY 2010, but our intention is to implement ASC quality reporting in a future rulemaking.

Comment: Commenters' expressed varied opinions regarding the feasibility of requiring ASCs to submit cost data to the Secretary. MedPAC's comments on CMS' solicitation in the CY 2010 OPPS/ASC proposed rule (74 FR 35391) stated that, although ASCs are generally small facilities that may have limited resources for collecting cost data, other small providers submit cost reports to CMS and, therefore, MedPAC did not believe that the resources involved in submitting cost data would be an insurmountable obstacle for ASCs. Further, MedPAC suggested that the scale of cost reporting for ASCs would be limited to the information that analysts would need to assess the adequacy of Medicare payments and evaluate the ASC update and may be satisfied by implementing either a streamlined cost report or a random survey. If a survey method is used, MedPAC recommended that CMS require ASCs to respond in order to ensure adequate data.

Other commenters, mostly those representing hospitals, believed that in light of the MedPAC recommendation that ASCs be required to submit cost data, ASCs should be required to do so even though ASC cost data are not used to set or revise the payment rates. They suggested that collection of ASC cost data could be accomplished through implementing an ASC cost reporting system or through the periodic collection of ASC cost data at the procedure level. On the other hand, some commenters (predominantly commenters who represented ASCs) opposed a requirement that ASCs submit cost data to CMS. The

commenters believed that a requirement to submit cost data would be both unnecessary and administratively burdensome for ASCs. Further, some commenters stated that requiring ASCs to submit cost data that would not be used to update or set payment rates would very likely result in submissions of data that would not be reliable.

In its comment on the statement in the proposed rule (74 FR 35391) that CMS has the authority under section 109(b) of the MIEA-TRHCA (Pub. L. 109-432) to implement ASC quality measure reporting, MedPAC noted that CMS was not required to implement that reporting as MedPAC recommended in its March 2009 Report to Congress. MedPAC expressed concern about CMS' proposal to delay implementation of ASC quality measurement reporting and argued it should be technically feasible for ASCs to report in CY 2010 on at least the five quality measures that were developed by the ASC industry-sponsored ASC Quality Collaboration and endorsed by the National Quality Forum. Briefly, these five facility-level measures are: patient being burned; patient fall in the ASC; errors related to wrong surgery, wrong patient, wrong side, wrong site or wrong implant; timing of prophylactic intravenous antibiotic; and hospital transfer/admission upon discharge from the ASC. MedPAC believed that ASCs could report these five measures without undue administrative burden and that CMS should require ASCs to report these measures without further delay.

Many other commenters also urged CMS to implement ASC quality reporting as soon as possible and reported that ASCs are anxious to begin the process. The commenters believed that CMS should ensure the availability of fair and accurate quality data from ASCs in order to promote transparency and allow beneficiaries to make meaningful comparisons across outpatient surgical settings. Some commenters believed that ASCs should be required to report quality data because ASCs should be held to the same accountability standards as hospitals with respect to the quality of care they deliver and that the ASC quality measures should be consistent, and where possible, identical to the measures reported by HOPDs.

Response: We thank all of the commenters for their thoughts regarding the feasibility and value of requiring ASCs to submit cost data that could be used to evaluate the adequacy of the Medicare ASC payment rates. We will keep the commenters' perspectives about collecting cost information from

ASCs in mind as we further consider the adequacy of the Medicare ASC payment rates.

We also appreciate the commenters' thoughtful remarks and suggestions regarding ASC quality reporting. We will be mindful of the opinions and information shared in the public comments as we move toward implementation of ASC quality reporting.

H. Revision to Terms of Agreements for Hospital-Operated ASCs

1. Background

The August 5, 1982 ASC final rule (47 FR 34082) established the initial Medicare ASC payment system and implementing Federal regulations under 42 CFR part 416. Under § 416.26 of our regulations, ASCs operated by hospitals, like other ASCs, must meet the applicable conditions for coverage and enter into an agreement with CMS in which CMS accepts the ASC as qualified to furnish ambulatory surgical services. Sections 416.30(a) through (g) of our regulations specify terms of agreement for ASCs. Section 416.30(f) specifies the following additional terms of agreement for an ASC operated by a hospital—

- The agreement is made effective on the first day of the next Medicare cost reporting period of the hospital that operates the ASC;
- The ASC participates and is paid only as an ASC, without the option of converting to or being paid as a hospital outpatient department, unless CMS determines there is good cause to do otherwise; and
- Costs incurred by the ASC are treated as a nonreimbursable cost center on the hospital's Medicare cost report.

In addition, § 416.35 provides guidance regarding the termination of ASC agreements with CMS. Voluntary terminations are those initiated by an ASC and as specified in § 416.35, an ASC may terminate its agreement either by sending written notice to CMS or by ceasing to furnish services to the community.

Although some sections of part 416 of the regulations governing ASCs have been revised since they were established in 1982, most recently for CY 2008 with the adoption of the revised ASC payment system, §§ 416.30(a) through 416.30(g) have not been changed or updated. At the time §§ 416.30 and 416.35 were promulgated, Medicare paid for hospital outpatient services on a reasonable cost basis. In contrast, Medicare initially paid ASCs for a small number of surgical procedures at one of only four prospective rates that were

developed for the ASC payment system using cost data obtained from surveys of ASCs. Since then, Medicare has adopted a prospective payment system for HOPDs (the OPPIs), the ASC list of covered surgical procedures and payment rates have been updated a number of times, and, beginning in CY 2008, the revised ASC payment system was introduced.

Under the revised ASC payment system, Medicare greatly increased the number and types of surgical procedures that are eligible for payment in ASCs. As a result, many more of the same surgical procedures may be paid under the OPPIs and ASC payment system, with the specific payment determined by whether the service is provided by a hospital or an ASC. Further, under the current, revised payment methodology, ASC payment rates have a direct relationship to the relative payment weights under the OPPIs for the same services. Today, hospital outpatient and ASC surgical procedures are paid based on the relative weights adopted for the OPPIs, and the difference between payments under the two systems is largely a reflection of the differences in capital and operating costs attributable to being an ASC or being an HOPD.

Another change that has taken place since the establishment of the Medicare ASC payment system and the implementing regulations at § 416.30 has been our effort to simplify the Medicare regulations to reduce the burden on providers and suppliers. As discussed in the August 1, 2002 IPPS final rule (67 FR 50084 through 50090), as part of that effort, we revised the provider-based status regulations at § 413.65 that outline the requirements for a determination that a facility or an organization has provider-based status as a department or entity of a hospital (main provider). The provider-based status rules generally apply to situations where there is a financial incentive for a facility or organization to claim affiliation with a main provider. The provider-based status rules establish criteria for a facility or organization to demonstrate that it is integrated with the main provider for payment purposes. We do not make provider-based status determinations for certain facilities, listed under § 413.65(a)(1)(ii) of the regulations, because the outcome of the determination (that is, whether a facility, unit, or department is found to be freestanding or provider-based) would not affect the methodology used to make Medicare or Medicaid payment, the scope of benefits available to a Medicare beneficiary in or at the facility, or the deductible or coinsurance

liability of a Medicare beneficiary in or at the facility. According to § 413.65(a)(1)(ii), we do not make provider-based determinations for ASCs or other suppliers that have active supplier agreements with Medicare because services provided in such entities are paid under other fee schedules, specifically in the case of ASCs regardless of whether the ASC is operated by a hospital.

In the August 1, 2002 IPPS final rule (67 FR 50084 through 50090), we revised the provider-based status rules where the main providers were no longer required to submit an attestation to CMS to demonstrate that their provider-based departments or entities met the provider-based status rules. However, the provider-based department or entity of a main provider must still meet the provider-based status rules in § 413.65 in order for the main provider to bill for services performed in the provider-based department or entity.

2. Change to the Terms of Agreements for ASCs Operated by Hospitals

In the CY 2010 OPPI/ASC proposed rule (74 FR 35392), in order to further streamline our regulations to reduce the administrative burden on providers and suppliers, we proposed to revise existing § 416.30(f)(2) to remove the language requiring a hospital-operated ASC to satisfy CMS that there is good cause for its request to become a provider-based department of a hospital prior to being recognized as such. Specifically, we proposed to remove the language, “without the option of converting to or being paid as a hospital outpatient department, unless CMS determines there is good cause to do otherwise.” We believe that this proposed revision to the requirements that apply to hospital-operated ASCs is consistent with our earlier regulation simplification activities related to the provider-based status rules under § 413.65. We believe that we would reduce the administrative burden on hospitals and ASCs that terminate their supplier agreements with Medicare and bring the requirements into closer alignment with the provider-based status rules for other facilities or organizations that wish to be integrated with the main provider for payment purposes. While an ASC participating in Medicare would continue to be paid only as an ASC, an ASC would also continue to be able to voluntarily terminate its agreements in accordance with § 416.35. Thus, if an ASC chooses to voluntarily terminate its agreement as an ASC and a main provider wants to consider the surgical facility a provider-

based department of that main provider, the facility must meet the provider-based status rules under § 413.65.

We did not receive any public comments on our proposal to revise § 416.30(f)(2) to remove the language, “without the option of converting to or being paid as a hospital outpatient department, unless CMS determines there is good cause to do otherwise.” Therefore, we are adopting as final our proposed revision of § 416.30(f)(2), without modification.

I. Calculation of the ASC Conversion Factor and ASC Payment Rates

1. Background

In the August 2, 2007 final rule (72 FR 42493), we established our policy to base ASC relative payment weights and payment rates under the revised ASC payment system on APC groups and relative payment weights. Consistent with that policy and the requirement at section 1833(i)(2)(D)(ii) of the Act that the revised payment system be implemented so that it would be budget neutral, the initial ASC conversion factor (CY 2008) was calculated so that estimated total Medicare payments under the revised ASC payment system in the first year would be budget neutral to estimated total Medicare payments under the prior (CY 2007) ASC payment system. That is, application of the ASC conversion factor was designed to result in aggregate Medicare expenditures under the revised ASC payment system in CY 2008 equal to aggregate Medicare expenditures that would have occurred in CY 2008 in the absence of the revised system, taking into consideration the cap on ASC payments in CY 2007 as required under section 1833(i)(2)(E) of the Act (72 FR 42521 through 42522).

We note that we consider the term “expenditures” in the context of the budget neutrality requirement under section 1833(i)(2)(D)(ii) of the Act to mean expenditures from the Medicare Part B Trust Fund. We do not consider expenditures to include beneficiary coinsurance and copayments. This distinction was important for the CY 2008 ASC budget neutrality model that considered payments across hospital outpatient, ASC, and MPFS payment systems. However, because coinsurance is almost always 20 percent for ASC services, this interpretation of expenditures has minimal impact for subsequent budget neutrality adjustments calculated within the revised ASC payment system.

In the CY 2008 OPPI/ASC final rule with comment period (72 FR 66857 through 66858), we set out a step-by-step illustration of the final budget

neutrality adjustment calculation based on the methodology finalized in the August 2, 2007 final rule (72 FR 42521 through 42531) and as applied to updated data available for the CY 2008 OPPS/ASC final rule with comment period. The application of that methodology to the data available for the CY 2008 OPPS/ASC final rule with comment period resulted in a budget neutrality adjustment of 0.65.

For CY 2008, we adopted the OPPS relative payment weights as the ASC relative payment weights for most services and, consistent with the final policy, we calculated the CY 2008 ASC payment rates by multiplying the ASC relative payment weights by the final CY 2008 ASC conversion factor of \$41,401. For covered office-based surgical procedures and covered ancillary radiology services, the established policy is to set the relative payment weights so that the national unadjusted ASC payment rate does not exceed the MPFS unadjusted nonfacility PE RVU amount. Further, as discussed in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66841 through 66847), we also adopted alternative ratesetting methodologies for specific types of services (for example, device-intensive procedures).

As discussed in the August 2, 2007 final rule (72 FR 42518) and as codified under § 416.172(c) of the regulations, the revised ASC payment system accounts for geographic wage variation when calculating individual ASC payments by applying the pre-floor and pre-reclassified hospital wage indices to the labor-related share, which is 50 percent of the ASC payment amount. Beginning in CY 2008, CMS accounted for geographic wage variation in labor cost when calculating individual ASC payments by applying the pre-floor and pre-reclassified hospital wage index values that CMS calculates for payment, using updated Core Based Statistical Areas (CBSAs) issued by the Office of Management and Budget in June 2003. The reclassification provision provided at section 1886(d)(10) of the Act is specific to hospitals. We believe the use of the most recent available raw pre-floor and pre-reclassified hospital wage indices results in the most appropriate adjustment to the labor portion of ASC costs. In addition, use of the unadjusted hospital wage data avoids further reductions in certain rural statewide wage index values that result from reclassification. We continue to believe that the unadjusted hospital wage indices, which are updated yearly and are used by many other Medicare payment systems, appropriately account

for geographic variation in labor costs for ASCs.

Comment: Several commenters recommended that CMS adopt for the ASC payment system the same wage index values used for hospital payment under the OPPS. They believed that applying different wage indices in the ASC payment system than are used in the OPPS is inequitable because, in many market areas, ASCs compete directly with hospitals for employees with skills and functions that are applicable in both settings. The commenters believed that, in all but a few instances, the adjusted wage index values used in the OPPS would be higher than the current wage index values used in the ASC payment system. Specifically, the commenters believed the adjustments that are applied to the wage indices used in the OPPS system also should be applied to the ASC wage indices. The adjustments that commenters requested be applied to the wage index values used in the ASC payment system are: an imputed statewide rural wage index for States with no counties outside of an urban area; a mechanism to prevent urban areas from having indices below the statewide rural wage index; a mechanism to prevent the wage index of urban areas that cross state lines from falling below the State-specific rural floor; and an adjustment for counties where a significant proportion of residents commute to other counties for work.

Response: We continue to believe that the unadjusted hospital wage indices, which are updated yearly and are used by almost all Medicare payment systems, appropriately account for geographic variance in labor costs for ASCs. The post-reclassification wage indices for 1886(d) hospitals include many statutory adjustments specific to 1886(d) hospitals and some regulatory adjustments for 1886(d) hospitals including, but not limited to, the areas requested by commenters: an imputed statewide rural wage index for States with no counties outside of an urban area; a "rural floor" mechanism to prevent urban areas from having indices below the statewide rural wage index; a mechanism to prevent the wage index of urban areas that cross state lines from falling below the State-specific rural floor; and an adjustment for counties where a significant proportion of residents commute to other counties. Because many of these adjustments are specified in statute for 1886(d) hospitals, we believe it is appropriate to apply these adjustments to 1886(d) hospitals. The OPPS adopts the post-reclassification wage indices (adjusted

hospital wage indices) because the majority of participating hospitals are section 1886(d) hospitals and, in these hospitals, the exact same personnel staff the ancillary departments of the hospital that simultaneously treat both inpatients and outpatients. For payments systems for other providers and suppliers for which there is no specific statutory provision for adjustments to the wage index values, CMS calculates and employs unadjusted hospital wage indices that reflect the reported cost of hospital labor in each area. Specifically, CMS uses some form of the unadjusted hospital wage indices to pay long-term care, psychiatric, and inpatient rehabilitation hospitals for inpatient care, as well as skilled nursing facilities, hospice programs, home health agencies, and ESRD facilities. CMS historically has only applied the adjusted, post-reclassification hospital wage indices to pay section 1886(d) hospitals for both inpatient and outpatient services for the reasons noted above. It is our policy to treat ASCs as we do all other providers and suppliers using hospital wage index values.

Further, adopting the post-reclassification hospital wage indices with rural floor and associated statewide budget neutrality adjustment would not increase overall ASC payment because we apply a budget neutrality adjustment for changes in the wage indices to the conversion factor. Therefore, any anticipated increases in aggregate ASC payment created by adopting the post-reclassification wage indices would lead to a comparable downward adjustment to the conversion factor to ensure that the only increase in payments to ASCs are those allowed by the update factor. We discuss our budget neutrality adjustment for changes to the wage indices below in section XVI.2.b. of this final rule with comment period.

2. Calculation of the ASC Payment Rates

a. Updating the ASC Relative Payment Weights for CY 2010 and Future Years

We update the ASC relative payment weights each year using the national OPPS relative payment weights (and MPFS nonfacility PE RVU amounts, as applicable) for that same calendar year and uniformly scale the ASC relative payment weights for each update year to make them budget neutral (72 FR 42531 through 42532). In the CY 2010 OPPS/ASC proposed rule (74 FR 35393), consistent with our established policy, we proposed to scale the CY 2010 relative payment weights for ASCs according to the following method. Holding ASC utilization and the mix of

services constant from CY 2008, for CY 2010, we proposed to compare the total payment weight using the CY 2009 ASC relative payment weights under the 50/50 blend (of the CY 2007 payment rate and the ASC payment rate calculated under the ASC standard ratesetting methodology) with the total payment weight using the CY 2010 ASC relative payment weights under the 25/75 blend (of the CY 2007 ASC payment rate and the ASC payment rate calculated under the ASC standard ratesetting methodology) to take into account the changes in the OPPS relative payment weights between CY 2009 and CY 2010. We proposed to use the ratio of CY 2009 to CY 2010 total payment weight (the weight scaler) to scale the ASC relative payment weights for CY 2010. The proposed CY 2010 ASC scaler was 0.9514 and scaling would apply to the ASC relative payment weights of the covered surgical procedures and covered ancillary radiology services for which the ASC payment rates are based on OPPS relative payment weights.

Scaling would not apply in the case of ASC payment for separately payable covered ancillary services that have a predetermined national payment amount (that is, their national ASC payment amounts are not based on OPPS relative payment weights), such as drugs and biologicals that are separately paid or services that are contractor-priced or paid at reasonable cost in ASCs. Any service with a predetermined national payment amount would be included in the ASC budget neutrality comparison, but scaling of the ASC relative payment weights would not apply to those services. The ASC payment weights for those services without predetermined national payment amounts (that is, those services with national payment amounts that would be based on OPPS relative payment weights if a payment limitation did not apply) would be scaled to eliminate any difference in the total payment weight between the current year and the update year.

The proposed weight scaler used to model CY 2010 ASC fully implemented payment rates in order to reflect our estimated of rates if there was no transition was equal to 0.9329. We applied this scaler to the payment weights subject to scaling, in order to estimate the ASC payment rates for CY 2010 without the transition, for purposes of the ASC impact analysis discussed in section XXI.C. of the CY 2010 OPPS/ASC proposed rule (74 FR 35418).

For any given year's ratesetting, we typically use the most recent full calendar year of claims data to model

budget neutrality adjustments. When we developed the CY 2010 OPPS/ASC proposed rule, we had available 98 percent of CY 2008 ASC claims data. In the CY 2010 OPPS/ASC proposed rule (74 FR 35393), we reported that we had 95 percent of the CY 2008 ASC claims data available to model proposed revisions to the CY 2010 ASC payment system, but we have since confirmed that we had a slightly higher percentage available at that time. For this final rule with comment period, we have close to 100 percent of all claims for CY 2008. CY 2010 is the first year that the claims data used for ratesetting include new covered surgical procedures and covered ancillary services under the revised ASC payment system. Because we had almost all of the CY 2008 claims data available when we calculated the conversion factor and budget neutrality adjustments for our proposed rule, for the final CY 2010 budget neutrality adjustments, we did not expect there would be significant changes in our calculated budget neutrality adjustments (the weight scaler or wage adjustment) that could be attributable to more utilization available from additional claims data for this CY 2010 OPPS/ASC final rule with comment period.

To create an analytic file to support calculation of the weight scaler and budget neutrality adjustment for the wage index (discussed below), we summarized available CY 2008 ASC claims by provider and by HCPCS code. We created a unique supplier identifier solely for the purpose of identifying unique ASCs within the CY 2008 claims data. We used the supplier zip code reported on the claim to associate State, county, and CBSA with each ASC. This file, available to the public as a supporting data file for the CY 2010 OPPS/ASC proposed rule, is posted on the CMS Web site at: http://www.cms.hhs.gov/ASCPayment/01_Overview.asp#TopOfPage.

Comment: Many commenters again expressed their opposition to scaling the ASC relative payment weights. Many of the commenters on the CY 2010 proposed rule offered the same views as the public commenters on the CY 2009 OPPS/ASC final rule with comment period, the year when CMS first applied the scaling policy that was finalized in the August 2, 2007 final rule. The commenters expressed many concerns, including that scaling is contrary to the intent of using the cost-based OPPS relative payment weights as the bases for determining the relative payments for the same services in ASCs and that scaling would continue to erode the payment relationship between the OPPS

and ASC payment system. Further, the commenters stated that increasing the difference between ASC and OPPS payments is in direct conflict with the goal of ensuring that patients have continued access to surgical care in the lowest priced setting appropriate to their clinical needs. They asserted that, although scaling is intended to maintain budget neutrality within the ASC payment system, it is instead creating increasingly large payment differentials between the ASC and OPPS payments for the same services, without evidence of growing differences in capital and operating costs between the two settings.

The commenters argued that CMS is not required to scale the ASC relative weights and that it should use its authority to suspend the application of scaling the ASC relative weights for CY 2010. They noted that CMS established at § 416.171(e)(2) of the regulations a process by which it *may* (emphasis added) make annual adjustment to the relative payment weights, *as needed* (emphasis added).

The commenters also expressed their continuing disagreement with aspects of the budget neutrality adjustment methodology used by CMS to establish the conversion factor. They provided the results of their comparison of actual volume and payment for services that were new to the ASC list in CY 2008. Based upon the results of their analyses of CY 2008 claims data, the commenters concluded that the migration estimates used by CMS to establish budget neutrality in CY 2008 were several times higher than the actual ASC spending for newly covered procedures and, therefore, that the resulting CY 2008 conversion factor was too low. They believed that these findings provide a further basis for CMS not to scale the ASC relative payment weights for CY 2010 after the weights are scaled under the OPPS.

In addition, many of the commenters reasoned that because the ASC payment system is based on the OPPS relative weights, the weights should be equal in both settings and because the weights are scaled to ensure budget neutrality under the OPPS, the weights should not be scaled again to ensure budget neutrality under the ASC system. The commenters believed that the CY 2010 OPPS relative payment weights reflected real growth in the relative costs of surgical services provided in HOPDs and that the ASC scaler should not reclaim dollars from the ASC payment system because there also has been real cost growth for the surgical services provided in ASCs. However, they acknowledged that suspending

application of the scaler for CY 2010 would result in an aggregate increase in ASC spending in that year.

The commenters expressed concern that other payment adjustments are depressing the ASC payments for many procedures, including the freeze on the ASC payment update through CY 2009 and the transition policy and that scaling further reduces rates to inappropriately low levels. Further, the commenters argued that scaling is forcing procedures for which the OPPS median cost increased from CY 2009 to CY 2010 to finance the transitional payment policies, and that the procedures the transition was intended to aid are the procedures financing the bulk of the scaler.

Response: Many of these comments are similar to public comments on the proposal for the revised ASC payment system that we responded to in the August 2, 2007 final rule (72 FR 42531 through 42533). For example, with regard to scaling, we addressed these same concerns raised by commenters “that annual rescaling would cause divergence of the relative weights between the OPPS and the revised ASC payment system for individual procedures” in the August 2, 2007 final rule (72 FR 42532). We refer the commenters to that discussion for our detailed response in promulgating the scaling policy that was initially applied in CY 2009 (72 FR 42531 through 42533). Below, we address new issues raised by the commenters and provide a general summary of some of the relevant responses from the August 2, 2007 final rule and the CY 2009 OPPS/ASC final rule with comment period (73 FR 68754 through 68755).

The ASC weight scaling methodology is entirely consistent with the OPPS methodology for scaling the relative payment weights and, for the most part, the increasing payment differentials between the ASC and OPPS payments for the same services are not attributable to scaling ASC relative payment weights. Considerations of differences between the capital and operating costs of ASCs and HOPDs are not part of the ASC standard ratesetting methodology, which relies only on maintaining the same relativity of payments for services under the two payment systems, as well as budget neutrality within each payment system. Furthermore, unlike HOPDs, we do not have information about the costs of ASC services in order to assess differences in capital and operating costs over time between the two settings. In order to maintain budget neutrality of the ASC payment system, we need to adjust for the effects of changes in relative weights. The ASC

payment system adopts the OPPS relative weights as the mechanism for apportioning total payments, after application of the update factor, among all of the services covered by the ASC payment system. The OPPS relative weights serve the same purpose in the OPPS. The OPPS relative weights do not represent an estimate of absolute cost of any given procedure; rather, they reflect our estimate of the cost of the procedure within the context of our cost estimation methodology for the OPPS. With the exception of services with a predetermined national payment amount, the use of a uniform scaling factor for changes in total weight between years in the ASC payment system does not alter the relativity of the OPPS payment weights as used in the ASC payment system. Differences in the relativity between the ASC relative payment weights and the OPPS relative payment weights are not driven by the application of the uniform scaling factor. The ASC weight scaling methodology is entirely consistent with the OPPS weight scaling methodology and the weights serve the same purpose in both systems, to apportion total budget neutral payment allowed under the update.

We do not believe that the application of the scaler will lead to beneficiary access problems. We believe that the fully implemented relative weights will be representative of relative costs across all ASC services and that payments will support the continued provision of high quality surgical procedures to Medicare beneficiaries in the most appropriate settings. We also expect that, over time, ASCs will provide an increased breadth of services. Appropriate beneficiary access to services in appropriate care settings is always an important concern and we will continue to monitor access under the revised ASC payment system.

As stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68754), with respect to the use of “as needed” in the text of § 416.171(e)(2) that commenters have interpreted to mean that CMS has the authority to suspend scaling the relative payment weights if it determines there is not a need to do so, the phrase does not mean that CMS will determine whether or not to adjust for budget neutrality. Rather, it means that CMS adjusts the relative payment weights as needed to ensure budget neutrality and, as acknowledged by the commenters on the CY 2010 OPPS/ASC proposed rule, if we were not to scale the ASC relative payment weights, we estimate that the CY 2010 revisions would not be budget neutral.

We agree that there are differences between the service volume estimates

CMS used to establish budget neutrality based on CY 2006 claims data and those reflected in the CY 2008 claims data. In the final regulations implementing the revised ASC payment system, we made our best actuarial estimate to ensure budget neutrality. We did not intend to revisit the actuarial budget neutrality regardless of whether or not it could be determined that there was a difference between actual experience and our underlying data assumptions and regardless of whether or not any difference that could be determined resulted in increased or decreased expenditures under the revised ASC payment system.

Establishing budget neutrality under the OPPS does not result in budget neutrality under the revised ASC payment system; it is only to maintain budget neutrality under the OPPS. Scaling the ASC relative payment weights is an integral and separate process for maintaining budget neutrality under the ASC prospective payment system. Scaling is the budget neutrality adjustment that ensures that changes in the relative weights do not, in and of themselves, change aggregate payment to ASCs. It ensures a specific amount of payment for ASCs in any given year. Without scaling, total ASC payment could increase or decrease relative to changes in hospital outpatient payment.

Although the commenters believed that scaling prevents increases in ASC spending that may be appropriate because ASC costs have increased over time, increases in cost in a prospective payment system are handled by the update factor. In a budget neutral system, we remove the independent effects of increases or decreases in payments as a result of changes in the relative payment weights or the wage indices and constrain increases to the allowed update factor. Therefore, changes in aggregate ASC expenditures related to payment rates should be determined by the update to the ASC conversion factor, the CPI-U.

Regarding commenters' concern that other payment adjustments, including the freeze on ASC payment updates and the transitional payment policy, are depressing the ASC payments for many procedures and that scaling has a disproportionate effect on some covered surgical procedures, we note that the statute set a zero percent update for CY 2008 and CY 2009. We implemented the 4-year transitional payment policy in response to public comments that persuaded us that ASCs would benefit from more gradual implementation of the revised ASC payment rates, especially for historically high volume

procedures because the prior rates for those procedures were disproportionately high compared to the prior rates for other ASC procedures. As explained in the August 2, 2007 final rule (72 FR 42542), a major effect of the revised ASC payment system is redistribution of payments across all ASC procedures. Historically, the highest volume ASC procedures had payment rates that were close to the payments in HOPDs and, as such, accounted for most of the total Medicare payments to ASCs. As a result, payments for many of those high volume services are the most adversely affected under the revised payment system as the relative weights across all ASC procedures become more closely aligned with those of the OPPS. We appreciate the commenters' concern that scaling is forcing procedures for which the OPPS median cost increased from CY 2009 to CY 2010 to finance the transitional payment policies, and that the procedures the transition was intended to aid are the procedures financing the bulk of the scaler. However, as already noted, the ASC payment system adopts the relativity of the OPPS weights, not the actual median costs or payments for OPPS services. It is fully consistent that a budget neutrality adjustment for differences in aggregate payment weight, specifically scaling, would change the amount of payment under the ASC payment system relative to the OPPS median cost and to the previous year's payment under the ASC payment system for the same service. It is critical that the amount of payment allowed under the ASC payment system, after application of the update factor, distributes the appropriate proportional payment amount to each service. The same statement is true for commenters' concerns that scaling is reducing payment for services explicitly designated as receiving a transition payment. Scaling ensures that the changes in the relative weights do not, in and of themselves, change aggregate payment to ASCs. The calculation of the transition weight over a fully implemented weight for any procedures paid in CY 2007 under the previous ASC payment system changes the relativity of the weight of those services relative to other services newly covered by the revised ASC payment system. This clearly changes the proportional resources distributed to services subject to the transition compared to what would be distributed under a fully implemented system. However, entitlement to a transition weight under a budget neutral system does not

guarantee a specific amount of payment in absolute dollar terms. A service that experienced an increase in the OPPS relative weight may very well experience a decline in payment relative to the previous year's actual payment rate because the scaling necessary to maintain equal weight in the system is greater than the proportional increase in the OPPS relative weight portion of the transition weight. Again, this outcome is fully consistent with implementation of a budget neutral prospective payment system with a specific update factor.

For this final rule with comment period, we used our proposed methodology described above to calculate the scaler adjustment using updated ASC claims data. The final CY 2010 scaler adjustment for the third year of the transition is 0.9567. This scaler adjustment is necessary to budget neutralize the difference in aggregate ASC payments calculated using the CY 2009 ASC transitional (50/50 blend) relative payment weights and the CY 2010 ASC transitional (75/25 blend) relative payment weights. We calculated the difference in aggregate payments due to the change in relative payment weights (including drugs and biologicals) holding constant the ASC conversion factor, the most recent CY 2008 ASC utilization from our claims data, and the CY 2009 wage index values. For this final CY 2010 calculation, we used the CY 2009 ASC conversion factor updated by the CY 2010 CPI-U, which is 1.2 percent.

After consideration of the public comments we received, we are finalizing our CY 2010 ASC relative payment weight scaling methodology, without modification. The final CY 2010 ASC payment weight scaler is 0.9567.

b. Updating the ASC Conversion Factor

Under the OPPS, we typically apply a budget neutrality adjustment for provider-level changes, most notably a change in the wage index values for the upcoming year, to the conversion factor. In the CY 2010 OPPS/ASC proposed rule (74 FR 35393), consistent with our final ASC payment policy, for the CY 2010 ASC payment system, we proposed to calculate and apply the pre-floor and pre-reclassified hospital wage indices that are used for ASC payment adjustment to the ASC conversion factor, just as the OPPS wage index adjustment is calculated and applied to the OPPS conversion factor (73 FR 41539). For CY 2010, we calculated this proposed adjustment for the ASC payment system by using the most recent CY 2008 claims data available and estimating the difference in total payment that would be created by

introducing the CY 2010 pre-floor and pre-reclassified hospital wage indices. Specifically, holding CY 2008 ASC utilization and service-mix and CY 2010 national payment rates after application of the weight scaler constant, we calculated the total adjusted payment using the CY 2009 pre-floor and pre-reclassified hospital wage indices and the total adjusted payment using the proposed CY 2010 pre-floor and pre-reclassified hospital wage indices. We used the 50-percent labor-related share for both total adjusted payment calculations. We then compared the total adjusted payment calculated with the CY 2009 pre-floor and pre-reclassified hospital wage indices to the total adjusted payment calculated with the proposed CY 2010 pre-floor and pre-reclassified hospital wage indices and applied the resulting ratio of 0.9996 (the proposed CY 2010 ASC wage index budget neutrality adjustment) to the CY 2009 ASC conversion factor to calculate the proposed CY 2010 ASC conversion factor.

Section 1833(i)(2)(C) of the Act requires that, if the Secretary has not updated the ASC payment amounts in a calendar year, the payment amounts shall be increased by the percentage increase in the CPI-U as estimated by the Secretary for the 12-month period ending with the midpoint of the year involved. However, section 1833(i)(2)(C)(iv) of the Act required that the increase of ASC payment amounts for CYs 2008 and 2009 equal zero percent. As discussed in the August 2, 2007 final rule, we adopted a final policy to update the ASC conversion factor using the CPI-U in order to adjust ASC payment rates for CY 2010 and subsequent years (72 FR 42518 through 42519 and § 416.171(a)(2)). In the CY 2010 OPPS/ASC proposed rule (74 FR 35394), we proposed to implement the annual updates through an adjustment to the ASC conversion factor beginning in CY 2010 when the statutory requirement for a zero update no longer applies.

For our proposed rule, for the 12-month period ending with the midpoint of CY 2010, the Secretary estimated that the CPI-U is 0.6 percent. Therefore, we proposed to apply to the ASC conversion factor a 0.6 percent increase for CY 2010.

Thus, for CY 2010, we proposed to adjust the CY 2009 ASC conversion factor (\$41.393) by the wage adjustment for budget neutrality of 0.9996 and the update of 0.6 percent, which resulted in a proposed CY 2010 ASC conversion factor of \$41.625.

Comment: Many commenters requested that CMS adopt the hospital

market basket to update the ASC payment system. They explained that not only is the CPI-U lower than the hospital market basket but it is not appropriate for updating health care providers because, unlike the hospital market basket which analyzes hospital spending, the CPI-U is designed to capture household spending. The commenters stated that in the most recent years, the CPI-U has been dominated by energy and housing costs rather than healthcare provider spending. Further, the commenters stated that CMS' use of the midyear CPI-U percent change is problematic because other federal agencies, such as the Bureau of Labor Statistics and the Congressional Budget Office, use an end-of-year timeframe. They believed that a negative consequence of the midyear timing for CMS' forecasted CPI-U percent change is that the CPI-U used to update the ASC payment system cannot be validated directly with an independent source.

The commenters argued that the difference between the ASC and OPSS conversion factors is not due to real differences in the growth of costs of goods and services furnished by ASCs and HOPDs and should not be perpetuated. The commenters asserted that CMS clearly has the authority to use an alternative update mechanism, and believed CMS should adopt a more appropriate update for the ASC payment system to prevent further increases in differential between the ASC and OPSS conversion factors.

Response: We understand the commenters' concerns regarding the update to the conversion factor for CY 2010, but note that we did not propose to change the conversion factor update methodology. We refer readers to the discussion in the August 2, 2007 final rule on this issue (72 FR 42518 through 42519).

After consideration of the public comments we received, we are applying our established methodology for determining the final CY 2010 ASC conversion factor. Using more complete CY 2008 data for this final rule with comment period than was available for the proposed rule, we calculated a wage index budget neutrality adjustment of 0.9996 and the updated CPI-U projected for the midpoint of CY 2010 is 1.2 percent. The final ASC conversion factor of \$41.873 is the product of the CY 2009 conversion factor of \$41.393 multiplied by 0.9996 and the 1.2 percent CPI-U.

3. Display of ASC Payment Rates

Addenda AA and BB to this CY 2010 OPSS/ASC final rule with comment

period display the updated ASC payment rates for CY 2010 for covered surgical procedures and covered ancillary services, respectively. These addenda contain several types of information related to the CY 2010 payment rates. Specifically, in Addendum AA, a "Y" in the column titled "Subject to Multiple Procedure Discounting" indicates that the surgical procedure would be subject to the multiple procedure payment reduction policy. As discussed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66829 through 66830), most covered surgical procedures are subject to a 50-percent reduction in the ASC payment for the lower-paying procedure when more than one procedure is performed in a single operative session. Display of the comment indicator "CH" in the column titled "Comment Indicator" indicates a final change in payment policy for the item or service, including identifying discontinued HCPCS codes, designating items or services newly payable under the ASC payment system, and identifying items or services with changes in the ASC payment indicator for CY 2010. Display of the comment indicator "NI" in the column titled "Comment Indicator" indicates that the code is new (or substantially revised) and that the payment indicator assignment is an interim assignment that is open to comment on this final rule with comment period.

The values displayed in the column titled "CY 2010 Third Year Transition Payment Weight" are the relative payment weights for each of the listed services for CY 2010, the third year of the 4-year transition period. The CY 2010 ASC payment rates for the covered surgical procedures subject to transitional payment (payment indicators "A2" and "H8" in Addendum AA) are based on a blend of 25 percent of the CY 2007 ASC payment rate for the procedure and 75 percent of the CY 2010 ASC rate calculated under the ASC standard ratesetting methodology before scaling for budget neutrality. The payment weights for all covered surgical procedures and covered ancillary services whose ASC payment rates are based on OPSS relative payment weights are scaled for budget neutrality. Thus, scaling was not applied to the device portion of the device-intensive procedures, services that are paid at the MPFS nonfacility PE RVU amount, separately payable covered ancillary services that have a predetermined national payment amount, such as drugs and biologicals that are separately paid under the OPSS,

or services that are contractor-priced or paid at reasonable cost in ASCs.

To derive the CY 2010 national unadjusted payment rate displayed in the "CY 2010 Third Year Transition Payment" column, each ASC payment weight in the "CY 2010 Third Year Transition Payment Weight" column is multiplied by the final CY 2010 ASC conversion factor of \$41.873. The conversion factor includes a budget neutrality adjustment for changes in the wage index values and the CPI-U percentage increase.

In Addendum BB, there are no relative payment weights displayed in the "CY 2010 Third Year Transition Payment Weight" column for items and services with predetermined national payment amounts, such as separately payable drugs and biologicals. The "CY 2010 Third Year Transition Payment" column displays the final CY 2010 national unadjusted ASC payment rates for all items and services. The CY 2010 ASC payment rates listed in the Addendum AA for separately payable drugs and biologicals are based on ASP data used for payment in physicians' offices in October 2009.

For informational purposes only, we also have posted on the CMS Web site the fully transitioned ASC payment rates for CY 2010. These rates do not represent what the payment rates would be once the transition is over, only what the CY 2010 rates would be if there were no transition. The Web site address is: <https://www.cms.hhs.gov/ASCPayment/>.

We did not receive any public comments regarding the continuation of our policy to provide CY 2010 ASC payment information as detailed in Addenda AA and BB. Therefore, Addenda AA and BB to this final rule with comment period display the updated ASC payment rates for CY 2010 for covered surgical procedures and covered ancillary services, respectively, and provide additional information related to the CY 2010 rates.

XVI. Reporting Quality Data for Annual Payment Rate Updates

A. Background

1. Overview

CMS has implemented quality measure reporting programs for multiple settings of care. These programs promote higher quality, more efficient health care for Medicare beneficiaries. The quality data reporting program for hospital outpatient care, known as the Hospital Outpatient Quality Data Reporting Program (HOP QDRP), has been generally modeled after the program for hospital inpatient services,

the Reporting Hospital Quality Data for Annual Payment Update (RHQDAPU) program. Both of these quality reporting programs for hospital services, as well as the program for physicians and other eligible professionals, known as the Physician Quality Reporting Initiative (PQRI), have financial incentives for reporting of quality data to CMS. CMS has also implemented quality reporting programs for home health agencies and skilled nursing facilities that are based on conditions of participation, and an end-stage renal disease quality reporting program that is based on conditions for coverage.

2. Hospital Outpatient Quality Data Reporting Under Section 109(a) of Pub. L. 109–432

Section 109(a) of the MIEA–TRHCA (Pub. L. 109–432) amended section 1833(t) of the Act by adding a new subsection (17) that affects the payment rate update applicable to OPSS payments for services furnished by hospitals in outpatient settings on or after January 1, 2009. Section 1833(t)(17)(A) of the Act, which applies to hospitals as defined under section 1886(d)(1)(B) of the Act, states that subsection (d) hospitals that fail to report data required for the quality measures selected by the Secretary in the form and manner required by the Secretary under section 1833(t)(17)(B) of the Act will receive a 2.0 percentage point reduction to their annual payment update factor. Section 1833(t)(17)(B) of the Act requires that hospitals submit quality data in a form and manner, and at a time, that the Secretary specifies. Section 1833(t)(17)(C)(i) of the Act requires the Secretary to develop measures appropriate for the measurement of the quality of care (including medication errors) furnished by hospitals in outpatient settings, that these measures reflect consensus among affected parties and, to the extent feasible and practicable, that these measures include measures set forth by one or more national consensus building entities.

The National Quality Forum (NQF) is a voluntary consensus standard-setting organization that is composed of a diverse representation of consumer, purchaser, provider, academic, clinical, and other health care stakeholder organizations. NQF was established to standardize health care quality measurement and reporting through its consensus development process. We generally prefer to adopt NQF-endorsed measures for CMS quality reporting programs. However, we believe that consensus among affected parties also can be reflected by other means,

including: consensus achieved during the measure development process; consensus shown through broad acceptance and use of measures; and consensus through public comment. We also note that section 1833(t)(17) of the Act does not require that each measure we adopt for the HOP QDRP be endorsed by a national consensus building entity, or by the NQF specifically.

Section 1833(t)(17)(C)(ii) of the Act authorizes the Secretary to select measures for the HOP QDRP that are the same as (or a subset of) the measures for which data are required to be submitted under section 1886(b)(3)(B)(viii) of the Act (the RHQDAPU program). Section 1833(t)(17)(D) of the Act gives the Secretary the authority to replace measures or indicators as appropriate, such as when all hospitals are effectively in compliance or when the measures or indicators have been subsequently shown not to represent the best clinical practice. Section 1833(t)(17)(E) of the Act requires the Secretary to establish procedures for making data submitted under the HOP QDRP available to the public. Such procedures must include giving hospitals the opportunity to review their data before these data are released to the public.

As we stated in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68758 through 68759), we continue to believe that it is most appropriate and desirable to adopt measures that specifically apply to the hospital outpatient setting for the HOP QDRP. In other words, we do not believe that we should simply, without further analysis, adopt the RHQDAPU program measures as the measures for the HOP QDRP. Nonetheless, we note that section 1833(t)(17)(C)(ii) of the Act allows the Secretary to “[select] measures that are the same as (or a subset of) the measures for which data are required to be submitted” under the RHQDAPU program.

3. Reporting ASC Quality Data for Annual Payment Update

Section 109(b) of the MIEA–TRHCA amended section 1833(i) of the Act by redesignating clause (iv) as clause (v) and adding new clause (iv) to paragraph (2)(D) and adding paragraph (7). These amendments may affect ASC payments for services furnished in ASC settings on or after January 1, 2009. Section 1833(i)(2)(D)(iv) of the Act authorizes the Secretary to implement the revised payment system for services furnished in ASCs (established under section 1833(i)(2)(D) of the Act), “so as to provide for a reduction in any annual

update for failure to report on quality measures.”

Section 1833(i)(7)(A) of the Act states that the Secretary may provide that any ASC that fails to report data required for the quality measures selected by the Secretary in the form and manner required by the Secretary under section 1833(i)(7) of the Act will incur a reduction in any annual payment update of 2.0 percentage points. Section 1833(i)(7)(A) of the Act also specifies that a reduction for one year cannot be taken into account in computing the ASC update for a subsequent calendar year.

Section 1833(i)(7)(B) of the Act provides that, “[e]xcept as the Secretary may otherwise provide,” the hospital outpatient quality data provisions of sections 1833(t)(17)(B) through (E) of the Act, summarized above, shall apply to ASCs. We did not implement an ASC quality reporting program for CY 2008 (72 FR 66875) or for CY 2009 (73 FR 68779).

We refer readers to section XVI.H. of this final rule with comment period for a discussion of our decision to implement ASC quality data reporting in a later rulemaking.

4. HOP QDRP Quality Measures for the CY 2009 Payment Determination

For the CY 2009 annual payment update, we required HOP QDRP reporting using seven quality measures—five Emergency Department (ED) AMI measures and two Perioperative Care measures. These measures address care provided to a large number of adult patients in hospital outpatient settings, across a diverse set of conditions, and were selected for the initial set of HOP QDRP measures based on their relevance as a set to all HOPDs.

Specifically, in order for hospitals to receive the full OPSS payment update for services furnished in CY 2009, in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66860), we required that subsection (d) hospitals paid under the OPSS submit data on the following seven measures for hospital outpatient services furnished on or after April 1, 2008: (1) ED–AMI–1: Aspirin at Arrival; (2) ED–AMI–2: Median Time to Fibrinolysis; (3) ED–AMI–3: Fibrinolytic Therapy Received within 30 Minutes of Arrival; (4) ED–AMI–4: Median Time to Electrocardiogram (ECG); (5) ED–AMI–5: Median Time to Transfer for Primary PCI; (6) PQRI #20: Perioperative Care—Timing of Antibiotic Prophylaxis; and (7) PQRI #21: Perioperative Care—Selection of Perioperative Antibiotic.

5. HOP QDRP Quality Measures for the CY 2010 Payment Determination

a. Background

In the CY 2009 OPPTS/ASC final rule with comment period, for the CY 2010 payment update, we required continued submission of data on the existing seven measures discussed above (73 FR 68761), and adopted four imaging measures (73 FR 68766). For CY 2010, we changed the measure designations for the existing seven measures, including a change to an “OP-X” format in order to maintain a consistent sequential designation system that we could expand as we add additional measures.

The four imaging measures that we adopted beginning with the CY 2010

payment determination (OP-8: MRI Lumbar Spine for Low Back Pain, OP-9: Mammography Follow-up Rates, OP-10: Abdomen CT—Use of Contrast Material, and OP-11: Thorax CT—Use of Contrast Material) are claims-based measures that CMS will calculate using Medicare Part B claims data without imposing upon hospitals the burden of additional chart abstraction. For purposes of the CY 2010 payment determination, we will calculate these measures using CY 2008 Medicare administrative claims data.

In the CY 2009 OPPTS/ASC proposed rule, OP-10 had two submeasures listed: OP-10a: CT Abdomen—Use of contrast material excluding calculi of the kidneys, ureter, and/or urinary tract, and OP-10b: CT Abdomen—Use of

contrast material for diagnosis of calculi in the kidneys, ureter, and or urinary tract. In the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68766), we finalized OP-10: Abdomen CT—Use of Contrast Material. To clarify, we are calculating OP-10 excluding patients with renal disease. This exclusion is described in greater detail in the *Specifications Manual for Hospital Outpatient Department Quality Measures* (HOPD Specifications Manual) located at the QualityNet Web site (<http://www.QualityNet.org>).

The complete set of measures to be used for the CY 2010 payment determination is set out below, and is shown with the CY 2010 measure designations as well as their ED-AMI and PQRI designations:

HOP QDRP measurement set to be used for CY 2010 payment determination	CY 2009 designation
OP-1: Median Time to Fibrinolysis	ED-AMI-2.
OP-2: Fibrinolytic Therapy Received Within 30 Minutes	ED-AMI-3.
OP-3: Median Time to Transfer to Another Facility for Acute Coronary Intervention	ED-AMI-5.
OP-4: Aspirin at Arrival	ED-AMI-1.
OP-5: Median Time to ECG	ED-AMI-4.
OP-6: Timing of Antibiotic Prophylaxis	PQRI #20.
OP-7: Prophylactic Antibiotic Selection for Surgical Patients	PQRI #21.
OP-8: MRI Lumbar Spine for Low Back Pain	NA.
OP-9: Mammography Follow-up Rates	NA.
OP-10: Abdomen CT—Use of Contrast Material	NA.
OP-11: Thorax CT—Use of Contrast Material	NA.

b. Maintenance of Technical Specifications for Quality Measures

Technical specifications for each HOP QDRP measure are listed in the HOPD Specifications Manual, which is posted on the CMS QualityNet Web site at <http://www.QualityNet.org>. We maintain the technical specifications for the measures by updating this HOPD Specifications Manual and include detailed instructions and calculation algorithms for hospitals to use when collecting and submitting data on required measures.

In the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68766), we established a subregulatory process for updates to the technical specifications that we use to calculate HOP QDRP measures. This process is used when changes to the measure specifications are necessary due to changes in scientific evidence or in the measure as endorsed by the consensus entity. Changes of this nature may not coincide with the timing of our regulatory actions, but nevertheless require inclusion in the measure specifications so that the HOP QDRP measures are calculated based on the most up-to-date scientific and consensus standards. We indicated that notification of changes to the measure specifications on the

QualityNet Web site, <http://www.QualityNet.org>, and in the HOPD Specifications Manual that occurred as a result of changes in scientific evidence or national consensus would occur no less than 3 months before any changes become effective for purposes of reporting under the HOP QDRP.

The HOPD Specifications Manual is released every 6 months and addenda are released as necessary, providing at least 3 months of advance notice for nonsubstantive changes such as changes to ICD-9, CPT, NUBC, and HCPCS codes, and at least 6 months notice for substantive changes to data elements that would require significant systems changes.

Comment: A few commenters indicated that they agreed with the maintenance of the outpatient measure technical specifications in a manner consistent with the inpatient measure technical specifications. They agreed that providing a 3-month notification period for code updates is sufficient. One commenter also agreed the OP-X designations along with short measure names are appropriate. Commenters indicated that CMS should ensure that the subregulatory process that it uses to update the technical specifications for

HOP QDRP measures is regular and transparent.

Response: We thank the commenters for their support of the subregulatory manual update process and timeframes. We will continue to make such updates on a regular semi-annual basis with addenda as necessary, and to issue notifications of updates via the QualityNet Web site, <http://www.QualityNet.org>, in order to maintain the transparency of the process. The HOPD Specifications Manual will continue to be released regularly and addenda will continue to be issued as necessary, providing at least 3 months of advance notice for nonsubstantive changes such as changes to ICD-9, CPT, NUBC, and HCPCS codes, and at least 6 months notice for substantive changes to data elements that would require significant systems changes.

c. Publication of HOP QDRP Data

Section 1833(t)(17)(E) of the Act requires that the Secretary establish procedures to make data collected under the HOP QDRP program available to the public. CMS also requires hospitals to complete and submit a registration form (“participation form”) in order to participate in the HOP QDRP. With submission of this form, participating

hospitals agree that they will allow CMS to publicly report the quality measures, including those that CMS calculates using Medicare claims, as required by the Act and the HOP QDRP.

In the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68778), we established that for CY 2010, hospitals sharing the same CMS Certification Number (CCN, previously known as the Medicare Provider Number (MPN)) must combine data collection and submission across their multiple campuses for the clinical measures for public reporting purposes. We finalized the policy that, under the HOP QDRP, we will publish quality data by the corresponding CCN. This approach is consistent with the approach taken under the RHQDAPU program. In that final rule with comment period, we also stated that we intend to indicate instances where data from two or more hospitals are combined to form the publicly reported measures on the Web site.

We discuss our CY 2010 policy regarding publication of HOP QDRP data in section XVI.F. of this final rule with comment period.

B. Quality Measures for the CY 2011 Payment Determination

1. Considerations in Expanding and Updating Quality Measures Under the HOP QDRP

In general, when selecting measures for the HOP QDRP program, we take into account several considerations and goals. These include: (a) Expanding the types of measures beyond process of care measures to include an increased number of outcome measures, efficiency measures, and patients' experience-of-care measures; (b) expanding the scope of hospital services to which the measures apply; (c) considering the burden on hospitals in collecting chart-abstracted data; (d) harmonizing the measures used in the HOP QDRP program with other CMS quality programs to align incentives and promote coordinated efforts to improve quality; (e) seeking to use measures based on alternative sources of data that do not require chart abstraction or that utilize data already being reported by many hospitals, such as data that hospitals report to clinical data registries, or all-payer claims data bases; and (f) weighing the relevance and utility of the measures compared to the burden on hospitals in submitting data under the HOP QDRP program. Specifically, we give priority to quality measures that assess performance on: (a) Conditions that result in the greatest mortality and morbidity in the Medicare population; (b) conditions that are high

volume and high cost for the Medicare program; and (c) conditions for which wide cost and treatment variations have been reported, despite established clinical guidelines. We have used and continue to use these criteria to guide our decisions regarding what measures to add to the HOP QDRP measure set.

Comment: Many commenters indicated that, although CMS is not required to adopt only measures that are endorsed by NQF, CMS should continue to rely on NQF evaluations to guide selection of measures, and to seek NQF approval for measures considered and adopted for the HOP QDRP in order to maintain consistency in the selection processes for quality measures across physician and hospital services. Many commenters indicated that they prefer that measures adopted for HOP QDRP first go through the rigorous, consensus-based assessment processes of both the NQF and HQA, and that given the number of NQF-endorsed and HQA-adopted measures currently available for use, it is both feasible and practicable for CMS to choose only NQF-endorsed and HQA-adopted measures. Other commenters indicated that although a consensus-based process may have been employed by CMS or CMS contractors to develop measures, it does not equal the rigor or broad stakeholder input of NQF endorsement and HQA adoption.

Response: Section 1833(t)(17)(C)(i) of the Act requires the Secretary to "develop measures that the Secretary determines to be appropriate for the measurement of the quality of care (including medication errors) furnished by hospitals in outpatient settings and that reflect consensus among affected parties and, to the extent feasible and practicable, shall include measures set forth by one or more national consensus building entities." This provision does not require that the measures we adopt for the HOP QDRP be endorsed by any particular entity, and we believe that consensus among affected parties can be reflected by means other than endorsement by a national consensus building entity, including consensus achieved during the measure development process, consensus shown through broad acceptance and use of measures, and consensus through public comment. Nevertheless, we have stated on numerous occasions that we prefer to adopt quality measures that have been endorsed by the NQF because the NQF uses a formal consensus development process and has been recognized as a voluntary consensus standards-setting organization as defined by the National Technology Transfer and Advancement Act of 1995 (NTTAA) and Office of

Management and Budget Circular A 119 (see http://www.qualityforum.org/Measuring_Performance/Consensus_Development_Process.aspx). We are unaware of any other organizations that qualify as an NTTAA consensus organization for the endorsement of quality measures. However, when we propose and adopt quality measures, we take into consideration the measures adopted by the HQA as well as an array of input from the public. We appreciate HQA's integral efforts to improve hospital quality of care by supporting CMS' public reporting programs.

Comment: Some commenters expressed concern regarding the accuracy of measures that rely solely on administrative (that is, claims) data and requested that CMS not consider these types of measures in the future. Several commenters questioned the value of measures based solely on claims data/administrative data for public reporting and pay-for-performance in terms of their capacity to improve care delivered to Medicare beneficiaries.

Response: We do not agree with these commenters' statements. We believe that claims data/administrative data are an appropriate data source upon which quality measures selected by the Secretary may be based. We note that many NQF-endorsed evidence-based quality measures that have been found appropriate for public reporting and quality improvement rely upon claims and administrative data as a data source. Furthermore, the use of claims-based measures reduces reliance upon chart abstraction and its associated burden for quality measurement.

Comment: Commenters submitted the following suggested measure selection criteria for the HOP QDRP:

- Potential for quality improvement;
- Processes measured are related to improved patient outcomes;
- Processes measured occur closer in time to patient outcomes of interest;
- Outcome measures are related to modifiable processes that affect patient outcomes;
- Minimal unintended adverse consequences;
- Alignment with national priorities as described in the NQF NPP project;
- Amenable to collection via alternative mechanisms such as electronic health records (EHRs), registries, and claims;
- Harmonizes with measures used for reporting programs in similar settings;
- Attributable to the facility rather than a prescribing physician;
- Data collection should not increase hospital operational burden; and
- Fully tested in a variety of outpatient settings.

Response: We thank the commenters for these suggestions, and we note that these suggestions were not submitted in reference to specific measures. In section XVI.B.1. of this final rule with comment period, we have set out the criteria that we use to guide our decisions regarding what measures to add to the HOP QDRP measure set. We determine the suitability of potential measures using consensus development processes, including, when appropriate, relying upon the NQF's voluntary consensus standards in addition to our rulemaking in determining the suitability of quality measures.

In the CY 2009 OPPTS/ASC final rule with comment period, we adopted four claims-based quality measures that do not require a hospital to submit chart-abstracted clinical data. This supports our goal of expanding the measures for the HOP QDRP while minimizing the burden upon hospitals and, in particular, without significantly increasing the chart abstraction burden. In addition to claims-based measures, we are considering registries¹ and EHRs as alternative ways to collect data from hospitals. Many hospitals submit data to and participate in existing registries. In addition, registries often capture outcome information and provide ongoing quality improvement feedback to registry participants. Instead of requiring hospitals to submit the same data to CMS that they are already submitting to registries, we could collect the data directly from the registries with the permission of the hospital, thereby enabling us to expand the HOP QDRP measure set without increasing the burden of data collection for those hospitals participating in the registries. The data that we would receive from registries would be used to calculate quality measures required under the HOP QDRP, and would be publicly reported like other HOP QDRP quality measures, encouraging improvements in the quality of care. In the CY 2010 OPPTS/ASC proposed rule (74 FR 35397), we invited public comment on such an approach.

Comment: Many commenters expressed concern about the potential use of registries as a source of data for the HOP QDRP. Many commenters indicated that the fees imposed by registries would be prohibitive for smaller hospitals and rural hospitals. Regarding registry-based data submission for the HOP QDRP, CMS was urged to do the following:

- Develop and test alternatives for hospitals choosing to submit data directly to CMS in lieu of participating in a registry (that is, chart abstraction, CART tool);
- Determine and articulate a process for validating data submitted through registries for completeness and accuracy;
- Determine and articulate a process to transmit registry-based data to the national data warehouse in a secure fashion and without violating HIPAA or other rules;
- Explore and determine the willingness and ability of the ORYX vendors to submit data for those hospitals not participating in a registry; and
- Require standardized, externally verifiable sampling for the measures.

Response: We are interested in minimizing the burden associated with quality measurement. If hospitals are participating in registries and submit the same data to those registries that they would otherwise have to submit for measures that are part of the HOP QDRP, we believe that the registry-based data would be an efficient alternative source from which to collect the data, and that this would prevent the hospital from having to report the same data twice. Many hospitals are currently participating in a number of registries that collect data on quality measures that are topics of interest to us. However, we acknowledge the commenters' concerns regarding the cost associated with participation in certain registries that may make this alternative mechanism for data submission less feasible for some hospitals, and the need for standardized validation strategies for registry-based data. We will take these considerations into account when considering registry-based measure submission options for this and other reporting programs in the future.

Comment: Some commenters strongly supported the use of registries as an alternative source of data for the HOP QDRP. These commenters stated that registries provide a substantial advantage over chart-abstracted data because registries provide regular feedback reports to participating hospitals on their performance, further minimize the reporting burden for physicians and facilities because registry-based data could be used for more than one reporting program, and aggregate clinical data from a provider's entire patient population and enable these data to be analyzed and tracked over time for adherence to evidence-based medicine and health outcomes. Commenters encouraged CMS to

continue to explore this mechanism and to develop the infrastructure standards needed to accurately capture such data as soon as practicable.

Response: We thank these commenters for their encouragement and will continue to investigate the feasibility of such an approach to the HOP QDRP and other quality data reporting programs.

Comment: One commenter recommended that CMS request legislative authority to base payments on pay-for-performance so that a portion of payments will depend on providers' performance on the selected quality measures, not simply on whether they report the specified data to CMS. This commenter also expressed support for CMS' efforts to collect data on measures of hospital quality as a valuable step toward pay-for-performance.

Response: We thank the commenter for sharing this suggestion for future program direction and for supporting current program operations.

In the CY 2009 OPPTS/ASC final rule with comment period, we also stated our intention to explore mechanisms for data submission using EHRs (73 FR 68769). Establishing such a system will require interoperability between EHRs and CMS data collection systems, additional infrastructure development on the part of hospitals and CMS, and the adoption of standards for the capturing, formatting, and transmission of data elements that make up the measures. However, once these activities are accomplished, the adoption of measures that rely on data obtained directly from EHRs will enable us to expand the HOP QDRP measure set with less cost and burden to hospitals.

Comment: Some commenters strongly supported the use of EHRs and other health information technology (IT). These commenters believed that such technology has the ability to capture, store, and readily report the types of clinical data not available from medical claims data, such as diagnostic laboratory test results and prescription drug dispensing data. Commenters commended CMS for encouraging the development and adoption of uniform data content and information technology standards across the health care industry that will support automated data collection and reporting of clinical data from EHR systems. These commenters believed that such efforts would streamline hospital data submission procedures and enable providers to view real-time measurement results to initiate their own improvement interventions in a more timely and efficient manner.

¹ A registry is a collection of clinical data for purposes of assessing clinical performance, quality of care, and opportunities for quality improvement.

Response: We appreciate these supportive comments regarding EHR-based data collection as an alternative data source for quality measures. We agree that EHR-based data submission may provide an alternative means of submitting quality data that would benefit hospitals by reducing their chart abstraction burden. We also agree that such systems may enable providers to implement more timely improvement efforts. Although we encourage adoption of EHRs, we also acknowledge the challenges that must be met both by hospitals and CMS to establish the infrastructure and interoperability necessary to collect data on quality measures via EHRs. We will continue to work collaboratively with health IT standard-setting and consensus development organizations to ensure that quality measures can be collected in a standardized manner.

Comment: Many commenters were concerned about the ability of EHRs to accurately capture the data required for meaningful and accurate quality measures for the HOP QDRP. Commenters indicated that, currently, all of the necessary information for measuring performance against essential metrics of quality (such as exclusion and inclusion criteria and contraindications) is not codified within EHRs, and that the need for such information will still require medical record review because the information cannot be adequately found in EHRs. Other commenters indicated that current products feature inconsistent communication standards and may pose privacy concerns. Several commenters indicated that small rural hospitals may not be able to enhance their health IT infrastructure to support EHR-based reporting. Several commenters supported one-way transmission of specific data elements from EHRs, but would not support providing access to the whole EHR to abstract clinical information for quality measures. Commenters encouraged CMS to consider postponing new measure implementation for CY 2012 until new measures can be verified to be structured for EHR data collection, especially given impending challenges of ICD-10 implementation.

Response: We do not agree with the commenters' belief that quality data produced from EHRs is not likely to accurately capture data elements needed for quality measurement. The data collected from the EHR would essentially be the same data that hospitals would otherwise have to manually abstract from a medical chart. These data are what we currently use for quality measure reporting. We

acknowledge that additional programming work may be needed in order to enable current EHR systems to collect and submit quality measure data. We are currently working with the Healthcare Information Technology Standards Panel (HITS-SP), a public-private partnership working to establish health IT interoperability standards under contract to the HHS Office of the National Coordinator on Health IT (ONC), to standardize the specifications of data elements used in several measure sets so that they may be collected and reported via EHRs. Standardization of the specifications allows software to convert clinical data of different types into a form that can be analyzed for quality measurement. We encourage collaboration among standard-setting organizations and measure developers on the creation of standards for electronic collection of data elements for other quality measures as well, particularly those used in our quality data reporting programs.

With regard to the commenters' concern about having to provide access to the entire EHR, we would only require that the hospital provide access to those data elements in the EHRs that are needed to calculate the measures. We also acknowledge the burden faced by hospitals in implementing multiple technological changes, including the ICD-10 coding system. We will carefully consider any additional burden that may be imposed by adopting additional measures for the HOP QDRP and will continue to consider other feasible alternatives to data collection such as registries.

Comment: Several commenters encouraged the collection of all-payer or multiple-payer claims information in order to calculate measures for the HOP QDRP as it would provide a more complete picture of care to consumers. Commenters also encouraged CMS to ensure the validity of any third party data used in the development or calculation of measures for public reporting.

Response: We thank the commenters for their encouragement of the collection of all-payer claims data, and we agree that all-payer claims data would enable us to provide consumers with more comprehensive claims-based quality measures that provide a comprehensive picture of the quality of care provided by a hospital. We currently collect other all-payer data where feasible for the hospital quality data reporting programs, and currently this is feasible for chart-abstracted data elements. It has been our policy to collect all-payer chart-abstracted data since the inception of both inpatient

(RHQDAPU program) and outpatient (HOP QDRP) quality data reporting. While we currently do not have the infrastructure in place to accept all-payer claims data, we intend to work with stakeholders to identify options, processes, and opportunities to collect all-payer claims data to supplement the Medicare claims data we currently use in many of our reporting programs.

Comment: Several commenters indicated that CMS should only concern itself with obtaining information and outcomes for Medicare beneficiaries, and should not collect information regarding patients for whom other payers are responsible.

Response: For the HOP QDRP, section 1833(t)(17)(C)(i) of the Act requires the Secretary to develop measures appropriate for the measurement of the quality of care (including medication errors) furnished by hospitals in outpatient settings. The collection and publication of quality measures based on all-payer data captures variations in the care delivered by a hospital to different populations and payers, and therefore allows us to obtain comprehensive information regarding the quality of care provided to its beneficiaries. Therefore, we are collecting all-payer data elements to calculate the chart-abstracted measures adopted into the HOP QDRP. We wish to eventually provide a similarly comprehensive picture of the quality of care provided by HOPDs with respect to the claims-based measures adopted into the HOP QDRP.

2. Retirement of HOP QDRP Quality Measures

In the FY 2010 IPPS/R Y 2010 LTCH PPS proposed rule, we proposed a process for immediate retirement of RHQDAPU program measures based on evidence that the continued use of the measure as specified raises patient safety concerns (74 FR 24168). As we explained in that proposed rule, in situations such as the one prompting immediate retirement of the AMI-6 measure from the RHQDAPU program in December 2008, we do not believe that it would be appropriate to wait for the annual rulemaking cycle to retire a measure. This proposal was later finalized for the RHQDAPU program in the FY 2010 IPPS/R Y 2010 LTCH PPS final rule (74 FR 43863). We proposed to adopt this same immediate retirement policy for the HOP QDRP (74 FR 35397). Specifically, in the CY 2010 OPPS/ASC proposed rule, we proposed that if we receive evidence that continued collection of a measure that has been adopted for the HOP QDRP raises patient safety concerns, we would

promptly retire the measure and notify hospitals and the public of the retirement of the measure and the reasons for its retirement through the usual means by which we communicate with hospitals, including but not limited to hospital e-mail blasts and the QualityNet Web site. We also proposed to confirm the retirement of the measure in the next OPDS rulemaking. In other circumstances, where we do not believe that continued use of a measure raises specific patient safety concerns, we stated that we intend to use the regular rulemaking process to retire a measure.

We invited public comment on this proposal allowing for immediate retirement of a HOP QDRP measure following evidence of a patient safety concern followed by confirmation in the next rulemaking cycle.

Comment: Several commenters applauded the proposal to immediately retire a HOP QDRP measure if CMS receives evidence that the continued collection of a measure raises patient safety concerns. They encouraged CMS to establish consistent and transparent processes that address changes in evidence-based guidelines more quickly and to establish channels to exchange this type of information between the agency and measure developers. The commenters also encouraged CMS to retire measures under the following conditions:

- A measure is no longer consistent with current clinical guidelines;
- Another indicator exists that better, or more accurately, assesses good quality care;
- Redundancy of measurement on a given topic or process; and
- The burden associated with data collection and reporting a measure outweighs the benefit of public reporting;

Response: We thank the commenters for their support for the proposed policy of prompt retirement when potential patient harm could result from the continued collection of a measure, and are finalizing our policy in this final rule with comment period. With respect to the suggestions we received, these criteria reflect examples of conditions that may warrant retirement via notice and comment rulemaking as opposed to prompt retirement because continued collection of the measure does not raise patient safety concerns. Another example of a nonurgent circumstance where we would use the rulemaking process to retire a measure would be when a measure is “topped out.” While we did not solicit public comments on criteria for retirement under circumstances other than potential patient harm, we will consider these

suggestions as we consider whether to propose to retire measures in nonurgent circumstances.

After consideration of the public comments we received, we are finalizing our proposal to promptly retire measures under circumstances in which we receive evidence that continued collection of a measure that has been adopted for the HOP QDRP raises patient safety concerns, to notify hospitals and the public of the retirement of the measure and the reasons for its retirement through the usual means by which we communicate with hospitals, including, but not limited to, hospital e-mail blasts and the QualityNet Web site, and to confirm the retirement of measures retired in this manner in the next rulemaking cycle.

3. HOP QDRP Quality Measures for the CY 2011 Payment Determination

For the CY 2011 payment determination, in the CY 2010 OPDS/ASC proposed rule (74 FR 35397), we proposed to continue requiring that hospitals submit data on the existing 11 HOP QDRP measures. These measures continue to address areas of topical importance regarding the quality of care provided in HOPDs, and reflect consensus among affected parties. Seven of these 11 measures are chart-abstracted measures in two areas of importance that are also measured for the inpatient setting: AMI care and surgical care. The remaining four measures address imaging efficiency in HOPDs.

For the CY 2011 payment determination, we proposed not to add any new HOP QDRP measures. Although we considered adding a number of chart-abstracted measures, we are sensitive to the burden upon HOPDs associated with chart abstraction and believe that adopting such measures at this time would not be consistent with our stated goal to minimize the collection burden associated with quality measurement. We will continue to assess whether we can collect data on additional quality measures through mechanisms other than chart abstraction, such as from Medicare administrative claims data and EHRs.

We invited public comment on our proposal to retain the existing 11 HOP QDRP measures and to not adopt additional measures for the CY 2011 payment determination.

Comment: Most commenters were pleased that CMS recognizes the burden that data collection and reporting places on facilities and did not propose to add new measures to the HOP QDRP measurement set for the CY 2011 payment determination. In particular,

some hospitals indicated that they have only one staff member performing chart abstraction for both the inpatient and outpatient quality data reporting programs, and that the burden of adding measures has a great impact under such circumstances.

Response: We thank the commenters for expressing their support for this proposal. We will continue to carefully weigh the burden associated with adding chart-abstracted measures to quality reporting programs such as the HOP QDRP against the benefit of adding such measures in the future.

We also received specific comments, discussed below, on some of the measures we proposed to retain.

- OP-3: Median Time To Transfer to Another Facility for Acute Coronary Intervention

Comment: One commenter recommended that CMS consider measuring the overall median time to percutaneous coronary intervention (PCI) in transferred patients because this captures the entire process of care and will encourage collaboration between transferring and receiving ST-segment elevation myocardial infarction (STEMI) centers.

Response: We thank the commenter for this suggestion. The current measure is meant to be one of accountability for the initial (transferring) facility rather than for both the transferring and receiving facility. Therefore, the outpatient measure that is currently in place (OP-3) focuses on the measurable time of arrival to time of physical departure from the first hospital, which is an important component of the total time to reperfusion. A modification to the measure as suggested would not currently be feasible to implement as it would require capturing information from medical records at two separate facilities.

- OP-4: Aspirin at Arrival & OP-5: Median Time to ECG

Comment: One commenter recommended that CMS consider excluding “Chest Pain NEC” from the list of eligible cases for these two measures because many of these cases are not “probable cardiac chest pain” as is the intent of the measures. This commenter also recommended only using the working diagnosis in the “final impression,” rather than working diagnoses used throughout the ED documentation forms, and recommended excluding patients in observation status, as patients believed to have “probable cardiac chest pain” or AMI will likely not be kept under observation status. The commenter believed implementing these

recommendations will eliminate many cases that these measures did not intend to capture. Another commenter noted that OP-4 has the potential to become "topped out" as the program matures.

Response: We will consider these suggestions as part of maintenance of the technical specifications for the measure. We also will evaluate the performance of OP-4 over time as we do with other measures that have been adopted for public reporting programs.

- **Imaging Efficiency Measures Generally**

Comment: Several commenters objected to CMS continuing to include the four imaging efficiency measures in the HOP QDRP. Many of these commenters objected because none of the four measures have been adopted by the HQA. Other commenters acknowledged that OP-8 and OP-11 are NQF-endorsed, and also acknowledged that NQF endorsement is not required, but recommended that CMS obtain endorsement for OP-9 and OP-10 in order to establish their credibility. Some commenters opined that the two non-NQF endorsed Imaging Efficiency measures, OP-9 and OP-10, are inappropriate for the HOP QDRP and could cause patient harm. One commenter cautioned that, because the protocols for reporting contrast media on claims have varied over the years, CMS should be aware that the use of contrast media may not be reliably documented in claims.

Response: Many of the concerns raised by the commenters about the imaging efficiency measures were also raised at the time the imaging measures were proposed. We responded to these concerns when we adopted the measures (74 FR 68762 through 68766). We stated that the measures meet the statutory definition of reflecting consensus among affected parties through their consensus-based development, and that the measures address important patient safety concerns related to exposure to unnecessary radiation and contrast materials. We also stated that the Secretary is not required to limit measures considered for selection to only those adopted by the HQA or to those that have been NQF-endorsed. We anticipate submitting OP-9 and OP-10 for NQF endorsement, along with national performance information and other supporting information, when an appropriate call for measures occurs. We note the cautionary advice regarding the varying requirements for reporting contrast media on claims. However, the OP-10 and OP-11 measures rely on procedure codes rather than on specific

material codes to determine whether a with-contrast procedure or without-contrast procedure was performed. In other words, these measures only consider whether contrast media was appropriately used during diagnostic imaging procedures, regardless of type.

- **OP-8: MRI Lumbar Spine for Low Back Pain**

Comment: Some commenters indicated that complete details of a patient treatment plan and history such as conservative therapy for the previous 60 days would often be unavailable to an imaging provider or outpatient center. Other commenters indicated that this measure would be appropriate for the PQRI program.

Response: While a HOPD may not have complete information about a patient's treatment plan and history, such as conservative therapy, HOPDs are in a position to consult and directly communicate with ordering physicians and the radiologists employed by the HOPD. HOPDs can also educate hospital medical staff and community physicians on the appropriate use of MRI for low back pain. We thank the commenters for suggesting that OP-8 may be appropriate for the PQRI program, and will consider this suggestion. We agree that the basis for the measure may be appropriately applied at the ordering physician level. However, we note that the measure has been endorsed by the NQF as appropriate for facility-level measurement.

- **OP-9: Mammography Follow-up Rates**

Comment: Some commenters objected to this measure because they believed that there is a lack of consensus as to what the appropriate recall rate should be, and that there is no established link between providers' recall rates and patient outcomes. Many commenters expressed concern that the measure implies that high follow-up rates are undesirable, leading to decreased access to these tests, and an increase in undiagnosed early cases of cancer. Other commenters supported this measure, stating that mammography is a life saving tool that is currently underutilized. In addition, some commenters suggested revisions that they believed would improve the current measure. These suggestions include:

- Extending the call back period to 3 months to allow adequate time for a patient to return;
- Counting breast MRI within 3 months of a screening examination as a call back;

- Revising the measure to be a proportion of screening mammograms interpreted as positive, or where the radiologist has recommended further evaluation.

Response: We do not believe that HOPDs should refuse access to mammograms when appropriate follow-up study is needed. We also do not believe that the measure encourages HOPDs to do so. The measure allows identification of facilities with abnormally high rates of "call-backs" from indeterminate or inadequate screening studies. We will evaluate the commenters' suggestions for improvements to the measure specifications as part of the maintenance process for the measure.

- **OP-10: Abdomen CT—Use of Contrast Material**

Comment: Some commenters indicated that there is a lack of evidence in the literature to determine the appropriate use of contrast material for these patients, and, thus, there is no accepted best practice. In addition, some commenters asserted that, because the measure contains a number of patient exclusions, the applicable patient population is unclear. Other commenters approved of the decision to exclude renal disease patients from this measure.

Response: We have incorporated existing clinical guidelines for appropriate use of combined imaging studies (with and without contrast) into the imaging efficiency measures. Nevertheless, imaging efficiency measures are not intended to define absolutes and should not be interpreted to mean that combined studies would never be considered appropriate. We believe that the measures will promote more careful consideration in individual cases as to whether, in the particular circumstance, a combined study is necessary and thus enhance the efficient use of combined studies. We also anticipate that the variation that exists will lessen and approaches to the use of combined studies will become more standardized.

By implementing the denominator exclusions, we seek to more clearly define the applicable patient population for the quality measure. We thank the commenters that supported the exclusion of renal disease patients from the denominator of this measure.

- **OP-11: Thorax CT—Use of Contrast Material**

Comment: One commenter objected to the inclusion of this measure, stating that current guidelines indicate that it is acceptable to perform a with-contrast

study followed by a without-contrast study as clinically indicated.

Response: We agree that, if clinically indicated, such dual studies are appropriate. As with the OP-10 measure, the intent of the OP-11 measure is not to reduce the use of

contrast studies or dual studies to zero, but to identify facilities utilizing dual study protocols in the majority of cases when not clinically appropriate.

After consideration of the public comments we received, we have decided to adopt as final our proposal

to retain the existing 11 HOP QDRP measures without adding new measures to the measure set for the CY 2011 payment determination. The measure set that will be used for the CY 2011 payment determination is displayed below.

HOP QDRP MEASUREMENT SET TO BE USED FOR THE CY 2011 PAYMENT DETERMINATION

- OP-1: Median Time to Fibrinolysis
- OP-2: Fibrinolytic Therapy Received Within 30 Minutes
- OP-3: Median Time to Transfer to Another Facility for Acute Coronary Intervention
- OP-4: Aspirin at Arrival
- OP-5: Median Time to ECG
- OP-6: Timing of Antibiotic Prophylaxis
- OP-7: Prophylactic Antibiotic Selection for Surgical Patients
- OP-8: MRI Lumbar Spine for Low Back Pain
- OP-9: Mammography Follow-up Rates
- OP-10: Abdomen CT—Use of Contrast Material
- OP-11: Thorax CT—Use of Contrast Material

C. Possible Quality Measures Under Consideration for CY 2012 and Subsequent Years

In previous years' rulemakings, we have provided lists of quality measures

that are under consideration for future adoption into the HOP QDRP measurement set. In the CY 2010 OPSP/ASC proposed rule (74 FR 35398), we set out a list of measures under

consideration for the CY 2012 payment determination and subsequent years. That list is displayed below.

QUALITY MEASURES UNDER CONSIDERATION FOR CY 2012 AND SUBSEQUENT YEARS' PAYMENT DETERMINATIONS

Topic	No.	Measure	Potential data sources
Cancer	1	Adjuvant Chemotherapy Is Considered or Administered Within 4 Months of Surgery to Patients Under Age 80 With AJCC III Colon Cancer. The measure specifications are similar to PQRI # 72 found at the PQRI manual Web site: http://www.cms.hhs.gov/apps/ama/license.asp?file=/PQRI/downloads/2009PQRIQualityMeasureSpecificationsManualandReleaseNotes.zip .	Registry.
	2	Adjuvant Hormonal Therapy for Patients with Breast Cancer	Claims, Registry.
	3	Needle Biopsy To Establish Diagnosis of Cancer Precedes Surgical Excision/Resection. The measure specifications can be found at: http://www.qualityforum.org/pdf/reports/Cancer_Nonmember_Report.pdf .	Claims, Registry.
ED Throughput	4	Median Time From ED Arrival to ED Departure for Discharged ED Patients. The measure specifications can be found at http://www.qualitynet.org/ in Appendix P of the specifications manual under Hospital—Out-patient.	Chart, EHR.
Diabetes	5	Low Density Lipoprotein Control in Type 1 or 2 Diabetes Mellitus	Claims, EHR.
	6	Urine protein screening or medical attention for nephrology during at least one office visit within last year for patient with diabetes mellitus. The measure specifications are similar to PQRI # 119 found at the PQRI manual Web site: http://www.cms.hhs.gov/apps/ama/license.asp?file=/PQRI/downloads/2009PQRIQualityMeasureSpecificationsManualandReleaseNotes.zip .	Claims, EHR.
	7	Eligible diabetes patients with documentation of an eye exam or referral for an eye exam within the last 24 months.	Claims, EHR.

QUALITY MEASURES UNDER CONSIDERATION FOR CY 2012 AND SUBSEQUENT YEARS' PAYMENT DETERMINATIONS—
Continued

Topic	No.	Measure	Potential data sources
		The measure specifications are similar to PQRI # 117 found at the PQRI manual Web site: http://www.cms.hhs.gov/apps/ama/license.asp?file=/PQRI/downloads/2009PQRIQualityMeasureSpecificationsManualandReleaseNotes.zip .	
	8	Patients who received at least one complete foot exam (visual inspection, sensory examination with monofilament and pulse exam within the last 12 months). The measure specifications are similar to PQRI # 126 found at the PQRI manual Web site: http://www.cms.hhs.gov/apps/ama/license.asp?file=/PQRI/downloads/2009PQRIQualityMeasureSpecificationsManualandReleaseNotes.zip .	Claims, EHR.
Medication Reconciliation	9	Medication Reconciliation The measure specifications are similar to PQRI # 46 found at the PQRI manual Web site: http://www.cms.hhs.gov/apps/ama/license.asp?file=/PQRI/downloads/2009PQRIQualityMeasureSpecificationsManualandReleaseNotes.zip .	Claims, EHR.
Immunization	10	Pneumococcal Vaccination Status—Overall Rate The measure specifications are available at http://www.qualityforum.org/pdf/reports/Immunization/4%2029%20Immunizations_Nonmembers.pdf .	Chart, EHR.
	11	Influenza Vaccination Status—Overall Rate The measure specifications are available at http://www.qualityforum.org/pdf/reports/Immunization/4%2029%20Immunizations_Nonmembers.pdf .	Chart, EHR.
Imaging Efficiency	12	SPECT MPI and Stress Echocardiography for Preoperative Evaluation for Low-Risk Non-Cardiac Surgery Risk Assessment. The measure specifications can be found at http://www.imagingmeasures.com/ .	Claims.
	13	Use of Stress Echocardiography or SPECT MPI Post-Revascularization Coronary Artery Bypass Graft. The measure specifications can be found at http://www.imagingmeasures.com/ .	Claims.
	14	Use of Computed Tomography in Emergency Department for Headache The measure specifications can be found at http://www.imagingmeasures.com/ .	Claims.
	15	Simultaneous Use of Brain Computed Tomography and Sinus Computed Tomography. The measure specifications can be found at http://www.imagingmeasures.com/ .	Claims.
Surgery	16	Appropriate surgical site hair removal The measure specifications are similar to Surgical Care Improvement Project Infection (SCIP)—6 which can be found at http://www.qualitynet.org/ under Hospital—Inpatient.	Chart, EHR.

We invited public comment on these quality measures and topics that we may consider proposing to adopt beginning with the CY 2012 payment determination. We also sought suggestions and rationales to support the adoption of measures and topics for the HOP QDRP which do not appear in the table above.

- Cancer (Potential Measures 1, 2, and 3)

Comment: A number of commenters supported the cancer measures because: (1) They align with national priorities

and CMS priority condition areas; (2) they provide insight into an area of care that is very relevant to the consumer; and (3) the measure set seems to address health care provided across settings. One commenter indicated that CMS should more clearly state whether CMS or the HOPD will collect information on chemotherapy within the 4-month timeframe stated in the measure, and how this information will be collected. Some commenters stated that, because some of the cancer measures are registry-based measures, the added costs of implementing measures that require

paying a fee to a nongovernmental entity would hinder small rural hospitals from being able to report data. Other commenters indicated that a process for validating registry-based data should be proposed prior to implementing quality measures based on registry data.

Response: We agree with the commenters who supported the cancer measures. We acknowledge that receiving data from registries presents additional issues, but believe that in circumstances where substantial timeframes are involved, registries may

provide the best data collection mechanism. We will take these comments into consideration in deciding whether to propose these measures in the future for the HOP QDRP, and would specify the form and manner for data submission required should we, in the future, adopt these measures.

- Emergency Department Throughput (Potential Measure 4)

Comment: A few commenters expressed strong support for the ED throughput Measure 4 (Median Time from ED Arrival to ED Departure for Discharged ED Patients) and recommended its inclusion in the HOP QDRP. Some commenters stated that a measure assessing delays in patient care is important as providers experience a growth in demand for ED services. Commenters saw the measure as making significant contributions to reducing overcrowding, and in turn increasing the quality of care delivered, particularly when public reporting occurs.

Response: We thank these commenters for their supportive statements. We agree with the commenters that this measure addresses the issue of timely emergency department care and delays which have an adverse impact on quality of care due to overcrowding.

Comment: One commenter indicated that the ED throughput measure is overly burdensome for hospitals to collect as it will require an arrival time to be noted for each patient, whether the patient is on observation, and will require sampling over 300 records per quarter. Other commenters indicated that, as currently structured, the measure includes the time spent receiving care in the ED in addition to the time spent waiting in the ED. These commenters indicated that the time spent receiving care in the ED should not be counted against the hospital, as it does not represent a delay in care. The commenters stated that, for patients discharged back into the community, and not admitted or transferred to another facility, there is no wait time in the ED after the patient has received the appropriate care. The commenters noted that, for these patients, any time spent waiting in the ED occurs before they see a provider. The commenters suggested that CMS modify the measure so that it reflects only the time spent waiting in the ED to see a provider. One commenter questioned whether the measure actually measures quality because fast care is not necessarily better care. Another commenter

indicated that it could not locate the specifications for this measure.

Response: We do not agree that the measure, as currently specified, would be overly burdensome to collect, because hospitals routinely collect the key information needed to calculate the median time (ED arrival date and time and ED departure date and time) for each emergency department patient. The current measure is an NQF-endorsed measure of quality, and feasibility of collection was among the considerations for its endorsement. Revising the measure in the manner suggested by the commenters to exclude active treatment times would be impractical, as it would impose a severe burden for hospitals to accurately track and collect the time spent in the ED not receiving care. We do not agree with the comment that prolonged ED throughput is solely due to time elapsed between arrival and first contact with a provider. The measure specifications are currently available in Appendix P of the HOPD Specifications Manual (versions 2.1b and 3.0) which is posted on the QualityNet Web site (<http://www.qualitynet.org/>).

- Diabetes (Potential Measures 5, 6, 7, and 8)

Comment: A few commenters indicated that they believed the three diabetes measures would be better suited to measure the quality of care in physician offices or physician-based clinics rather than in HOPDs. Other commenters indicated that, if finalized, CMS should consider HOPD participation in a disease management registry, or recognition and/or certification in disease management, as substitutes for the requirement of submitting diabetes-related quality measures to the CMS for the HOP QDRP. One commenter indicated that, for the LDL Control measure, CMS should account for the fact that some patients may not reach the goal but their risk may be mitigated by high HDL. One commenter indicated that the timeframe for the eye examination should be 24 months if there is no retinopathy and 12 months if retinopathy is known to be present.

Response: We agree on the suitability of such measures for the physician office setting and note that these measures are currently part of the PQRI program. We would anticipate that these measures would be appropriate for reporting by those HOPDs that function as a primary care provider. We would not view participation in a registry or disease management program certification/recognition as a substitute for reporting quality measures for the

HOP QDRP because it would not allow us to achieve the goal of providing comparative quality information on HOPDs to Medicare beneficiaries. These measures are currently being specified for the HOPD setting, and we will consider the suggestions for enhancements submitted by commenters.

- Medication Reconciliation (Potential Measure 9)

Comment: A few commenters supported the medication reconciliation measure but urged CMS to clarify its expectation of medication reconciliation in the ED. Some commenters indicated that, although medication reconciliation measures were recently NQF-endorsed, implementing a quality measure in multiple outpatient settings may result in more medication errors, and recommended that the measure be implemented in primary care settings.

Response: We are interested in medication reconciliation in all settings of care because medication errors may result in serious avoidable complications, and receiving the appropriate medications throughout the continuum of care may prevent the onset or worsening of serious medical conditions. Thus, the reduction of medication errors would contribute to overall improvements in patient outcomes and quality of life and would reduce mortality and hospital readmissions. We would expect that, prior to administration of or prescription of drugs in an ED setting, a patient's current medications, drug allergies, current acute condition, and chronic conditions would be assessed to the extent possible in order to prevent adverse drug-drug interactions and drug-disease interactions. We will take these comments into consideration in determining whether to propose this measure for the HOP QDRP in the future.

- Immunization (Potential Measures 10 and 11)

Comment: One commenter stated that the influenza and pneumococcal vaccination measures will contribute to ED overcrowding, and that the measures are not appropriate for the HOPD setting as administering influenza and pneumococcal vaccinations are not part of routine emergency care protocols like administering a tetanus vaccine would be for wound care. The commenter believed that the measures will work against the ED throughput measures and would be more appropriate for physician offices and community public health departments.

Response: These measures are currently being specified for HOPDs and are not intended for EDs. These measures are intended to apply to the facility under circumstances where the HOPD serves as a primary care provider. We will consider these comments in deciding whether to propose this measure for the HOP QDRP.

- Imaging Efficiency—SPECT MPI and Stress Echocardiography (Potential Measures 12 and 13)

Comment: Some commenters indicated that the two measures on SPECT MPI and Stress Echocardiography should not be considered for the following reasons:

- Lack of benchmarks;
- Preoperative or postoperative period difficult for provider of test to track;
- Lack of medical history makes it difficult for a provider to determine if a test is appropriate for a patient;
- Not clear how the purpose of test (preoperative evaluation) is captured in Medicare claims; and
- Medicare claims provide an incomplete picture of facility performance.

In addition, for Measure 13 (Use of Stress Echocardiography or SPECT MPI Post-Revascularization Coronary Artery Bypass Graft (CABG)), the commenters indicated that the measure's long time span of a 5-year period post-CABG hinders its usability as the information will be unavailable for a number of years and will be irrelevant by the time it becomes available.

Response: These measures are currently under development, and we will take these comments into consideration as the measures are developed further.

Comment: Some commenters applauded CMS' effort to obtain consensus among affected parties as evidenced by hosting of a public comment period during the measure development process, and supported Measure 12 (SPECT MPI and Stress Echocardiography for Preoperative Evaluation of Low Risk Non-Cardiac Surgery) and Measure 13. However, these commenters also recommended stratification of the measures by imaging procedure.

Response: We appreciate the supportive comments regarding our consensus-based measure development process. We will consider these suggestions for these measures as we continue measure development.

- Imaging Efficiency—Computed Tomography (Potential Measures 14 and 15)

Comment: For both measures, several commenters indicated that the measures target important areas where overuse of diagnostic imaging may be detrimental to patient care, and the measures appear valid and usable. However, the commenters believed that because the measures are based on Medicare claims, they would provide an incomplete picture of facility performance. One commenter suggested excluding "sign of meningeal irritation (stiff neck)" from Measure 14 (Use of Computed Tomography in Emergency Department for Headache).

Response: Both of these measures are under development, and both address overutilization of CT scans in the outpatient setting which have implications for patient safety due to radiation exposure. The goal of these measures is not to reduce outpatient diagnostic CT imaging in these circumstances to zero, but to encourage its use only in circumstances where it is clinically indicated. Though all-payer claims are not currently included in these measures, due to the high volume of these services in the Medicare population relative to other populations, we believe that calculation of these measures based on Medicare claims only will target performance improvement where it is most needed: in the population that is at high risk for inappropriate imaging studies. We appreciate the supportive comments, and will consider these suggestions in the continuing development of these measures.

- Surgery (Potential Measure 16)

Comment: Some commenters indicated that Measure 16 (Appropriate surgical site hair removal) is an unnecessary measure, as performance on the measure in the inpatient setting is already in the high 90 percent range for the Nation. One commenter also indicated that SCIP officials may retire the measure because it is "topped out" and no longer distinguishes between high performers and low performers. Other commenters suggested that CMS not use this measure for quality reporting or, at the very least, exclude cases for which there is no supporting evidence that the use of razors results in lesser quality of care, and cases in which razors would prevent wound bandages from falling off, thus decreasing the chance of infection.

Response: While hospitals may perform highly on this measure in the inpatient setting, we currently do not

know if this is the case for the outpatient setting. We will take these comments into consideration in determining whether to propose this measure for the HOP QDRP in the future.

- Other Suggested Measures or Measurement Areas

Comment: Commenters suggested several measures or measurement areas for CMS to consider for future development and adoption. The suggestions include:

- Heart Failure: ACE or ARB for LVSD (NQF #0137);
- Pneumonia: Empiric antibiotic for CAP (NQF #0096);
- Diabetes: Hemoglobin A1c poor control in type 1 or 2 diabetes mellitus (NQF #0059);
- Outcome-based measures;
- Radiation therapy administered within 1 year of diagnosis for women under 70 receiving breast conserving surgery for breast cancer;
- Patient centeredness;
- Total lipid treatment;
- ED throughput;
- Orthopedic procedures;
- Diagnostic Mammography Positive Predictive Value;
- Screening Mammography Positive Predictive Value;
- Cancer Detection Rate (CDR);
- Abnormal Interpretation Rate;
- Emergency Department AMI mortality;
- Emergency Department-related nonmortality outcome measures (that is, NQF Sepsis measures);
- Overall cardiac care;
- Use and overuse of Cardiac CT;
- Percutaneous Cardiac Interventions ("PCI");
- Care transitions/care coordination;
- AMI-2: Aspirin prescribed at discharge;
- AMI-5: Beta Blocker prescribed at discharge;
- HF-1: Discharge instructions;
- PN-3b: Blood culture performed before first antibiotic received in hospital;
- COPD management;
- NQF-endorsed ASC quality measures;
- Rate of surgical infections in outpatient surgery centers; and
- Rate of infection outbreaks related to contaminated scopes, syringes, and other medical equipment.

Response: We thank the commenters for these suggestions for quality measures and measurement areas for the HOP QDRP, and we will consider them for the future. Some of the topics are reflected in the current list of measures and topics for future consideration.

Some of the specific measures suggested were considered in the past for the HOP QDRP but, upon evaluation, were either found not to be appropriate measures for HOPD services or were found to be overly burdensome. Other measures and measure topics on this list are currently under consideration as future areas of measurement for inpatient quality measure reporting, and we will examine the appropriateness of these measures for the HOP QDRP as well.

D. Payment Reduction for Hospitals That Fail To Meet the HOP QDRP Requirements for the CY 2010 Payment Update

1. Background

Section 1833(t)(17)(A) of the Act, which applies to hospitals as defined under section 1886(d)(1)(B) of the Act, requires that hospitals that fail to report data required for the quality measures selected by the Secretary, in the form and manner required by the Secretary under section 1833(t)(17)(B) of the Act, incur a 2.0 percentage point reduction to their OPD fee schedule increase factor, that is, the annual payment update factor. Section 1833(t)(17)(A)(ii) of the Act specifies that any reduction would apply only to the payment year involved and would not be taken into account in computing the applicable OPD fee schedule increase factor for a subsequent payment year.

In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68769 through 68772), we discussed how the payment reduction for failure to meet the administrative, data collection, and data submission requirements of the HOP QDRP affected the CY 2009 payment update applicable to OPPS payments for HOPD services furnished by the hospitals defined under section 1886(d)(1)(B) of the Act to which the program applies. The application of a reduced OPD fee schedule increase factor results in reduced national unadjusted payment rates that apply to certain outpatient items and services provided by hospitals that are required to report outpatient quality data and that fail to meet the HOP QDRP requirements. All other hospitals paid under the OPPS receive the full OPPS payment update without the reduction.

The national unadjusted payment rates for many services paid under the OPPS equal the product of the OPPS conversion factor and the scaled relative weight for the APC to which the service is assigned. The OPPS conversion factor, which is updated annually by the OPD fee schedule increase factor, is used to calculate the OPPS payment rate for services with the following status

indicators (listed in Addendum B to this final rule with comment period): “P,” “Q1,” “Q2,” “Q3,” “R,” “S,” “T,” “V,” “U,” or “X.” In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68770), we adopted a policy that payment for all services assigned these status indicators would be subject to the reduction of the national unadjusted payment rates for applicable hospitals, with the exception of services assigned to New Technology APCs, assigned status indicator “S” or “T,” and brachytherapy sources, assigned status indicator “U,” which were paid at charges adjusted to cost in CY 2009. We excluded services assigned to New Technology APCs from the list of services subject to the reduced national unadjusted payment rates because the OPD fee schedule increase factor is not used to update the payment rates for these APCs.

In addition, section 1833(t)(16)(C) of the Act, as amended by section 142 of Public Law 110–275, specifically required that brachytherapy sources be paid during CY 2009 on the basis of charges adjusted to cost, rather than under the standard OPPS methodology. Therefore, the reduced conversion factor also was not applicable to CY 2009 payment for brachytherapy sources because payment would not be based on the OPPS conversion factor and, consequently, the payment rates for these services were not updated by the OPD fee schedule increase factor. However, in accordance with section 1833(t)(16)(C) of the Act, as amended by section 142 of Public Law 110–275, payment for brachytherapy sources at charges adjusted to cost is set to expire on January 1, 2010. For CY 2010, in the CY 2010 OPPS/ASC proposed rule (74 FR 35399), we proposed to pay prospectively for brachytherapy sources. Therefore, we proposed that the CY 2010 payment for brachytherapy sources would be based on the conversion factor and the quality reporting reduction policy would be applicable to brachytherapy sources, which are assigned status indicator “U.”

We did not receive any public comments on our proposal to apply the reporting reduction to payment for brachytherapy sources, effective for services furnished on and after January 1, 2010. Therefore, we are finalizing our CY 2010 proposal, without modification, to apply the reduction to payment for brachytherapy sources to hospitals that fail to meet the quality data reporting requirements of the HOP QDRP for the CY 2010 OPD fee schedule increase factor.

The OPD fee schedule increase factor, or market basket update, is an input into

the OPPS conversion factor, which is used to calculate OPPS payment rates. To implement the requirement to reduce the market basket update for hospitals that fail to meet reporting requirements, in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68770 through 68771), we calculated two conversion factors: a full market basket conversion factor (that is, the full conversion factor), and a reduced market basket conversion factor (that is, the reduced conversion factor). We then calculated a reduction ratio by dividing the reduced conversion factor by the full conversion factor. We refer to this reduction ratio as the “reporting ratio” to indicate that it applies to payment for hospitals that fail to meet their reporting requirements. Applying this reporting ratio to the OPPS payment amounts results in reduced national unadjusted payment rates that are mathematically equivalent to the reduced national unadjusted payment rates that would result if we multiplied the scaled OPPS relative weights by the reduced conversion factor. To determine the reduced national unadjusted payment rates that applied to hospitals that failed to meet their quality reporting requirements for the CY 2009 OPPS, we multiplied the final full national unadjusted payment rate in Addendum B to the CY 2009 OPPS/ASC final rule with comment period by the CY 2009 OPPS final reporting ratio of 0.981 (73 FR 68771).

In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68771 through 68772), we established a policy that the Medicare beneficiary’s minimum unadjusted copayment and national unadjusted copayment for a service to which a reduced national unadjusted payment rate applies would each equal the product of the reporting ratio and the national unadjusted copayment or the minimum unadjusted copayment, as applicable, for the service. We applied the reporting ratio to both the minimum unadjusted copayment and national unadjusted copayment for those hospitals that received the payment reduction for failure to meet the HOP QDRP reporting requirements. This application of the reporting ratio to the national unadjusted and minimum unadjusted copayments was calculated according to § 419.41 of the regulations, prior to any adjustment for hospitals’ failure to meet the quality reporting standards according to § 419.43(h). Beneficiaries and secondary payers thereby share in the reduction of payments to these hospitals.

In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68772), we

established the policy that all other applicable adjustments to the OPPS national unadjusted payment rates apply in those cases when the OPD fee schedule increase factor is reduced for hospitals that fail to meet the requirements of the HOP QDRP. For example, the following standard adjustments apply to the reduced national unadjusted payment rates: the wage index adjustment; the multiple procedure adjustment; the interrupted procedure adjustment; the rural sole community hospital adjustment; and the adjustment for devices furnished with full or partial credit or without cost. We believe that these adjustments continue to be equally applicable to payments for hospitals that do not meet the HOP QDRP requirements. Similarly, outlier payments will continue to be made when the criteria are met. For hospitals that fail to meet the quality data reporting requirements, the hospitals' costs are compared to the reduced payments for purposes of outlier eligibility and payment calculation. This policy conforms to current practice under the IPPS. For a complete discussion of the OPPS outlier calculation and eligibility criteria, we refer readers to section II.F. of this CY 2010 OPPS/ASC final rule with comment period.

2. Reporting Ratio Application and Associated Adjustment Policy for CY 2010

In the CY 2010 OPPS/ASC proposed rule (74 FR 35400), we proposed to continue our established policy of applying the reduction of the OPD fee schedule increase factor through the use of a reporting ratio for those hospitals that fail to meet the HOP QDRP requirements for the full CY 2010 annual payment update factor. For the CY 2010 OPPS, the proposed reporting ratio was 0.980, calculated by dividing the reduced conversion factor of \$66.118 by the full conversion factor of \$67.439. The final CY 2010 OPPS reporting ratio is 0.980, calculated by dividing the reduced conversion factor of \$66.086 by the full conversion factor of \$67.406. We proposed to continue to apply the reporting ratio to all services calculated using the OPPS conversion factor. For the CY 2010 OPPS, we proposed to apply the reporting ratio, when applicable, to all HCPCS codes to which we have assigned status indicators "P," "Q1," "Q2," "Q3," "R," "S," "T," "V," or "X" and, effective for services furnished on or after January 1, 2010, to also apply it to the HCPCS codes for brachytherapy sources, to which we have assigned status indicator "U." Under our established policy, we

would continue to exclude services paid under New Technology APCs. We proposed to continue to apply the reporting ratio to the national unadjusted payment rates and the minimum unadjusted and national unadjusted copayment rates of all applicable services for those hospitals that fail to meet the HOP QDRP reporting requirements. We also proposed to continue to apply all other applicable standard adjustments to the OPPS national unadjusted payment rates for hospitals that fail to meet the requirements of the HOP QDRP. Similarly, we proposed to continue to calculate OPPS outlier eligibility and outlier payment based on the reduced payment rates for those hospitals that fail to meet the reporting requirements.

We did not receive any public comments on our CY 2010 proposal to apply the HOP QDRP reduction in the manner described in the paragraph above and, therefore, are finalizing our proposal, without modification. For the CY 2010 OPPS, we are applying a reporting ratio of 0.980 to the national unadjusted payments, minimum unadjusted copayments, and national unadjusted copayments for all applicable services reported by those hospitals failing to meet the HOP QDRP reporting requirements. This reporting ratio applies to lines with HCPCS codes assigned status indicators "P," "Q1," "Q2," "Q3," "R," "S," "T," "U," "V," or "X," excluding services paid under New Technology APCs. All other applicable standard adjustments to the OPPS national unadjusted payment rates will continue to apply. This includes the OPPS outlier eligibility and payment calculations, which are determined using the reduced payment rates.

E. Requirements for HOPD Quality Data Reporting for CY 2011 and Subsequent Years

In order to participate in the HOP QDRP, hospitals must meet administrative, data collection and submission, and data validation requirements (if applicable). Hospitals that do not meet the requirements of the HOP QDRP, as well as hospitals not participating in the program and hospitals that withdraw from the program, will not receive the full OPPS payment rate update. Instead, in accordance with section 1833(t)(17)(A) of the Act, those hospitals will receive a reduction of 2.0 percentage points in their updates for the applicable payment year.

For payment determinations affecting the CY 2011 payment update, in the CY 2010 OPPS/ASC proposed rule (74 FR

35400), we proposed to implement the requirements listed below. Most of these requirements are the same as the requirements we implemented for the CY 2010 payment determination, with some proposed modifications.

1. Administrative Requirements

To participate in the HOP QDRP, several administrative steps must be completed. These steps require the hospital to:

- Identify a QualityNet administrator who follows the registration process located on the QualityNet Web site (<http://www.QualityNet.org>) and submits the information to the appropriate CMS-designated contractor. All CMS-designated contractors will be identified on the QualityNet Web site. The same person may be the QualityNet administrator for both the RHQDAPU program and the HOP QDRP. From our experience, we believe that the QualityNet administrator typically fulfills a variety of tasks related to the hospital's ability to participate in the HOP QDRP, such as: creating, approving, editing and/or terminating QualityNet user accounts within the organization; monitoring QualityNet usage to maintain proper security and confidentiality measures; and serving as a point of contact for information regarding QualityNet and the HOP QDRP.

In the past, we have required not only that the hospital designate a QualityNet administrator for purposes of registering the hospital to participate in the HOP QDRP, but also that the hospital continually maintain a QualityNet administrator for as long as the hospital participates in the program. We have become aware that the required maintenance of the QualityNet administrator is creating an undue technical burden for some hospitals and that, in some cases, is preventing the hospital from meeting all of the HOP QDRP requirements. Therefore, we proposed to no longer require that a hospital maintain current designation of a QualityNet administrator. We invited public comment on this proposed change. Nevertheless, we strongly urged hospitals to maintain current designation of a QualityNet administrator, regardless of whether the hospital submits data directly to the CMS-designated contractor or uses a vendor for transmission of data.

Comment: Many commenters agreed with CMS' proposal to remove the requirement to maintain current designation of a QualityNet administrator. Some of these commenters expressed their belief that it is in a hospital's best interest to

maintain a QualityNet administrator if possible. One commenter greatly appreciated the proposal to no longer require hospitals to maintain a QualityNet administrator to oversee the collection of HOP QDRP data because of the undue technical burden, particularly for hospitals in rural areas. This commenter believed that giving hospitals the option will lead to better quality data collection by lessening this burden on certain rural hospitals.

Response: We thank these commenters for their support; we agree that removing this program requirement will relieve a technical burden especially for some small or rural hospitals. However, due to information systems security requirements, we have now determined that we are prohibited from removing the requirement for a QualityNet Security Administrator at this time. We remind hospitals that are submitting their own data without the use of a vendor that the hospital must have at least one active QualityNet account with the appropriate role assigned in order to submit data. We note that those hospitals with QualityNet accounts (Security Administrator and non-Security Administrator) that are in danger of lapsing receive multiple e-mail notifications that contain reminders that they must sign in or the account will be deactivated.

After consideration of the public comments we received, due to systems requirements, we are not adopting our proposal to no longer require that a hospital maintain current designation of a QualityNet Administrator. Instead, hospitals must continue to maintain a QualityNet Security Administrator as part of the HOP QDRP requirements.

- Register with QualityNet, regardless of the method used for data submission.
- Complete and submit an online participation form if this form (or a paper Notice of Participation form) has not been previously completed, if a hospital has previously withdrawn, or if the hospital acquires a new CCN. For HOP QDRP decisions affecting the CY 2011 payment determination, hospitals that share the same CCN must complete a single online participation form. In the CY 2009 OP/ASC final rule with comment period (73 FR 68772), we implemented an online registration form and eliminated the paper form. At this time, the participation form for the HOP QDRP is separate from the RHQDAPU program and completing a form for each program is required. Agreeing to participate includes acknowledging that the data submitted to the CMS-designated contractor will be submitted to CMS and may also be shared with

one or more other CMS contractors that support the implementation of the HOP QDRP and be publicly reported.

Under our current requirements, the deadline for submitting the participation form is 30 days following receipt of a CCN form from CMS (73 FR 68772). In the CY 2010 OP/ASC proposed rule (74 FR 35400), we proposed to change this requirement as follows:

Hospitals with Medicare acceptance dates on or after January 1, 2010: For the CY 2011 payment update, we proposed that any hospital that has a Medicare acceptance date on or after January 1, 2010 (including a new hospital and hospitals that have merged) must submit a completed participation form no later than 180 days from the date identified as its Medicare acceptance date on the CMS Online System Certification and Reporting (OSCAR) system. Hospitals typically receive a package notifying them of their new CCN after they receive their Medicare acceptance date. The Medicare acceptance date is the earliest date that a hospital can receive Medicare payment for the services that it furnishes. Completing the participation form includes supplying the name and address of each hospital campus that shares the same CCN.

The use of the Medicare acceptance date as beginning the timeline for HOP QDRP participation will allow CMS to monitor more effectively hospital compliance with the requirement to complete a participation form because a hospital's Medicare acceptance date is readily available to CMS through its data systems. In addition, providing an extended time period to register for the program will allow newly functioning hospitals sufficient time to get their operations up and running before having to collect and submit quality data. We invited public comment on these proposed changes.

Hospitals with Medicare acceptance dates before January 1, 2010, that want to participate or withdraw: For the CY 2011 payment update, we proposed that any hospital that has a Medicare acceptance date on or before December 31, 2009 that wants to withdraw from participation in the CY 2011 HOP QDRP or that is not currently participating in the HOP QDRP and wishes to participate in the CY 2011 HOP QDRP must submit a participation form by March 31, 2010. We proposed a deadline of March 31, 2010, because we believe it will give hospitals sufficient time to decide whether they wish to participate in the HOP QDRP, as well as put into place the necessary staff and resources to timely report data for first

quarter CY 2010 services. This requirement applies to all hospitals whether or not the hospital has billed for payment under the OP/ASC. We invited public comment on these proposed changes.

Comment: Several commenters agreed with CMS' proposal to provide additional time for hospitals to submit an HOP QDRP participation form. One commenter believed that, for hospitals with Medicare acceptance dates prior to January 2010, a 3-month window ending March 31, 2010, is reasonable in which to make a decision regarding participation.

Response: We thank the commenters for their support.

Comment: One commenter stated that, based on its experience with hospital mergers, the surviving facility does not typically receive a new CCN and asked that the requirement that merged facilities submit a new participation form be confirmed.

Response: Annual payment update decisions are made for a hospital's CCN. If a hospital's CCN does not change in a merger situation and the hospital is currently participating in the HOP QDRP, the hospital with that CCN would continue to be subject to HOP QDRP requirements, so a new participation form would not be required. However, the participation form requests that hospitals submit National Provider Identifier (NPI) information for any facilities connected to the hospital that bill under the hospital's CCN. The hospital may want to update its participation form to include any facilities added due to a merger, but there is no HOP QDRP requirement to do so at this time. If the hospital's CCN did not change in a merger situation and it was not participating in the HOP QDRP and now wishes to do so, or was participating and now wishes to withdraw, it must comply with the March 31, 2010 timeframe for completing a participation form.

After consideration of the public comments we received, we are adopting as final our proposed administrative requirements with one exception. We are not adopting our proposal to no longer require that a hospital maintain current designation of a QualityNet Administrator. Instead, hospitals must continue to maintain a QualityNet Security Administrator as part of HOP QDRP requirements.

2. Data Collection and Submission Requirements

a. General Data Collection and Submission Requirements

In the CY 2010 OPSS/ASC proposed rule (74 FR 35401), we proposed that, to be eligible for the full CY 2011 OPSS payment update, hospitals must:

- **Submit data:** Hospitals that are participating in the HOP QDRP must submit data for each applicable quarter by the deadline posted on the QualityNet Web site; there must be no lapse in data submission. For the CY 2011 annual payment update, the applicable quarters will be as follows: 3rd quarter CY 2009, 4th quarter CY 2009, 1st quarter CY 2010, and 2nd quarter CY 2010. Hospitals that did not participate in the CY 2010 HOP QDRP, but would like to participate in the CY 2011 HOP QDRP, and that have a Medicare acceptance date on the OSCAR system before January 1, 2010, must begin data submission for 1st quarter CY 2010 services using the CY 2011 measure set that we are finalizing in this final rule with comment period. For those hospitals with Medicare acceptance dates on or after January 1, 2010, data submission must begin with the first full quarter following the submission of a completed online participation form. For the four claims-based measures, we will calculate the measures using the hospital's Medicare claims data. For the CY 2011 payment update, we will utilize paid Medicare fee-for-service (FFS) claims submitted prior to January 1, 2010, to calculate these four measures.

Sampling and Case Thresholds: It will not be necessary for a hospital to submit data for all eligible cases for some measures if sufficient eligible case thresholds are met. Instead, for those measures where a hospital has a sufficiently large number of cases, it can sample cases and submit data for these sampled cases rather than submitting data from all eligible cases. This sampling scheme which includes the minimum number of cases based upon case volume will be set out in the HOPD Specifications Manual at least 4 months in advance of the required data collection. Hospitals must meet the sampling requirements for required quality measures each reporting quarter.

In addition, in order to reduce the burden on hospitals that treat a low number of patients but otherwise meet the submission requirements for a particular quality measure, hospitals that have five or fewer claims (both Medicare and non-Medicare) for any measure included in a measure topic in a quarter will not be required to submit

patient level data for the entire measure topic for that quarter. Even if hospitals are not required to submit patient level data because they have five or fewer claims (both Medicare and non-Medicare) for any measure included in a measure topic in a quarter, they may voluntarily do so.

Hospitals must submit all required data according to the data submission schedule that will be available on the QualityNet Web site (<http://www.QualityNet.org>). This Web site meets or exceeds all current Health Insurance Portability and Accountability Act requirements. Submission deadlines will, in general, be four months after the last day of each calendar quarter. Thus, for example, the submission deadline for data for services furnished during the first quarter of CY 2010 (January–March 2010) will be on or around August 1, 2010. The actual submission deadlines will be posted on the <http://www.QualityNet.org> Web site.

Hospitals must submit data to the OPSS Clinical Warehouse using either the CMS Abstraction and Reporting Tool for Outpatient Department (CART-OPD) measures or the tool of a third-party vendor that meets the measure specification requirements for data transmission to QualityNet.

Hospitals must submit quality data through My QualityNet, the secure portion of the QualityNet Web site, to the OPSS Clinical Warehouse. The OPSS Clinical Warehouse, which is maintained by a CMS-designated contractor, will submit the OPSS Clinical Warehouse data to CMS. OPSS Clinical Warehouse data are not currently considered to be Quality Improvement Organization (QIO) data; rather, we consider such data to be CMS data. However, it is possible that the information in the OPSS Clinical Warehouse may at some point become QIO information. If this occurs, these data would also become protected under the stringent QIO confidentiality regulations in 42 CFR part 480.

Hospitals must collect HOP QDRP data from outpatient episodes of care to which the required measures apply. For the purposes of the HOP QDRP, an outpatient “episode of care” is defined as care provided to a patient who has not been admitted as an inpatient, but who is registered on the hospital's medical records as an outpatient and receives services (rather than supplies alone) directly from the hospital. Every effort will be made to ensure that data elements common to both inpatient and outpatient settings are defined consistently for purposes of quality reporting (such as “time of arrival”).

Hospitals are to submit required quality data using the CCN under which the care was furnished.

To be accepted into the OPSS Clinical Warehouse, data submissions, at a minimum, must be timely, complete, and accurate. Data submissions are considered to be “timely” when data are successfully accepted into the OPSS Clinical Warehouse on or before the reporting deadline. A “complete” submission is determined based on whether the data satisfy the sampling criteria that are published and maintained in the HOPD Specifications Manual, and must correspond to both the aggregate number of cases submitted by a hospital and the number of Medicare claims the hospital submits for payment. We are aware of “data lags” that occur due to when hospitals submit claims, then cancel and correct those claims; efforts will be made to take such events into account that can change the aggregate Medicare case counts. To be considered “accurate,” submissions must pass validation, if applicable.

CMS strongly recommends that hospitals review OPSS Clinical Warehouse feedback reports and the HOP QDRP Provider Participation Reports that are accessible through their QualityNet accounts. These reports enable hospitals to verify whether the data they or their vendor submitted was accepted into the OPSS Clinical Warehouse and the date/time that such acceptance occurred. We also note that irrespective of whether a hospital submits data to the OPSS Clinical Warehouse itself or uses a vendor to complete the submissions, the hospital is responsible for ensuring that HOP QDRP requirements are met.

Finally, although not required, hospitals may submit, on a voluntary basis, the aggregate numbers of outpatient episodes of care which are eligible for submission under the HOP QDRP and sample size counts. These aggregated numbers of outpatient episodes represent the number of outpatient episodes of care in the universe of all possible cases eligible for data reporting under the HOP QDRP. We do not wish to require this submission at this time because we continue to see evidence that some hospitals would not be able to meet this requirement. However, as it is vital for quality data reporting for hospitals to be able to determine their population sizes, we believe it is highly beneficial for hospitals to develop systems that can determine whether or not they have furnished services or billed for five or fewer cases for a particular measure topic on a quarterly basis. CMS strongly

recommends that all hospitals work to develop systems that can accurately determine their population and sample sizes for purposes of quality reporting.

In the future, we plan to use the aggregate population and sample size data to assess data submission completeness and adherence to sampling requirements for Medicare and non-Medicare patients.

For the reporting of aggregate numbers of outpatient episodes of care and sample size counts, we proposed that the deadlines for this reporting will be the same as they are for the reporting of quality measures, and these deadlines will be posted on the data submission schedule that will be available on the QualityNet Web site.

We invited public comment on these proposed changes.

Comment: One commenter appreciated CMS' clear and concise definition of an outpatient episode of care. Another commenter asked for a clear definition of what constitutes an outpatient setting.

Response: We thank the first commenter for its support for our definition of an outpatient episode of care. This definition is drawn from the CMS Claims Processing Manual, Chapter 1, Section 50.3.1 of the CMS Claims Processing Manual (issued 06-23-09; Effective Date: 07-01-09) states "Outpatient" means a person who has not been admitted as an inpatient but who is registered on the hospital or critical access hospital (CAH) records as an outpatient and receives services (rather than supplies alone) directly from the hospital or CAH." Thus, based upon the definition of outpatient, an outpatient setting would be a health care setting where a person is registered on the hospital or CAH records as an outpatient and receives services from the hospital or CAH. Under the HOP QDRP, hospitals are defined as including all the facilities connected to a hospital that are operating and billing for OPSS services under the same CCN. We note that the above definition does not restrict the outpatient setting to the hospital or CAH itself.

Comment: Several commenters stated that many of the program's processes were specified in detail for the first time and that this specificity was appreciated, as it is helpful for hospitals to have clear direction on both the requirements and the process of the program.

Response: We thank the commenters for their feedback.

Comment: One commenter supported CMS' proposal that all hospitals sharing the same CCN be required to combine data across multiple campuses for all

clinical data submissions. The commenter believed reporting by CCN is appropriate to align clinical and financial reporting. This commenter also supported this proposal because this approach is similar to the approach being taken by the RHQDAPU program and noted that consistency between administrative aspects of the two programs is appreciated. One commenter stated that while in situations where a new facility is opened or remains under the same ownership, the statement that hospitals are to submit required quality data using the CCN under which the care was furnished is true. However, the commenter believed that, for mergers, there will be a tremendous resource burden placed on the hospital and measure vendor if abstracted data must be separated according to the CCN that applied to the hospital at the time the care was furnished. This commenter stated that because data are submitted and published for public view on a quarterly basis, there is tremendous concern that the public could select a hospital for patient care based on data that does not represent a full quarter. The commenter recommended that if a merger does not occur at the very beginning/end of a quarter, the data be combined for both facilities under the parent/surviving facility CCN because the child or absorbed facility, under its old CCN, no longer exists once the merger occurs.

Response: We thank the commenters that supported our proposal to have quality measure data reported by the CCN that applies to the hospital at the time the care is furnished. We understand that there could be issues regarding burden and completeness of data reporting in the case of mergers. We point out that hospitals operating under separate CCNs participating in the HOP QDRP would be collecting quality measure data separately prior to a merger, and that the data could be kept separate after the merger. Regarding the issue of incomplete data, we acknowledge that the surviving hospital would have less data for the quarter in which the merger took place than if the absorbed hospital's data were included, but the surviving hospital will have all the data for care furnished under its CCN. Because the CCN is the financial and certification identifier for hospitals and is the identifier used by the HOP QDRP to monitor completeness of data reporting, we believe it is important that all quality measure data be reported under the CCN that was in place when the care was delivered. We urge hospitals and vendors to have data

reporting systems sufficiently robust to be able to efficiently handle data by the CCN under which the care was delivered.

Comment: One commenter strongly agreed that hospitals with five or fewer claims (Medicare and non-Medicare) for a specific measure should not be required to submit patient-level data for the entire measure topic while being allowed to report data voluntarily, but believed that the allowable time period to not report data was a year. Another commenter urged CMS to modify this provision so that it would apply to hospitals with five or fewer Medicare claims, not five or fewer claims across all payers.

Response: We thank the first commenter for supporting our policy not to require hospitals with five or fewer claims for a specific measure to submit data while allowing these hospitals to report data voluntarily. However, we are clarifying that hospitals that have five or fewer claims (both Medicare and non-Medicare) for any measure included in a measure topic in a quarter will not be required to submit patient-level data for the entire measure topic only for that quarter.

With respect to the second commenter's suggestion that we modify our policy to apply to five or fewer Medicare claims (rather than five or fewer Medicare and non-Medicare claims), we selected more than 5 cases per quarter (more than 20 cases per year) as the minimum threshold to ensure that the vast majority of hospitals with sufficient caseload would be required to submit data, while easing the burden on hospitals whose patient counts were too small to reliably predict hospital performance. Because we collect quality measure data on both Medicare and non-Medicare patients, we believe it is appropriate to set our case thresholds using the population for which we are collecting data, which includes both Medicare and non-Medicare patients.

Comment: One commenter questioned the proposal under which hospitals that have five or fewer claims (both Medicare and non-Medicare) for any measure included in a measure topic in a quarter will not be required to submit patient-level data for the entire measure topic for that quarter. The commenter stated that one of the goals of the CMS quality improvement programs is to improve the care given to Medicare beneficiaries and that, by allowing hospitals with five or fewer cases in a quarter to not report, the very hospitals that need improvement the most may be missed. The commenter also believed

that the burden of reporting is minor for hospitals that would abstract five or fewer charts as this would take less than 30 minutes in most cases, which would not be much of a burden over a 3-month period.

Response: We agree with the commenter that abstracting five charts over a quarter would not be overly burdensome. In implementing the reporting of quality measure data where there are five or fewer cases in a quarter, we are also addressing the reported excessive burden associated with determining the individual cases and submitting the data for those hospitals with small case numbers for a measure. We continue to strive to collect quality of care data while limiting burden. We acknowledge that it is possible that quality of care concerns may exist with hospitals with small case numbers for a particular quality measure. We note that quality of care is also monitored through other mechanisms, including, but not limited to, the Medicare beneficiary complaint process through QIOs and the survey and certification process.

Comment: One commenter agreed that data abstraction processes for outpatient services are sufficiently different from inpatient services as to require the hospital to spend time creating processes to ensure that they capture the accurate population and abstracted data. One commenter agreed with keeping the submission deadline for population and sampling counts the same as the deadline for case-level data, and stated that last minute updates to the population occur because of coding changes. The commenter also stated that synchronizing the deadlines aided hospitals in capturing the most accurate information possible. One commenter agreed with CMS' proposal to keep the submission of population and sampling counts voluntary, and noted that this can be very time consuming for hospitals.

Response: We thank these commenters for confirming our views on the ability of some hospitals to meet a requirement to submit population and sampling data at this time. We reiterate that it is vital, for quality data reporting, that hospitals are able to determine their population sizes and that all hospitals work to develop systems that can accurately determine their population and sample sizes for purposes of quality reporting.

Comment: One commenter urged immediate adoption of an effective mechanism that allows hospitals and their vendors to resubmit quality measure data if an error is discovered and emphasized that the point of public reporting is to put accurate and useful

information into the hands of the public and this is facilitated by allowing known reporting mistakes to be corrected.

Response: We agree with the commenter that publicly reporting accurate and useful information is important and that a mechanism or process for correcting errors should be implemented. While a proposal addressing this concern was not included in this current rulemaking, we will consider ways to address this concern in a future rulemaking.

Comment: One commenter disagreed with CMS' approach that sampling requirements should apply based on both Medicare and non-Medicare cases and believed that CMS should focus only on the population of patients for which the agency is responsible.

Response: We believe the commenter is arguing that we should not apply sampling criteria to non-Medicare claims and should focus only on Medicare claims. As we collect quality measure data on both Medicare and non-Medicare patients, we believe it is appropriate to set our sampling criteria so that they apply to the same population, which includes both Medicare and non-Medicare patients.

Comment: One commenter supported the concept of sampling and the development of eligible thresholds and recommended that CMS distribute sampling criteria when new measures are implemented (that is, any measures proposed in a calendar year should include the sampling criteria as part of the OPPI/ASC proposed rule).

Response: We thank the commenter for this suggestion and will consider it in future planning. We note that sampling criteria are included in each release of the HOPD Specifications Manual.

After consideration of the public comments we received, we are adopting as final our proposals regarding HOP QDRP data collection and submission requirements for the CY 2011 payment determination.

b. Extraordinary Circumstance Extension or Waiver for Reporting Quality Data

In our experience, there have been times when hospitals have been unable to submit required quality data due to extraordinary circumstances that are not within their control. It is our goal to not penalize hospitals for such circumstances and we do not want to unduly increase their burden during these times. Therefore, in the CY 2010 OPPI/ASC proposed rule (74 FR 35402), we proposed a process for hospitals to request and for CMS to grant extensions

or waivers with respect to the reporting of required quality data when there are extraordinary circumstances beyond the control of the hospital.

Under the proposed process, in the event of extraordinary circumstances not within the control of the hospital, for the hospital to receive consideration for an extension or waiver of the requirement to submit quality data for one or more quarters, a hospital must—

(1) Submit to CMS a request form that will be made available on the QualityNet Web site. The following information should be noted on the form:

- Hospital CCN;
- Hospital Name;
- CEO and any other designated personnel contact information, including name, e-mail address, telephone number, and mailing address (must include a physical address, a post office box address is not acceptable);
- Identified reason for requesting an extension or waiver;
- Hospital's reason for requesting an extension or waiver;
- Evidence of the impact of the extraordinary circumstances, including but not limited to photographs, newspaper and other media articles; and
- A date when the hospital will again be able to submit HOP QDRP data, and a justification for the proposed date.

The request form must be signed by the hospital's CEO. A request form must be submitted within 30 days of the date that the extraordinary circumstance occurred.

Following receipt of such a request, CMS will—

(1) Provide a written acknowledgement using the contact information provided in the request, to the CEO and any additional designated hospital personnel, notifying them that the hospital's request has been received; and

(2) Provide a formal response to the CEO and any additional designated hospital personnel using the contact information provided in the request notifying them of our decision.

We invited public comment on these proposed procedures for requesting an extraordinary circumstance extension or waiver of the requirement to submit quality data for one or more quarters.

Comment: Several commenters expressed their appreciation for CMS' recognition that hospitals facing certain extraordinary circumstances should be granted an extension or waiver. The commenters believed that, while decisions on granting an extension or waiver would best be made on a case-by-case basis depending on each hospital's unique situation, they

suggested that CMS adopt some general criteria to apply when it determines whether such extensions or waivers would be granted. The commenters also expressed concern that it might not be feasible for a hospital to file a request form for an extraordinary circumstances waiver within 30 days of such an event and urged a creative and flexible approach to working with hospitals in these situations to ensure that an undue burden is not placed on hospitals during a time of hardship.

Response: We will consider these comments as we further develop program procedures for extraordinary circumstance extensions or waivers. We are mindful that many hospitals operating in these adverse situations cannot access the Internet or mail service. We note that we currently use a variety of means to communicate with hospitals in these circumstances, including using our HOP QDRP support contractor and both national and State hospital associations, and will continue to do so. Regarding the ability to file a request form for an extraordinary circumstances waiver within 30 days of such an event, we believe that 30 days is sufficient in the vast majority of circumstances. However, we agree that additional time may be warranted in some extreme circumstances.

After consideration of the public comments we received, we are adopting our proposals regarding extraordinary circumstance extensions or waivers for the reporting of quality data under the HOP QDRP, with a modification that the request form must be submitted within 45 calendar days of the date that the extraordinary circumstance occurred, rather than the 30 days we proposed.

3. HOP QDRP Validation Requirements

In the CY 2009 OPSS/ASC final rule with comment period (73 FR 68776), we announced a voluntary test validation program, the results of which would not affect the CY 2010 payment update for any hospital. Due to resource constraints, we were not able to implement this test validation plan.

a. Data Validation Requirements for CY 2011

Validation, as discussed in the CY 2008 OPSS/ASC final rule with comment period (72 FR 66871), is intended to provide assurance of the accuracy of the hospital abstracted data. For the CY 2011 payment determination, in the CY 2010 OPSS/ASC proposed rule (74 FR 35402), we proposed to implement a validation program that will require hospitals to supply requested medical documentation to a CMS contractor for purposes of being validated. However,

the results of the validation will not affect the CY 2011 payment update for any hospital. We believe that it is important for hospitals to have some experience and knowledge of the HOP QDRP validation process before payment determinations are made based upon validation results. We proposed to implement a validation program that will both limit burden upon hospitals, especially small hospitals, as well as provide feedback to all hospitals on validation performance.

Specifically, we proposed to select a random sample of 7,300 cases from all cases successfully submitted to the OPSS Clinical Warehouse by all participating hospitals for the relevant time period described below and validate those data. Based upon the quality data submitted for the CY 2009 payment update and our methodology for drawing the sample, we estimate that the sample will include up to 20 cases per participating hospital; the same number of cases sampled on an annual basis for validation under the RHQDAPU program. A sample size of 7,300 was chosen because it will enable us to detect a relative difference of 10 percent in the measured overall accuracy rate with a 95 percent (two-tailed) confidence interval and should provide sufficient data to conduct post-hoc stratified analyses that provide meaningful feedback. These figures are based upon a power analysis assuming a population measure mismatch rate of five percent with the outcomes being either a match or a mismatch between what the hospital submitted versus what was determined by validation. We intend to supply feedback on the validation results to all hospitals.

We proposed to request medical documentation from hospitals for April 1, 2009 through March 31, 2010 episodes of care, which will allow us to gather one full year of submitted data for validation purposes.

Once we have completed the random selection, a designated CMS contractor will use certified mail to request that each selected hospital send to the CMS contractor supporting medical record documentation that corresponds to each selected episode of care. Each hospital must submit this documentation to the designated CMS contractor within 45 calendar days of the date of the request (as documented on the request letter). If the hospital fails to comply within 30 days of the initial medical documentation request, the designated CMS contractor will send a second certified letter to the hospital reminding the hospital that the requested documentation must be received within 45 calendar days following the date of

the initial request. If the hospital still fails to comply, a "zero" score will be assigned to each data element for each selected case and the case will fail for all measures in the same topic (for example, OP-6 and OP-7 measures for a surgical care case).

Once the CMS contractor receives the requested medical documentation, the CMS contractor will independently reabstract the same quality measure data elements that the hospital previously abstracted and submitted and compare the two sets of data to determine whether they match. Specifically, the CMS contractor will conduct a measures level validation by calculating each measure within a submitted record using the independently reabstracted data and then comparing this to the measure reported by the hospital; a percent agreement will then be calculated.

As we stated above, the results of the validation will not affect a hospital's CY 2011 annual payment update because we want to give hospitals time to gain experience with the medical documentation requests and the validation process before these results are used in payment determinations. However, hospitals must supply the medical documentation for each requested case; failure to provide this documentation may result in a 2.0 percentage point reduction in a hospital's CY 2011 annual payment update.

Comment: Many commenters expressed their support for a validation approach where they could receive feedback on their validation results without having their payment affected for the CY 2011 payment update. Several commenters applauded the approach, stating that hospitals will benefit from the program as they continue to gain experience with outpatient quality reporting. One of these commenters specifically agreed with requiring hospitals to submit charts for validation for the CY 2011 payment determination. One commenter expressed support for a 12-month validation period in CY 2011. One commenter expressed support for a different validation process than the one in use for FY 2010 under the inpatient RHQDAPU program.

Response: We thank these commenters and appreciate their support. We agree that it is important that hospitals gain experience with validation for HOP QDRP data collection prior to using validation results to make payment determinations. We also believe that hospitals will benefit from the results of our proposed validation plan, both by

reviewing results on selected individual cases as well as the aggregate results.

Comment: One commenter stated that hospitals must incur the additional burden of paying to copy and ship medical records to CMS because this funding was not incorporated into the outpatient project.

Response: The issue of costs for copying and shipping medical records to CMS was not discussed in the CY 2010 OPPS/ASC proposed rule, and we are currently considering our policy approach regarding this issue. We are aware that these costs are not insignificant.

Comment: One commenter believed that it is unrealistic to conduct validation and calculate reliable measures from collected HOP QDRP quality measure data because there have been so many changes in the abstraction instructions over the past year. The commenter argued that validation should not be part of the determination for payment decisions until the entire measure set is tested and proven to be reliable and valid. Another commenter stated that the two statements “the results of the validation will not affect a hospital’s CY 2011 annual payment update” and “failure to provide this documentation may result in a 2.0 percentage point reduction in the hospital’s CY 2011 payment update” were inconsistent.

Response: We agree that there have been issues with the HOP QDRP data, including changes in abstraction instructions over the past year. However, we believe that the validation approach that we have proposed addresses these concerns. For CY 2011 payment determinations, we have proposed to conduct validation on April 1, 2009 to March 31, 2010 episodes of care reported to the OPPS Clinical Warehouse; this is one year after the initial data reported, which was for April 1, 2008 to June 30, 2008 episodes of care. With one exception, most of the significant changes in abstraction instructions during the program’s life were incorporated by April 1, 2009. The exception, the exclusion of cancelled surgery cases from the Surgical measures, went into effect with July 1, 2009 episodes of care. We intend to determine whether a selected Surgical Care measure case was cancelled based upon the submitted medical documentation rather than drop April 1, 2009 to June 30, 2009 cases from selection in order to maintain a 12-month sampling frame for validation. We intend to either drop any selected April to June 2009 cancelled surgery cases or otherwise account for this factor.

While it would be optimal to use the preliminary results of the validation effort for CY 2011 in the final design of the validation process for the CY 2012 payment determinations, due to resource constraints, we will not obtain the results of the CY 2011 validation before we must put forth our proposal for CY 2012. We do intend to conduct further analyses of collected HOP QDRP data and will utilize these results in developing our CY 2012 proposals.

Regarding our statement that the results of this validation will not affect a hospital’s annual payment update, we wish to clarify that when we referred to “results,” we meant the results of the validation process where what is independently abstracted from a hospital’s submitted documentation is compared to what the hospital self-reported. Therefore, while the validation “results” will not affect a hospital’s CY 2011 payment update factor, to ensure that we receive an adequate supply of records for our test validation, a hospital must submit required medical record documentation for the selected cases and if it fails to do so, that failure would affect its CY 2011 payment update factor.

Comment: Several commenters supported CMS’ proposal to document the validation medical record request process. Specifically, the commenters supported CMS’ plans to send two certified letter requests for medical records for data validation in case the hospital does not respond to the first request. The commenters suggested that phone calls be placed to hospitals that do not respond to the first letter to ensure that every effort is made to communicate the request to the appropriate staff; and suggested that this phone call should be placed to the QualityNet Administrator for those hospitals that have a person serving in this role. One commenter suggested that a telephone call to the hospital chief executive officer be made before assigning a “zero” score for validation, as there may be circumstances in which the CEO is unaware of an insufficient response to a request for records.

Response: We thank the commenters and agree that certified letters provide hospitals with multiple written documented notification and reminder attempts; we believe that this alone is sufficient notification. However, we note that the planned contractor for this work as standard practice attempts to call hospitals at least three times about 30 calendar days after it sends the initial medical record request. As a practice, we intend to continue attempting to call hospitals at least three times around the 30th calendar day following the initial

request, in addition to sending written certified letters. We believe that these attempted calls at different time periods around the 30th calendar day following the initial request demonstrate our commitment to notify hospitals of medical record requests using multiple communication modes.

Comment: One commenter expressed concern that a random sampling of 7,300 cases is not sufficient to provide hospitals meaningful feedback on the accuracy of their data collection and recommended that CMS select a larger number of cases over a longer period of time. The commenter also suggested that individual feedback on validation results be provided to those hospitals that submit records for this initial validation process.

Response: The major purpose of our proposed sampling scheme for CY 2011 payment determinations is to provide aggregate level feedback to hospitals on data elements abstracted and to validate the quality measure data collected while limiting hospital burden. However, we do intend to provide individual feedback on validation results to those hospitals that submit records for this initial validation effort. Regarding the suggestion to select cases over a longer time period, we have selected the April 1, 2009 to March 31, 2010 timeframe because this time period provides the most recent 12-month period possible. We believe that it is important to use the most recent data possible while maintaining at least a full year’s worth of data. We have not included any data from the April 1, 2008 to March 31, 2009 time period because we believe that this will minimize the use for validation purposes of HOP QDRP data that may be unreliable because of changes in data abstraction guidelines that occurred, or because it was collected in the early stages of the program when hospitals were still developing HOP QDRP data collection systems. We believe that hospitals will be able to gain meaningful information from aggregate results produced under our validation sampling scheme, although we agree that it would be more useful to select an increased number of cases. Selecting an increased number of cases is not possible with present resource constraints and we note that this approach would increase hospital burden.

Comment: Several commenters disagreed with randomly selecting only a subset of hospitals for validation because they believe that all hospitals should be held to the same threshold/expectation. One commenter believed that “random validation” would not produce accurate results, and another

commenter expressed concern that the proposed validation approach is not robust enough given the ever increasing scope of measures included in the measure set.

Response: We appreciate the concern that these commenters express for treating all hospitals similarly, both in costs and benefits of the validation process and for maintaining hospital performance in regard to data validation.

Under the HOP QDRP, all hospitals are responsible for meeting reliability levels for submitted abstracted data. Because hospitals will not know in advance whether they will be selected for validation and because selection will be random, we believe that hospitals will have sufficient incentive to maintain data quality.

Comment: Several commenters suggested that CMS build safeguards into the sampling process to ensure that no more than 20 patient cases are selected from each hospital.

Response: We appreciate the commenters' concerns about the number of cases selected under the proposed sampling design. However, it is highly unlikely, given the number of hospitals and the cases submitted, that any hospital will have more than 20 cases selected. In addition, building in a threshold for the number of cases selected would take away the "random" element of the sampling process.

Comment: One commenter suggested that rather than randomly selecting hospitals each year, CMS adopt a validation process that results in those hospitals not selected in one year having a greater likelihood of being selected in the next/future years.

Response: While we did not propose this approach in our validation plan for CY 2011, in the CY 2010 OPPTS/ASC proposed rule (74 FR 35403 through 35404), we discussed additional data validation conditions under consideration for CY 2012 and subsequent years, including the use of targeting criteria. Examples of possible targeting criteria include considering whether a hospital had not been previously selected for validation for 2 or more consecutive years.

Comment: One commenter urged CMS to set strict timelines so that the public has access to data that have undergone the test validation process as soon as possible by publicly reporting validated 2nd and 3rd quarter 2009 data no later than June 2010.

Response: We agree that it is important to make validated data publicly available as soon as possible and will make every effort to do so. However, with present resource

constraints, it is not possible to meet the commenter's suggested timeline.

Comment: One commenter urged that State QIOs be supportive not only during the validation appeals process, but proactively during data collection and reporting.

Response: The HOP QDRP was implemented separately from the QIO program and State QIOs have not been involved with the HOP QDRP to date. State QIOs, however, are involved in the RHQDAPU program. We note that QIOs are available for quality of care concerns for individual Medicare cases and that their purview includes the outpatient setting.

Comment: One commenter preferred a validation approach that would select five cases per quarter stratified by measure/topic for all hospitals. The commenter argued that such an approach provides hospitals an opportunity to focus on mismatches by measure set.

Response: We interpret the commenter's statement to mean that the commenter prefers to have five cases that are stratified by measure/topic be selected each quarter for all hospitals so that information on each hospital's individual abstraction accuracy can be assessed by measure/topic. We acknowledge the commenter's belief that such an approach would continually improve data abstraction, but we believe that the improved reliability under the proposed validation process coupled with the reduction of overall hospital burden associated with validation participation will outweigh the potential benefits of validating a smaller number of records for all hospitals. Regardless of whether a hospital was included in any validation selection, we intend to provide aggregate validation information to all hospitals that submit quality measure data.

Comment: Several commenters preferred that CMS give continued attention to individual data element level analysis for validation for a variety of reasons, including: Increasing the denominator and minimizing the impact of a small number of errors; and looking at the individual data elements with a threshold based on sample size. Some of these commenters doubted CMS' statement that higher accuracy rates are possible using the proposed measure level match approach versus a data element level approach and believed that the proposed approach appeared to place hospitals at high risk for not receiving the full CY 2012 payment update. The commenters recommended a period or process where any changes in the validation process be tested

without penalty against any payment update prior to broad implementation.

Response: We agree that there should be continued attention to the data element level as the individual data elements are used to calculate quality measures. We proposed a measure level match approach to replace the data element match approach because of what we have observed in the RHQDAPU program for inpatient quality measure data reporting. The intent of the measure match approach is to minimize the impact of errors for noncritical data elements. As we explain in more detail below, we are not finalizing our validation proposal for CY 2012. Instead, we will be conducting further analyses on collected HOP QDRP data and considering all public comments we received on validation before we propose a validation process for the CY 2012 payment determination.

We agree that hospitals should be allowed to gain some experience with validation under the HOP QDRP before we consider such results toward payment determinations, and we are doing so through our validation approach for the CY 2011 payment determination.

Comment: Many commenters agreed with the adoption of a measure score match for validation instead of a data element matching approach. Several commenters believed that it is appropriate to focus on the hospital's measure rate, as opposed to individual data elements, because the measure rate captures the information that is truly important to patient care. The commenters observed that, for data validation in the RHQDAPU (inpatient reporting) program, there have been several instances in which a mismatch between single data elements unrelated to the quality of care provided by a hospital, such as the patient's birth date, have caused hospitals to fail validation and that validating the hospital's measure rate should eliminate these unfortunate incidents.

Response: We thank these commenters for their support. The proposed validation process focuses on validating whether hospital abstracted data results in accurate measure rates and measure denominator counts. We will continue to use the data elements to calculate the measure values and, subsequently, the validation scores.

Comment: One commenter urged CMS to extend the turnaround time for chart selection to 60 days and believed that hospitals should have the option to submit validation cases electronically rather than printed copies because this would avoid shipping delays and allow faster turnaround time.

Response: We understand the commenter's concern about the deadline for hospitals to submit requested medical records. However, extending this time period increases the amount of time between when services are furnished, initial hospital data are submitted, data are validated, and, ultimately, when the results can be compiled for program purposes. We will consider accepting electronic submission of validation cases using compact disc and electronic health record submission in future years. We must consider both the cost to accept and review these submissions and the added benefit to the hospitals using electronic methods to store medical record information.

After consideration of the public comments we received, we are adopting as final, without modification, our proposals for validation for the CY 2011 payment determination.

b. Data Validation Approach for CY 2012 and Subsequent Years

Similar to our proposal for the FY 2012 RHQDAPU program (74 FR 24178), in the CY 2010 OPSS/ASC proposed rule (74 FR 35403), we proposed to validate data from 800 randomly selected hospitals (approximately 20 percent of all participating HOP QDRP hospitals) each year, beginning with the CY 2012 payment determination. We note that because the 800 hospitals will be selected randomly, every HOP QDRP-participating hospital will be eligible each year for validation selection. For each selected hospital, we proposed to randomly validate per year up to 48 patient episodes of care (12 per quarter) from the total number of cases that the hospital successfully submitted to the OPSS Clinical Warehouse. However, if a selected hospital has submitted less than 12 cases in one or more quarters, only those cases available will be validated. For each selected episode of care, a designated CMS contractor will request that the hospital submit the supporting medical record documentation that corresponds to the episode. We will not be selecting cases stratified by measure or topic; our interest is whether the data submitted by hospitals accurately reflect the care delivered and documented in the medical record, not what the accuracy is by measure or whether there are differences by measure or topic. We proposed to sample data for April 1, 2010 to March 31, 2011 services because this will provide a full year of the most recent data possible to use for purposes of completing the validation in time to make the CY 2012 payment determination.

For the CY 2012 and subsequent years' payment determinations, we proposed to use the validation methodology proposed for the CY 2011 payment update with validation being done for each selected hospital. Specifically, we would conduct a measures level validation by calculating each measure within a submitted record using the independently reabstracted data and then comparing this to the measure reported by the hospital; a percent agreement will then be calculated.

To receive the full OPSS payment update, we proposed that hospitals must attain at least a 90 percent reliability score, based upon our validation process, for the designated time period. We will use the lower bound of a two-tailed 95 percent confidence interval to estimate the validation score. If the calculated upper limit is above the required 90 percent reliability threshold, we will consider a hospital's data to be "validated" for payment purposes. We believe that hospitals will be able to attain higher accuracy rates based on the proposed measure level match approach versus a data element level approach; therefore, we proposed to implement a higher threshold for accuracy than we currently use (and are using) for validation purposes under the RHQDAPU program. We believe that a hospital will be able to achieve a higher accuracy rate under this validation process because we are not calculating whether each data element matches. Instead, we are determining whether or not the reabstracted measure result (for example, was aspirin given at arrival as part of an episode of care that was properly included in the reported data) matches the measure result that was submitted by the hospital. In other words, we are more interested in whether the measure as a whole has been accurately reported than we are in whether each data element that makes up the measure has been accurately reported. Thus, we are focusing on whether the quality measure as a whole that a hospital reports matches what is in the medical record as determined by our re-abstractation. The reason we proposed to implement a measure level match for the HOP QDRP, rather than a data element match, is that in our experience with the RHQDAPU program, hospitals sometimes receive low validation scores due to data element mismatching and not because the care furnished did not match what was documented in the medical record.

We believe that validating a larger number of cases per hospital, but only for 800 randomly selected hospitals, and validating these cases at the measure

level (rather than at the data element level) has several benefits. We believe that this approach is suitable for the HOP QDRP because it will: produce a more reliable estimate of whether a hospital's submitted data have been abstracted accurately; provide more statistically reliable estimates of the quality of care delivered in each selected hospital as well as at a national level; and reduce overall hospital burden because most hospitals will not be selected to undergo validation each year.

We solicited public comments on this proposed validation methodology. The public comments we received and our responses are discussed below.

Comment: Several commenters stated that CMS' proposed process for validating hospitals' quality data beginning with CY 2012 holds promise as a reasonable approach to ensure the accuracy of the quality data and improves upon the deficiencies in the current inpatient program validation process.

Response: We agree with the commenters that the proposed new validation process beginning with CY 2012 is an improved approach, and we thank these commenters for their support. However, we have decided that we want to further evaluate and refine the approach and have decided to not finalize a validation approach for CY 2012 in this final rule with comment period. We intend to put forth a proposal for a CY 2012 HOP QDRP validation process in the CY 2011 OPSS/ASC proposed rule.

Comment: One commenter supported the proposed validation process for CY 2012 and subsequent years; however, this commenter believed that the data from the CY 2011 test year should be reviewed to ensure the process is functioning as it was intended and that CMS should make modifications to the process if necessary.

Response: We thank the commenter for supporting our proposed validation process. We agree that it would be helpful to review the data from the CY 2011 test year to evaluate the extent to which the process is functioning as it was intended and to use the results to assist us in determining whether to propose modifications to the validation process for CY 2012 and subsequent years. However, due to resource constraints, we will not receive the results of the CY 2011 validation before we must put forth a proposed validation plan for CY 2012. Instead, we will be conducting further analyses on collected HOP QDRP data and intend to utilize these results as well as all comments we

received in developing our CY 2012 proposals.

Comment: Several commenters recommended that if validation is limited to randomly selected hospitals each year, a hospital that is selected in one year should be excluded from the selection process for some period, which could be an indeterminate number of subsequent years, the following year, or the next 2 years, or, alternatively, CMS could limit the number of times during a 5-year period a hospital could be randomly selected. Many of these commenters suggested that such an exemption could apply to all hospitals that pass validation, those that pass with high reliability scores, or all hospitals regardless if they pass. Some of these commenters based their suggestions on limiting burden to hospitals and/or rewarding hospitals for participation or for achieving a high reliability or accuracy score. Some commenters believed that exempting a hospital from subsequent validation for some time period for high reliability scores would encourage hospitals to ensure that their data are as accurate as possible and would increase data quality.

Response: We understand the commenters' desire to limit hospital burden on a guaranteed basis and/or to reward hospitals for performance. However, we do not agree that exempting hospitals from validation because they were selected in a previous year or achieved a high reliability score will encourage increased data quality or that it should be a "reward" for meeting a program requirement. It is our belief that any guaranteed exclusion from participating in the validation process also has the potential to undermine a hospital's incentive to maintain data quality.

Comment: One commenter asserted that random selection of hospitals for validation will reduce the hospitals' focus on accuracy because hospitals will have the chance of not being chosen in a given year.

Response: We understand and appreciate the commenter's concern for data accuracy. We believe that each hospital having a chance at selection for validation each year will provide incentive to hospitals to maintain data quality.

Comment: Several commenters made suggestions regarding the sampling scheme for validation. One commenter suggested a random selection of fewer hospitals while increasing the number of records selected and that a random selection of hospitals be done quarterly to reduce the demands on any one hospital while increasing CMS' ability

to monitor performance throughout the industry. One commenter supported a random sample of 200 hospitals per quarter with a minimum number of 20 charts reviewed with hospitals not to be selected for validation any more frequently than one time per year.

Another commenter agreed with having a separate random selection process for small and rural hospitals that have five or less cases per condition each quarter.

Response: We chose 800 as the number of hospitals we would select for validation each year because this comprises about 25 percent of the total number of HOP QDRP participating hospitals and will provide us with enough data to be statistically assured that we have obtained a representative sample of all hospital data. Regarding randomly sampling hospitals quarterly, we have increased the sample size to gain increased statistical reliability for individual hospital data; lowering the number of cases per hospital or sampling different hospitals each quarter would not enable us to achieve the same result. We agree with the commenters that stratifying sampling by quarter is a possible approach and will consider this as we develop our proposal for a validation process for the CY 2012 payment determination.

Regarding having a separate random selection process for small and rural hospitals that have five or less cases per condition each quarter, we did not propose to stratify by hospital size or by a threshold number of cases per condition. However, we will strongly consider this approach when we develop our CY 2012 validation proposal.

Comment: One commenter supported the random sampling of 800 hospitals (which would require selected hospitals to submit supporting medical records for 48 randomly selected patient episodes of care (12 per quarter)) if the sampling methodology to select the 800 hospitals considers the proportion of rural to urban hospitals. This commenter also believed that this sample must take into account the large number of hospitals that have sample sizes that are too small to make justified decisions based on gathered data. One commenter argued that random validation on a larger number of cases per hospital is excessive for small PPS hospitals.

Response: We agree that hospital size and urban/rural status are important considerations regarding burden and representation of hospital type in any sampling scheme utilized for validation and view these as possible characteristics to stratify sampling of hospitals for CY 2012 and subsequent

year's validation. We intend to consider these factors as we further evaluate any proposed validation methodology for CY 2012. Regarding the commenter's belief that this sample must take into account the large number of hospitals that have sample sizes that are too small to make justified decisions based on gathered data, we interpret this statement to mean that the commenter believes these hospitals would not have a sufficient number of cases for us to reliably determine that the hospitals have submitted valid data. We will further assess the numbers of cases submitted by hospitals and, as discussed here in this final rule with comment period, we will be considering whether we should refine our proposed validation methodology to take into account hospital size or a threshold number of case counts in any proposed sampling scheme.

Comment: Several commenters argued that the 90 percent reliability proposal is too stringent for the first year of data validation. Many of these commenters suggested that CMS establish a lower reliability threshold (for example, 70 percent, 75 percent, or 80 percent). Commenters suggesting a 75 percent reliability threshold for the HOP QDRP noted that a 75 percent threshold will be used in the RHQDAPU program for FY 2012 when that program adopts a similar validation approach. One commenter recommended that CMS use data and experience from the CY 2011 test year to determine what an appropriate threshold should be, and until that is determined, the threshold should be the same as the 75 percent threshold that will be required in the inpatient setting for FY 2012. One commenter believed that more analysis of validation results is necessary before establishing the threshold at 90 percent.

Response: After consideration of the public comments regarding the 90-percent threshold, we agree with these commenters that this level is too stringent for the first year of validation where the results may affect a hospital's annual payment update. We appreciate the suggestion that the experience from the CY 2011 test year should be utilized to determine an appropriate rate. However, due to resource constraints, we will not be able to determine any results of the CY 2011 validation prior to proposing and finalizing validation requirements for the CY 2012 payment update factor. However, we will be analyzing collected HOP QDRP data and will take any analyses we complete, as well as the public comments we have received on this proposal, into account as we develop a new proposed

validation process for CY 2012 and subsequent years.

Comment: One commenter suggested that no hospital be penalized in terms of its payment update if it fails the validation requirement for a single quarter.

Response: We interpret this comment to apply to the proposed validation methodology for CY 2012 and subsequent years because the results of the proposed CY 2011 validation methodology will not affect the CY 2011 payment determination. We did not address whether a hospital would be penalized in terms of its payment update if it fails the validation requirement for a single quarter in the CY 2010 OPPS/ASC proposed rule with comment period. We will take the commenter's suggestion regarding this aspect of validation into consideration as we develop our new proposal for validation for CY 2012 and subsequent years.

We appreciate all the public comments we received regarding the validation process we proposed for CY 2012 and subsequent years and will take them into account as we develop our validation proposals for these years.

c. Additional Data Validation Conditions Under Consideration for CY 2012 and Subsequent Years

In the CY 2010 OPPS/ASC proposed rule (74 FR 35403), we stated that we are considering building upon what we proposed as a validation approach for CY 2012 and subsequent years. We are considering, in addition to selecting a random sample of hospitals for validation purposes, selecting targeted hospitals based on criteria designed to measure whether the data they have reported raises a concern regarding data accuracy. Because little data have been collected under the HOP QDRP at this point, we are considering this approach for possible use beginning with the CY 2012 payment determination. Examples of targeting criteria could include:

- Abnormal data patterns identified such as consistently high HOP QDRP measure denominator exclusion rates resulting in unexpectedly low denominator counts;
- Whether a hospital had previously failed validation; and/or
- Whether a hospital had not been previously selected for validation for 2 or more consecutive years.

Another example of a possible targeting criterion would involve some combination of the some or all of the criteria discussed above.

We solicited public comments on whether these criteria, or another approach, should be applied in future

years. We especially solicited suggestions for additional criteria that could be used to target hospitals for validation.

Comment: One commenter opposed the targeting criteria that might supplement random validation as the commenter opposed random validation and instead preferred that all hospitals have cases selected for validation.

Response: We appreciate the commenter's desire for us to validate data from all hospitals. However, we believe that the increased reliability that will be achieved by increasing the number of cases that we validate per selected hospital, coupled with the overall reduction of burden on hospitals that our proposed validation methodology will achieve, outweighs any potential benefit from requiring all HOP QDRP participating hospitals to undergo validation each year of a small number of cases. We chose the sample size of 800 hospitals because this comprises about 25 percent of the total number of HOP QDRP participating hospitals and will provide us with sufficient data to be statistically assured that we have obtained a representative sample of all hospital data.

Comment: One commenter recommended that CMS use similar statistical processes to those used by the Joint Commission to identify outliers in scoring, as well as low denominators compared to population sizes, as these are processes that many hospitals already know.

Response: We thank the commenter for its suggestions, and we will take them into account for our validation proposals for CY 2012.

We greatly appreciate all the public comments we received regarding the validation process proposed for CY 2012. However, as we stated above, we are not finalizing a validation process for CY 2012 at this time. We will take all of the comments we received into account when we develop our validation proposals for CY 2012.

F. 2010 Publication of HOP QDRP Data

In the CY 2009 OPPS/ASC final rule with comment period, we stated our intention to make the information collected under the HOP QDRP available to the public in 2010 (74 FR 68778). In the CY 2008 OPPS/ASC final rule with comment period, we stated that "[i]nformation from non-validated data, including the initial reporting period (April–June 2008) will not be posted" (72 FR 66874). However, section 1833(t)(17)(E) of the Act requires that the Secretary establish procedures to make data collected under the HOP QDRP available to the public, and does

not require that such data be validated before it is made public. Moreover, under existing procedures for the RHQDAPU program, data submitted by hospitals are publicly reported regardless of whether those data are successfully validated for payment determination purposes. For these reasons, in the CY 2010 OPPS/ASC proposed rule (74 FR 35404), we proposed to make data collected for quarters beginning with the third quarter of CY 2008 (July–September 2008) under the HOP QDRP publicly available, regardless of whether those data have been validated for payment determination purposes. We invited public comment on this proposal.

Comment: Many commenters opposed the public reporting of unvalidated HOP QDRP data. The commenters stated that these data would not be accurate and may mislead the public. The commenters also argued that because data for the inpatient reporting program were validated before they were publicly posted, the outpatient data should be as well. The commenters stated that making these data available on Hospital Compare and in the downloadable public use file accompanying a Hospital Compare release may lead to inappropriate use of the data in research and policy deliberations, and may result in inaccurate portrayals of the data on various other Web sites that currently utilize Hospital Compare data. The commenters argued that public reporting of unvalidated data may cause confusion among consumers utilizing these data on different Web sites. The commenters were concerned that it is not known now how the non-validated data compare to the validated data. The commenters argued that because data for cases from quarters earlier than the second quarter of 2009 will not have been validated, HOP QDRP data should not be publicly reported prior to this time period.

Response: The validation process for both the RHQDAPU program and the HOP QDRP pertains only to chart-abstracted measures. Validation under these programs for the purposes of payment determination seeks to validate the accurate application of the abstraction rules described in the HOPD Specifications Manual when data are abstracted from medical records and submitted to CMS. Neither the HOP QDRP nor the RHQDAPU program has any validation process for claims data. Thus, the context of our discussion of the public reporting of unvalidated HOP QDRP data in previous rulemakings, our proposal in the CY 2010 OPPS/ASC proposed rule, and our discussion of

public comments we received regarding whether to post unvalidated data and which quarters of data would be appropriate to post on Hospital Compare pertain only to chart-abstracted measures.

We note that the Secretary is required under section 1833(t)(17)(E) of the Act to establish procedures to make the data submitted under the HOP QDRP available to the public, and we intend to use generally the same procedures that we currently use for the RHQDAPU program. In the RHQDAPU program, we currently publicly report the data as they have been submitted by the hospitals, and we report these data regardless of whether the hospital passed validation. Also, no changes are made to quality data that have been submitted by hospitals that fail validation in the inpatient RHQDAPU program. However, for the RHQDAPU program, we have suppressed data from display on Hospital Compare in circumstances where we have become aware of inaccuracies in the calculation of the measure rates due to systematic issues with the data submitted. We believe that the finalized methodological improvements in the validation process for the CY 2011 HOP QDRP will allow CMS to better assess the overall accuracy of the chart-abstracted data that are submitted by hospitals to CMS. We also believe that our approach will encourage hospitals to optimize their chart abstraction processes and improve the accuracy of their data because it is data that hospitals are responsible for that are ultimately posted on Hospital Compare.

Although we appreciate the concerns raised about the public reporting of unvalidated data, prior to public reporting hospitals are given an opportunity to preview the results to be reported. Additionally, should our consumer testing suggest a different approach to public reporting, or should our validation process for CY2011 reveal a low reliability of self-reported data, we will reconsider our approach for future public reporting.

Comment: Many commenters were opposed to the publication of data beginning with 3rd quarter 2008 services because, during part of that period: (1) Some antibiotics needed to be updated in the specifications but these updates were not present at that time; (2) some procedures needed to be removed from the specifications but these exclusions were not present at that time; and (3) canceled procedures were not able to be excluded from the calculation of certain measures reported at that time. Because these changes were later put into effect, many commenters

suggested suppressing earlier quarters of data, and beginning public reporting with the 1st quarter of 2009 data when these changes had been made, and the data specifications were stabilized.

Response: The issues raised by the commenters apply only to the chart-abstracted HOP QDRP surgical care measures OP-6 and OP-7. We have considered the issues raised by the commenters, and because there was an issue with surgeries that were cancelled prior to incision that was not resolved in the specifications until July 1, 2009, we have decided not to report either of the surgery-related process measures (OP-6 & OP-7) for any quarter before the 3rd quarter of 2009.

Comment: Some commenters expressed concern that the 1 year to 2 year time lag in the public reporting of administrative claims-based measures would not be useful to the public.

Response: In the CY 2009 OPPS/ASC final rule with comment period we adopted the 4 claims based imaging efficiency measures for the CY 2010 payment determination and indicated that we would calculate the claims-based imaging efficiency measures for that payment determination using CY 2008 claims (73 FR 68761-68763). We recognize that the time lag for the claims-based measures is a concern, but because of claims submission and claims processing timeframes, this time lag ensures that the most complete and accurate paid claims are used in the measure calculations. Part of the time lag is also due to the time needed for data extraction, data processing, calculation of the measures for the payment determination and subsequent public reporting, quality assurance processes, and the Hospital Compare preview and update schedule. We intend to publicly report the claim-based measures as soon as possible, and the earliest we anticipate being able to make these claims-based measures available to the public is June 2010.

Comment: Some commenters commended CMS for its efforts to publicly report hospital outpatient measures on Hospital Compare in 2010, and encouraged CMS to continue to engage multi-stakeholder groups in testing and preparing the measures for display on Hospital Compare, as is done with inpatient measures.

Response: We appreciate the supportive comments regarding public reporting of hospital outpatient measures on Hospital Compare in 2010. We began stakeholder and consumer focus group testing of the HOPD measures in the Fall of 2009. We will continue to engage in consumer testing and obtain stakeholder input regarding

public reporting of HOP QDRP measures.

Comment: Commenters made numerous suggestions for enhancing the public reporting of HOP QDRP data including:

- Reporting comparative hospital outpatient (OP) and ambulatory surgical center (ASC) quality information on surgical services.
- Providing a mechanism for providers to raise concerns with information to be posted prior to its publication.
- Including a provider narrative section allowing the providers to advise consumers of any concerns regarding reliability or accuracy of information presented.
- Including information on facility accreditation status, state licensure and Medicare certification.
- Creating clear and explicit names and explanations for the measures geared toward consumers.
- Grouping like measures by condition and distinguishing them by care setting.
- Communicating efficiency measures in a manner that clearly interprets the differences among providers, and explains how consumers should integrate this information into decision making.
- Conducting consumer testing and multi-stakeholder vetting of changes in the Hospital Compare architecture, navigation, display and language.
- Considering how best to display hospital outpatient data in the context of current and anticipated future public reporting efforts for ASCs.
- Adding an identifier to the CCN to enable the reporting and display of data for individual hospitals rather than combining results from two or more hospitals sharing the same CCN.
- Displaying measures in a way that allows greater range and detail in categorizations.

Response: We thank the commenters for these suggestions and will consider them as we further develop our procedures for the public reporting of HOP QDRP quality data.

After consideration of the public comments we received, we have decided to finalize our proposal to publicly report HOP QDRP data on Hospital Compare in 2010 with some modifications in the periods of time to be reported. For measures OP-1 through OP-5, we will publicly report data periods beginning with the 3rd quarter of 2008. For measures OP-6 and OP-7, we will publicly report data periods beginning with the 3rd quarter of 2009. For measures OP-8 through OP-11, we will report CY 2010 payment

determination calculations using CY 2008 claims.

As we noted in section XVI.A.5.c. of this final rule with comment period, in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68778), we established that, for CY 2010, hospitals sharing the same CCN must combine data collection and submission across their multiple campuses for the clinical measures for public reporting purposes and that we will publish quality data by CCN under the HOP QDRP. This approach is consistent with the approach taken under the RHQDAPU program. In that final rule with comment period, we also stated that we intend to indicate instances where data from two or more hospitals are combined to form the publicly reported measures on the Web site.

Comment: One commenter suggested that combining the results from two or more hospitals that share the same CCN is misleading and will not allow consumers and health care payers to assess and use the information, reducing the effectiveness of Hospital Compare. This commenter stated that hospitals should be required to report at the unit of the hospital rather than the CCN. Another commenter suggested that CMS add an identifier to the CCN in order to enable the reporting and display of data of quality measure data for individual hospitals rather than combining results from two or more hospitals that share the same CCN.

Response: We believe that we should publicly report combined data from hospitals with the same CCN because it is important to align clinical reporting with financial reporting. We proposed to report data by CCN for several reasons. First, the unit affected by the OPPS annual update and that must meet all HOP QDRP requirements is the CCN, not an individual hospital or facility that falls under that CCN. Second, hospitals are obligated to comply with applicable survey and certification requirements by CCN, and not by any other individual facility identifier. Third, the additional Medicare identifier for facilities, the National Provider Identifier (NPI), is not a uniform identifier. Fourth, reporting by CCN aligns the reporting of quality of care data with the reporting of financial data. For these reasons, we consider the CCN to be representative of the entire hospital entity for purposes of public reporting under the HOP QDRP. However, as resources permit, we will evaluate whether the benefits the commenter believes would flow from its approach outweigh the reasons outlined above for using the current CCN and whether this suggestion is feasible.

After consideration of the public comments, we have decided to finalize our proposal to publicly report HOP QDRP data on Hospital Compare starting as early as 2010 with some modifications in the periods of time to be reported. Should our consumer testing suggest a different approach to public reporting, or should our validation process for CY 2011 reveal a low reliability of self-reported data, we will reconsider our approach for future public reporting. For measures OP-1 through OP-5, we will publicly report data periods beginning with the 3rd quarter of 2008. For measures OP-6 and OP-7, we will publicly report data periods beginning with the 3rd quarter of 2009. For measures OP-8 through OP-11, we will report 2010 payment determination calculations using calendar year 2008 claims.

G. HOP QDRP Reconsideration and Appeals Procedures

When the RHQDAPU program was initially implemented, it did not include a reconsideration process for hospitals. Subsequently, we received many requests for reconsideration of those payment decisions and, as a result, established a process by which participating hospitals would submit requests for reconsideration. We anticipated similar concerns with the HOP QDRP and, therefore, in the CY 2008 OPPS/ASC final rule with comment period (72 FR 66875), we stated our intent to implement for the HOP QDRP a reconsideration process modeled after the reconsideration process we implemented for the RHQDAPU program. In the CY 2009 OPPS/ASC final rule with comment period (73 FR 68779), we adopted a mandatory reconsideration process that will apply to the CY 2010 payment decisions. In the CY 2010 OPPS/ASC proposed rule (74 FR 35404), we proposed to continue this process for the CY 2011 payment update. Under this proposed process, the hospitals must—

(1) Submit to CMS, via QualityNet, a Reconsideration Request form that will be made available on the QualityNet Web site; this form must be submitted by February 3, 2011, and must contain the following information:

- Hospital CCN.
- Hospital Name.
- CMS-identified reason for failure (as provided in any CMS notification of failure to the hospital).
- Hospital basis for requesting reconsideration. This must identify the hospital's specific reason(s) for believing it met the HOP QDRP

requirements and should receive a full annual payment update.

- CEO and any additional designated hospital personnel contact information, including name, e-mail address, telephone number, and mailing address (must include physical address, not just a post office box).

- A copy of all materials that the hospital submitted in order to receive the full payment update for CY 2011. Such material would include, but may not be limited to, the applicable Notice of Participation form or completed online registration form, and quality measure data that the hospital submitted via QualityNet.

The request must be signed by the hospital's CEO.

(2) Following receipt of a request for reconsideration, CMS will—

- Provide an e-mail acknowledgement, using the contact information provided in the reconsideration request, to the CEO and any additional designated hospital personnel notifying them that the hospital's request has been received.
- Provide a formal response to the hospital CEO and any additional designated hospital personnel, using the contact information provided in the reconsideration request, notifying the hospital of the outcome of the reconsideration process.

If a hospital is dissatisfied with the result of a HOP QDRP reconsideration decision, the hospital may file an appeal under 42 CFR part 405, subpart R (PRRB appeal).

Comment: Several commenters supported the proposed hospital reconsideration and appeals process. Some commenters suggested establishing a timeline for CMS to respond to reconsiderations and appeals. One of these commenters suggested a timeline for CMS to respond so that hospitals can better plan in the event the payment rate update is reduced by 2 percentage points. One commenter urged that the full payment rate update not be reduced for hospitals until the reconsideration and appeals process is complete. One commenter believed that this mandatory reconsideration and appeals process should be a permanent component to the quality reporting program and, therefore, not proposed or renewed each calendar year.

Response: We thank these commenters for their support of a hospital reconsideration and appeals process. We plan to complete any CY 2010 reconsideration reviews and communicate the results of these determinations within 60 to 90 days following the date we receive the

request for reconsideration. While we recognize the commenter's concern about possibly withholding the full payment rate update before the reconsideration and appeals process is complete, we need to consider that if we waited to reduce the payment, the agency could encounter issues with recouping funds that were improperly paid to a hospital that did not meet the HOP QDRP requirements.

Regarding making the reconsideration and appeals process a permanent component to the quality reporting program and, therefore, not proposing or renewing it each calendar year, we have customarily proposed most of the HOP QDRP requirements and procedures, even those we propose to continue with only minor modifications, through the annual rulemaking process in order to afford the public additional opportunities to comment on them. In the case of the reconsideration and appeals procedures, each year we also propose the date by which hospitals must request for reconsiderations (for the CY 2011 payment update, these requests must be submitted by February 3, 2011).

After consideration of the public comments we received, we are adopting as final the proposed HOP QDRP reconsideration and appeals procedures for CY 2010.

H. Reporting of ASC Quality Data

As discussed above, section 109(b) of the MIEA-TRHCA amended section 1833(i) of the Act by redesignating clause (iv) as clause (v) and adding new clause (iv) to paragraph (2)(D) and new paragraph (7) to the Act. These amendments authorize the Secretary to require ASCs to submit data on quality measures and to reduce the annual payment update in a year by 2.0 percentage points for ASCs that fail to do so. These provisions permit, but do not require, the Secretary to require ASCs to submit such data and to reduce any annual increase for noncompliant ASCs.

In the CY 2008 OPPTS/ASC final rule with comment period (72 FR 66875) and in the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68780), we indicated that we intended to implement the provisions of section 109(b) of the MIEA-TRHCA in a future rulemaking. While promoting high quality care in the ASC setting through quality reporting is highly desirable and fully in line with our efforts under other payment systems, the transition to the revised payment system in CY 2008 posed significant challenges to ASCs, and we determined that it would be most appropriate to allow time for ASCs

to gain some experience with the revised payment system before introducing other new requirements. Further, by implementing quality reporting under the OPPTS prior to establishing quality reporting for ASCs, CMS would gain experience with quality measurement in the ambulatory setting in order to identify the most appropriate measures for quality reporting in ASCs prior to the introduction of the requirement in ASCs. Finally, we are sensitive to the potential burden on ASCs associated with chart abstraction and believe that adopting such measures at this time is in contrast with our desire to minimize collection burden, particularly when measures may be reported via EHRs in the future.

We continue to believe that promoting high quality care in the ASC setting through quality reporting is highly desirable and fully in line with our efforts under other payment systems. However, we continue to have the concerns outlined above for CY 2010 and, therefore, in the CY 2010 OPPTS/ASC proposed rule (74 FR 35405), we stated that we intend to implement the provisions of section 109(b) of the MIEA-TRHCA in a future rulemaking. We invited public comment on this deferral of quality data reporting for ASCs and invited suggestions for quality measures geared toward the services provided by ASCs. We again sought public comment on potential reporting mechanisms for ASC quality data, including electronic submission of these data.

Comment: Several commenters agreed with CMS' proposal and reasons for deferring quality data reporting for ASCs. Some commenters supported the CMS rationale for the proposed approach, that is, enabling ASCs to gain experience with the recently launched payment system and permitting CMS to gain experience in the HOPD setting before implementing quality data reporting requirements for ASCs. Several commenters supported CMS' decision to move with caution in expanding quality data reporting to the ASC setting and appreciated CMS' sensitivity to administrative burdens faced by ASCs. Commenters stated that it would be beneficial to allow extra time in order to assess implementation challenges and identify appropriate measures. These commenters also indicated that issues such as preventability, risk adjustment, unintended consequences, coding accuracy, burden, and effect on overall health care costs need to be carefully examined before starting a reporting program for a new setting. Other

commenters indicated it would be better to wait until ICD-10 implementation to begin measurement in a new setting in order to allow for more accurate coding and measurement and POA coding. One commenter agreed with the continued delay in implementing a quality data reporting program for ASCs based upon CMS' rationale set forth in the proposed rule and suggested that CMS discuss implementation of the requirements, including when ASC reporting will occur and the potential effects on ASC staff. One commenter argued that requiring ASCs to conduct quality data reporting is unnecessary because quality improvement is a key requirement for ASCs to obtain and maintain accreditation and such reporting would result in additional costs to ASC operations.

Response: We thank these commenters for acknowledging the many operational issues that must be addressed prior to implementing a quality data reporting program for ASCs and supporting our decision to defer ASC quality reporting to a future time. However, with regard to the commenter's suggestion that an ASC quality data reporting program is unnecessary in light of ASC quality improvement accreditation requirements, we believe that quality measure data reporting for ASCs would provide consumers with quality of care information for this type of health care provider as well as support our quality improvement efforts. Therefore, notwithstanding the current issues that have led to our determination to implement an ASC quality reporting program in a future rulemaking, we believe it is important and necessary to require ASCs to submit quality data.

Comment: Numerous commenters advocated that CMS move forward with an ASC quality data reporting program as soon as possible. Many commenters indicated that the collection and reporting of quality data is a common practice for ASC facilities, and that the industry is eager to make quality data available to consumers in a manner that allows direct comparisons between equivalent surgical care delivered in HOPDs and ASCs, particularly as the percentage of outpatient surgical services being provided in ASC settings has grown. Some commenters urged CMS to implement a quality data reporting system for ASCs for CY 2010. One commenter was concerned about the continued delay in quality measurement for the rapidly growing ASC setting and indicated that by now it should be technically feasible for ASCs to report on at least the set of five quality measures that were developed

by the industry-sponsored ASC Quality Collaboration. Several commenters argued that any quality data reporting system implemented would not create significant administrative burden on ASCs. Some of these commenters recommended the use of administrative claims as a means for quality reporting as both chart abstraction and Internet-based reporting would impose major disadvantages for ASCs, most of which are classified as small businesses. Some commenters suggested beginning with a set of six ASC quality measures that have already been developed. Commenters also suggested that CMS consider claims-based reporting for ASCs, which would eliminate the chart abstraction burden, would capitalize on existing data collection infrastructure, and would be most feasible for the industry at this time. Another commenter indicated that specialty-specific measures for ASCs should be implemented in such a reporting program in order to ensure broad opportunity for participation, including those ASCs that specialize in a few services or procedures. One commenter indicated that at least 35 States collect data on ASCs.

Response: We agree that it would be beneficial for consumers to be able to compare the quality of surgical care across HOPDs and ASCs. Currently, in addition to the reasons we outlined in the proposed rule, we do not have the resources needed to implement a quality data reporting program for ASCs. We are aware of the set of five quality measures that were developed by the ASC Quality Collaboration as well as six NQF-endorsed quality measures. While some of the measures may be feasible to collect using claims data, others (such as the patient safety-related measures) may not be meaningful to report unless all patients treated were captured, and hence all-payer claims were collected to generate the measures. We will evaluate the suggested measures for ASC quality data reporting, as well as the feasibility of claims-based measure reporting for ASCs and the need for specialty-specific measures for ASCs in the future.

Comment: One commenter encouraged CMS to focus on electronic submission of data for quality reporting once it has been determined to move forward with the ASC quality program. One commenter recommended that any ASC reporting program build on the public reporting programs in place for the inpatient and outpatient settings and that the measures reported in the ASC setting be consistent and, where possible, identical to the outpatient department program as the consistency will minimize confusion, simplify data

collection, and assure greater comparability across sectors.

Response: We thank the commenters for these suggestions. We will consider them in the planning process for ASC quality measure data reporting.

After consideration of the public comments we received, we are finalizing our proposal to implement quality measure reporting in the ASC setting in a future rulemaking. We continue to believe that promoting high quality care in the ASC setting through quality data reporting is highly desirable and fully in line with our efforts under other payment systems.

I. Electronic Health Records

As stated above, CMS is actively seeking alternatives to manual chart abstraction for the collection of quality measures for its quality data reporting programs. Among these alternatives are claims-based measure calculation, collection of data from systematic registries widely used by hospitals, and electronic submission of quality measures using EHRs. In the CY 2009 OP/ASC final rule with comment period, we discussed public commenters' suggestions that we adopt measures that can be collected via EHRs (73 FR 68769). We agree with the commenters about the importance of actively working to move to a system of data collection based on submission from EHRs. We have been engaged with health IT standard-setting organizations to promote the adoption of the necessary standards regarding data capture to facilitate data collection via EHRs, and have been collaborating with such organizations on standards for a number of quality measures. We encourage hospitals to take steps toward the adoption of EHRs that will allow for reporting of clinical quality data from the EHR directly to a CMS data repository. We also encourage hospitals that are implementing, upgrading, or developing EHR systems to ensure that such systems conform to standards adopted by HHS. In the CY 2010 OP/ASC proposed rule (74 FR 35405), we invited public comment on the future direction of EHR-based quality measure submission with respect to the HOP QDRP.

Comment: One commenter strongly encouraged CMS to consider aligning the HOP QDRP with the ONC measures for "meaningful use," and to provide clarity on those measures that appear to be similar to those identified as measures for meaningful use, such as the OP/ASC CY 2011 "OP-4: Aspirin at Arrival" and the meaningful use matrix measure for CY 2011, "Improve quality, safety, efficiency, and reduce health

disparities: Percentage patients at high-risk for cardiac events on aspirin prophylaxis [OP]."

Response: The measure matrix referenced by the commenter is a list of criteria developed by the Health IT Policy Council, an advisory committee to ONC, for consideration by the Department as it develops the initial criteria for determining whether an eligible hospital or eligible professional is a meaningful user of certified EHR technology. Eligible hospitals and eligible professionals who demonstrate that they meaningfully use certified EHR technology will be eligible for payment incentives under Medicare, as authorized under the HITECH Act. To be considered a meaningful user of the certified EHR technology, section 1886(n)(3)(A)(iii) of the Act, as added by section 4102(a) of the HITECH Act, requires that eligible hospitals submit to CMS, using certified EHR technology, the clinical quality measures and such other measures selected by the Secretary. Section 1886(n)(3)(B)(iii) of the Act provides that in selecting these reporting measures, the Secretary shall seek to avoid redundant or duplicative reporting with the reporting otherwise required, including reporting under RHQDAPU. CMS will establish the initial meaningful use criteria in future rulemaking, including the selection of quality measures for hospital reporting purposes under this separate incentive program. Some of the clinical quality measures included on the Health IT Policy Council's matrix are similar to measures adopted into the HOP QDRP. As we stated in the "considerations for measurement" section of this final rule with comment period, because we seek to harmonize applicable measures across settings, and many of the measures for the HOP QDRP that apply to HOPDs have been adapted from the RHQDAPU program, some of the measures that appear on the Health IT Policy Council's matrix are similar to measures adopted into the RHQDAPU program. We thank the commenters and will take these comments into consideration as we consider the future direction of EHR-based quality measure submission with respect to the HOP QDRP.

XVII. Healthcare-Associated Conditions

A. Background

1. Preventable Medical Errors and Hospital-Acquired Conditions (HACs) Under the IPPS

As noted in its landmark 1999 report "To Err is Human: Building a Safer Health System," the Institute of Medicine found that medical errors are

a leading cause of morbidity and mortality in the United States. Total national costs of these errors due to lost productivity, disability, and health care costs were estimated at \$17 billion to \$29 billion.² As one approach to combating healthcare-associated conditions, in 2005, Congress authorized CMS to adjust Medicare IPPS hospital payments to encourage the prevention of these conditions. Section 1886(d)(4)(D) of the Act (as added by section 5001(c) of the Deficit Reduction Act (DRA) of 2005, Pub. L. 109–171) required the Secretary to select by October 1, 2007, at least two conditions that are: (1) High cost, high volume, or both; (2) assigned to a higher paying diagnosis-related group (DRG) when present as a secondary diagnosis; and (3) could reasonably have been prevented through the application of evidence-based guidelines. CMS has titled this initiative “Hospital-Acquired Conditions (HAC) and Present on Admission (POA) Indicator Reporting.” Since October 1, 2008, Medicare no longer assigns a hospital inpatient discharge to a higher paying Medicare Severity Diagnosis-Related Group (MS-DRG) if a selected HAC is not present on admission. That is, if there is an HAC, the case is paid as though the secondary diagnosis was not present. However, if any nonselected complications or comorbidities appear on the claim, the claim will be paid at the higher MS-DRG rate; to cause a lower MS-DRG payment, all complications or comorbidities on the claim must be selected conditions for the HAC payment provision. Since October 1, 2007, CMS has required hospitals to submit information on Medicare hospital inpatient claims specifying whether diagnoses were POA.

2. Expanding the Principles of the IPPS HACs Payment Provision to the OPSS

In the CY 2009 OPSS/ASC proposed rule and final rule with comment period (73 FR 41547 and 68781, respectively), we discussed whether the principle of Medicare not paying more for preventable HACs during inpatient stays paid under the IPPS could be applied more broadly to other Medicare payment systems in other settings for conditions that occur or result from health care delivered in those settings. We also acknowledged that implementation of this concept would be different for each setting, as each Medicare payment system is unique. As

we have used in past rulemaking and general notices, in the following discussion in this final rule with comment period, we refer to conditions that occur in the hospital inpatient setting as “hospital-acquired conditions (HACs),” to conditions that occur in HOPDs as “hospital outpatient healthcare-associated conditions (HOP-HACs),” and to conditions that result from care in settings other than the hospital inpatient and HOPD settings as “healthcare-associated conditions.”

In both the CY 2009 OPSS/ASC proposed rule and final rule with comment period, we specifically presented our rationale for considering the HOPD as a possible appropriate setting for Medicare to extend to the OPSS the concept of not paying more for preventable healthcare-associated conditions that occur as a result of care provided during a hospital encounter. For example, hospitals provide a broad array of services in their HOPDs that may overlap or precede the inpatient activities of the hospital, including many surgical procedures and diagnostic tests that are commonly performed on both hospital inpatients and outpatients. Similarly, individuals who are eventually admitted as hospital inpatients often initiate their hospital encounter in the HOPD, where they receive care during clinic or emergency department visits or observation care that precede their inpatient hospital admission. In addition, like the IPPS, the OPSS is also subject to the “pay-for-reporting” provision that affects the hospital outpatient annual payment update by the authority of section 1833(t)(17) of the Act (as amended by section 109(a) of Pub. L. 109–432 (MIEA–TRHCA)). (We refer readers to section XVI. of this final rule with comment period for a discussion of the HOP QDRP provisions for hospitals that fail to meet the reporting requirements established for the hospital outpatient payment update.)

The risks of preventable medical errors leading to the occurrence of healthcare-associated conditions are likely to be high in outpatient settings, given the large number of encounters and exposures that occur in these settings. Approximately 530,000 preventable drug-related injuries are estimated to occur each year among Medicare beneficiaries in outpatient clinics.³ These statistics clearly point to the significant magnitude of the problem of healthcare-associated

conditions in outpatient settings. Recent trends have shown a shift in services from the inpatient setting to the HOPD, and we expect the occurrence of healthcare-associated conditions stemming from outpatient care to grow directly as a result of this shift in sites of service.

For the CY 2009 OPSS, we did not adopt any new Medicare policy in our discussion of healthcare-associated conditions as they relate to the OPSS. Instead, in the CY 2009 OPSS/ASC proposed rule, we solicited public comments on options and considerations, including the statutory authority related to expanding the IPPS HAC provision to the OPSS. Our discussion addressed the following areas:

- Criteria for possible candidate OPSS conditions;
- Collaboration process;
- Potential OPSS HOP-HACs, including object left in during surgery; air embolism; blood incompatibility; and falls and trauma, including fractures, dislocations, intracranial injuries, crushing injuries, and burns; and
- OPSS infrastructure and payment for encounters resulting in healthcare-associated conditions, including the necessity of POA reporting for hospital outpatient services, methods for risk stratification, and potential methods for adjusting hospital payment.

3. Discussion in the CY 2009 OPSS/ASC Final Rule With Comment Period

In the CY 2009 OPSS/ASC final rule with comment period (73 FR 68784 through 68787), we responded to the public comments we received on healthcare-associated conditions in the context of the OPSS. Several commenters fully supported expanding the IPPS HAC policy to other settings such as HOPDs and ASCs, but many commenters stated that CMS should not implement a related policy in other settings without gaining implementation experience with the IPPS HACs. A number of commenters addressed concerns regarding some of the potential specific HOP-HACs discussed in the CY 2009 OPSS/ASC proposed rule (73 FR 41549), and some commenters suggested other conditions that should be considered or identified those that should not be considered. Many commenters stated that the attribution of HOP-HACs in the HOPD setting is difficult and stated that there was a need to develop risk adjustment techniques to account for differences in patient severity of illness or other patient characteristics. Many commenters asserted that the POA

² Institute of Medicine: To Err Is Human: Building a Safer Health System, November 1999. Available at: <http://www.iom.edu/Object.File/Master/4/117/ToErr-8pager.pdf>.

³ Asplen, P., Wolcott, J., Bootman, J.L., Cronenwett, L.R. (editors): Preventing Medication Errors: Quality Chasm Series, The National Academy Press, 2007. Available at: http://www.nap.edu/catalog.php?record_id=11623.

indicators may need to be modified for use in the HOPD or ASC setting. Some commenters suggested that a “present on encounter” indicator or another form of incorporation of preexisting conditions into an episode-of-care might be more useful than a POA indicator. Several commenters believed that, without changes to the existing OPSS payment structure, there would be no straightforward methodology for adjusting hospital payment.

While we acknowledged these challenges in the CY 2009 OPSS/ASC final rule with comment period (73 FR 68787), we noted that we view addressing the ongoing problem of preventable healthcare-associated conditions in outpatient settings, including the HOPD, as a key value-based purchasing strategy to sharpen the focus on such improvements beyond hospital inpatient care to those settings where the majority of Medicare beneficiaries receive most of their health care services. We also noted that we looked forward to continuing to work with stakeholders to improve the quality, safety, and value of health care provided to Medicare beneficiaries, beginning with a joint IPPS/OPSS listening session.

B. Public Comments and Recommendations on Issues Regarding Healthcare-Associated Conditions From the Joint IPPS/OPSS Listening Session

Subsequent to the issuance of the CY 2009 OPSS/ASC final rule with comment period, we held a joint Hospital-Acquired Conditions and Hospital Outpatient Healthcare-Associated Conditions Listening Session on December 18, 2008. (The listening session was announced in a notice published in the **Federal Register** on October 30, 2008 (73 FR 64618).) During the listening session, we provided an overview of the HAC program under the IPPS and our previous discussions of extending the underlying concepts to the HOPD, including OPSS infrastructure concerns such as the lack of a POA indicator and the need to address current ICD-9-CM POA reporting guidelines, attribution of conditions in the HOPD, and payment adjustment considerations. In addition to the initial candidate HOP-HACs that we had previously identified based on their adoption under the IPPS, we discussed other potential HOP-HACs, such as medication errors, conditions related to complications of hospital outpatient surgery or other procedures, and infections related to HOPD care. A transcript of the listening session is available on the CMS Web site at: [http://](http://www.cms.hhs.gov/HospitalAcqCond/07-EducationalResources.asp#TopOfPage)

www.cms.hhs.gov/HospitalAcqCond/07-EducationalResources.asp#TopOfPage.

Of the many public comments presented orally at the listening session or submitted in writing, approximately one-half commented on expansion of the IPPS HAC payment provision to other settings. Some commenters were in favor of an expansion to the HOPD and other settings. Many commenters requested that CMS delay any expansion, citing the short duration of experience with HACs and POA indicator reporting for inpatient hospitalizations and the need to evaluate the current program prior to its expansion to other settings.

We appreciate these commenters' perspectives and note that now that we have early data on the HAC program, in the near future we plan to evaluate the impact of the HAC payment provision through a joint program evaluation with CDC, AHRQ, and the Office of Public Health and Science.

Many commenters pointed to the need to define the boundaries of an episode-of-care for healthcare-associated conditions in the HOPD and other settings in order to define when, how, and to whom an expanded policy would apply. These commenters also noted that hospital outpatients have frequently received care from numerous practitioners and providers over an extended period of time and the hospitals' or clinics' role would be supportive, rather than prescriptive, with respect to that patient care. They requested that CMS develop a comprehensive and accurate definition of an episode-of-care in order to appropriately attribute responsibility and the additional costs associated with HOP-HACs.

We have previously acknowledged that short-term consideration of HOP-HACs would necessarily be limited to conditions that occur during and result from care provided in a single hospital outpatient encounter because a broader definition of an episode-of-care has not yet been developed.

Many commenters believed that detailed information should be gathered and analyzed from the IPPS POA indicator reporting experience before an expansion of the HAC payment provision and POA indicator reporting to the HOPD. Other commenters pointed out that the initial four conditions under consideration for HOPDs based on their adoption under the IPPS would likely require emergency admission for treatment of the event. Though secondary to an initial encounter in the HOPD, they indicated that these conditions would be coded as POA for the IPPS according to current reporting

guidelines and would not be captured as HOP-HACs. Several commenters stated that, in the HOPD, it would be particularly important to make an assessment over an entire episode-of-care; thus, POA might be better defined in terms of “present on encounter” for this purpose. Other commenters pointed to the need for the development of new codes and determinations of when the codes should apply in order to capture POA conditions under the OPSS, an activity that would potentially significantly increase hospitals' administrative burden. Some commenters suggested waiting to expand the HAC payment provision to other settings until implementation of the ICD-10 classification system, which would provide more precise coding to identify preexisting conditions.

We have acknowledged a number of these challenges already, and we will continue to consider these reporting issues as we refine our views regarding potential HOP-HACs.

Many commenters highlighted that patients receiving hospital outpatient care may receive care in multiple departments of the hospital, both during a single outpatient encounter and longitudinally over many outpatient encounters of relatively short duration. These commenters stated that, because of these common patterns of care, the timely identification of HOP-HACs and their provider attribution would be particularly challenging. In addition, the commenters pointed out that patient factors may play a role in the development of potential HOP-HACs, such as adverse drug events. Several of these commenters suggested targeting the HOP-HAC policy to specific APCs, specific HCPCS codes, or specific HOPD settings, such as the emergency department. In the CY 2009 OPSS/ASC proposed rule and final rule with comment period (73 FR 41549 through 41550 and 68785 through 68787, respectively), we discussed the challenge of provider attribution under the OPSS, particularly for conditions that may develop over time and involve multiple encounters and other care settings. We understand the importance of this issue and will continue to be cognizant of it in future policy development.

Several commenters asserted that CMS should consider risk adjustment models that incorporate population risk adjustments to avoid creating barriers to access for more complex patients or to avoid unduly placing providers treating more complex patients at higher risk for payment consequences due to HOP-HACs. A number of commenters endorsed the use of rate-based measures

of conditions on a provider-specific level so that the level of preventability of specific clinical conditions could be determined and compared. Several commenters stated that, under the best of circumstances, falls may not be "reasonably preventable," particularly in the HOPD. Many commenters also believed that adverse drug events would require further definition in order to appropriately address medication errors that were not directly under the control of the hospital providing the treatment of the medication-related problem and were, therefore, not "reasonably preventable." Similarly, some commenters stated that it would be difficult to appropriately attribute metabolic derangements in the HOPD to the hospital treating the resulting clinical problem.

We appreciate these public comments and will use our collaborative process with CDC, AHRQ, and the Office of Public Health and Science to help define potential HOP-HACs that are clinically meaningful for patient safety, as well as attributable to care furnished by providers.

Numerous commenters urged CMS to generally proceed with care, to promote the use of evidence-based guidelines and care coordination, and to ensure that any HOP-HAC program is aligned with other CMS quality programs. Many commenters believed that the challenges involved might be better addressed operationally within a full-scale value-based purchasing program. We appreciate these suggestions and will consider them as we advance policies that will ensure paying for the highest quality, safest, and most effective health care for Medicare beneficiaries.

C. CY 2010 Approach to Healthcare-Associated Conditions Under the OPSS

For CY 2010, we did not propose to expand the principles behind the IPPS HAC payment provision to the OPSS through a HOP-HAC program (74 FR 35407). We stated that we continue to believe that it may be appropriate to expand the principles of the IPPS HAC payment provision to the OPSS in the future. However, we acknowledged that, at this time, there are many operational challenges to such an expansion that will require further consideration and infrastructure development. We appreciate the input and guidance provided by the many public commenters to date on how to approach these challenges. Most stakeholders have strongly encouraged CMS to evaluate the impact of the IPPS HAC payment provision before further considering any expansion to other settings. We explained that we are

evaluating the impact of the HAC and POA indicator reporting initiative on Medicare payment and that we plan to consider any relevant findings as part of our future decision making regarding any expansion of the HAC payment provision to other settings. We welcomed additional suggestions and comments from stakeholders on potential HOP-HACs as additional information becomes available and health care delivery continues to evolve.

Comment: Several commenters commended CMS for considering an extension of the current IPPS HAC policy to other care settings and payment systems, including the OPSS. These commenters suggested that it would be reasonable to begin with patient safety-related conditions such as an object left in during surgery, air embolism, blood incompatibility, falls and trauma, including fractures, dislocations, intracranial injuries, crushing injuries, and burns.

However, the majority of commenters supported CMS' position in not proposing to expand the IPPS HAC payment provision to the OPSS at this time. They agreed with the plan to consider any relevant findings from the agency's evaluation of the impact of the HAC and POA indicator reporting initiatives (74 FR 35407) as part of CMS' future decision-making regarding expansion of the IPPS HAC policy to other settings. Several commenters reiterated their concerns related to technical challenges in expanding the IPPS HAC program to the OPSS, with particular emphasis on the need to develop a type of POA coding for hospital outpatient services and the fact that current POA guidelines designate conditions that develop during an outpatient encounter, such as clinic and emergency department visits, observation services, or outpatient surgery, as POA for hospital inpatient reporting. Other commenters encouraged CMS to develop a comprehensive definition of an episode-of-care, with the potential for inclusion of related care settings to appropriately attribute accountability. Some commenters urged CMS to evaluate the role of ICD-10 classification in the HAC program and to consider postponing implementation of HOP-HACs until the adoption of ICD-10. A number of commenters recommended that CMS focus on areas of patient safety in a future HOP-HACs program, including outpatient surgery and outpatient procedures that are correlated with the potential for injury and medication errors. Finally, several commenters encouraged CMS to ensure that any future HOP-HAC program provides an

incentive for care coordination, aligns with other CMS quality initiatives, and has no detrimental effect on patient access to care.

Response: We appreciate the thoughtful and detailed suggestions from commenters. We will continue to consider the concerns and suggestions from commenters as we evaluate the impact of the HAC and POA indicator reporting initiative and as part of our future decision making regarding any expansion of the HAC payment provision to other settings.

XVIII. Files Available to the Public Via the Internet

A. Information in Addenda Related to the CY 2010 Hospital OPSS

Addenda A and B to this final rule with comment period provide various data pertaining to the CY 2010 payment for items and services under the OPSS. Addendum A, which includes a list of all APCs payable under the OPSS, and Addendum B, which includes a list of all active HCPCS codes with their CY 2010 OPSS payment status and comment indicators, are available to the public by clicking "Hospital Outpatient Regulations and Notices" on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientOPSS/>.

For the convenience of the public, we also are including on the CMS Web site a table that displays the HCPCS code data in Addendum B sorted by APC assignment, identified as Addendum C.

Addendum D1 defines the payment status indicators that are used in Addenda A and B. Addendum D2 defines the comment indicators that are used in Addendum B. Addendum E lists the HCPCS codes that are only payable to hospitals as inpatient procedures and are not payable under the OPSS. Addendum L contains the out-migration wage adjustment for CY 2010. Addendum M lists the HCPCS codes that are members of a composite APC and identifies the composite APC to which each is assigned. This addendum also identifies the status indicator for the HCPCS code and a comment indicator if there is a change in the code's status with regard to its membership in the composite APC. Each of the HCPCS codes included in Addendum M has a single procedure payment APC, listed in Addendum B, to which it is assigned when the criteria for assignment to the composite APC are not met. When the criteria for payment of the code through the composite APC are met, one unit of the composite APC payment is paid, thereby providing packaged payment for all services that are assigned to the composite APC

according to the specific I/OCE logic that applies to the APC. We refer readers to the discussion of composite APCs in section II.A.2.e. of this final rule with comment period for a complete description of the composite APCs.

These addenda and other supporting OPSS data files are available on the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/>.

B. Information in Addenda Related to the CY 2010 ASC Payment System

Addenda AA and BB to this final rule with comment period provide various data pertaining to the CY 2010 payment for the covered surgical procedures and covered ancillary services for which ASCs may receive separate payment. Addendum AA lists the CY 2010 ASC covered surgical procedures, their payment indicators, and their payment rates. Addendum BB displays the CY 2010 ASC covered ancillary services, their payment indicators, and their payment rates. All ASC relative payment weights and payment rates for CY 2010 are a result of applying the revised ASC payment system methodology established in the final rule for the revised ASC payment system published in the **Federal Register** on August 2, 2007 (72 FR 42470 through 42548) to the CY 2010 OPSS and MPFS ratesetting information.

Addendum DD1 defines the payment indicators that are used in Addenda AA and BB. Addendum DD2 defines the comment indicators that are used in Addenda AA and BB.

Addendum EE (available only on the CMS Web site) lists the surgical procedures that are excluded from Medicare payment if furnished in ASCs. The excluded procedures listed in Addendum EE are surgical procedures that are assigned to the OPSS inpatient list, are not covered by Medicare, are reported using a CPT unlisted code, or have been determined to pose a significant safety risk or are expected to require an overnight stay when performed in ASCs.

These addenda and other supporting ASC data files are included on the CMS Web site at: <http://www.cms.hhs.gov/ASCPayment/>. The MPFS data files are located at: <http://www.cms.hhs.gov/PhysicianFeeSched/>.

The links to all of the FY 2010 IPSS wage index-related tables (that are used

for the CY 2010 OPSS) that were published in the FY 2010 IPSS/LTCH PPS final rule (74 FR 44032 through 44125) are accessible on the CMS Web site at: <http://www.cms.hhs.gov/AcuteInpatientPPS/WIFN>.

XIX. Collection of Information Requirements

A. Legislative Requirement for Solicitation of Comments

Under the Paperwork Reduction Act of 1995, we are required to provide 30-day notice in the **Federal Register** and solicit public comment before a collection of information requirement is submitted to the Office of Management and Budget (OMB) for review and approval. In order to fairly evaluate whether an information collection should be approved by OMB, section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 requires that we solicit comment on the following issues:

- The need for the information collection and its usefulness in carrying out the proper functions of our agency.
- The accuracy of our estimate of the information collection burden.
- The quality, utility, and clarity of the information to be collected.
- Recommendations to minimize the information collection burden on the affected public, including automated collection techniques.

The CY 2010 OPSS/ASC proposed rule and this final rule with comment period do not specify any information collection requirements through regulatory text. However, in the proposed rule and in this final rule with comment period, we make reference to associated information collection requirements that are not discussed in the regulation text contained in this document. The following is a discussion of those requirements, for which we solicited public comment in the CY 2010 OPSS/ASC proposed rule (74 FR 35232).

B. Associated Information Collections Not Specified in Regulatory Text

1. Hospital Outpatient Quality Data Reporting Program (HOP QDRP)

As previously stated in section XVI. of this final rule with comment period, the quality data reporting program for hospital outpatient care, known as the Hospital Outpatient Quality Data Reporting Program (HOP QDRP), has been generally modeled after the quality

data reporting program for hospital inpatient services, the Reporting Hospital Quality Data for Annual Payment Update (RHQDAPU) program. Section 109(a) of the MIEA-TRHCA (Pub. L. 109-432) amended section 1833(t) of the Act by adding a new subsection (17) that affects the payment rate update applicable to OPSS payments for services furnished by hospitals in outpatient settings on or after January 1, 2009. Section 1833(t)(17)(A) of the Act, which applies to hospitals as defined under section 1886(d)(1)(B) of the Act, states that subsection (d) hospitals that fail to report data required for the quality measures selected by the Secretary in the form and manner required by the Secretary under section 1833(t)(17)(B) of the Act will receive a 2.0 percentage point reduction to their annual payment update factor. Section 1833(t)(17)(B) of the Act requires that hospitals submit quality data in a form and manner, and at a time, that the Secretary specifies. Section 1833(t)(17)(C)(i) of the Act requires the Secretary to develop measures appropriate for the measurement of the quality of care (including medication errors) furnished by hospitals in outpatient settings, that these measures reflect consensus among affected parties and, to the extent feasible and practicable, that these measures include measures set forth by one or more national consensus building entities.

2. HOP QDRP Quality Measures for the CY 2010 and CY 2011 Payment Determinations

In the CY 2009 final rule with comment period (73 FR 68766), we adopted 4 claims-based imaging measures for use in CY 2010, bringing the total number to 11 measures. For the CY 2010 payment update, we are requiring hospitals to submit data related to the seven data abstracted measures; we will calculate the claims-based measures using administrative paid claims data and do not require additional hospital data submissions. Similarly, as we proposed, we are using the same 11 measures and the same data submission requirements related to the seven data abstracted measures for CY 2011 payment determinations.

HOP QDRP measurement set to be used for CY 2010 and CY 2011 payment determination

OP-1: Median Time to Fibrinolysis

OP-2: Fibrinolytic Therapy Received Within 30 Minutes

OP-3: Median Time to Transfer to Another Facility for Acute Coronary Intervention

OP-4: Aspirin at Arrival

HOP QDRP measurement set to be used for CY 2010 and CY 2011 payment determination

OP-5: Median Time to ECG
OP-6: Timing of Antibiotic Prophylaxis
OP-7: Prophylactic Antibiotic Selection for Surgical Patients
OP-8: MRI Lumbar Spine for Low Back Pain
OP-9: Mammography Follow-up Rates
OP-10: Abdomen CT—Use of Contrast Material
OP-11: Thorax CT—Use of Contrast Material

As part of the data submission process pertaining to the 11 measures listed above, hospitals must also complete and submit a notice of participation in the HOP QDRP. By submitting this document, hospitals agree that they will allow CMS to publicly report the quality measures as required by the HOP QDRP.

The burden associated with this section is the time and effort associated with completing the notice of participation as well as collecting and submitting the data on the seven data abstracted measures. We estimate that there will be approximately 3,500 respondents per year. For hospitals to collect and submit the information on the required measures, we estimate it will take 30 minutes per sampled case. We estimate there will be a total of 1,800,000 cases per year, approximately 514 cases per respondent. The estimated annual burden associated with the aforementioned submission requirements is 900,000 hours $((1,800,000 \text{ cases/year}) \times (0.5 \text{ hours/case}))$.

We did not receive any public comments on the burden associated with these information collection requirements.

3. HOP QDRP Validation Requirements

In addition to requirements for submitting of quality data, hospitals must also comply with the requirements for data validation in CY 2011. As specified in detail in section XVI.E. of this final rule with comment period, for the CY 2011 payment determination, as we proposed, we are implementing a validation program that will require hospitals to supply requested medical documentation to a CMS contractor for purposes of validating those data. However, the results of the validation will not affect the CY 2011 payment update for any hospital, although the payment update may be affected if a hospital fails to submit the requested data. We believe that it is important for hospitals to have some experience and knowledge of the HOP QDRP validation process before payment determinations are made based upon validation results. As we proposed, we are implementing a validation program that will both limit burden upon hospitals, especially small

hospitals, as well as provide feedback to all hospitals on validation performance. We are requesting medical documentation from hospitals for April 1, 2009 through March 31, 2010 episodes of care, which, with a modification for two of the quality measures, will allow us to gather one full year of submitted data for validation purposes.

The burden associated with the CY 2011 requirement is the time and effort necessary to submit validation data to a CMS contractor. We estimate that it will take each hospital approximately 38 minutes to comply with these data submission requirements. To comply with the requirements, we estimate each hospital must submit between 2 to 3 cases on average for review. We estimate that 3,200 hospitals will need to comply with these requirements in order for us to collect a total of 7,300 charts across all sampled hospitals. The estimated annual burden associated with the data validation process for CY 2011 is 2,026 hours.

Similar to our policy for the FY 2012 RHQDAPU program (74 FR 43884 through 43889), we proposed (74 FR 35403) to validate data from 800 randomly selected hospitals each year under the HOP QDRP, beginning with the CY 2012 payment determination. We note that, because the 800 hospitals will be selected randomly, every HOP QDRP-participating hospital will be eligible each year for validation selection. For each selected hospital, we will randomly validate per year up to 48 patient episodes of care (12 per quarter) from the total number of cases that the hospital successfully submitted to the OPSS Clinical Warehouse during the applicable time period. However, if a selected hospital has submitted less than 12 cases in one or more quarters, only those cases available will be validated. However, we did not adopt that proposal in this final rule with comment period. Instead, we indicated that we would take into account results of further analyses of collected data, as well as public comments we received on our proposal and propose a validation process for the CY 2012 payment rate update in next year's rulemaking process.

The burden associated with the proposed CY 2012 requirement, if we adopt it next year is the time and effort necessary to submit validation data to a CMS contractor. We estimate that it will take each of the 800 sampled hospitals approximately 12 hours to comply with these data submission requirements. To comply with the requirements, we estimate each hospital must submit 48 cases for the affected year for review. We estimate that 800 hospitals must comply with these requirements to submit a total of 38,400 charts across all sampled hospitals. The estimated annual burden associated with the data validation process for CY 2012 and subsequent years is 9,600 hours.

We discuss public comments on this information collection requirement in section XVI.E.3.b. of this final rule with comment period.

4. HOP QDRP Reconsideration and Appeals Procedures

In the CY 2009 OPSS/ASC final rule with comment period (73 FR 68779), we adopted a mandatory reconsideration process that will apply to the CY 2010 payment decisions. As we proposed in the CY 2010 OPSS/ASC proposed rule, we will continue this process for the CY 2011 payment update. Under this process, the hospitals must meet all of the requirements specified in section XVI.G. of this final rule with comment period. We did not assign burden to the aforementioned information collection requirements in the CY 2010 OPSS/ASC proposed rule because we believed the associated information collection requirements were exempt under 5 CFR 1320.4 (that is, information collected subsequent to an administrative action is not subject to the PRA). However, upon further review, in this final rule with comment period, we are assigning burden to the reconsideration and appeals procedures. The burden associated with the reconsideration and appeals procedures is the time and effort necessary to gather the required information and submit it to CMS. The required information, as specified in section XVI.G. of this final rule with comment period, involves the submission of a completed reconsideration request form that is

signed by the hospital's chief financial officer. We estimate that 25 hospitals will avail themselves of the reconsideration and appeals procedures on an annual basis. We estimate that it will take each hospital approximately 40 minutes to gather the required information, complete the required reconsideration request form, obtain the signature of the chief financial officer, and forward the documentation to CMS. The total annual estimated burden associated with these requirements is 1,000 minutes.

We did not receive any public comments on these information collection requirements.

5. Additional Topics

While we are seeking OMB approval for the information collection requirements associated with the HOP QDRP and the data validation processes, in the CY 2010 OPSS/ASC proposed rule (74 FR 35232), we also sought public comment on several issues that have the potential to ultimately affect the burden associated with HOP QDRP and the data validation processes. Specifically, that proposed rule listed the possible quality measures under consideration for CY 2012 and subsequent years. We also solicited public comments to explore the use of registries to comply with the HOP QDRP submission requirements, the use of EHRs as a data submission tool, the use of a standardized process for the retirement of HOP QDRP quality measures, the use of an extraordinary circumstance extension or waiver for reporting quality data, and the implementation of additional data validation conditions. We discussed the comments we received on these issues in section XVI. of the preamble of this final rule with comment period.

XX. Response to Comments

Because of the large number of public comments we normally receive on **Federal Register** documents, we are not able to acknowledge or respond to them individually. We will consider all comments we receive by the date and time specified in the **DATES** section of this final rule with comment period, and, when we proceed with a subsequent document(s), we will respond to those comments in the preamble to that document(s).

XXI. Regulatory Impact Analysis

A. Overall Impact

We have examined the impacts of this final rule with comment period as required by Executive Order 12866 (September 1993, Regulatory Planning

and Review), the Regulatory Flexibility Act (RFA) (September 19, 1980, Pub. L. 96-354), section 1102(b) of the Social Security Act, the Unfunded Mandates Reform Act of 1995 (Pub. L. 104-4), Executive Order 13132 on Federalism, and the Congressional Review Act (5 U.S.C. 804(2)).

1. Executive Order 12866

Executive Order 12866 directs agencies to assess all costs and benefits of available regulatory alternatives and, if regulation is necessary, to select regulatory approaches that maximize net benefits (including potential economic, environmental, public health and safety effects, distributive impacts, and equity). A regulatory impact analysis (RIA) must be prepared for major rules that have economically significant effects (\$100 million or more in any 1 year) or adversely affect in a material way the economy, a sector of the economy, productivity, competition, jobs, the environment, public health or safety, or State, local, or tribal government or communities (58 FR 51741).

We estimate that the effects of the OPSS provisions that are being implemented in this final rule with comment period will result in expenditures exceeding \$100 million in any 1 year. We estimate the total increase (from changes in this final rule with comment period as well as enrollment, utilization, and case-mix changes) in expenditures under the OPSS for CY 2010 compared to CY 2009 to be approximately \$1.9 billion. Because this final rule with comment period for the OPSS is "economically significant" as measured by the \$100 million threshold and also a major rule under the Congressional Review Act, we have prepared a regulatory impact analysis that, to the best of our ability, presents the costs and benefits of this rulemaking. Table 73 of this final rule with comment period displays the redistributive impact of the CY 2010 changes on OPSS payment to various groups of hospitals.

We estimate that the effects of the ASC provisions that are being implemented in this final rule with comment period for the ASC payment system will not exceed \$100 million in any 1 year and, therefore, are not economically significant. We estimate the total increase (from changes in this final rule with comment period as well as enrollment, utilization, and case-mix changes) in expenditures under the ASC payment system for CY 2010 compared to CY 2009 to be approximately \$80 million. However, because this final rule with comment period for the ASC

payment system substantially affects ASCs, we have prepared a regulatory impact analysis of changes to the ASC payment system that, to the best of our ability, presents the costs and benefits of this rulemaking. Table 75 and Table 76 of this final rule with comment period display the redistributive impact of the CY 2010 changes on ASC payment, grouped by specialty area and then by procedures with the greatest ASC expenditures, respectively.

2. Regulatory Flexibility Act (RFA)

The RFA requires agencies to analyze options for regulatory relief of small businesses if a rule has a significant impact on a substantial number of small entities. For purposes of the RFA, small entities include small businesses, nonprofit organizations, and small governmental jurisdictions. Many hospitals, other providers, ASCs, and other suppliers are considered to be small entities, either by being nonprofit organizations or by meeting the Small Business Administration (SBA) definition of a small business (hospitals having revenues of \$34.5 million or less in any 1 year and ASCs having revenues of \$10 million or less in any 1 year). (For details on the latest standards for health care providers, we refer readers the SBA's Web site at: http://sba.gov/idc/groups/public/documents/sba_homepage/serv_sstd_tablepdf.pdf (refer to the 620000 series).)

For purposes of the RFA, we have determined that many hospitals and most ASCs would be considered small entities according to the SBA size standards. Individuals and States are not included in the definition of a small entity. Therefore, the Secretary has determined that this final rule with comment period will have a significant impact on a substantial number of small entities. Because we acknowledge that many of the affected entities are small entities, the analyses presented throughout this final rule with comment period constitute our regulatory flexibility analysis. Therefore, in the CY 2010 OPSS/ASC proposed rule (74 FR 35410), we solicited public comments on our estimates and analyses of the impact of the proposed rule on those small entities.

3. Small Rural Hospitals

In addition, section 1102(b) of the Act requires us to prepare a regulatory impact analysis if a rule may have a significant impact on the operations of a substantial number of small rural hospitals. This analysis must conform to the provisions of section 603 of the RFA. With the exception of hospitals located in certain New England

counties, for purposes of section 1102(b) of the Act, we now define a small rural hospital as a hospital that is located outside an urban area and has fewer than 100 beds. Section 601(g) of the Social Security Amendments of 1983 (Pub. L. 98-21) designated hospitals in certain New England counties as belonging to the adjacent urban areas. Thus, for OPPS purposes, we continue to classify these hospitals as urban hospitals. We believe that the changes to the OPPS in this final rule with comment period will affect both a substantial number of rural hospitals as well as other classes of hospitals and that the effects on some may be significant. Also, the changes to the ASC payment system in this final rule with comment period will affect rural ASCs. Therefore, the Secretary has determined that this final rule with comment period will have a significant impact on the operations of a substantial number of small rural hospitals.

4. Unfunded Mandates

Section 202 of the Unfunded Mandates Reform Act of 1995 (UMRA) also requires that agencies assess anticipated costs and benefits before issuing any rule whose mandates require spending in any 1 year of \$100 million in 1995 dollars, updated annually for inflation. That threshold level is currently approximately \$133 million. This final rule with comment period will not mandate any requirements for State, local, or tribal governments, nor will it affect private sector costs.

5. Federalism

Executive Order 13132 establishes certain requirements that an agency must meet when it promulgates a proposed rule (and subsequent final rule) that imposes substantial direct costs on State and local governments, preempts State law, or otherwise has Federalism implications.

We have examined the OPPS and ASC provisions included in this final rule with comment period in accordance with Executive Order 13132, Federalism, and have determined that they will not have a substantial direct effect on State, local or tribal governments, preempt State law, or otherwise have a Federalism implication. As reflected in Table 73 below, we estimate that OPPS payments to governmental hospitals (including State and local governmental hospitals) will increase by 1.8 percent under this final rule with comment period. While we do not know the number of ASCs with government ownership, we anticipate that it is small. We believe

that the provisions related to payments to ASCs in CY 2010 will not affect payments to any ASCs owned by government entities.

The following analysis, in conjunction with the remainder of this document, demonstrates that this final rule with comment period is consistent with the regulatory philosophy and principles identified in Executive Order 12866, the RFA, and section 1102(b) of the Act.

This final rule with comment period will affect payments to a substantial number of small rural hospitals and a small number of rural ASCs, as well as other classes of hospitals and ASCs, and some effects may be significant.

B. Effects of OPPS Changes in This Final Rule With Comment Period

We are making several changes to the OPPS that are required by the statute. We are required under section 1833(t)(3)(C)(ii) of the Act to update annually the conversion factor used to determine the APC payment rates. We also are required under section 1833(t)(9)(A) of the Act to revise, not less often than annually, the wage index and other adjustments, including pass-through payments and outlier payments. In addition, we must review the clinical integrity of payment groups and weights at least annually. Accordingly, in this final rule with comment period, we are updating the conversion factor and the wage index adjustment for hospital outpatient services furnished beginning January 1, 2010, as we discuss in sections II.B. and II.C., respectively, of this final rule with comment period. We also are revising the relative APC payment weights using claims data for services furnished from January 1, 2008, through December 31, 2008, and updated cost report information. We are continuing the current payment adjustment for rural SCHs, including EACHs. Finally, we list the 6 drugs and biologicals in Table 30 of this final rule with comment period that we are removing from pass-through payment status for CY 2010.

Under this final rule with comment period, we estimate that the update change to the conversion factor and other adjustments as provided by the statute will increase total OPPS payments by 2.1 percent in CY 2010. The changes to the APC weights, the changes to the wage indices, and the continuation of a payment adjustment for rural SCHs, including EACHs, will not increase OPPS payments because these changes to the OPPS are budget neutral. However, these updates do change the distribution of payments within the budget neutral system as

shown in Table 73 below and described in more detail in this section. We also estimate that the total change in payments between CY 2010 and CY 2009, considering all payments, including changes in estimated total outlier payments and expiration of additional money for specified wages indices outside of budget neutrality, will increase total OPPS payments by 1.9 percent.

1. Alternatives Considered

Alternatives to the changes we are making and the reasons that we have chosen the options are discussed throughout this final rule with comment period. Some of the major issues discussed in this final rule with comment period and the options considered are discussed below.

a. Alternatives Considered for Pass-Through Payment for Implantable Biologicals

We are finalizing our proposal to change the way we evaluate transitional pass-through applications for implantable biologicals and the way we pay for implantable biologicals newly eligible for transitional pass-through status beginning in CY 2010. As discussed in detail in section V.A.4. of this final rule with comment period, we are finalizing a policy that the pass-through evaluation process and pass-through payment methodology for implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) and that are newly approved for pass-through payment beginning on or after January 1, 2010, be the device pass-through process and payment methodology only. As a result, implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) will no longer be eligible to submit biological pass-through applications and to receive biological pass-through payment at ASP+6 percent. Rather, implantable biologicals that are eligible for device pass-through payment will be paid based on the charges-adjusted-to-cost methodology used for all pass-through device categories.

We considered three alternatives for the pass-through evaluation process and payment methodology for implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice), as indicated in the CY 2010 OPPS/ASC proposed rule (74 FR 35411). The first alternative we considered was to make no change to the current pass-through evaluation process and payment methodology for

implantable biologicals that are surgically inserted or implanted. We did not select this alternative because this approach would continue the separate pass-through evaluation processes and payment methodologies for implantable biologicals and implantable nonbiological devices that are sometimes used for the same clinical indications, and where the implantable biologicals are often FDA-approved as devices. Moreover, under our current policy, implantable biologicals could potentially have two periods of pass-through payment, one as a biological and one as a device. We believe that it is most appropriate for a product to be eligible for a single period of OPPS pass-through payment, rather than a period of device pass-through payment and a period of drug or biological pass-through payment.

The second alternative we considered was to add a criterion requiring the demonstration of substantial clinical improvement to the biological pass-through evaluation process in order for a biological to be approved for pass-through payment. This alternative would provide pass-through payment only for those biologicals that demonstrate clinical superiority, consistent with the pass-through evaluation process for devices, and ensure that a product could receive only one period of pass-through payment. We did not choose this alternative because this approach would continue the different pass-through payment methods for implantable biological and nonbiological devices. Pass-through payment for biologicals is made at ASP+6 percent as required for drug and biological pass-through payment, while pass-through devices are paid at charges adjusted to cost. Therefore, this second alternative would result in continued inconsistent pass-through payment methodologies for biological and nonbiological devices that may substitute for one another.

The third alternative we considered and the one we are adopting for CY 2010 is to provide that, beginning in CY 2010, the pass-through evaluation process and pass-through payment methodology for implantable biologicals that are surgically inserted or implanted (through a surgical incision or a natural orifice) be the device pass-through process and payment methodology only. As we discuss in section V.A.4. of this final rule with comment period, after consideration of the public comments we received on the proposed rule, we are adopting this alternative because we believe that a consistent pass-through payment policy is to evaluate all such devices, both biological and

nonbiological, under the device pass-through process. We believe that implantable biologicals that function as and may be substitutes for implantable devices are most similar to devices because of their required surgical insertion or implantation, and that it would be appropriate to evaluate them as devices because they share significant clinical similarity with implantable nonbiological devices.

b. Alternatives Considered for Payment of the Acquisition and Pharmacy Overhead Costs of Drugs and Biologicals That Do Not Have Pass-Through Status

For CY 2010, we are finalizing a transition payment for separately payable drugs and biologicals at ASP+4 percent, which will continue to represent combined payment for both the acquisition and pharmacy overhead costs of separately payable drugs and biologicals. As discussed in detail in section V.B.3. of this final rule with comment period, we are redistributing \$200 million total of packaged drug cost (\$150 million of the pharmacy overhead cost currently attributed to coded packaged drugs and biologicals with an ASP and \$50 million of the estimated pharmacy overhead cost currently attributed to uncoded packaged drugs and biologicals) to separately payable drugs and biologicals to provide an adjustment for the pharmacy overhead costs of these separately payable products. As a result, we are proportionally reducing the cost of packaged drugs and biologicals that is included in the separate payment for procedural APCs to offset the \$200 million adjustment to provide payment for separately payable drugs and biologicals at ASP+4 percent. We received favorable public comments on our proposal to redistribute cost within drugs and biologicals to adjust payment for separately payable drugs and biologicals. The public commenters also agreed that our estimated total cost for all drugs and biologicals in our claims data is accurate. Therefore, we are redistributing a portion of pharmacy overhead cost in the CY 2010 final rule claims data from some packaged drugs and biologicals to separately payable drugs and biologicals. A redistribution within drug cost maintains the estimated total cost of drugs and biologicals under the OPPS.

We considered three alternatives for payment of the acquisition and pharmacy overhead costs of drugs and biologicals that do not have pass-through status for CY 2010. The first alternative we considered was to continue our standard policy of comparing the estimated aggregate cost

of separately payable drugs and biologicals in our claims data to the estimated aggregate ASP dollars for separately payable drugs and biologicals, using the ASP as a proxy for average acquisition cost, to calculate the estimated percent of ASP that would serve as the proxy for the combined acquisition and pharmacy overhead costs of separately payable drugs and biologicals (70 FR 68642). Under this standard methodology, using July 2009 ASP information and updated final rule costs derived from CY 2008 OPPS claims data, we estimated the combined acquisition and pharmacy overhead costs of separately payable drugs and biologicals to be ASP-3 percent. We also determined the combined acquisition and pharmacy overhead costs of coded packaged drugs and biologicals with an ASP to be 258 percent of ASP. As discussed in section V.B.3. of this final rule with comment period, we did not choose this alternative because we believe that this analysis indicates that our standard drug payment methodology has the potential to "compress" the calculated costs of separately payable drugs and biologicals and inflate the calculated costs of packaged drugs and biologicals to some degree. Further, we recognize that the attribution of pharmacy overhead costs to packaged or separately payable drugs and biologicals through our standard drug payment methodology of a combined payment for acquisition and pharmacy overhead costs also depends on the determination of separate or packaged payment for all drugs and biologicals each year based on our annual drug packaging threshold. Changes to the packaging threshold and the packaged status of drugs or biologicals may result in changes to the estimated combined acquisition and pharmacy overhead costs of drugs and biologicals that do not reflect actual changes in hospital pharmacy overhead cost for those products.

The second alternative we considered was to adopt the February 2009 APC Panel recommendation to accept the pharmacy stakeholders' recommended methodology for payment of drugs and biologicals that do not have pass-through status. This recommended methodology would establish ASP+6 percent as the cost of packaged drugs and biologicals, including all pharmacy overhead costs; establish ASP+6 percent as the acquisition cost of separately payable drugs and biologicals with some overhead cost included; and reallocate the residual cost of packaged drugs and biologicals currently reflected in the claims data across three categories of

pharmacy overhead cost that would then be paid separately for each administration of separately payable drugs and biologicals in CY 2010. The pharmacy stakeholders recommended that we pay the pharmacy overhead amount specific to the overhead category to which a drug or biological is assigned, in addition to the ASP+6 percent payment for the separately payable drug or biological, each time a separately payable drug or biological is administered. We refer readers to section V.B.3. of this final rule with comment period for a more detailed discussion of the pharmacy stakeholders' recommended methodology. We did not choose this alternative because we do not believe that ASP+6 percent would pay appropriately for the acquisition and pharmacy overhead costs of packaged drugs. We believe the amount of redistribution of pharmacy overhead costs from packaged to separately payable drugs and biologicals incorporated in the recommendation of the pharmacy stakeholders would be too great. In addition, we do not believe that it would be appropriate to establish separate payment for pharmacy overhead costs, thereby unbundling payment for the acquisition and overhead costs of separately payable drugs and biologicals when hospitals report a single charge for these products that represents both types of costs. For these reasons, we are not accepting the APC Panel recommendation to adopt the pharmacy stakeholders' recommended methodology.

The third alternative we considered and the one we selected for CY 2010 is to make a transition payment for nonpass-through separately payable drugs and biologicals at ASP+4 percent, which will continue to represent a combined payment for both the acquisition costs of separately payable drugs and biologicals and the pharmacy overhead costs applicable to these products. We also are reducing the cost of packaged drugs and biologicals that is included in the payment for procedural APCs to offset the \$200 million adjustment to payment for separately payable drugs and biologicals. The \$200 million consists of \$150 million (one-third of the pharmacy overhead cost) from the cost of coded packaged drugs and biologicals with an ASP and \$50 million from the uncoded packaged drug and biological cost. To model this policy for the CY 2010 final rule with comment period, we reduced the cost of coded packaged drugs and biologicals with an ASP by 24 percent (based on final rule data; the reduction was 27

percent based on proposed rule data) and the cost of uncoded packaged drugs and biologicals by 8 percent when we calculated the median costs of the CY 2010 procedural APCs. We chose this transition alternative because we believe that it provides an appropriate redistribution of pharmacy overhead costs associated with drugs and biologicals and is consistent with the principles of a prospective payment system.

c. Alternatives Considered for the Physician Supervision of Hospital Outpatient Services

We are revising or further defining several policies related to the physician supervision of services in the HOPD for CY 2010. We refer readers to section XII.D. of this final rule with comment period for the full discussion of these policies. Specifically, for the CY 2010 OPSS, we are revising our existing policy that requires direct supervision to be provided by a physician to allow, when statutorily permitted under the Social Security Act, specified nonphysician practitioners to supervise the hospital outpatient therapeutic services that they are able to personally perform within their State scope of practice and hospital-granted privileges. We note that section 144(a)(1) of Public Law 100-275 imposes strict requirements for the direct physician supervision of PR, CR, and ICR services and gives us no flexibility to modify the requirement beyond direct physician supervision by a doctor of medicine or a doctor of osteopathy. We also are establishing a policy for hospital outpatient therapeutic services furnished in the main hospital buildings or in on-campus provider-based departments (PBDs) that "direct supervision" means that the supervisory practitioner must be on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. In addition, we are establishing in regulations a policy that applies the MPFS physician supervision requirements for diagnostic tests to all hospital outpatient diagnostic tests performed directly by the hospital or under arrangement.

We considered three alternatives for the physician supervision of hospital outpatient services for CY 2010 in the CY 2010 OPSS/ASC proposed rule (74 FR 35412 through 35413). The first alternative we considered was to make no changes to the existing supervision policies for hospital outpatient therapeutic and diagnostic services and to provide no new policy guidance in this area. This approach would require

hospitals to ensure that only physicians supervise services that may currently be ordered or performed by nonphysician practitioners within their State scope of practice and hospital-granted privileges. Hospitals would not receive payment for outpatient services for which they were unable to provide supervision by a physician. In addition, there could continue to be confusion regarding what "direct supervision" means for services provided in an area of the hospital that may not be a PBD of the hospital. Lastly, there would be potential for misunderstanding regarding the appropriate level of physician supervision required for hospital outpatient diagnostic services without a clearly stated policy, codified in regulations, that would apply the same level of physician supervision to all hospital outpatient diagnostic services, whether provided directly or under arrangement, as applies to those services currently furnished in physicians' offices and independent diagnostic testing facilities. We did not choose this alternative because we believe that it is important to address the issues outlined above, including areas of potential confusion or limited current policy guidance, to ensure that hospitals are able to comply with the hospital outpatient supervision requirements while providing access to care for Medicare beneficiaries.

The second alternative we considered was to permit specified nonphysician practitioners to supervise the hospital outpatient therapeutic services that they are able to personally perform within their State scope of practice and hospital-granted privileges, but to make no changes that would provide clearer statements of policy regarding other concerns raised by hospitals regarding physician supervision for hospital outpatient therapeutic and diagnostic services. We did not choose this alternative because we believe it is important to clearly specify the policies that apply to the supervision of both therapeutic and diagnostic services in all hospital outpatient settings in order to ensure the safety and effectiveness of hospital outpatient services furnished to Medicare beneficiaries.

The third alternative we considered and the one we selected for CY 2010 was to revise our existing policy to permit specified nonphysician practitioners, when statutorily permitted in the Social Security Act, to supervise the services that they are able to personally perform within their State scope of practice and hospital-granted privileges; to establish a specific definition of "direct supervision" for hospital outpatient therapeutic services

furnished in the hospital or in on-campus PBDs that was consistent for services furnished by the hospital on-campus; and to apply the MPFS supervision requirements for diagnostic tests to all hospital outpatient diagnostic tests provided directly by the hospital or under arrangement. We selected this alternative because we believe that it is appropriate that, unless the Act imposes strict requirements for the direct supervision of certain services, such as PR, CR, and ICR services, a licensed nonphysician practitioner who may bill and be paid by Medicare for the practitioner's professional services should be able to supervise the therapeutic services that he or she may personally perform within his or her State scope of practice and hospital-granted privileges. Furthermore, we believe that it is necessary and appropriate to establish consistent and operationally feasible policies regarding the supervision requirements for hospital outpatient therapeutic and diagnostic services in order to ensure safe and effective health care services for Medicare beneficiaries. We refer readers to section XII.D. of this final rule with comment period for a complete discussion of the final physician supervision policies.

2. Limitations of Our Analysis

The distributional impacts presented here are the projected effects of the CY 2010 policy changes on various hospital groups. We post on the CMS Web site our hospital-specific estimated payments for CY 2010 with the other supporting documentation for this final rule with comment period. To view the hospital-specific estimates, we refer readers to the CMS Web site at: <http://www.cms.hhs.gov/HospitalOutpatientPPS/>. Select "regulations and notices" from the left side of the page and then select "CMS-1414-P" from the list of regulations and notices. The hospital-specific file layout and the hospital-specific file are listed with the other supporting documentation for this final rule with comment period. We show hospital-specific data only for hospitals whose claims were used for modeling the impacts shown in Table 73 below. We do not show hospital-specific impacts for hospitals whose claims we were unable to use. We refer readers to section II.A.2. of this final rule with comment period for a discussion of the hospitals whose claims we do not use for ratesetting and impact purposes.

We estimate the effects of the individual policy changes by estimating payments per service, while holding all other payment policies constant. We use

the best data available, but do not attempt to predict behavioral responses to our policy changes. In addition, we do not make adjustments for future changes in variables such as service volume, service mix, or number of encounters. As we have done in previous rules, in the CY 2010 OPFS/ASC proposed rule (74 FR 35413), we solicited public comment and information about the anticipated effects of our proposed changes on providers and our methodology for estimating them.

We received many public comments on the proposed changes to payment policies and to proposed payment rates for the CY 2010 OPFS. We have summarized these public comments and provided our responses to them in other sections of this final rule with comment period as part of our discussions of the specific topics to which the comments pertained. We did not receive any public comments on our methodology for estimating the anticipated effects of our proposed changes on providers or other parties.

3. Estimated Effects of This Final Rule With Comment Period on Hospitals

Table 73 below shows the estimated impact of this final rule with comment period on hospitals. Historically, the first line of the impact table, which estimates the change in payments to all hospitals, has always included cancer and children's hospitals, which are held harmless to their pre-BBA payment-to-cost ratio. We also are including CMHCs in the first line that includes all providers because we included CMHCs in our weight scaler estimate.

We present separate impacts for CMHCs in Table 73 because CMHCs are paid under only two APCs for services under the OPFS: APC 0172 (Level 1 Partial Hospitalization (3 services)) and APC 0173 (Level II Partial Hospitalization (4 or more services)). We note that CMHCs are also a different provider type. We discuss the impact on CMHCs in section XXI.B.4. of this final rule with comment period.

The estimated increase in the total payments made under the OPFS is limited by the increase to the conversion factor set under the methodology in the statute. The distributional impacts presented do not include assumptions about changes in volume and service mix. The enactment of Public Law 108-173 on December 8, 2003, provided for the additional payment outside of the budget neutrality requirement for wage indices for specific hospitals reclassified under section 508. Public Law 108-173 extended section 508 reclassifications

through September 30, 2008. Section 124 of Public Law 110-275 further extended section 508 reclassifications through September 30, 2009. The amounts attributable to these reclassifications are incorporated into the CY 2009 estimates in Table 73.

Table 73 shows the estimated redistribution of hospital and CMHC payments among providers as a result of APC reconfiguration and recalibration; wage indices; the combined impact of the APC recalibration, wage effects, and the market basket update to the conversion factor; and, finally, estimated redistribution considering all payments for CY 2010 relative to all payments for CY 2009, including the impact of changes in the outlier threshold, expiring section 508 wage indices, and changes to the pass-through payment estimate. We did not model an explicit budget neutrality adjustment for the rural adjustment for SCHs because we are not making any changes to the policy for CY 2010. Because the updates to the conversion factor, including the update of the market basket and the subtraction of additional money dedicated to pass-through payment for CY 2010, are applied uniformly across services, observed redistributions of payments in the impact table for hospitals largely depend on the mix of services furnished by a hospital (for example, how the APCs for the hospital's most frequently furnished services will change), and the impact of the wage index changes on the hospital. However, total payments made under this system and the extent to which this final rule with comment period will redistribute money during implementation also will depend on changes in volume, practice patterns, and the mix of services billed between CY 2009 and CY 2010 by various groups of hospitals, which CMS cannot forecast.

Overall, the final OPFS rates for CY 2010 will have a positive effect for providers paid under the OPFS, resulting in a 1.9 percent estimated increase in Medicare payments. Removing cancer and children's hospitals, because their payments are held harmless to the pre-BBA ratio between payment and cost, and CMHCs suggests that these changes also will result in a 1.9 percent estimated increase in Medicare payments to all other hospitals.

To illustrate the impact of the final CY 2010 changes, our analysis begins with a baseline simulation model that uses the final CY 2009 weights, the FY 2009 final post-reclassification OPFS wage indices, and the final CY 2009 conversion factor. Column 2 in Table 73

shows the independent effect of the changes resulting from the reclassification of services among APC groups and the recalibration of APC weights, based on 12 months of CY 2008 OPSS hospital claims data and the most recent cost report data. We modeled the effect of the APC recalibration changes for CY 2010 by varying only the weights (the final CY 2009 weights versus the final CY 2010 weights calculated using the service mix and volume in the CY 2008 claims used for this final rule with comment period) and calculating the percent difference in weight. Column 2 also reflects the effect of the changes resulting from the APC reclassification and recalibration changes and any changes in multiple procedure discount patterns or conditional packaging that occur as a result of the changes in the relative magnitude of payment weights.

Column 3 reflects the independent effects of the updated wage indices, including the application of budget neutrality for the rural floor policy on a statewide basis. We did not model a budget neutrality adjustment for the rural adjustment for SCHs because we are making no changes to the policy for CY 2010. We modeled the independent effect of updating the wage indices by varying only the wage indices, holding APC relative weights, service mix, and the rural adjustment constant and using the CY 2010 scaled weights and a CY 2009 conversion factor that included a budget neutrality adjustment for the effect of changing the wage indices between CY 2009 and CY 2010.

Column 4 demonstrates the combined "budget neutral" impact of APC recalibration (that is, Column 2), the wage index update (that is, Column 3), as well as the impact of updating the conversion factor with the market basket update. We modeled the independent effect of the budget neutrality adjustments and the market basket update by using the weights and wage indices for each year, and using a CY 2009 conversion factor that included the market basket update and a budget neutrality adjustment for differences in wage indices.

Finally, Column 5 depicts the full impact of the CY 2010 policies on each hospital group by including the effect of all the changes for CY 2010 (including the APC reconfiguration and recalibration shown in Column 2) and comparing them to all estimated payments in CY 2009 (these CY 2009 estimated payments include the payments resulting from the non-budget neutral increases to wage indices under section 508 of Pub. L. 108–173 as extended by Pub. L. 110–275). Column 5 shows the combined budget neutral

effects of Columns 2 through 4, plus the impact of the change to the fixed-dollar outlier threshold from \$1,800 to \$2,175; the impact of the expiration of section 508 reclassifications; the change in the HOP QDRP payment reduction for the small number of hospitals in our impact model that failed to meet the reporting requirements; and the impact of increasing the estimate of the percentage of total OPSS payments dedicated to transitional pass-through payments. We discuss our CY 2010 change to the outlier threshold in section II.F. of this final rule with comment period. Of the 85 hospitals that failed to meet the HOP QDRP reporting requirements for the full CY 2009 update (and assumed, for modeling purposes, to be the same number for CY 2010), we included 28 in our model because they had both CY 2008 claims data and recent cost report data. We estimate that the cumulative effect of all changes for CY 2010 will increase payments to all providers by 1.9 percent for CY 2010. We modeled the independent effect of all changes in Column 5 using the final weights for CY 2009 and the final weights for CY 2010. We used the final conversion factor for CY 2009 of \$66.059 and the final CY 2010 conversion factor of \$67.406. Column 5 also contains simulated outlier payments for each year. We used the charge inflation factor used in the FY 2010 IPPS/RV 2010 LTCH PPS final rule of 6.86 percent (1.0686) to increase individual costs on the CY 2008 claims, and we used the most recent overall CCR in the July 2009 Outpatient Provider-Specific File (OPSF) (74 FR 44010). Using the CY 2008 claims and a 6.86 percent charge inflation factor, we currently estimate that outlier payments for CY 2009, using a multiple threshold of 1.75 and a fixed-dollar threshold of \$1,800, will be approximately 1.03 percent of total payments. Outlier payments of 1.03 percent are incorporated in the CY 2009 comparison in Column 5. We used the same set of claims and a charge inflation factor of 14.18 percent (1.1418) and the CCRs in the July 2009 OPSF, with an adjustment of 0.9880 to reflect relative changes in cost and charge inflation between CY 2008 and CY 2010, to model the CY 2010 outliers at 1.0 percent of total payments using a multiple threshold of 1.75 and a fixed-dollar threshold of \$2,175.

Column 1: Total Number of Hospitals

The first line in Column 1 in Table 73 shows the total number of providers (4,222), including cancer and children's hospitals and CMHCs for which we were able to use CY 2008 hospital outpatient claims to model CY 2009 and

CY 2010 payments, by classes of hospitals. We excluded all hospitals for which we could not accurately estimate CY 2009 or CY 2010 payment and entities that are not paid under the OPSS. The latter entities include CAHs, all-inclusive hospitals, and hospitals located in Guam, the U.S. Virgin Islands, Northern Mariana Islands, American Samoa, and the State of Maryland. This process is discussed in greater detail in section II.A. of this final rule with comment period. At this time, we are unable to calculate a disproportionate share (DSH) variable for hospitals not participating in the IPPS. Hospitals for which we do not have a DSH variable are grouped separately and generally include freestanding psychiatric hospitals, rehabilitation hospitals, and long-term care hospitals. We show the total number (3,942) of OPSS hospitals, excluding the hold-harmless cancer and children's hospitals and CMHCs, on the second line of the table. We excluded cancer and children's hospitals because section 1833(t)(7)(D) of the Act permanently holds harmless cancer hospitals and children's hospitals to a proportion of their pre-BBA payment relative to their pre-BBA costs and, therefore, we removed them from our impact analyses. We show the isolated impact on 221 CMHCs in the last row of the impact table and discuss that impact separately below.

Column 2: APC Changes Due to Reassignment and Recalibration

This column shows the combined effects of the reconfiguration, recalibration, and other policies (such as setting payment for separately payable drugs and biologicals at ASP+4 percent with an accompanying reduction in the amount of cost associated with packaged drugs and biologicals, payment for brachytherapy sources based on median unit cost, and changes in payment for PHP services. Overall, we estimate that changes in APC reassignment and recalibration across all services paid under the OPSS will increase payments to urban hospitals by 0.1 percent. We estimate that both large and other urban hospitals will see an increase of 0.1 percent, all attributable to recalibration. We estimate that urban hospitals billing fewer than 11,000 lines for OPSS services will experience decreases of 0.2 to 1.0 percent, while urban hospitals billing 11,000 or more lines for OPSS services will see no change or an increase of 0.1 percent.

Overall, we estimate that rural hospitals will experience a decrease of 0.1 percent as a result of changes to the APC structure. We estimate that rural

hospitals of all bed sizes will experience no change or decreases of up to 0.2 percent as a result of APC recalibration. We estimate that rural hospitals that report fewer than 5,000 lines for OPSS services will experience a decrease of 0.8 percent, while rural hospitals that report more than 5,000 lines for OPSS services will see decreases of 0.1 percent to 0.4 percent.

Among teaching hospitals, we estimate that the largest observed impact resulting from APC recalibration will include an increase of 0.2 percent for major teaching hospitals and a increase of 0.1 percent for minor teaching hospitals.

Classifying hospitals by type of ownership suggests that proprietary and governmental hospitals will see no change, and voluntary hospitals will see an estimated increase of 0.1 percent.

Finally, we estimate that hospitals for which DSH payments are not available will experience decreases of 2.5 to 2.7 percent that are largely attributable to the reduction in PHP payment for APC 0172. We estimate that most other classes of hospitals will not experience any change from CY 2009 to CY 2010 or will experience a modest increase.

Column 3: New Wage Indices and the Effect of the Rural Adjustment

This column estimates the impact of applying the FY 2010 IPPS wage indices for the CY 2010 OPSS. We are not changing the rural payment adjustment for CY 2010. We estimate that the combination of updated wage data and statewide application of rural floor budget neutrality will redistribute payment among regions. We also updated the list of counties qualifying for the section 505 out-migration adjustment. Overall, we estimate that urban hospitals will not experience any change from CY 2009 to CY 2010, and that rural hospitals will experience a decrease of 0.1 percent as a result of the updated wage indices. However, we estimate that hospitals in rural New England States and rural West South Central States will experience decreases of 0.9 and 0.7 percent, respectively. We estimate that urban and rural Mountain States will experience increases of 0.6 percent.

Column 4: All Budget Neutrality Changes and Market Basket Update

We estimate that the addition of the market basket update of 2.1 percent will mitigate any negative impacts on hospital payments for CY 2010 created by the budget neutrality adjustments made in Columns 2 and 3, with the exception of hospitals not paid under the IPPS, including freestanding

psychiatric, rehabilitation, and long-term care hospitals, that we estimate will continue to experience decreases of between -0.6 and -0.7 percent. In general, Column 4 shows that all hospitals will experience an estimated increase of 2.1 percent, attributable to the 2.1 percent market basket increase.

Overall, we estimate that these changes will increase payments to urban hospitals by 2.2 percent. We estimate that large urban hospitals will experience an increase of 2.3 percent, and "other" urban hospitals will experience a 2.1 percent increase.

Overall, we estimate that rural hospitals will experience a 1.9 percent increase as a result of the market basket update and other budget neutrality adjustments. We estimate that rural hospitals that bill less than 5,000 lines of OPSS services will experience an increase of 1.5 percent and that rural hospitals that bill more than 5,000 lines of OPSS services will experience increases of 1.8 to 1.9 percent.

Among teaching hospitals, we estimate that the observed impacts resulting from the market basket update and other budget neutrality adjustments will include an increase of 2.4 and 2.2 percent, respectively, for major and minor teaching hospitals.

Classifying hospitals by type of ownership suggests that voluntary, proprietary, and governmental hospitals will experience estimated increases of 2.2 percent, 2.1 percent, and 2.0 percent, respectively.

Column 5: All Changes for CY 2010

Column 5 compares all estimated changes for CY 2010 to estimated final payment for CY 2009, including the expiration of the reclassifications under section 508, the change in the outlier threshold, payment reductions for hospitals that failed to meet the HOP QDRP reporting requirements, and the difference in pass-through estimates that are not included in the combined percentages shown in Column 4. This column includes estimated payment for a handful of hospitals receiving reduced payment because they did not meet their hospital outpatient quality measure reporting requirements; however, we estimate that the anticipated change in payment between CY 2009 and CY 2010 for these hospitals will be negligible. Overall, we estimate that providers will experience an increase of 1.9 percent under this final rule with comment period in CY 2010 relative to total spending in CY 2009. The projected 1.9 percent increase for all providers in Column 5 of Table 73 reflects the 2.1 percent market basket increase, less 0.03 percent for the

change in the pass-through estimate between CY 2009 and CY 2010, less 0.03 percent for the difference in estimated outlier payments between CY 2009 (1.03 percent) and CY 2010 (1.0 percent), and less 0.14 percent due to the expiration of the special, non-budget neutral wage index payments made under section 508. We estimate that when we exclude cancer and children's hospitals (which are held harmless to their pre-OPSS costs) and CMHCs, the increase remains at 1.9 percent.

We estimate that the combined effect of all changes for CY 2010 will increase payments to urban hospitals by 2.0 percent. We estimate that large urban hospitals will experience a 2.1 percent increase, while "other" urban hospitals will experience an increase of 1.9 percent. We estimate that urban hospitals that bill less than 5,000 lines of OPSS services will experience an increase of 1.2 percent, and we estimate that all urban hospitals that bill more than 5,000 lines of OPSS services will experience increases between 1.9 percent and 2.0 percent.

Overall, we estimate that rural hospitals will experience a 1.6 percent increase as a result of the combined effects of all changes for CY 2010. We estimate that rural hospitals that bill less than 5,000 lines of OPSS services will experience an increase of 1.3 percent and rural hospitals that bill greater than 5,000 lines of OPSS services will experience increases ranging from 1.4 percent to 1.8 percent.

Among teaching hospitals, we estimate that the impacts resulting from the combined effects of all changes will include an increase of 2.0 percent for major teaching hospitals and an increase of 1.9 percent for minor teaching hospitals.

Classifying hospitals by type of ownership, we estimate that proprietary hospitals will gain 2.0 percent, governmental hospitals will experience an increase of 1.8 percent, and voluntary hospitals will experience an increase of 1.9 percent.

4. Estimated Effects of This Final Rule With Comment Period on CMHCs

The last row of the impact analysis in Table 73 demonstrates the impact on CMHCs. We modeled this impact assuming that CMHCs will continue to provide the same number of days of PHP care, with each day having either three services or four or more services, as seen in the CY 2008 claims data. We excluded days with one or two services. Using these assumptions, we estimate that there would be a 5.0 percent decrease in payments to CMHCs due to these APC policy changes (shown in

Column 2). The relative weight for low intensity partial hospitalization APC 0172 (Level 1 Partial Hospitalization (3 services)) declines between CY 2009 and CY 2010 under this final rule with comment period. CMHCs perform a greater proportion of low intensity partial hospitalization days than freestanding psychiatric hospitals. Table 73 demonstrates our estimate that hospitals not paid under the IPPS for which a disproportionate patient percentage is not available (DSH Not

Available), consisting largely of freestanding psychiatric hospitals, will experience a more moderate decline in payments of 2.7 percent. Psychiatric hospitals provide a greater proportion of APC 0173 (Level II Partial Hospitalization (4 or more services)) for which the relative weight increases between CY 2009 and CY 2010 under this final rule with comment period.

Column 3 shows that the estimated impact of adopting the CY 2010 wage index values will be no change in

payments to CMHCs. We note that all providers paid under the OPPIs, including CMHCs, will receive a 2.1 percent market basket increase. Combining this market basket increase, along with changes in APC policy for CY 2010 and the CY 2010 wage index updates, and changes in outlier payments, we estimate that the combined impact on CMHCs for CY 2010 will be a 3.0 percent decrease.

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**TABLE 73.—ESTIMATED IMPACT OF THE FINAL CY 2010 HOSPITAL
OUTPATIENT PROSPECTIVE PAYMENT SYSTEM**

		Number of Hospitals (1)	APC Recalibration (2)	New Wage Index and Rural Adjustment (3)	Comb (cols 2,3) with Market Basket Update (4)	All Changes (5)
ALL PROVIDERS *		4,222	0.0	0.0	2.1	1.9
ALL HOSPITALS		3,942	0.0	0.0	2.1	1.9
(excludes hospitals permanently held harmless and CMHCs)						
URBAN HOSPITALS						
	LARGE URBAN (GT 1 MILL.)	1,608	0.1	0.1	2.3	2.1
	OTHER URBAN (LE 1 MILL.)	1,343	0.1	0.0	2.1	1.9
RURAL HOSPITALS						
	SOLE COMMUNITY	391	-0.1	0.0	2.0	1.6
	OTHER RURAL	600	-0.1	-0.2	1.7	1.6
BEDS (URBAN)						
	0 - 99 BEDS	1,005	0.0	0.1	2.1	2.0
	100-199 BEDS	881	0.0	0.0	2.1	1.8
	200-299 BEDS	460	0.1	0.0	2.2	2.1
	300-499 BEDS	407	0.1	0.0	2.1	1.9
	500 + BEDS	198	0.2	0.1	2.4	2.1
BEDS (RURAL)						
	0 - 49 BEDS	359	-0.2	0.0	1.8	1.6
	50- 100 BEDS	375	0.0	-0.1	1.9	1.7
	101- 149 BEDS	148	-0.2	-0.2	1.7	1.6
	150- 199 BEDS	64	-0.1	-0.1	1.8	1.5
	200 + BEDS	45	-0.1	-0.1	1.9	1.4
VOLUME (URBAN)						
	LT 5,000 Lines	604	-1.0	0.1	1.2	1.2
	5,000 - 10,999 Lines	174	-0.2	0.1	2.0	1.9
	11,000 - 20,999 Lines	250	0.0	0.0	2.0	1.9

		Number of Hospitals (1)	APC Recalibration (2)	New Wage Index and Rural Adjustment (3)	Comb (cols 2,3) with Market Basket Update (4)	All Changes (5)
	21,000 - 42,999 Lines	514	0.0	0.1	2.2	2.0
	GT 42,999 Lines	1,409	0.1	0.0	2.2	2.0
VOLUME (RURAL)						
	LT 5,000 Lines	80	-0.8	0.2	1.5	1.3
	5,000 - 10,999 Lines	89	-0.3	0.1	1.9	1.8
	11,000 - 20,999 Lines	186	-0.4	0.1	1.8	1.4
	21,000 - 42,999 Lines	323	-0.1	-0.2	1.9	1.7
	GT 42,999 Lines	313	-0.1	-0.2	1.9	1.6
REGION (URBAN)						
	NEW ENGLAND	150	0.1	0.6	2.8	2.4
	MIDDLE ATLANTIC	370	-0.1	0.3	2.3	1.9
	SOUTH ATLANTIC	455	0.1	-0.3	1.9	1.8
	EAST NORTH CENT.	470	0.0	-0.2	2.0	1.7
	EAST SOUTH CENT.	188	0.1	-0.2	2.0	1.9
	WEST NORTH CENT.	197	0.2	0.1	2.4	2.3
	WEST SOUTH CENT.	486	0.1	-0.2	2.0	1.9
	MOUNTAIN	192	0.2	0.6	3.0	2.9
	PACIFIC	393	0.1	0.0	2.3	2.1
	PUERTO RICO	50	0.2	-0.2	2.1	2.3
REGION (RURAL)						
	NEW ENGLAND	24	-0.1	-0.9	1.1	1.1
	MIDDLE ATLANTIC	68	-0.1	0.5	2.5	2.2
	SOUTH ATLANTIC	169	-0.2	-0.3	1.6	1.5
	EAST NORTH CENT.	130	0.0	-0.1	2.0	1.6
	EAST SOUTH CENT.	177	-0.1	-0.1	1.9	1.8
	WEST NORTH CENT.	104	0.0	0.1	2.2	1.5
	WEST SOUTH CENT.	216	-0.4	-0.7	1.0	0.9
	MOUNTAIN	72	0.1	0.6	2.8	2.5
	PACIFIC	31	0.1	0.0	2.2	1.7
TEACHING STATUS						
	NON-TEACHING	2,952	0.0	-0.1	2.0	1.9

		Number of Hospitals (1)	APC Recalibration (2)	New Wage Index and Rural Adjustment (3)	Comb (cols 2,3) with Market Basket Update (4)	All Changes (5)
	MINOR	706	0.1	0.0	2.2	1.9
	MAJOR	284	0.2	0.1	2.4	2.0
DSH PATIENT PERCENT						
	0	9	0.1	0.1	2.3	2.2
	GT 0 - 0.10	401	0.2	0.1	2.3	2.1
	0.10 - 0.16	434	0.1	0.0	2.2	1.8
	0.16 - 0.23	763	0.1	0.0	2.1	1.9
	0.23 - 0.35	974	0.0	0.0	2.1	1.9
	GE 0.35	744	0.0	0.1	2.2	2.1
	DSH NOT AVAILABLE **	617	-2.7	-0.1	-0.7	-0.8
URBAN TEACHING/DSH						
	TEACHING & DSH	894	0.1	0.1	2.3	2.0
	NO TEACHING/DSH	1,468	0.1	0.0	2.1	2.0
	NO TEACHING/NO DSH	8	0.1	0.1	2.3	2.2
	DSH NOT AVAILABLE**	581	-2.5	-0.2	-0.6	-0.6
TYPE OF OWNERSHIP						
	VOLUNTARY	2,100	0.1	0.0	2.2	1.9
	PROPRIETARY	1,257	0.0	-0.1	2.1	2.0
	GOVERNMENT	585	0.0	-0.1	2.0	1.8
CMHCs		221	-5.0	0.0	-2.9	-3.0

Column (1) shows total hospitals.

Column (2) shows the impact of changes resulting from the reclassification of HCPCS codes among APC groups and the recalibration of APC weights based on 2008 hospital claims data.

Column (3) shows the budget neutral impact of updating the wage index by applying the FY 2010 hospital inpatient wage index. We did not make any changes to the rural adjustment.

Column (4) shows the impact of all budget neutrality adjustments and the addition of the market basket update.

Column (5) shows the additional adjustments to the conversion factor resulting from a change in the pass-through estimate and adds outlier payments. This column also shows the expiration of section 508 wages on September 30, 2009.

*These 4,222 providers include children and cancer hospitals, which are held harmless to pre-BBA payments, and CMHCs.

**Complete DSH numbers are not available for providers that are not paid under IPPS, including rehabilitation, freestanding psychiatric, and long-term care hospitals.

5. Estimated Effect of This Final Rule With Comment Period Rule on Beneficiaries

For services for which the beneficiary pays a copayment of 20 percent of the payment rate, the beneficiary share of payment will increase for services for which the OPSS payments will rise and will decrease for services for which the OPSS payments will fall. For example, for a service assigned to Level IV Needle Biopsy/Aspiration Except Bone Marrow (APC 0037) in the CY 2009 OPSS, the national unadjusted copayment is \$228.76, and the minimum unadjusted copayment is \$178.60. For CY 2010, the national unadjusted copayment for APC 0037 will be \$228.76, the same rate in effect for CY 2009. The minimum unadjusted copayment for APC 0037 will be \$208.97 or 20 percent of the CY 2010 national unadjusted payment rate for APC 0037 of \$1,044.81. The minimum unadjusted copayment will rise because the payment rate for APC 0037 will rise for CY 2010. In all cases, the statute limits beneficiary liability for copayment for a procedure to the hospital inpatient deductible for the applicable year. The CY 2009 hospital inpatient deductible is \$1,068. The CY 2010 hospital inpatient deductible is \$1,100.

In order to better understand the impact of changes in copayment on

beneficiaries, we modeled the percent change in total copayment liability using CY 2008 claims. We estimate, using the claims of the 4,222 hospitals and CMHCs on which our modeling is based, that total beneficiary liability for copayments will decline as an overall percentage of total payments, from 23.0 percent in CY 2009 based on updated claims data for this final rule with comment period to 22.6 percent in CY 2010.

6. Conclusion

The changes in this final rule with comment period will affect all classes of hospitals and CMHCs. We estimated that some classes of hospitals will experience significant gains and others less significant gains, but all classes of hospitals will experience positive updates in OPSS payments in CY 2010 with one exception. We estimate that hospitals not paid under the IPPS will see an overall decrease in payment of 0.6 to 0.8 percent because they are largely freestanding psychiatric hospitals that bill mostly PHP services. As we discuss in substantial detail in section X. of this final rule with comment period, payment for APC 0172 will decline for CY 2010 and, therefore, we estimate that payments to CMHCs and hospitals that furnish mostly PHP services will also decline. In general, we

estimate that CMHCs will experience an overall decline of 3.0 percent in total payment due to the recalibration of the payment rates.

Table 73 demonstrates the estimated distributional impact of the OPSS budget neutrality requirements that will result in a 1.9 percent increase in payments for all services paid under the OPSS in CY 2010, after considering all changes to APC reconfiguration and recalibration, as well as the market basket increase, wage index changes, estimated payment for outliers, and changes to the pass-through payment estimate. The accompanying discussion, in combination with the rest of this final rule with comment period, constitutes a regulatory impact analysis.

7. Accounting Statement

As required by OMB Circular A-4 (available at <http://www.whitehouse.gov/omb/circulars/a004/a-4.pdf>), in Table 74, we have prepared an accounting statement showing the CY 2010 estimated hospital OPSS incurred benefit impact associated with the CY 2010 hospital outpatient market basket update shown in this final rule with comment period based on the baseline for the 2009 Trustees Report. All estimated impacts are classified as transfers.

TABLE 74—ACCOUNTING STATEMENT: CY 2010 ESTIMATED HOSPITAL OPSS INCURRED BENEFIT IMPACT ASSOCIATED WITH THE CY 2010 HOSPITAL OUTPATIENT MARKET BASKET UPDATE

Category	Transfers
Annualized Monetized Transfers	\$0.5 billion.
From Whom to Whom	Federal Government to outpatient hospitals and other providers who received payment under the hospital OPSS.
Total	\$0.5 billion.

C. Effects of ASC Payment System Changes in This Final Rule With Comment Period Rule

On August 2, 2007, we published in the **Federal Register** the final rule for the revised ASC payment system, effective January 1, 2008 (72 FR 42470). In that final rule, we adopted the methodologies to set payment rates for covered ASC services to implement the revised payment system so that it would be designed to result in budget neutrality as required by section 626 of Public Law 108-173; established that the OPSS relative payment weights would be the basis for payment and that we would update the system annually as part of the OPSS rulemaking cycle; and provided that the revised ASC payment rates would be phased-in over 4 years. During the 4-year transition to

full implementation of the ASC payment rates, payments for surgical procedures paid in ASCs in CY 2007 are made using a blend of the CY 2007 ASC payment rate and the ASC payment rate calculated according to the ASC standard ratesetting methodology for the applicable transitional year. In CY 2009, we are paying ASCs using a 50/50 blend, in which payment is calculated by adding 50 percent of the CY 2007 ASC rate for a surgical procedure on the CY 2007 ASC list of covered surgical procedures and 50 percent of the CY 2009 ASC rate calculated according to the ASC standard ratesetting methodology for the same procedure. For CY 2010, we are transitioning the blend to a 25/75 blend of the CY 2007 ASC rate and the ASC payment rate calculated according to the ASC

standard ratesetting methodology. Beginning in CY 2011, we will pay ASCs for all covered surgical procedures, including those on the CY 2007 ASC list, at the ASC payment rates calculated according to the ASC standard ratesetting methodology. Payment for procedures that were not included on the ASC list of covered surgical procedures in CY 2007 is not subject to the transitional payment methodology.

ASC payment rates are calculated by multiplying the ASC conversion factor by the ASC relative payment weight. As discussed fully in section XV. of this final rule with comment period, we set the CY 2010 ASC relative payment weights by scaling CY 2010 ASC relative payment weights by the ASC scaler of 0.9567. These weights take into consideration the 25/75 blend for the

third year of transitional payment for certain services. If there were no transition, the scaler for the CY 2010 relative payment weights would be 0.9338. The estimated effects of the updated relative payment weights on payment rates during this transitional period are varied and are reflected in the estimated payments displayed in Tables 75 and 78 below.

The CY 2010 ASC conversion factor was calculated by adjusting the CY 2009 ASC conversion factor to account for changes in the pre-floor and pre-reclassified hospital wage indices between CY 2009 and CY 2010 and by applying the CY 2010 CPI-U of a 1.2 percent increase. The CY 2010 ASC conversion factor is \$41,873.

1. Alternatives Considered

Alternatives to the changes we are making and the reasons that we have chosen the options are discussed throughout this final rule with comment period. Some of the major ASC issues discussed in this final rule with comment period and the options considered are discussed below.

a. Alternatives Considered for Office-Based Procedures

According to our final policy for the revised ASC payment system, we designate as office-based those procedures that are added to the ASC list of covered surgical procedures in CY 2008 or later years and that we determine are predominantly performed in physicians' offices based on consideration of the most recent available volume and utilization data for each individual procedure HCPCS code and/or, if appropriate, the clinical characteristics, utilization, and volume of related HCPCS codes. We establish payment for procedures designated as office-based at the lesser of the MPFS nonfacility PE RVU amount or the ASC rate developed according to the standard methodology of the revised ASC payment system.

In developing this final rule with comment period, we reviewed the full CY 2008 utilization data for all surgical procedures added to the ASC list of covered surgical procedures in CY 2008 or later years and for those procedures for which the office-based designation is temporary in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68730 through 68733). Based on that review, and as discussed in section XV.C.1.b. of this final rule with comment period, we are newly designating six existing surgical procedures as office-based and making permanent the office-based designations of four existing surgical procedures that

have temporary office-based designations in CY 2009. We also are providing temporary office-based designations for 16 CY 2010 procedures reported with new or substantially revised CPT codes and continuing the temporary office-based designations for 6 procedures that were temporarily office-based in CY 2009. We considered two alternatives in developing this policy.

The first alternative we considered was to make no change to the procedure payment designations. This would mean that we would pay for the 6 procedures we are designating as permanently office-based and the 16 procedures we are newly designating as temporarily office-based at an ASC payment rate calculated according to the standard ratesetting methodology of the revised ASC payment system and for the 10 procedures with temporary office-based designations for 2009 according to the office-based methodology. We did not select this alternative because our analysis of the data and our clinical review indicated that all 10 procedures we are designating permanently office-based as well as the 22 procedures that we are designating temporarily office-based could be considered to be predominantly performed in physicians' offices. Consistent with our final policy adopted in the August 2, 2007 final rule (72 FR 42509), we were concerned that making payments at the standard ASC payment rate for the 6 procedures newly designated as office-based and 16 new procedures designated as temporarily office-based could create financial incentives for the procedures to shift from physicians' offices to ASCs for reasons unrelated to clinical decisions regarding the most appropriate setting for surgical care. Further, consistent with our policy, we believe that when adequate data become available to make permanent determinations about procedures with temporary office-based designations, maintaining the temporary designation is no longer appropriate.

The second alternative we considered and the one we selected for CY 2010 is to designate six additional procedures as office-based for CY 2010 and to make permanent the office-based designations of four of the procedures with temporary office-based designations in CY 2009. We also are designating 16 new procedures described by new or substantially revised CPT codes for CY 2010 as temporarily office-based and continuing to designate 6 procedures as temporarily office-based in CY 2010. We chose this alternative because our claims data and clinical review indicate that these procedures could be considered to be predominantly

performed in physicians' offices. We believe that designating these procedures as office-based, which results in the CY 2010 ASC payment rate for these procedures potentially being capped at the CY 2010 physicians' office rate (that is, the MPFS nonfacility PE RVU amount), if applicable, is an appropriate step to ensure that Medicare payment policy does not create financial incentives for such procedures to shift unnecessarily from physicians' offices to ASCs, consistent with our final policy adopted in the August 2, 2007 final rule.

b. Alternatives Considered for Covered Surgical Procedures

According to our final policy for the revised ASC payment system, we designate as covered all surgical procedures that we determine would not be expected to pose a significant risk to beneficiary safety or would not be expected to require an overnight stay when performed on Medicare beneficiaries in an ASC.

In developing this final rule with comment period, we reviewed the clinical characteristics and full CY 2008 utilization data, if applicable, for all procedures reported by Category III CPT codes implemented July 1, 2009, and surgical procedures that were excluded from ASC payment for CY 2009. In response to public comments received on the CY 2009 OPPS/ASC proposed rule, we stated in the CY 2009 OPPS/ASC final rule with comment period (73 FR 68724) that, as we developed the CY 2010 OPPS/ASC proposed rule and final rule with comment period, we would perform a comprehensive review of the APCs in order to identify potentially inconsistent ASC treatment of procedures assigned to a single APC under the OPPS. Thus, for this final rule with comment period, we examined surgical procedures that were excluded from the CY 2009 ASC list of covered surgical procedures and the APCs to which they were assigned under the OPPS. Based on this review, we identified 26 surgical procedures that meet the criteria for inclusion on the ASC list of covered surgical procedures, and we are adding those procedures to the list for CY 2010 payment, in addition to the 2 new surgical procedures described by Category III CPT codes that were new for July 2009, and that we determined were appropriate for addition to the ASC list. We considered two alternatives in developing this policy.

The first alternative we considered was to make no change to the ASC list of covered surgical procedures for CY 2010. We did not choose this alternative because our analysis of data and clinical

review indicated that the 28 procedures we are designating as covered surgical procedures for CY 2010 would not be expected to pose a significant risk to beneficiary safety in ASCs and would not be expected to require an overnight stay. Consistent with our final policy, we were concerned that by continuing to exclude them from the list of ASC covered surgical procedures, we may unnecessarily limit beneficiaries' access to the services in the most clinically appropriate settings.

The second alternative we considered and the one we selected for CY 2010 was to designate 28 additional procedures as ASC covered surgical procedures for CY 2010. We chose this alternative because our claims data and clinical review indicate that these procedures would not be expected to pose a significant risk to beneficiary safety and would not be expected to require an overnight stay, and thus they meet the criteria for inclusion on the list of ASC covered surgical procedures. We believe that adding these procedures to the list of covered surgical procedures is an appropriate step to ensure that beneficiary access to services is not limited unnecessarily.

2. Limitations of Our Analysis

Presented here are the projected effects of the changes for CY 2010 on Medicare payment to ASCs. A key limitation of our analysis is our inability to predict changes in ASC service mix between CY 2008 and CY 2010 with precision. We believe that the net effect on Medicare expenditures resulting from the CY 2010 changes will be small in the aggregate for all ASCs. However, such changes may have differential effects across surgical specialty groups as ASCs continue to adjust to the payment rates based on the policies of the revised ASC payment system. We are unable to accurately project such changes at a disaggregated level. Clearly, individual ASCs will experience changes in payment that differ from the aggregated estimated impacts presented below.

3. Estimated Effects of This Final Rule With Comment Period on Payments to ASCs

Some ASCs are multispecialty facilities that perform the gamut of surgical procedures, from excision of lesions to hernia repair to cataract extraction; others focus on a single specialty and perform only a limited range of surgical procedures, such as eye, digestive system, or orthopedic procedures. The combined effect on an individual ASC of the update to the CY 2010 payments will depend on a

number of factors, including, but not limited to, the mix of services the ASC provides, the volume of specific services provided by the ASC, the percentage of its patients who are Medicare beneficiaries, and the extent to which an ASC provides different services in the coming year. The following discussion presents tables that display estimates of the impact of the CY 2010 update to the revised ASC payment system on Medicare payments to ASCs, assuming the same mix of services as reflected in our CY 2008 claims data. Table 75 depicts the estimated aggregate percent change in payment by surgical specialty or ancillary items and services group by comparing estimated CY 2009 payments to estimated CY 2010 payments, and Table 76 shows a comparison of estimated CY 2009 payments to estimated CY 2010 payments for procedures that we estimate will receive the most Medicare payment in CY 2010.

Table 75 shows the estimated effects on aggregate Medicare payments under the revised ASC payment system by surgical specialty or ancillary items and services group. We have aggregated the surgical HCPCS codes by specialty group, grouped all HCPCS codes for covered ancillary items and services into a single group, and then estimated the effect on aggregated payment for surgical specialty and ancillary items and services groups, considering separately the CY 2010 transitional rates and the ASC payment rates calculated according to the ASC standard ratesetting methodology that would apply in CY 2010 if there were no transition. The groups are sorted for display in descending order by estimated Medicare program payment to ASCs. The following is an explanation of the information presented in Table 75.

- **Column 1—Surgical Specialty or Ancillary Items and Services Group** indicates the surgical specialty into which ASC procedures are grouped or the ancillary items and services group which includes all HCPCS codes for covered ancillary items and services. To group surgical procedures by surgical specialty, we used the CPT code range definitions and Level II HCPCS codes and Category III CPT codes, as appropriate, to account for all surgical procedures to which the Medicare program payments are attributed.

- **Column 2—Estimated ASC Payments** were calculated using CY 2008 ASC utilization (the most recent full year of ASC utilization) and CY 2009 ASC payment rates. The surgical specialty and ancillary items and services groups are displayed in

descending order based on estimated CY 2009 ASC payments.

- **Column 3—Estimated CY 2010 Percent Change With Transition (25/75 Blend)** is the aggregate percentage increase or decrease, compared to CY 2009, in Medicare program payment to ASCs for each surgical specialty or ancillary items and services group that is attributable to updates to the ASC payment rates for CY 2010 under the scaled, 25/75 blend of the CY 2007 ASC payment rates and the CY 2010 ASC payment rates calculated according to the ASC standard ratesetting methodology.

- **Column 4—Estimated CY 2010 Percent Change Without Transition (Fully Implemented)** is the aggregate percentage increase or decrease in Medicare program payment to ASCs for each surgical specialty or ancillary items and services group that would be attributable to updates to ASC payment rates for CY 2010 compared to CY 2009 if there were no transition period to the fully implemented payment rates. The percentages appearing in Column 4 are presented only as illustrative comparisons to the percentage changes under the transition policy in Column 3. We are not eliminating or modifying the policy for a 4-year transition that was finalized in the August 2, 2007 final rule (72 FR 42519).

As seen in Table 75, we estimate that the update to ASC rates for CY 2010 will result in no change in aggregate payment amounts for eye and ocular adnexa procedures and in aggregate decreases of 4 percent in payment amounts for both digestive system and nervous system procedures. As shown in Column 4 in the table, we estimate that if there were no transitional payment for these three surgical specialty groups in CY 2010, aggregate payments would decrease by 1 percent for eye and ocular adnexa procedures and by 10 and 6 percent for digestive and nervous system procedures, respectively.

Generally, for the surgical specialty groups that account for less ASC utilization and spending, we estimate that the payment effects of the CY 2010 update are positive. We estimate that ASC payments for procedures in those surgical specialties will increase in CY 2010 with the 25/75 transitional payment rates and, in the absence of the transition, will increase even more. For instance, we estimate that, in the aggregate, payment for integumentary system procedures will increase by 13 percent under the CY 2010 rates and by 20 percent if there were no transition. We estimate similar effects for genitourinary, cardiovascular,

musculoskeletal, respiratory, hematologic and lymphatic systems, and auditory system procedures as well.

An estimated increase in aggregate payment for the specialty group does not mean that all procedures in the group will experience increased payment rates. For example, the substantial estimated increase for CY 2010 for integumentary procedures is likely due to the significant median cost increase for APC 0137 (Level V Skin Repair) under the OPSS. The highest volume procedure in the integumentary surgical specialty group, described by CPT code 15823 (Blepharoplasty, upper eyelid; with excessive skin weighting

down lid), is assigned to that APC under the OPSS. In contrast, the estimated increased payments at the surgical specialty group level may be due to decreased payments for some of the most frequently provided procedures in the group and the moderating effect of the sometimes substantial payment increases for the less frequently performed procedures within the surgical specialty group.

Also displayed in Table 75 is a separate estimate of Medicare ASC payments for the group of separately payable covered ancillary items and services. We estimate that aggregate payments for these items and services

will remain the same for CY 2010. The payment estimates for the covered surgical procedures include the costs of packaged ancillary items and services. In prior years' rules, we did not have ASC payment data for covered ancillary items and services because prior to CY 2008, they were paid under other fee schedules or packaged into payment for the covered surgical procedures. Beginning with the CY 2010 proposed rule and this final rule with comment period, we have utilization data for those services as well as for all of the covered surgical procedures provided in ASCs under the revised payment system.

TABLE 75—ESTIMATED IMPACT OF THE FINAL CY 2010 ASC PAYMENT SYSTEM ON AGGREGATE CY 2010 MEDICARE PROGRAM PAYMENTS UNDER THE 25/75 TRANSITION BLEND AND WITHOUT A TRANSITION, BY SURGICAL SPECIALTY OR ANCILLARY ITEMS AND SERVICES GROUP

Surgical specialty group	Estimated CY 2009 ASC payments (in millions)	Estimated CY 2010 percent change with transition (25/75 blend)	Estimated CY 2010 percent change without transition (fully implemented)
(1)	(2)	(3)	(4)
Total	3,077	1	1
Eye and ocular adnexa	1,405	0	-1
Digestive system	731	-4	-10
Nervous system	365	-4	-5
Musculoskeletal system	285	15	29
Genitourinary system	112	10	17
Integumentary system	106	13	20
Respiratory system	27	24	37
Cardiovascular system	20	17	27
Ancillary items and services	15	0	-1
Auditory system	8	9	17
Hematologic & lymphatic systems	3	22	40

Table 76 below shows the estimated impact of the updates to the revised ASC payment system on aggregate ASC payments for selected surgical procedures during CY 2010 with and without the transitional blended rate. The table displays 30 of the procedures receiving the greatest estimated CY 2009 aggregate Medicare payments to ASCs. The HCPCS codes are sorted in descending order by estimated CY 2009 program payment.

- Column 1—HCPCS code.
- Column 2—Short Descriptor of the HCPCS code.
- Column 3—Estimated CY 2009 Allowed Charges were calculated using CY 2008 ASC utilization (the most recent full year of ASC utilization) and the CY 2009 ASC payment rates. The estimated CY 2009 allowed charges are expressed in millions of dollars.
- Column 4—Estimated CY 2010 Percent Change with Transition (25/75 Blend) reflects the percent differences between the estimated ASC payment for

CY 2009 and the estimated payment for CY 2010 based on the update, incorporating a 25/75 blend of the CY 2007 ASC payment rate and the CY 2010 ASC payment rate calculated according to the ASC standard ratesetting methodology.

- Column 5—Estimated CY 2010 Percent Change without Transition (Fully Implemented) reflects the percent differences between the estimated ASC payment for CY 2009 and the estimated payment for CY 2010 based on the update if there were no transition period to the fully implemented payment rates. The percentages appearing in Column 5 are presented as a comparison to the percentage changes under the transition policy in Column 4 for informational purposes only. We are not eliminating or modifying the policy for the 4-year transition that was finalized in the August 2, 2007 final rule (72 FR 42519). As displayed in Table 76, 24 of the 30 procedures with the greatest estimated aggregate CY 2009 Medicare payment

are included in the 3 surgical specialty groups that are estimated to account for the most Medicare payment to ASCs in CY 2009, specifically eye and ocular adnexa, digestive system, and nervous system surgical groups. Consistent with the estimated payment effects on the surgical specialty groups displayed in Table 75, the estimated effects of the CY 2010 update on ASC payment for individual procedures in year 3 of the transition shown in Table 76 are varied. Aggregate ASC payments for many of the most frequently furnished ASC procedures will decrease as the transitional rates more closely align the individual procedure relative ASC payment weights with the relativity of payments under the OPSS.

The ASC procedure for which the most Medicare payment is estimated to be made in CY 2009 is the cataract removal procedure reported with CPT code 66984 (Extracapsular cataract removal with insertion of intraocular lens prosthesis (one stage procedure),

manual or mechanical technique (e.g., irrigation and aspiration or phacoemulsification)). We estimate that the update to the ASC rates will result in a negligible payment decrease for this procedure in CY 2010. The estimated payment effects on the three other eye and ocular adnexa procedures included in Table 76 will be slightly positive or negative, but for CPT code 66821 (Discission of secondary membranous cataract (opacified posterior lens capsule and/or anterior hyaloid); laser surgery (e.g., YAG laser) (one or more stages)), the estimated CY 2010 payment decrease will be 10 percent, significantly greater than the decreases estimated for any of the other eye and ocular adnexa procedures shown.

We estimate that the transitional payment rates for all but 1 of the 9 digestive system procedures included in Table 76 will decrease by 5 to 8 percent in CY 2010. Those estimated decreases are consistent with decreases in the previous 2 years under the revised ASC payment system and are expected because, under the previous ASC payment system, the payment rates for many high volume endoscopy

procedures were almost the same as the payments for the procedures under the OPSS.

The estimated effects of the CY 2010 update on the 9 nervous system procedures for which the most Medicare ASC payment is estimated to be made in CY 2009 will be variable. Our estimates indicate that the CY 2010 update will result in payment decreases of 4 percent or less for 4 of the 9 procedures and in more substantial decreases for 2 others. We estimate that the greatest decreases will be for the add-on procedure described by CPT code 64484 (Injection, anesthetic agent and/or steroid, transforaminal epidural; lumbar or sacral, each additional level) and for the procedure described by CPT code 63685 (Insertion or replacement of spinal neurostimulator pulse generator or receiver, direct or inductive coupling), which we estimate to have 19 and 9 percent payment decreases, respectively, in CY 2010. In contrast, the three nervous system procedures for which we estimate positive effects on CY 2010 payments, CPT code 63650 (Percutaneous implantation of neurostimulator electrode array,

epidural); CPT code 64721 (Neuroplasty and/or transposition; median nerve at carpal tunnel); and CPT code 64622 (Destruction by neurolytic agent, paravertebral facet joint nerve; lumbar or sacral, single level), are estimated to have substantial payment increases of 10, 13, and 6 percent, respectively.

The estimated payment effects for most of the remaining procedures listed in Table 76 will be positive. For example, the CY 2010 transitional payment rates for musculoskeletal CPT codes 29880 (Arthroscopy, knee, surgical; with meniscectomy (medial AND lateral, including any meniscal shaving)) and 29881 (Arthroscopy, knee, surgical; with meniscectomy (medial OR lateral, including any meniscal shaving)) are estimated to increase 17 percent over the CY 2009 transitional payment rates. We estimate that musculoskeletal procedures will account for a greater percentage of CY 2010 Medicare ASC spending as we estimate that payment for procedures in that surgical specialty group will increase under the revised payment system in CY 2010.

TABLE 76—ESTIMATED IMPACT OF THE FINAL CY 2010 ASC PAYMENT SYSTEM ON AGGREGATE PAYMENTS FOR SELECTED PROCEDURES

HCPCS code *	Short descriptor	Estimated CY 2009 allowed charges (in mil)	Estimated CY 2010 percent change with transition (25/75 blend)	Estimated CY 2010 percent change without transition (fully implemented)
(1)	(2)	(3)	(4)	(5)
66984	Cataract surg w/iol, 1 stage	1,064	0	-2
43239	Upper gi endoscopy, biopsy	164	-6	-13
45380	Colonoscopy and biopsy	133	-5	-11
45378	Diagnostic colonoscopy	124	-5	-11
45385	Lesion removal colonoscopy	96	-5	-11
66821	After cataract laser surgery	71	-10	-20
62311	Injct spine l/s (cd)	69	-4	-8
66982	Cataract surgery, complex	62	0	-2
64483	Inj foramen epidural l/s	57	-4	-8
15823	Revision of upper eyelid	35	15	21
45384	Lesion remove colonoscopy	33	-5	-11
G0105	Colorectal scrn; hi risk ind	33	-8	-17
G0121	Colon ca scrn not hi rsk ind	32	-8	-16
29881	Knee arthroscopy/surgery	25	16	30
63650	Implant neuroelectrodes	25	10	14
43235	Uppr gi endoscopy, diagnosis	24	2	1
64721	Carpal tunnel surgery	23	13	24
52000	Cystoscopy	22	-5	-9
29880	Knee arthroscopy/surgery	20	17	30
63685	Insrt/redo spine n generator	18	-9	-8
29826	Shoulder arthroscopy/surgery	17	28	54
62310	Injct spine c/t	15	-4	-8
67904	Repair eyelid defect	15	0	2
28285	Repair hammertoe	15	14	25
29827	Arthroscop rotator cuff repr	14	22	42
64622	Destr paravertebrl nerve l/s	14	6	14
64484	Inj foramen epidural add-on	13	-19	-38
43248	Uppr gi endoscopy/guide wire	12	-6	-13
64623	Destr paravertebral n add-on	12	-4	-8

TABLE 76—ESTIMATED IMPACT OF THE FINAL CY 2010 ASC PAYMENT SYSTEM ON AGGREGATE PAYMENTS FOR SELECTED PROCEDURES—Continued

HCPCS code *	Short descriptor	Estimated CY 2009 allowed charges (in mil)	Estimated CY 2010 percent change with transition (25/75 blend)	Estimated CY 2010 percent change without transition (fully implemented)
(1)	(2)	(3)	(4)	(5)
26055	Incise finger tendon sheath	12	12	21

* Note that HCPCS codes deleted for CY 2010 are not displayed in this table.

The previous ASC payment system served as an incentive to ASCs to focus on providing procedures for which they determined Medicare payments would support their continued operation. We note that, historically, the ASC payment rates for many of the most frequently performed procedures in ASCs were similar to the OPPS payment rates for the same procedures. Conversely, procedures with ASC payment rates that were substantially lower than the OPPS rates have historically been performed least often in ASCs. We believed that the revised ASC payment system will encourage greater efficiency in ASCs and will promote significant increases in the breadth of surgical procedures performed in ASCs because it distributes payments across the entire spectrum of covered surgical procedures based on a coherent system of relative weights that are related to the clinical and facility resource requirements of those procedures.

The CY 2008 claims data that we used to develop the CY 2010 ASC payment system relative weights and rates reflect the first year of utilization under the revised payment system. Although the changes in the claims data are not large, the data reflect increased Medicare ASC spending for procedures that were newly added to the ASC list in CY 2008. Our estimates based on CY 2008 data indicate that for CY 2010 there would be especially noticeable increases in spending for genitourinary and cardiovascular procedures, compared to the previous ASC payment system.

4. Estimated Effects of This Final Rule With Comment Period on Beneficiaries

We estimate that the CY 2010 update to the ASC payment system will be generally positive for beneficiaries with respect to the new procedures that we are adding to the ASC list of covered surgical procedures and for those that we are designating as office-based for CY 2010. First, except for screening colonoscopy and flexible sigmoidoscopy procedures, the ASC coinsurance rate

for all procedures is 20 percent. This contrasts with procedures performed in HOPDs, where the beneficiary is responsible for copayments that range from 20 percent to 40 percent of the procedure payment. Second, ASC payment rates under the revised payment system are lower than payment rates for the same procedures under the OPPS; therefore, the beneficiary coinsurance amount under the ASC payment system almost always will be less than the OPPS copayment amount for the same services. (The only exceptions will be if the ASC coinsurance amount exceeds the inpatient deductible. The statute requires that copayment amounts under the OPPS not exceed the inpatient deductible.) For new procedures that we are adding to the ASC list of covered surgical procedures in CY 2010, as well as for procedures already included on the list, and that are furnished in an ASC rather than the HOPD setting, the beneficiary coinsurance amount will be less than the OPPS copayment amount. Furthermore, the additions to the ASC list of covered surgical procedures will provide beneficiaries access to more surgical procedures in ASCs. Beneficiary coinsurance for services migrating from physicians' offices to ASCs may decrease or increase under the revised ASC payment system, depending on the particular service and the relative payment amounts for that service in the physician's office compared to the ASC. However, for those additional procedures that we are designating as office-based in CY 2010, the beneficiary coinsurance amount will be no greater than the beneficiary coinsurance in the physician's office.

In addition, as finalized in the August 2, 2007 final rule (72 FR 42520), in CY 2010, the third year of the 4-year transition to the ASC payment rates calculated according to the ASC standard ratesetting methodology of the revised ASC payment system, ASC payment rates for a number of

commonly furnished ASC procedures will continue to be reduced, resulting in lower beneficiary coinsurance amounts for these ASC services in CY 2010.

5. Conclusion

The updates to the ASC payment system for CY 2010 will affect each of the approximately 5,000 ASCs currently approved for participation in the Medicare program. The effect on an individual ASC will depend on its mix of patients, the proportion of the ASC's patients that are Medicare beneficiaries, the degree to which the payments for the procedures offered by the ASC are changed under the revised payment system, and the extent to which the ASC provides a different set of procedures in the coming year.

The CY 2010 update to the revised ASC payment system includes a payment update of 1.2 percent that we estimate will result in a greater amount of Medicare expenditures in CY 2010 than was estimated to be made in CY 2009. We estimate that the update to the revised ASC payment system, including the addition of surgical procedures to the list of covered surgical procedures, will have a modest effect on Medicare expenditures compared to the estimated level of Medicare expenditures in CY 2009.

6. Accounting Statement

As required by OMB Circular A-4 (available at <http://www.whitehouse.gov/omb/circulars/a004/a-4.pdf>), in Table 77 below, we have prepared an accounting statement showing the classification of the expenditures associated with the statutorily authorized 1.2 percent update to the CY 2010 revised ASC payment system, based on the provisions of this final rule with comment period and the baseline spending estimates for ASCs in the 2009 Medicare Trustees Report. This table provides our best estimate of Medicare payments to suppliers as a result of the final update to the CY 2010 ASC payment system, as presented in this

final rule with comment period. All expenditures are classified as transfers.

TABLE 77—ACCOUNTING STATEMENT: CLASSIFICATION OF ESTIMATED EXPENDITURES FROM CY 2009 TO CY 2010 AS A RESULT OF THE CY 2010 UPDATE TO THE REVISED ASC PAYMENT SYSTEM

Category	Transfers
Annualized Monetized Transfers. From Whom to Whom.	\$33 Million. Federal Government to Medicare Providers and Suppliers.
Total	\$33 Million.

D. Effects of Requirements for Hospital Reporting of Quality Data for Annual Hospital Payment Update

In section XVI. of the CY 2009 OPPTS/ASC final rule with comment period (73 FR 68758), we discussed our requirements for subsection (d) hospitals to report quality data under the HOP QDRP in order to receive the full payment update for CY 2010. In section XVI. of this final rule with comment period, we established additional policies affecting the CY 2010 and CY 2011 HOP QDRP. We estimate that about 83 hospitals may not receive the full payment update in CY 2010. Most of these hospitals are either small rural or small urban hospitals. However, at this time, information is not available to determine the precise number of hospitals that do not meet the requirements for the full hospital market basket increase for CY 2010. We also estimate that 83 hospitals may not receive the full payment update in CY 2011.

In section XVI.E.3.a. of this final rule with comment period, for the CY 2011 payment update, as part of the validation process, we are requiring hospitals to submit paper copies of requested medical records to a designated contractor within the required timeframe. Failure to submit requested documentation can result in a 2 percentage point reduction in a hospital's update, but the failure to pass the validation itself would not. Of the 83 hospitals that we estimate will not receive the full payment update for CY 2011, we estimate that no more than 20 hospitals would fail the validation documentation submission requirement for the CY 2011 payment update.

For the CY 2011 payment update, our validation sample size is estimated to be about 7,300 medical records. We estimate that this requirement will cost

hospitals approximately 12 cents per page for copying and approximately \$4.00 per chart for postage. We have found, based on experience, that an average sized outpatient medical chart is approximately 30 pages. We estimate that the total cost to the impacted hospitals will be approximately \$55,480, with a maximum expected cost of \$152 for an individual hospital based upon an expected maximum of 20 selected records; the expected minimum will be \$0.00 if no records were selected from a hospital. We believe that this cost is minimal, compared with the 2.0 percentage point HOP QDRP component of the annual payment update at risk. CMS does not plan to reimburse hospitals for copying and mailing costs. This validation requirement is necessary so that CMS has all the information it needs to validate the accuracy of hospital submitted data abstracted from paper medical records.

In section XVI.E.3.b. of this final rule with comment period, we did not, at this time, adopt our proposal in the CY 2010 OPPTS/ASC proposed rule (74 FR 35403) to expand the CY 2011 validation requirement for the CY 2012 payment update. Instead, we will consider the public comments we received on that proposal, as well as any analyses we conduct of the CY 2011 validation process, and propose a CY 2012 validation process as a part of the CY 2011 OPPTS/ASC rulemaking. We believe that this approach will give HOP QDRP hospitals additional experience with the validation process and allow these hospitals sufficient time to prepare for the CY 2012 validation.

However, we noted in the CY 2010 OPPTS/ASC proposed rule (74 FR 35424) that we expected our proposal to validate data submitted by 800 hospitals for purposes of the CY 2012 HOP QDRP payment determination would not have changed the number of hospitals that fail the validation requirement from CY 2011. For CY 2011, and under our proposal for CY 2012 in the CY 2010 OPPTS/ASC proposed rule, we stated that we would calculate the validation matches for CY 2011 (we note, however, that the validation results will not affect the CY 2011 payment update) and CY 2012 by assessing whether the overall measure data submitted by the hospital matches the independently reabstracted measure data. We believe that this methodology will make it easier for hospitals to satisfy the validation requirement than if we calculate the percent agreement between what the hospital submitted and what the CMS-designated contractor independently reabstracted for each individual data element that was submitted. In addition,

for the CY 2012 payment update, in the CY 2010 OPPTS/ASC proposed rule, we proposed to validate data for only 800 hospitals out of the approximately 3,400 HOP QDRP participating hospitals. As a result, we believe that the effect of our proposed validation process for CY 2012 would have been minimal in terms of the number of hospitals that would not meet all program requirements. In the CY 2010 OPPTS/ASC proposed rule (74 FR 35424), we stated that of the 83 hospitals that we estimated would not have received the full payment update for CY 2012, we estimated that approximately 20 hospitals would have failed to meet our proposed CY 2012 validation requirements.

E. Executive Order 12866

In accordance with the provisions of Executive Order 12866, this final rule with comment period rule was reviewed by the OMB.

List of Subjects

42 CFR Part 410

Health facilities, Health professions, Laboratories, Medicare, Rural areas, X-rays.

42 CFR Part 416

Health facilities, Health professions, Medicare, Reporting and recordkeeping requirements.

42 CFR Part 419

Hospitals, Medicare, Reporting and recordkeeping requirements.

■ For reasons stated in the preamble of this document, the Centers for Medicare & Medicaid Services is amending 42 CFR chapter IV as set forth below:

PART 410—SUPPLEMENTARY MEDICAL INSURANCE (SMI) BENEFITS

■ 1. The authority citation for part 410 continues to read as follows:

Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

■ 2. Section 410.27 is amended by—

- a. Revising the section heading.
 - b. Revising the introductory text of paragraph (a) and paragraph (a)(1).
 - c. Revising paragraph (e).
 - d. Revising paragraph (f).
 - e. Adding new paragraph (g).
- The revisions and additions read as follows:

§ 410.27 Outpatient hospital or CAH services and supplies incident to a physician or nonphysician practitioner service: Conditions.

(a) Medicare Part B pays for hospital or CAH services and supplies furnished

incident to a physician or nonphysician practitioner service to outpatients, including drugs and biologicals that cannot be self-administered, if—

(1) They are furnished—

(i) By or under arrangements made by the participating hospital or CAH, except in the case of a SNF resident as provided in § 411.15(p) of this chapter;

(ii) As an integral though incidental part of a physician's or nonphysician practitioner's services;

(iii) In the hospital or CAH or in a department of the hospital or CAH, as defined in § 413.65 of this subchapter; and

(iv) Under the direct supervision of a physician or a nonphysician practitioner as specified in paragraph (f) of this section. Nonphysician practitioners may directly supervise services that they may personally furnish in accordance with State law and all additional requirements, including those specified in §§ 410.71, 410.73, 410.74, 410.75, 410.76, and 410.77.

(A) For services furnished in the hospital or CAH or in an on-campus outpatient department of the hospital or CAH, as defined in § 413.65 of this subchapter, "direct supervision" means that the physician or nonphysician practitioner must be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician or nonphysician practitioner must be present in the room when the procedure is performed. For pulmonary rehabilitation, cardiac rehabilitation, and intensive cardiac rehabilitation services, direct supervision must be furnished by a doctor of medicine or osteopathy, as specified in §§ 410.47 and 410.49, respectively.

(B) For services furnished in an off-campus outpatient department of the hospital or CAH, as defined in § 413.65 of this subchapter, "direct supervision" means the physician or nonphysician practitioner must be present in the off-campus provider-based department of the hospital or CAH and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician or nonphysician practitioner must be present in the room when the procedure is performed. For pulmonary rehabilitation, cardiac rehabilitation, and intensive cardiac rehabilitation services, direct supervision must be furnished by a doctor of medicine or osteopathy, as specified in §§ 410.47 and 410.49, respectively.

* * * * *

(e) Services furnished by an entity other than the hospital or CAH are subject to the limitations specified in § 410.42(a).

(f) For purposes of this section, "nonphysician practitioner" means a clinical psychologist, licensed clinical social worker, physician assistant, nurse practitioner, clinical nurse specialist, or certified nurse-midwife.

(g) For purposes of this section, "in the hospital or CAH" means areas in the main building(s) of the hospital or CAH that are under the ownership, financial, and administrative control of the hospital or CAH; that are operated as part of the hospital or CAH; and for which the hospital or CAH bills the services furnished under the hospital's or CAH's CMS Certification Number.

■ 3. Section 410.28 is amended by revising paragraph (e) to read as follows:

§ 410.28 Hospital or CAH diagnostic services furnished to outpatients: Conditions.

* * * * *

(e) Medicare Part B makes payment under section 1833(t) of the Act for diagnostic services furnished by or under arrangements made by the participating hospital, only when the diagnostic services are furnished under the appropriate level of physician supervision specified by CMS in accordance with the definitions in § 410.32(b)(3)(i), (b)(3)(ii), and (b)(3)(iii). Under general supervision, the training of the nonphysician personnel who actually perform the diagnostic procedure and the maintenance of the necessary equipment and supplies are the continuing responsibility of the facility. In addition—

(1) For services furnished directly or under arrangement in the hospital or in an on-campus outpatient department of the hospital, as defined in § 413.65 of this subchapter, "direct supervision" means that the physician must be present on the same campus and immediately available to furnish assistance and direction throughout the performance of the procedure. It does not mean that the physician must be present in the room when the procedure is performed. For this purpose, the definition of "in the hospital" is as specified in § 410.27(g).

(2) For services furnished directly or under arrangement in an off-campus outpatient department of the hospital, as defined in § 413.65 of this subchapter, "direct supervision" means the physician must be present in the off-campus provider-based department of the hospital and immediately available to furnish assistance and direction throughout the performance of the

procedure. It does not mean that the physician must be present in the room when the procedure is performed.

(3) For services furnished under arrangement in nonhospital locations, "direct supervision" means the definition specified in § 410.32(b)(3)(ii).

* * * * *

PART 416—AMBULATORY SURGICAL SERVICES

■ 4. The authority citation for Part 416 continues to read as follows:

Authority: Secs. 1102 and 1871 of the Social Security Act (42 U.S.C. 1302 and 1395hh).

■ 5. Section 416.30 is amended by revising paragraph (f)(2) to read as follows:

§ 416.30 Terms of the agreement with CMS.

* * * * *

(f) * * *

(2) The ASC participates and is paid only as an ASC.

* * * * *

PART 419—PROSPECTIVE PAYMENT SYSTEM FOR HOSPITAL OUTPATIENT DEPARTMENT SERVICES

■ 6. The authority citation for part 419 continues to read as follows:

Authority: Secs. 1102, 1833(t), and 1871 of the Social Security Act (42 U.S.C. 1302, 1395(t), and 1395(hh)).

■ 7. Section 419.64 is amended by adding new paragraphs (a)(4)(iii) and (a)(4)(iv), to read as follows:

§ 419.64 Transitional pass-through payments: Drugs and biologicals.

(a) * * *

(4) * * *

(iii) A biological that is not surgically implanted or inserted into the body.

(iv) A biological that is surgically implanted or inserted into the body, for which pass-through payment as a biological is made on or before December 31, 2009.

* * * * *

■ 8. Section 419.66 is amended by revising paragraph (b)(4)(iii) to read as follows:

§ 419.66 Transitional pass-through payments: Medical devices.

* * * * *

(b) * * *

(4) * * *

(iii) A material that may be used to replace human skin (for example, a biological skin replacement material or synthetic skin replacement material).

* * * * *

■ 9. Section 419.70 is amended by revising the heading of paragraph (d)(5) to read as follows:

§ 419.70 Transitional adjustments to limit decline in payments.

* * * * *

(d) * * *

(5) *Temporary treatment for small sole community hospitals on or after*

January 1, 2009 and through December 31, 2009. * * *

* * * * *

(Catalog of Federal Domestic Assistance Program No. 93.773, Medicare—Hospital Insurance; and Program No. 93.774, Medicare—Supplementary Medical Insurance Program)

Dated: October 26, 2009.

Charlene Frizzera,

Acting Administrator, Centers for Medicare & Medicaid Services.

Dated: October 29, 2009.

Kathleen Sebelius,

Secretary.

BILLING CODE 4120-01-P

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0056	Level II Foot Musculoskeletal Procedures	T	52.5258	\$3,540.55		\$708.11
0057	Bunion Procedures	T	31.5534	\$2,126.89	\$475.91	\$425.98
0058	Level I Strapping and Cast Application	S	1.0537	\$71.03		\$14.21
0060	Manipulation Therapy	S	0.3808	\$25.67		\$5.14
0061	Laminectomy, Laminectomy, or Incision for Implantation of Neurostimulator Electrode	S	86.5171	\$5,831.77		\$1,166.36
0062	Level I Treatment Fracture/Dislocation	T	25.8446	\$1,742.08	\$372.87	\$348.42
0063	Level II Treatment Fracture/Dislocation	T	45.4678	\$3,064.80		\$612.96
0064	Level III Treatment Fracture/Dislocation	T	65.4752	\$4,413.42		\$882.69
0065	Level I Stereotactic Radiosurgery, MRgFUS, and MEG	S	14.2808	\$962.61		\$192.53
0066	Level II Stereotactic Radiosurgery, MRgFUS, and MEG	S	36.9119	\$2,488.08		\$497.62
0067	Level III Stereotactic Radiosurgery, MRgFUS, and MEG	S	52.9891	\$3,571.78		\$714.36
0069	Thoracoscopy	T	34.0084	\$2,292.37	\$591.64	\$458.48
0070	Thoracotomy	T	5.5521	\$374.24		\$74.85
0071	Level I Endoscopy Upper Airway	T	0.8007	\$53.97	\$11.14	\$10.80
0072	Level II Endoscopy Upper Airway	T	1.8425	\$124.20		\$24.84
0073	Level III Endoscopy Upper Airway	T	4.4119	\$297.39	\$69.15	\$59.48
0074	Level IV Endoscopy Upper Airway	T	21.6572	\$1,459.83	\$292.26	\$291.97
0075	Level I Endoscopy Lower Airway	T	29.2718	\$1,973.09	\$445.92	\$384.62
0076	Level II Endoscopy Lower Airway	T	10.3707	\$699.05	\$189.82	\$139.81
0077	Level I Pulmonary Treatment	S	0.4058	\$27.35	\$7.74	\$5.47
0078	Level III Pulmonary Treatment	S	1.4142	\$95.33		\$19.07
0079	Ventilation Initiation and Management	S	3.0865	\$208.05		\$41.61
0080	Diagnostic Cardiac Catheterization	T	39.8099	\$2,683.43	\$838.92	\$536.69
0082	Coronary or Non-Coronary Atherectomy	T	93.3244	\$6,290.62		\$1,258.13
0083	Coronary or Non-Coronary Angioplasty and Percutaneous Valveoplasty	T	50.6809	\$3,416.20		\$683.24
0084	Level I Electrophysiologic Procedures	S	10.8110	\$728.73		\$145.75
0085	Level II Electrophysiologic Procedures	T	52.3882	\$3,531.28		\$706.26
0086	Level III Electrophysiologic Procedures	T	106.9978	\$7,212.29		\$1,442.46
0088	Thrombectomy	T	40.9003	\$2,756.93	\$655.22	\$551.39
0089	Insertion/Replacement of Permanent Pacemaker and Electrodes	T	118.8826	\$8,013.40	\$1,682.28	\$1,602.68
0090	Insertion/Replacement of Pacemaker Pulse Generator	T	98.1576	\$6,616.41	\$1,597.43	\$1,323.29
0091	Level II Vascular Ligation	T	45.0292	\$3,035.24		\$607.05
0092	Level I Vascular Ligation	T	26.5713	\$1,791.07		\$358.21
0093	Vascular Reconstruction/Fistula Repair without Device	S	35.5969	\$2,399.44		\$479.89
0094	Level I Resuscitation and Cardioversion	S	2.4553	\$165.50	\$46.29	\$33.10
0095	Cardiac Rehabilitation	S	0.5691	\$38.36	\$13.86	\$7.88
0096	Level II Noninvasive Physiologic Studies	S	1.6075	\$108.36	\$37.42	\$21.68
0097	Level I Noninvasive Physiologic Studies	S	0.9848	\$66.38	\$23.79	\$13.28
0099	Electrocardiograms	S	0.3940	\$26.56		\$5.32
0100	Cardiac Stress Tests	X	2.6136	\$176.17	\$41.44	\$35.24

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0001	Level I Photochemotherapy	S	0.5302	\$35.74		\$7.15
0002	Fine Needle Biopsy/Aspiration	T	1.5111	\$101.86		\$20.38
0003	Bone Marrow Biopsy/Aspiration	T	3.0998	\$208.95		\$41.79
0004	Level I Needle Biopsy/Aspiration Except Bone Marrow	T	4.5991	\$310.01		\$62.01
0005	Level II Needle Biopsy/Aspiration Except Bone Marrow	T	7.8145	\$526.74		\$105.35
0006	Level I Incision & Drainage	T	1.4557	\$98.12		\$19.63
0007	Level II Incision & Drainage	T	12.6217	\$850.78		\$170.16
0008	Level III Incision and Drainage	T	19.4063	\$1,308.10		\$261.62
0012	Level I Debridement & Destruction	T	0.4436	\$29.90		\$5.98
0013	Level II Debridement & Destruction	T	0.8789	\$59.24		\$11.85
0015	Level III Debridement & Destruction	T	1.5412	\$103.89		\$20.78
0016	Level IV Debridement & Destruction	T	2.7982	\$188.62		\$37.73
0017	Level VI Debridement & Destruction	T	21.2653	\$1,433.41		\$286.69
0019	Level I Excision/ Biopsy	T	4.3625	\$294.06	\$64.51	\$58.82
0020	Level II Excision/ Biopsy	T	8.2028	\$552.92		\$110.59
0021	Level III Excision/ Biopsy	T	17.4975	\$1,179.44		\$235.89
0022	Level IV Excision/ Biopsy	T	23.9880	\$1,576.49	\$354.45	\$315.30
0028	Level II Breast Surgery	T	24.7516	\$1,668.41		\$333.69
0029	Level III Breast Surgery	T	34.1654	\$2,302.95	\$581.52	\$460.59
0030	Level III Breast Surgery	T	41.9997	\$2,831.03	\$747.07	\$566.21
0031	Smoking Cessation Services	X	0.3448	\$23.24		\$4.65
0034	Mental Health Services Composite	S	3.1286	\$210.89		\$42.18
0035	Vascular Puncture and Minor Diagnostic Procedures	X	0.2320	\$15.64		\$3.13
0037	Level IV Needle Biopsy/Aspiration Except Bone Marrow	T	15.5003	\$1,044.81	\$228.76	\$208.97
0039	Level I Implantation of Neurostimulator Generator	S	206.1011	\$13,892.45		\$2,778.49
0040	Percutaneous Implantation of Neurostimulator Electrodes	S	65.7095	\$4,423.21		\$885.85
0041	Level I Arthroscopy	T	29.9198	\$2,016.77		\$403.36
0042	Level II Arthroscopy	T	48.8176	\$3,290.60	\$804.74	\$658.12
0045	Bone/Joint Manipulation Under Anesthesia	T	15.2922	\$1,030.79	\$268.47	\$206.16
0047	Arthroplasty without Prosthesis	T	39.8877	\$2,888.67		\$537.74
0048	Level I Arthroplasty or Implantation with Prosthesis	T	58.0838	\$3,915.20		\$783.04
0049	Level I Musculoskeletal Procedures Except Hand and Foot	T	22.0149	\$1,483.94		\$296.79
0050	Level II Musculoskeletal Procedures Except Hand and Foot	T	31.7717	\$2,141.60		\$428.32
0051	Level III Musculoskeletal Procedures Except Hand and Foot	T	45.5786	\$3,139.68		\$627.94
0052	Level IV Musculoskeletal Procedures Except Hand and Foot	T	88.6521	\$5,975.68		\$1,195.14
0053	Level I Hand Musculoskeletal Procedures	T	17.0420	\$1,148.73	\$253.49	\$229.75
0054	Level II Hand Musculoskeletal Procedures	T	28.2376	\$1,903.38		\$380.66
0055	Level I Foot Musculoskeletal Procedures	T	21.8439	\$1,472.41	\$355.34	\$294.49

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0153	Peritoneal and Abdominal Procedures	T	27.1855	\$1,832.47	\$376.05	\$366.50
0154	Hernia/Hydrocele Procedures	T	31.9682	\$2,154.85	\$464.85	\$430.97
0155	Level II Anal/Rectal Procedures	T	14.1140	\$951.37		\$190.28
0156	Level III Urinary and Anal Procedures	T	3.0214	\$203.66		\$40.74
0157	Colorectal Cancer Screening: Barium Enema	S	1.2511	\$84.33		\$16.87
0158	Colorectal Cancer Screening: Colonoscopy	T	8.0912	\$545.40		\$136.95
0159	Colorectal Cancer Screening: Flexible Sigmoidoscopy	S	3.7227	\$250.93		\$62.73
0160	Level I Cystourethroscopy and other Genitourinary Procedures	T	7.1342	\$480.89		\$96.18
0161	Level II Cystourethroscopy and other Genitourinary Procedures	T	17.0344	\$1,148.22		\$229.65
0162	Level III Cystourethroscopy and other Genitourinary Procedures	T	25.5223	\$1,720.36		\$344.08
0163	Level IV Cystourethroscopy and other Genitourinary Procedures	T	36.2009	\$2,440.16		\$488.04
0164	Level II Urinary and Anal Procedures	T	2.0194	\$136.12		\$27.23
0165	Level IV Urinary and Anal Procedures	T	20.0243	\$1,349.76		\$269.96
0166	Level I Urethral Procedures	T	20.3374	\$1,370.86		\$274.18
0168	Level II Urethral Procedures	T	31.3770	\$2,115.00		\$423.00
0169	Lithotripsy	T	41.3626	\$2,788.09		\$557.62
0170	Dialysis	S	6.8212	\$459.79		\$91.96
0172	Level I Partial Hospitalization (3 services)	P	2.2230	\$149.84		\$29.97
0173	Level II Partial Hospitalization (4 or more services)	P	3.1286	\$210.89		\$42.18
0174	Level IV Laparoscopy	T	109.9237	\$7,409.52	\$2,069.30	\$1,481.91
0181	Level II Male Genital Procedures	T	34.8837	\$2,351.37	\$621.82	\$470.28
0183	Level I Male Genital Procedures	T	23.2838	\$1,569.47		\$313.90
0184	Prostate Biopsy	T	12.4313	\$837.94		\$167.89
0188	Level II Female Reproductive Proc	T	3.4085	\$229.75		\$45.95
0189	Level III Female Reproductive Proc	T	22.5926	\$1,522.88		\$304.58
0190	Level I Hysterectomy	T	1.5277	\$102.96		\$20.60
0191	Level I Female Reproductive Proc	T	0.1320	\$8.90		\$1.78
0192	Level IV Female Reproductive Proc	T	6.7949	\$458.02		\$91.61
0193	Level V Female Reproductive Proc	T	20.0452	\$1,351.17		\$270.24
0195	Level VI Female Reproductive Procedures	T	35.2666	\$2,377.18	\$483.80	\$475.44
0202	Level VII Female Reproductive Procedures	T	44.9894	\$3,032.56	\$981.50	\$606.82
0203	Level IV Nerve Injections	T	13.2439	\$892.72	\$225.98	\$178.55
0204	Level I Nerve Injections	T	2.5558	\$172.28	\$40.13	\$34.46
0206	Level II Nerve Injections	T	3.7221	\$250.89	\$51.76	\$50.18
0207	Level III Nerve Injections	T	7.2002	\$485.34		\$97.07
0208	Laminotomies and Laminectomies	T	49.2256	\$3,318.10		\$663.62
0209	Level II Extended EEG, Sleep, and Cardiovascular Studies	S	11.4315	\$770.55	\$288.73	\$154.11
0213	Level I Extended EEG, Sleep, and Cardiovascular Studies	S	2.4043	\$162.06	\$53.58	\$32.42

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0101	Till Table Evaluation	S	4.3705	\$294.60	\$100.24	\$58.92
0102	Level II Pulmonary Treatment	S	0.7488	\$50.46		\$10.10
0103	Miscellaneous Vascular Procedures	T	17.8436	\$1,202.77		\$240.56
0104	Transcatheter Placement of Intracoronary Stents	T	84.7773	\$5,714.50	\$1,142.90	\$1,142.90
0105	Repair/Revision/Removal of Pacemakers, AICDs, or Vascular Devices	T	22.9412	\$1,546.37	\$309.28	\$309.28
0106	Insertion/Replacement of Pacemaker Leads and/or Electrodes	T	48.9381	\$3,298.59	\$659.72	\$659.72
0107	Insertion of Cardioverter-Defibrillator	T	325.8288	\$21,962.82	\$4,392.57	\$4,392.57
0108	Insertion/Replacement/Repair of Cardioverter-Defibrillator Leads	T	412.3722	\$27,796.36	\$5,569.28	\$5,569.28
0110	Transfusion	S	3.3809	\$227.89		\$45.58
0111	Blood Product Exchange	S	11.9424	\$804.99	\$198.40	\$161.00
0112	Apheresis and Stem Cell Procedures	S	33.3206	\$2,246.01	\$449.21	\$449.21
0113	Excision Lymphatic System	T	24.6146	\$1,659.17	\$331.84	\$331.84
0114	Thyroid/Lymphadenectomy Procedures	T	48.7561	\$3,286.45	\$657.20	\$657.20
0115	Cannula/Access Device Procedures	T	30.9784	\$2,088.13	\$417.63	\$417.63
0121	Level I Tube or Catheter Changes or Repositioning	T	6.3742	\$429.66	\$85.94	\$85.94
0126	Level II Urinary and Anal Procedures	T	1.0958	\$73.86	\$16.21	\$14.78
0127	Level IV Stereotactic Radiosurgery, MRgFUS, and MEG	S	108.9558	\$7,344.27	\$1,488.86	\$1,488.86
0128	Echocardiogram with Contrast	S	9.6604	\$651.17	\$216.29	\$190.24
0129	Level I Closed Treatment Fracture Finger/Toe/Trunk	T	1.6576	\$111.73	\$22.35	\$22.35
0130	Level II Laparoscopy	T	38.0540	\$2,565.07	\$659.53	\$513.02
0131	Level III Laparoscopy	T	48.8400	\$3,157.30	\$1,001.89	\$631.46
0132	Level III Laparoscopy	T	72.9582	\$4,917.82	\$1,239.22	\$983.57
0133	Level I Skin Repair	T	1.3542	\$91.28	\$25.67	\$18.26
0134	Level II Skin Repair	T	3.1508	\$212.38	\$42.48	\$42.48
0135	Level III Skin Repair	T	4.4366	\$298.19	\$59.84	\$59.84
0136	Level IV Skin Repair	T	16.0542	\$1,082.15	\$216.43	\$216.43
0137	Level V Skin Repair	T	23.9317	\$1,613.14	\$322.63	\$322.63
0138	Level III Closed Treatment Fracture Finger/Toe/Trunk	T	4.7946	\$323.18	\$64.64	\$64.64
0139	Level III Closed Treatment Fracture Finger/Toe/Trunk	T	18.3962	\$1,240.01	\$248.01	\$248.01
0140	Esophageal Dilation without Endoscopy	T	5.9622	\$401.89	\$84.79	\$80.38
0141	Level I Upper GI Procedures	T	8.7462	\$589.55	\$143.38	\$117.91
0142	Small Intestine Endoscopy	T	9.8503	\$663.97	\$152.78	\$132.80
0143	Lower GI Endoscopy	T	9.1051	\$613.74	\$186.06	\$122.75
0146	Level I Sigmoidoscopy and Anoscopy	T	5.7747	\$389.25	\$77.85	\$77.85
0147	Level II Sigmoidoscopy and Anoscopy	T	9.2450	\$623.17	\$124.64	\$124.64
0148	Level III Anal/Rectal Procedures	T	5.3650	\$361.63	\$72.33	\$72.33
0149	Level III Anal/Rectal Procedures	T	23.9703	\$1,615.74	\$323.15	\$323.15
0150	Level IV Anal/Rectal Procedures	T	32.1812	\$2,169.21	\$437.12	\$433.85
0151	Endoscopic Retrograde Cholangio-Pancreatography (ERCP)	T	22.6111	\$1,524.12	\$304.83	\$304.83
0152	Level I Percutaneous Abdominal and Biliary Procedures	T	30.7430	\$2,072.26	\$414.46	\$414.46

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0270	Level III Echocardiogram Without Contrast	S	8.8425	\$596.04	\$141.32	\$119.21
0272	Fluoroscopy	X	1.2693	\$95.56	\$31.15	\$17.12
0274	Myelography	S	7.2260	\$487.08	-	\$97.42
0275	Arthrography	S	3.9361	\$266.32	\$69.07	\$53.07
0276	Level I Digestive Radiology	S	1.2985	\$87.53	\$34.74	\$17.51
0277	Level II Digestive Radiology	S	2.1001	\$141.56	\$54.04	\$28.32
0278	Diagnostic Urography	S	2.5460	\$171.62	\$58.59	\$32.48
0279	Level II Angiography and Venography	S	29.1126	\$1,962.36	-	\$392.48
0280	Level III Angiography and Venography	S	45.8793	\$3,092.54	-	\$618.51
0282	Miscellaneous Computed Axial Tomography	S	1.6291	\$109.81	\$37.81	\$21.97
0283	Computed Tomography with Contrast	S	4.4066	\$297.03	\$96.86	\$69.41
0284	Magnetic Resonance Imaging and Magnetic Resonance Angiography with Contrast	S	6.2901	\$423.99	\$147.21	\$94.90
0288	Bone Density-Axial Skeleton	S	1.0755	\$72.50	\$28.66	\$14.50
0293	Level V Anterior Segment Eye Procedures	T	101.0161	\$6,809.09	-	\$1,361.92
0299	Hyperthermia and Radiation Treatment Procedures	S	5.6461	\$380.58	-	\$76.12
0300	Level I Radiation Therapy	S	1.3764	\$92.78	-	\$18.56
0301	Level II Radiation Therapy	S	2.3030	\$155.24	-	\$31.05
0303	Treatment Device Construction	X	2.8279	\$190.62	\$66.95	\$38.13
0304	Level I Therapeutic Radiation Treatment Preparation	X	1.5271	\$102.94	\$34.71	\$20.59
0305	Level II Therapeutic Radiation Treatment Preparation	X	3.9510	\$266.32	\$91.38	\$53.27
0307	Myocardial Positron Emission Tomography (PET) Imaging	S	21.2573	\$1,432.87	-	\$286.58
0308	Non-Myocardial Positron Emission Tomography (PET) Imaging	S	15.3895	\$1,037.34	-	\$207.47
0310	Level III Therapeutic Radiation Treatment Preparation	X	13.7576	\$927.34	\$325.27	\$185.47
0312	Radiation Applications	S	4.4846	\$302.29	-	\$60.46
0313	Brachytherapy	S	11.5353	\$777.55	\$283.30	\$155.51
0315	Level II Implantation of Neurostimulator Generator	S	274.7397	\$18,519.10	-	\$3,703.92
0317	Level II Miscellaneous Radiology Procedures	X	5.6012	\$377.55	-	\$75.51
0320	Electroconvulsive Therapy	S	5.8620	\$395.13	\$80.06	\$79.03
0322	Brief Individual Psychotherapy	S	1.1958	\$80.60	-	\$16.12
0323	Extended Individual Psychotherapy	S	2.6680	\$112.50	-	\$22.50
0324	Family Psychotherapy	S	2.2103	\$148.99	-	\$29.80
0325	Group Psychotherapy	S	0.8905	\$60.03	\$12.81	\$12.01
0330	Dental Procedures	S	10.3227	\$695.81	-	\$139.17
0332	Computed Tomography without Contrast	S	2.8940	\$195.07	\$75.24	\$39.02
0333	Computed Tomography without Contrast followed by Contrast	S	4.9479	\$333.52	\$116.41	\$66.71
0336	Magnetic Resonance Imaging and Magnetic Resonance Angiography without Contrast	S	5.1855	\$349.53	\$137.40	\$69.91
0337	Magnetic Resonance Imaging and Magnetic Resonance Angiography without Contrast followed by Contrast	S	7.9432	\$535.42	\$198.32	\$107.09
0340	Minor Ancillary Procedures	X	0.6693	\$45.11	-	\$9.03

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0215	Level I Nerve and Muscle Tests	S	0.6135	\$41.35	-	\$8.27
0216	Level III Nerve and Muscle Tests	S	2.8831	\$180.89	-	\$36.18
0218	Level II Nerve and Muscle Tests	S	1.1965	\$80.65	-	\$16.13
0220	Level I Nerve Procedures	T	18.7200	\$1,261.84	\$252.37	\$129.21
0221	Level II Nerve Procedures	T	37.2806	\$2,512.94	\$502.59	\$252.59
0224	Implantation of Catheter/Reservoir/Shunt	T	41.0288	\$2,765.99	\$553.12	\$276.59
0225	Implantation of Neurostimulator Electrodes, Cranial Nerve	S	157.9390	\$10,646.04	\$2,129.21	\$1,064.10
0227	Implantation of Drug Infusion Device	T	198.6572	\$13,390.69	\$2,678.14	\$1,339.07
0229	Transcatheter Placement of Intravascular Shunts	T	97.2910	\$6,558.00	\$1,311.60	\$655.80
0230	Level I Eye Tests & Treatments	S	0.9945	\$40.07	-	\$8.02
0231	Level III Eye Tests & Treatments	S	1.9575	\$131.95	\$26.39	\$13.19
0232	Level I Anterior Segment Eye Procedures	T	4.5074	\$303.83	\$76.12	\$38.06
0233	Level II Anterior Segment Eye Procedures	T	16.2485	\$1,095.25	\$263.77	\$131.88
0234	Level III Anterior Segment Eye Procedures	T	24.4021	\$1,644.85	\$311.31	\$155.65
0235	Level I Posterior Segment Eye Procedures	T	5.8498	\$394.31	-	\$78.87
0237	Level II Posterior Segment Eye Procedures	T	20.7145	\$1,396.28	\$279.26	\$139.63
0238	Level I Repair and Plastic Eye Procedures	T	3.2267	\$217.50	\$43.50	\$21.75
0239	Level II Repair and Plastic Eye Procedures	T	7.6421	\$515.12	\$103.03	\$51.51
0240	Level III Repair and Plastic Eye Procedures	T	18.1580	\$1,223.96	\$256.93	\$128.46
0241	Level IV Repair and Plastic Eye Procedures	T	26.8396	\$1,809.15	\$363.45	\$181.72
0242	Level V Repair and Plastic Eye Procedures	T	39.0115	\$2,623.61	\$597.36	\$298.68
0243	Strabismus/Muscle Procedures	T	23.4694	\$1,581.98	\$418.00	\$209.00
0244	Corneal and Amniotic Membrane Transplant	T	37.5388	\$2,530.34	\$803.26	\$401.63
0245	Level I Cataract Procedures without IOL Insert	T	15.8085	\$1,065.59	\$214.11	\$107.05
0246	Cataract Procedures with IOL Insert	T	24.2879	\$1,637.15	\$495.96	\$247.98
0247	Laser Eye Procedures	T	5.3014	\$357.35	\$104.31	\$52.15
0249	Level II Cataract Procedures without IOL Insert	T	30.3293	\$2,044.38	\$518.26	\$259.13
0250	Level I ENT Procedures	T	1.1522	\$77.67	\$25.10	\$12.55
0251	Level II ENT Procedures	T	3.4250	\$230.87	\$46.18	\$23.09
0252	Level III ENT Procedures	T	7.6196	\$513.61	\$109.16	\$54.58
0253	Level IV ENT Procedures	T	17.1879	\$1,158.57	\$282.29	\$141.14
0254	Level V ENT Procedures	T	24.9637	\$1,682.70	\$336.54	\$168.27
0256	Level VI ENT Procedures	T	42.9827	\$2,897.29	\$579.46	\$289.73
0259	Level VII ENT Procedures	T	428.8363	\$28,906.14	\$5,543.66	\$2,771.83
0260	Level I Plain Film Except Teeth	X	0.6661	\$44.90	-	\$8.98
0261	Level II Plain Film Except Teeth Including Bone Density Measurement	X	1.1161	\$75.23	-	\$15.05
0262	Plain Film of Teeth	X	0.4469	\$30.12	-	\$6.03
0263	Level I Miscellaneous Radiology Procedures	X	3.1192	\$210.25	-	\$42.05
0265	Level I Diagnostic and Screening Ultrasound	S	0.9267	\$62.47	\$22.34	\$11.17
0266	Level II Diagnostic and Screening Ultrasound	S	1.4434	\$97.29	\$37.61	\$19.46
0267	Level III Diagnostic and Screening Ultrasound	S	2.3005	\$155.07	\$60.50	\$30.25
0269	Level II Echocardiogram Without Contrast	S	6.6903	\$450.97	-	\$90.20

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0402	Level II Nervous System Imaging	S	8.5845	\$578.65		\$115.73
0403	Level I Nervous System Imaging	S	2.9181	\$196.70	\$72.42	\$39.94
0404	Renal and Genitourinary Studies	S	4.8221	\$325.04	\$83.10	\$65.01
0406	Level I Tumor/Infection Imaging	S	4.3034	\$290.07	\$88.22	\$58.02
0407	Level I Radionuclide Therapy	S	3.2566	\$218.10	\$78.13	\$43.62
0408	Level III Tumor/Infection Imaging	S	14.2874	\$963.06		\$192.62
0409	Red Blood Cell Tests	X	0.1161	\$7.83	\$2.20	\$1.57
0412	IMRT Treatment Delivery	X	6.2490	\$421.22		\$64.25
0413	Level II Radionuclide Therapy	S	5.2007	\$350.56		\$70.12
0414	Level II Tumor/Infection Imaging	S	7.5722	\$510.41		\$102.09
0415	Level II Endoscopy Lower Airway	T	25.9024	\$1,745.98	\$459.92	\$349.20
0418	Insertion of Left Ventricular Pacing Elect.	T	204.1256	\$13,759.29		\$2,751.86
0422	Level II Upper GI Procedures	T	24.2703	\$1,635.96	\$437.96	\$227.20
0423	Level II Percutaneous Abdominal and Biliary Procedures	T	51.3618	\$3,462.09		\$692.42
0425	Level II Arthroplasty or Implantation with Prosthesis	T	118.7670	\$8,005.61		\$1,601.13
0426	Level II Strapping and Cast Application	S	2.3457	\$158.11		\$31.63
0427	Level II Tube or Catheter Changes or Repositioning	T	15.3103	\$1,032.01		\$206.41
0428	Level III Sigmoidoscopy and Anoscopy	T	22.8208	\$1,538.26		\$307.66
0429	Level V Cystourethroscopy and other Genitourinary Procedures	T	46.6800	\$3,146.51		\$629.31
0432	Health and Behavior Services	S	0.6013	\$40.53		\$8.11
0433	Level II Pathology	X	0.2462	\$16.73	\$5.17	\$3.95
0434	Cardiac Defect Repair	T	147.3728	\$9,933.81		\$1,986.77
0436	Level I Drug Administration	S	0.3809	\$25.67		\$5.14
0437	Level II Drug Administration	S	0.5555	\$37.44		\$7.49
0438	Level III Drug Administration	S	1.1229	\$75.69		\$15.14
0439	Level IV Drug Administration	S	1.8808	\$126.78		\$25.96
0440	Level V Drug Administration	S	3.2632	\$219.96		\$44.00
0442	Dosimetric Drug Administration	S	25.7674	\$1,736.88		\$347.38
0604	Level 1 Hospital Clinic Visits	V	0.8893	\$57.92		\$11.59
0605	Level 2 Hospital Clinic Visits	V	1.0337	\$69.68		\$13.94
0606	Level 3 Hospital Clinic Visits	V	1.3222	\$89.12		\$17.83
0607	Level 4 Hospital Clinic Visits	V	1.6830	\$113.44		\$22.69
0608	Level 5 Hospital Clinic Visits	V	2.4853	\$167.52		\$33.51
0609	Level 1 Type A Emergency Visits	V	0.7886	\$53.16	\$12.70	\$10.64
0613	Level 2 Type A Emergency Visits	V	1.3033	\$87.85	\$21.06	\$17.57
0614	Level 3 Type A Emergency Visits	V	2.0796	\$140.18	\$34.50	\$28.04
0615	Level 4 Type A Emergency Visits	V	3.1109	\$223.17	\$48.49	\$44.64
0616	Level 5 Type A Emergency Visits	V	4.8917	\$329.73	\$72.86	\$65.95
0617	Critical Care	S	7.3492	\$495.38	\$111.59	\$99.08
0618	Trauma Response with Critical Care	S	12.3717	\$833.93		\$166.79
0621	Level I Vascular Access Procedures	T	11.1660	\$752.66		\$150.54
0622	Level II Vascular Access Procedures	T	25.3344	\$1,707.69		\$341.54

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0341	Skin Tests	X	0.0799	\$5.39	\$2.09	\$1.03
0342	Level I Pathology	X	0.1546	\$10.42		\$2.09
0343	Level III Pathology	X	0.5301	\$35.73	\$10.84	\$7.15
0344	Level IV Pathology	X	0.8028	\$54.11	\$15.60	\$10.83
0345	Level II Transfusion Laboratory Procedures	X	0.2195	\$14.80		\$2.96
0347	Level III Transfusion Laboratory Procedures	X	0.7384	\$25.17		\$5.04
0350	Administration of flu and PPV vaccine	X	0.3809	\$25.67		\$9.96
0360	Level I Allimentary Tests	S	1.5203	\$102.48	\$33.88	\$20.50
0361	Level II Allimentary Tests	X	4.0839	\$275.28	\$83.23	\$55.06
0363	Level I Otorhinolaryngological Function Tests	X	0.9061	\$61.08	\$17.10	\$12.22
0364	Level II Audiometry	X	0.4700	\$31.68	\$7.06	\$6.34
0366	Level III Audiometry	X	1.6275	\$85.44	\$18.52	\$17.09
0367	Level I Pulmonary Test	X	0.5968	\$40.23	\$25.00	\$21.90
0368	Level II Pulmonary Tests	X	0.8495	\$57.26	\$20.93	\$11.46
0369	Level III Pulmonary Tests	X	2.6228	\$176.79	\$42.29	\$35.36
0370	Allergy Tests	X	1.4598	\$96.40		\$19.68
0373	Level I Neuropsychological Testing	X	0.9758	\$65.77		\$13.16
0375	Ancillary Outpatient Services When Patient Expires	S	88.5075	\$5,965.94		\$1,193.19
0377	Level II Cardiac Imaging	S	11.4989	\$775.09	\$158.84	\$155.02
0378	Level II Pulmonary Imaging	S	4.8032	\$323.76	\$124.94	\$84.76
0379	Injection adenosine 6 MG	K		\$9.50		\$1.90
0381	Single Allergy Tests	X	0.4312	\$29.07		\$5.82
0382	Level II Neuropsychological Testing	X	2.6683	\$179.86		\$35.98
0383	Cardiac Computed Tomographic Imaging	S	3.9929	\$269.15	\$107.66	\$53.83
0384	GI Procedures with Stents	T	26.4913	\$1,785.67	\$357.14	
0385	Level I Prosthetic Urological Procedures	S	98.0867	\$6,811.63		\$1,322.33
0386	Level II Prosthetic Urological Procedures	S	164.3200	\$11,076.15		\$2,215.23
0387	Level II Hysteroscopy	T	37.2199	\$2,508.84	\$655.55	\$501.77
0388	Discography	S	25.8604	\$1,743.15		\$348.63
0389	Level I Non-imaging Nuclear Medicine	S	1.6695	\$112.53	\$30.02	\$22.51
0390	Level II Endocrine Imaging	S	2.1595	\$145.56	\$62.15	\$29.12
0391	Level III Endocrine Imaging	S	3.2894	\$221.73	\$66.13	\$44.35
0392	Level II Non-imaging Nuclear Medicine	S	2.6817	\$180.76	\$43.95	\$36.16
0393	Hematologic Processing & Studies	S	5.7873	\$390.10	\$79.97	\$78.02
0394	Hepatobiliary Imaging	S	4.3139	\$290.78	\$89.32	\$58.16
0396	Bone Imaging	S	3.6702	\$247.39	\$89.73	\$49.48
0397	Vascular Imaging	S	2.9679	\$200.05	\$46.29	\$40.01
0398	Level I Cardiac Imaging	S	4.5366	\$305.79	\$97.94	\$61.16
0400	Hematopoietic Imaging	S	3.8342	\$258.45	\$91.75	\$51.69
0401	Level I Pulmonary Imaging	S	3.0625	\$206.43	\$74.63	\$41.29

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0698	Level II Eye Tests & Treatments	S	0.9553	\$64.39		\$12.88
0699	Level IV Eye Tests & Treatments	T	15.5407	\$1,047.54		\$209.51
0701	Sr89 strontium	K		\$647.67		\$129.54
0726	Dexrazoxane HCl injection	K		\$340.03		\$68.01
0728	Fligrastrin 300 mcg injection	K		\$208.54		\$41.71
0730	Pamidronate disodium	K		\$18.42		\$3.69
0731	Sargramostim injection	K		\$23.31		\$4.67
0732	Mesna injection	K		\$4.34		\$0.87
0735	Ampho b cholesteryl sulfate	K		\$13.74		\$2.75
0736	Amphotericin b liposome inj	K		\$14.96		\$3.00
0738	Rasburicase	K		\$164.00		\$32.80
0747	Chlorothiazide sodium inj	K		\$292.02		\$58.41
0751	Mechlorethamine hcl inj	K		\$144.56		\$28.92
0752	Dactinomycin injection	K		\$533.21		\$106.65
0759	Naltrexone, depot form	K		\$2.14		\$0.43
0760	Andilungin injection	K		\$1.21		\$0.25
0800	Leuprolide acetate	K		\$480.20		\$96.04
0802	Etoposide oral	K		\$0.45		\$0.09
0807	Aldesleukin injection	K		\$931.49		\$186.30
0809	Bcg live intravesical vac	K		\$111.08		\$22.22
0810	Goserelin acetate implant	K		\$193.02		\$38.61
0812	Carmustine injection	K		\$173.73		\$34.75
0814	Asparaginase injection	K		\$56.92		\$11.39
0820	Daunorubicin injection	K		\$14.95		\$2.99
0821	Daunorubicin citrate inj	K		\$55.27		\$11.06
0823	Docetaxel injection	K		\$16.95		\$3.39
0825	Nelarabine injection	K		\$101.28		\$20.26
0827	Floxuridine injection	K		\$46.60		\$9.32
0828	Gemcitabine hcl injection	K		\$139.10		\$27.82
0830	Irinotecan injection	K		\$13.18		\$2.64
0831	Ifosfomide injection	K		\$29.39		\$5.88
0832	Idarubicin hcl injection	K		\$96.70		\$19.34
0834	Interferon alfa-2a inj	K		\$10.60		\$2.12
0835	Cosyntropin injection NOS	K		\$91.84		\$18.37
0836	Interferon alfa-2b inj	K		\$15.54		\$3.11
0838	Interferon gamma 1-b inj	K		\$294.03		\$58.81
0840	inj malphalian hydrochl	K		\$1,622.81		\$324.57
0842	Fludrabine phosphate inj	K		\$151.36		\$30.28
0843	Pegaspargase injection	K		\$2,695.67		\$539.14
0844	Pentostatin injection	K		\$1,399.56		\$279.92
0849	Pitumab injection	K		\$562.70		\$110.54
0850	Streptozocin injection	K		\$278.35		\$55.67
0851	Thiotepa injection	K		\$97.69		\$19.54

ADDENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0623	Level III Vascular Access Procedures	T	30.4396	\$2,051.81		\$410.37
0624	Phlebotomy and Minor Vascular Access Device Procedures	X	0.6132	\$41.33	\$12.65	\$6.27
0626	Level 1 Type B Emergency Visits	V	0.6796	\$45.81		\$9.17
0627	Level 2 Type B Emergency Visits	V	0.9229	\$62.21		\$12.45
0628	Level 3 Type B Emergency Visits	V	1.4572	\$99.22		\$19.65
0629	Level 4 Type B Emergency Visits	V	2.1041	\$141.83		\$28.37
0630	Level 5 Type B Emergency Visits	V	3.4466	\$232.32	\$52.26	\$46.47
0648	Level IV Breast Surgery	T	58.2278	\$3,924.90		\$784.98
0651	Complex Intraarterial Radiation Source Application	S	13.2555	\$893.30		\$178.70
0652	Insertion of Intraoperative and Pleural Catheters	T	30.4602	\$2,053.20		\$410.64
0653	Vascular Reconstruction/Fistula Repair with Device	T	46.6863	\$3,147.07		\$629.42
0654	Insertion/Replacement of a permanent dual chamber pacemaker	T	108.1692	\$7,291.25		\$1,458.25
0655	Insertion/Replacement/Conversion of a permanent dual chamber pacemaker	T	142.1623	\$9,582.59		\$1,916.52
0656	Transcatheter Placement of Intracoronary Drug-Eluting Stents	T	110.7870	\$7,467.71		\$1,493.55
0659	Hyperbaric Oxygen	S	1.5880	\$107.04		\$21.41
0660	Level II Otorhinolaryngologic Function Tests	X	1.5019	\$101.24	\$27.16	\$20.25
0661	Level V Pathology	X	2.4450	\$164.81	\$57.69	\$32.97
0662	CT Angiography	S	5.0511	\$340.47	\$115.03	\$68.10
0664	Level I Proton Beam Radiation Therapy	S	13.9796	\$942.31		\$188.47
0665	Bone Density Appendicular/Skeleton	S	0.4318	\$29.11	\$11.63	\$5.83
0667	Level II Proton Beam Radiation Therapy	S	18.2873	\$1,232.57		\$246.54
0672	Level I Angiography and Venography	S	10.2978	\$694.13		\$138.83
0673	Level III Posterior Segment Eye Procedures	T	39.9643	\$2,693.83		\$538.77
0674	Level IV Anterior Segment Eye Procedures	T	41.8840	\$2,823.23	\$649.56	\$564.65
0676	Prostate Cryoblation	T	113.8626	\$7,675.02		\$1,535.01
0678	Thrombolysis and Other Device Revisions	T	2.3954	\$161.46		\$32.30
0679	External Counterpulsation	T	1.5463	\$104.23		\$20.85
0680	Insertion of Patient Activated Event Recorders	S	5.5047	\$371.05	\$95.30	\$74.21
0683	Level III Photochemotherapy	S	78.0661	\$5,262.12		\$1,052.43
0685	Level III Needle Biopsy/Aspiration Except Bone Marrow	T	2.6790	\$180.58		\$36.12
0687	Revision/Removal of Neurostimulator Electrodes	T	9.6666	\$651.59		\$130.32
0688	Revision/Removal of Neurostimulator Pulse Generator Receiver	T	19.6381	\$1,323.73	\$397.37	\$284.75
0689	Level II Electronic Analysis of Devices	S	28.6636	\$1,932.10	\$770.83	\$386.42
0690	Level I Electronic Analysis of Devices	S	0.5680	\$38.29		\$7.86
0691	Level IV Electronic Analysis of Devices	S	0.3506	\$23.63	\$8.67	\$4.73
0692	Level III Electronic Analysis of Devices	S	2.5406	\$171.25	\$50.49	\$34.25
0694	Mohs Surgery	S	1.6000	\$107.85		\$21.57
0697	Level I Echocardiogram Without Contrast	S	4.9337	\$332.56	\$91.69	\$66.52
		S	3.9223	\$264.39		\$52.88

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010						
APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0947	Fibrogamma injection	K		\$36.51		\$7.31
0948	Gamma globulin injection	K		\$36.71		\$7.35
0949	Frozen plasma, pooled, sd	R	0.7589	\$51.15		\$10.23
0950	Whole blood for transfusion	R	3.0598	\$206.25		\$41.25
0951	Reclast injection	K		\$218.59		\$43.72
0952	Cryoprecipitate each unit	R	0.6910	\$46.58		\$9.32
0954	RBC leukocytes reduced	R	2.7703	\$186.75		\$37.35
0955	Plasma, frz between 8-24hour	R	1.1491	\$77.46		\$15.50
0956	Plasma protein fract,5%,50ml	R	0.9754	\$65.75		\$13.15
0957	Platelets, each unit	R	0.9882	\$66.61		\$13.33
0958	Platelet rich plasma unit	R	2.0294	\$136.79		\$27.36
0959	Red blood cells unit	R	2.1027	\$141.73		\$28.35
0960	Washed red blood cells unit	R	3.6495	\$246.00		\$49.20
0961	Albumin (human),5%, 50ml	K		\$16.89		\$3.38
0963	Albumin (human), 25%, 250 ml	K		\$60.58		\$12.12
0964	Albumin (human), 25%, 20 ml	K		\$25.67		\$5.14
0965	Albumin (human), 25%, 50ml	K		\$62.05		\$12.41
0966	Plasmaprotein fract,5%,250ml	R	1.6017	\$107.96		\$21.60
0967	Blood split unit	R	1.2965	\$87.39		\$17.48
0968	Platelets leukoreduced irradiated	R	1.9576	\$131.95		\$26.39
0969	RBC leukoreduced irradiated	R	3.6350	\$245.02		\$49.01
1009	Cryoprecipitate/reduced plasma	R	1.4035	\$94.60		\$18.92
1010	Blood, fr, cmv-neg	R	2.0076	\$135.32		\$27.07
1011	Platelets, hla-m, fr, unit	R	10.9290	\$736.68		\$147.34
1013	Platelets leukocytes reduced	R	1.5541	\$104.76		\$20.96
1015	Injection glatiramer acetate	K		\$81.23		\$16.25
1016	Blood, fr, froz/dg/wash	R	1.5973	\$103.62		\$20.73
1017	Plt, apher, fr, cmv-neg	R	6.2195	\$419.23		\$93.85
1018	Blood, fr, irradiated	R	2.4502	\$165.16		\$33.04
1019	Plate pheres leukoredu irradiated	R	10.0373	\$676.57		\$135.32
1020	Plt, pher, fr, cmv-neg, irr	R	9.7428	\$656.72		\$131.35
1021	RBC, frz/dg/wash, fr, irradiated	R	5.3658	\$363.04		\$72.61
1022	RBC, fr, cmv-neg, irradiated	R	4.3604	\$293.92		\$58.79
1023	Pralixime choride inj	K		\$85.83		\$17.77
1052	Injection, voriconazole	K		\$5.26		\$1.06
1064	I131 iodide cap, rx	K		\$16.51		\$3.31
1083	Adalimumab injection	K		\$357.53		\$71.51
1084	Denileukin difitox inj	K		\$1,448.32		\$289.67
1086	Temozolomide	K		\$8.59		\$1.72
1138	Hepagam b intravenous, inj	G		\$50.04		\$9.82
1139	Protein c concentrate	K		\$11.96		\$2.40
1142	Supprelin LA implant	G		\$14,875.43		\$2,918.95
1150	I131 iodide sol, rx	K		\$11.02		\$2.21

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010						
APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0852	Topotecan injection	K		\$983.88		\$197.78
0856	Porfimer sodium injection	K		\$2,745.46		\$549.10
0858	Inj cladribine	K		\$25.15		\$5.03
0861	Leuprolide acetate injection	K		\$5.29		\$1.06
0864	Mitoxantrone hydrochl	K		\$65.51		\$13.11
0865	Inferferon alfa-n3 inj	K		\$17.89		\$3.58
0868	Hydral/supatz inj per dose	K		\$5.42		\$1.09
0873	Oral apreplant	K		\$91.87		\$18.38
0874	Synvisc or synvisc-one	K		\$11.47		\$2.30
0875	Euflexxa inj per dose	K		\$113.96		\$22.80
0877	Orthovisc inj per dose	K		\$177.68		\$35.54
0878	Gallium nitrate injection	K		\$1.71		\$0.35
0884	Rho d immune globulin inj	K		\$84.39		\$16.88
0887	Azathioprine parenteral	K		\$90.64		\$18.13
0890	Lymphocyte immune globulin	K		\$453.67		\$90.74
0891	Facrolimus oral	K		\$3.96		\$0.80
0900	Alglucerase injection	K		\$41.19		\$8.24
0901	Alpha 1 proteinase inhibitor	K		\$3.63		\$0.73
0902	Injection, onabotulinumtoxinA	K		\$5.40		\$1.08
0903	Cytomegalovirus imm IV/Mal	K		\$82.24		\$17.45
0904	Gamma globulin 4 CC inj	K		\$60.20		\$12.04
0910	Inferferon beta-1b / 25 MG	K		\$166.90		\$33.78
0913	Ganciclovir long act implant	K		\$16,640.00		\$3,328.00
0916	Injection rimigucrase /unit	K		\$4.12		\$0.83
0917	Adenosine injection	K		\$76.42		\$15.29
0920	Gamma globulin 6 CC inj	K		\$90.35		\$18.07
0921	Gamma globulin 7 CC inj	K		\$105.27		\$21.06
0922	Gamma globulin 8 CC inj	K		\$120.40		\$24.08
0923	Gamma globulin 9 CC inj	K		\$150.50		\$30.10
0924	Gamma globulin 10 CC inj	K		\$150.50		\$30.10
0925	Factor viii	K		\$0.84		\$0.17
0927	Factor viii recombinant	K		\$1.08		\$0.22
0928	Factor ix complex	K		\$0.85		\$0.17
0929	Anti-inhibitor	K		\$1.53		\$0.31
0931	Factor IX non-recombinant	K		\$0.88		\$0.18
0932	Factor IX recombinant	K		\$1.06		\$0.22
0933	Gamma globulin > 10 CC inj	K		\$150.50		\$30.10
0934	Capecitabine, oral	K		\$18.73		\$3.75
0935	Clonidine hydrochloride	K		\$109.75		\$21.95
0943	Octagam injection	K		\$37.03		\$7.41
0944	Gammagard liquid injection	K		\$37.85		\$7.57
0945	Rhophylac injection	K		\$5.13		\$1.03
0946	Hepagam b lim injection	G		\$50.04		\$9.82

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
1263	Anthrithrombin iii injection	K		\$2.28		\$0.46
1266	Interferon alfacon-1 inj	K		\$6.75		\$1.95
1268	Xyntha inj	K		\$1.06		\$0.22
1270	Alloderm skin sub	K		\$31.72		\$6.35
1271	Cholera vaccine, injectable	K		\$0.16		\$0.04
1272	Acetylcysteine injection	K		\$2.29		\$0.48
1273	Dimecaprol Injection	K		\$26.81		\$5.37
1274	Eccelate calcium disodium inj	K		\$78.86		\$15.78
1275	Vivaglobin, inj	K		\$7.05		\$1.41
1276	Fondaparinux sodium	K		\$5.98		\$1.20
1277	Insulin for insulin pump use	K		\$3.34		\$0.67
1279	Factor VIII (porcine)	K		\$2.00		\$0.40
1280	Carticortropin injection	K		\$2,394.93		\$478.99
1281	Bevacizumab injection	K		\$1.41		\$0.29
1282	Gamma globulin 2 CC inj	K		\$30.10		\$9.03
1283	Gamma globulin 3 CC inj	K		\$45.14		\$9.03
1284	Gamma globulin 5 CC inj	K		\$75.26		\$15.06
1285	Nandrolone decanoate 50 MG	K		\$7.00		\$1.40
1286	Nandrolone decanoate 200 MG	K		\$14.74		\$2.95
1287	Visaliskin skin sub	K		\$9.36		\$1.88
1288	Visualization adjunct	K		\$4.11		\$0.83
1289	Abobotulinumtoxin type A	K		\$9.23		\$1.65
1290	Human fibrinogen conc inj	G		\$96.46		\$18.93
1291	Rilonacept injection	K		\$23.64		\$4.73
1292	Cyclosporine oral 100 mg	K		\$3.22		\$0.65
1293	Bladder calculi irrig sol	K		\$29.28		\$5.86
1294	Cyclosporine oral 25 mg	K		\$0.82		\$0.17
1295	Sm 153 lexidronam	K		\$4,671.66		\$934.34
1296	Degarelix injection	G		\$2.23		\$0.44
1297	Ferumoxytol, non-esrd	G		\$0.82		\$0.16
1298	Cosyntropin cortrosyn inj	K		\$91.84		\$18.37
1299	Gandofosvetet trisodium inj	K		\$1.29		\$0.00
1436	Etidronate disodium inj	K		\$70.06		\$14.02
1491	New Technology - Level IA (\$0-\$10)	S		\$5.00		\$1.00
1492	New Technology - Level IB (\$10-\$20)	S		\$15.00		\$3.00
1493	New Technology - Level IC (\$20-\$30)	S		\$25.00		\$5.00
1494	New Technology - Level ID (\$30-\$40)	S		\$35.00		\$7.00
1495	New Technology - Level IE (\$40-\$50)	S		\$45.00		\$9.00
1496	New Technology - Level IA (\$0-\$10)	T		\$5.00		\$1.00
1497	New Technology - Level IB (\$10-\$20)	T		\$15.00		\$3.00
1498	New Technology - Level IC (\$20-\$30)	T		\$25.00		\$5.00
1499	New Technology - Level ID (\$30-\$40)	T		\$35.00		\$7.00
1500	New Technology - Level IE (\$40-\$50)	T		\$45.00		\$9.00

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
1166	Cytarabine liposome inj	K		\$480.19		\$96.04
1167	inj, eprubicin hcl	K		\$2.95		\$0.51
1168	inj, temsirolimus	K		\$47.93		9.59
1178	Busulfan injection	K		\$14.18		\$2.84
1203	Verteporfin injection	K		\$9.31		\$1.87
1204	Cyclosporin parenteral	K		\$21.24		\$4.25
1207	Octreotide injection, depot	K		\$105.27		\$21.06
1209	Diethylstilbestrol injection	K		\$1,257.36		\$251.48
1213	Antihemophilic viiawf comp	K		\$0.84		\$0.17
1214	inj IVIG privitygen 500 mg	G		\$35.05		\$6.88
1216	Lyme disease vaccine, im	K		\$0.93		\$0.19
1220	Calcitonin salmon injection	K		\$48.37		\$9.68
1221	Dimethyl sulfoxide 50%	K		\$67.46		\$13.50
1222	Penistatarch 10% solution	K		\$1,270.88		\$254.18
1225	Somatrelin injection	K		\$43.99		\$8.80
1226	inj streptokinase (2500000 IU	K		\$73.00		\$15.60
1232	Mitomycin 5 MG inj	K		\$17.74		\$3.55
1233	Mitomycin 20 MG inj	K		\$70.98		\$14.20
1234	Mitomycin 40 MG inj	K		\$141.95		\$28.39
1235	Valrubicin injection	K		\$953.16		\$190.64
1236	Levofloxacin injection	G		\$0.99		\$0.19
1237	inj iron dextran	K		\$14.11		\$2.83
1238	Topotecan oral	G		\$71.35		\$14.00
1239	Rotavirus vacc 2 dose oral	K		\$106.60		\$21.32
1240	Apilgraf skin sub	K		\$32.16		\$6.44
1241	Oasis wound matrix skin sub	K		\$4.12		\$0.83
1242	Oasis burn matrix skin sub	K		\$4.12		\$0.83
1243	Integra BMWD skin sub	K		\$11.77		\$2.36
1244	Integra DRT skin sub	K		\$11.77		\$2.36
1245	Demagraft skin sub	K		\$39.25		\$7.85
1246	Integra matrix skin sub	K		\$69.23		\$17.85
1247	Integra matrix skin sub	K		\$17.98		\$3.60
1248	Primatrix skin sub	K		\$33.99		\$6.80
1249	Cymetra allograft	K		\$327.47		\$65.50
1250	Graftackett express allograft	K		\$327.47		\$65.50
1251	Integra, flowable wound mairi	G		\$907.36		\$178.05
1252	Gammagraft skin sub	K		\$7.13		\$1.43
1253	Triamcinolone A inj PRS-free	K		\$3.20		\$0.64
1254	Adenovirus vaccine, type 4	K		\$72.17		\$14.44
1255	Rotavirus vacc, 3 dose, oral	K		\$72.37		\$14.48
1256	Bronphentramine maleate inj	K		\$0.75		\$0.15
1257	Entuviridine injection	K		\$0.47		\$0.10
1260	Nandrolone decanoate 100 MG	K		\$7.00		\$1.40

APPENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
1546	New Technology - Level IX (\$700 - \$800)	T		\$750.00		\$150.00
1547	New Technology - Level X (\$800 - \$900)	T		\$850.00		\$170.00
1548	New Technology - Level XI (\$900 - \$1000)	T		\$950.00		\$190.00
1549	New Technology - Level XII (\$1000 - \$1100)	T		\$1,050.00		\$210.00
1550	New Technology - Level XIII (\$1100 - \$1200)	T		\$1,150.00		\$230.00
1551	New Technology - Level XIV (\$1200 - \$1300)	T		\$1,250.00		\$250.00
1552	New Technology - Level XV (\$1300 - \$1400)	T		\$1,350.00		\$270.00
1553	New Technology - Level XVI (\$1400 - \$1500)	T		\$1,450.00		\$290.00
1554	New Technology - Level XVII (\$1500-\$1600)	T		\$1,550.00		\$310.00
1555	New Technology - Level XVIII (\$1600-\$1700)	T		\$1,650.00		\$330.00
1556	New Technology - Level XIX (\$1700-\$1800)	T		\$1,750.00		\$350.00
1557	New Technology - Level XX (\$1800-\$1900)	T		\$1,850.00		\$370.00
1558	New Technology - Level XXI (\$1900-\$2000)	T		\$1,950.00		\$390.00
1559	New Technology - Level XXII (\$2000-\$2500)	T		\$2,250.00		\$450.00
1560	New Technology - Level XXIII (\$2500-\$3000)	T		\$2,750.00		\$550.00
1561	New Technology - Level XXIV (\$3000-\$3500)	T		\$3,250.00		\$650.00
1562	New Technology - Level XXV (\$3500-\$4000)	T		\$3,750.00		\$750.00
1563	New Technology - Level XXVI (\$4000-\$4500)	T		\$4,250.00		\$850.00
1564	New Technology - Level XXVII (\$4500-\$5000)	T		\$4,750.00		\$950.00
1565	New Technology - Level XXVIII (\$5000-\$5500)	T		\$5,250.00		\$1,050.00
1566	New Technology - Level XXIX (\$5500-\$6000)	T		\$5,750.00		\$1,150.00
1567	New Technology - Level XXX (\$6000-\$6500)	T		\$6,250.00		\$1,250.00
1568	New Technology - Level XXXI (\$6500-\$7000)	T		\$6,750.00		\$1,350.00
1569	New Technology - Level XXXII (\$7000-\$7500)	T		\$7,250.00		\$1,450.00
1570	New Technology - Level XXXIII (\$7500-\$8000)	T		\$7,750.00		\$1,550.00
1571	New Technology - Level XXXIV (\$8000-\$8500)	T		\$8,250.00		\$1,650.00
1572	New Technology - Level XXXV (\$8500-\$9000)	T		\$8,750.00		\$1,750.00
1573	New Technology - Level XXXVI (\$9000-\$9500)	T		\$9,250.00		\$1,850.00
1574	New Technology - Level XXXVII (\$9500-\$10000)	T		\$9,750.00		\$1,950.00
1605	Abciximab injection	K		\$459.36		\$91.88
1607	Eptifibatid injection	K		\$3.72		\$3.72
1608	Etanercept injection	K		\$183.61		\$36.73
1609	Rho(D) immune globulin h, sd	K		\$18.39		\$3.68
1612	Dacizumab, parenteral	K		\$378.20		\$75.64
1613	Trastuzumab injection	K		\$63.51		\$12.71
1630	Hep b ig, im	K		\$111.20		\$22.24
1631	Backlen intrathecal trial	K		\$71.24		\$14.25
1633	Alfacept	K		\$30.02		\$6.01
1643	Y90 Ibritumomab, rx	K		\$15,532.15		\$3,106.43
1645	I131 tositumomab, rx	K		\$9,929.05		\$1,985.81
1670	Tetanus immune globulin inj	K		\$199.91		\$39.99
1675	P32 Na phosphate	K		\$196.49		\$39.30
1676	P32 Chromic phosphate	K		\$113.44		\$22.69

APPENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
1502	New Technology - Level II (\$50 - \$100)	S		\$75.00		\$15.00
1503	New Technology - Level III (\$100 - \$200)	S		\$150.00		\$30.00
1504	New Technology - Level IV (\$200 - \$300)	S		\$250.00		\$50.00
1505	New Technology - Level V (\$300 - \$400)	S		\$350.00		\$70.00
1506	New Technology - Level VI (\$400 - \$500)	S		\$450.00		\$90.00
1507	New Technology - Level VII (\$500 - \$600)	S		\$550.00		\$110.00
1508	New Technology - Level VIII (\$600 - \$700)	S		\$650.00		\$130.00
1509	New Technology - Level IX (\$700 - \$800)	S		\$750.00		\$150.00
1510	New Technology - Level X (\$800 - \$900)	S		\$850.00		\$170.00
1511	New Technology - Level XI (\$900 - \$1000)	S		\$950.00		\$190.00
1512	New Technology - Level XII (\$1000 - \$1100)	S		\$1,050.00		\$210.00
1513	New Technology - Level XIII (\$1100 - \$1200)	S		\$1,150.00		\$230.00
1514	New Technology - Level XIV (\$1200 - \$1300)	S		\$1,250.00		\$250.00
1515	New Technology - Level XV (\$1300 - \$1400)	S		\$1,350.00		\$270.00
1516	New Technology - Level XVI (\$1400 - \$1500)	S		\$1,450.00		\$290.00
1517	New Technology - Level XVII (\$1500-\$1600)	S		\$1,550.00		\$310.00
1518	New Technology - Level XVIII (\$1600-\$1700)	S		\$1,650.00		\$330.00
1519	New Technology - Level XIX (\$1700-\$1800)	S		\$1,750.00		\$350.00
1520	New Technology - Level XX (\$1800-\$1900)	S		\$1,850.00		\$370.00
1521	New Technology - Level XXI (\$1900-\$2000)	S		\$1,950.00		\$390.00
1522	New Technology - Level XXII (\$2000-\$2500)	S		\$2,250.00		\$450.00
1523	New Technology - Level XXIII (\$2500-\$3000)	S		\$2,750.00		\$550.00
1524	New Technology - Level XXIV (\$3000-\$3500)	S		\$3,250.00		\$650.00
1525	New Technology - Level XXV (\$3500-\$4000)	S		\$3,750.00		\$750.00
1526	New Technology - Level XXVI (\$4000-\$4500)	S		\$4,250.00		\$850.00
1527	New Technology - Level XXVII (\$4500-\$5000)	S		\$4,750.00		\$950.00
1528	New Technology - Level XXVIII (\$5000-\$5500)	S		\$5,250.00		\$1,050.00
1529	New Technology - Level XXIX (\$5500-\$6000)	S		\$5,750.00		\$1,150.00
1530	New Technology - Level XXX (\$6000-\$6500)	S		\$6,250.00		\$1,250.00
1531	New Technology - Level XXXI (\$6500-\$7000)	S		\$6,750.00		\$1,350.00
1532	New Technology - Level XXXII (\$7000-\$7500)	S		\$7,250.00		\$1,450.00
1533	New Technology - Level XXXIII (\$7500-\$8000)	S		\$7,750.00		\$1,550.00
1534	New Technology - Level XXXIV (\$8000-\$8500)	S		\$8,250.00		\$1,650.00
1535	New Technology - Level XXXV (\$8500-\$9000)	S		\$8,750.00		\$1,750.00
1536	New Technology - Level XXXVI (\$9000-\$9500)	S		\$9,250.00		\$1,850.00
1537	New Technology - Level XXXVII (\$9500-\$10000)	S		\$9,750.00		\$1,950.00
1539	New Technology - Level II (\$50 - \$100)	T		\$75.00		\$15.00
1540	New Technology - Level III (\$100 - \$200)	T		\$150.00		\$30.00
1541	New Technology - Level IV (\$200 - \$300)	T		\$250.00		\$50.00
1542	New Technology - Level V (\$300 - \$400)	T		\$350.00		\$70.00
1543	New Technology - Level VI (\$400 - \$500)	T		\$450.00		\$90.00
1544	New Technology - Level VII (\$500 - \$600)	T		\$550.00		\$110.00
1545	New Technology - Level VIII (\$600 - \$700)	T		\$650.00		\$130.00

APPENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
2731	Immune globulin, powder	K		\$29.83		\$5.97
2770	Quinupristin/dalfopristin	K		\$144.08		\$28.82
3030	Sumatriptan succinate	K		\$55.49		\$11.10
3041	Bivalirudin	K		\$2.40		\$0.48
3043	Gamma globulin 1 CC inj	K		\$15.05		\$3.01
3050	Sermorelin acetate injection	K		\$350.07		\$70.02
7000	Amifostine	K		\$176.89		\$35.38
7005	Gonadorelin hydroch	K		\$242.16		\$48.44
7011	Oprelvekin injection	K		\$53.47		\$10.70
7034	Somatropin injection	K		\$319.43		\$63.89
7035	Teniposide	K		\$449.09		\$89.82
7036	Monoclonal antibodies	K		\$1,109.45		\$221.89
7038	Urokinase 250,000 IU inj	K		\$7.83		\$1.57
7041	Tirofiban HCl	K		\$5.68		\$1.14
7042	Capecitabine, oral	K		\$57.60		\$11.52
7043	Infliximab injection	K		\$124.80		\$24.96
7045	inj trimetrexate glucuronate	K		\$450.51		\$90.11
7046	Doxorubicin hcl liposome inj	K		\$324.44		\$64.89
7048	Alteplase recombinant	K		\$35.03		\$7.01
7049	Fingolimod 480 mcg injection	K		\$324.44		\$64.89
7051	Leuprolide acetate implant	K		\$4,728.88		\$945.78
7308	Annulevulmic acid hcl top	K		\$127.60		\$25.52
8000	Cardiac Electrophysiologic Evaluation and Ablation Composite	T	150.1090	\$10,118.25		\$2,023.65
8001	LDR Prostate Brachytherapy Composite	T	46.1770	\$3,112.61		\$622.53
8002	Level I Extended Assessment & Management Composite	V	5.6573	\$381.34		\$76.27
8003	Level II Extended Assessment & Management Composite	V	10.4630	\$705.27		\$141.06
8004	Ultrasound Composite	S	2.8400	\$191.43		\$38.29
8005	CT and CTA without Contrast Composite	S	6.2228	\$419.45		\$83.89
8006	CT and CTA with Contrast Composite	S	9.3240	\$628.49		\$125.70
8007	MRI and MRA without Contrast Composite	S	10.5672	\$712.29		\$142.46
8008	MRI and MRA with Contrast Composite	S	14.7694	\$995.55		\$199.11
9001	Linectoplas injection	K		\$29.37		\$5.88
9002	Tenecteplase injection	K		\$40.10		\$8.02
9003	Palivizumab	K		\$937.29		\$187.46
9004	Gemtuzumab ozogamicin inj	K		\$2,572.82		\$514.57
9005	Pelteplase injection	K		\$1,230.80		\$246.16
9006	Tacrolimus injection	K		\$136.82		\$27.37
9012	Arsenic trioxide injection	K		\$36.73		\$7.35
9015	Mycophenolate mofetil oral	K		\$2.45		\$0.49
9018	inj, imabotulinumtoxinB	K		\$10.36		\$2.08
9019	Caspofungin acetate	K		\$11.52		\$2.31
9020	Sirolimus, oral	K		\$9.44		\$1.89

APPENDUM A.—FINAL OPPTS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
1682	Apronin, 10,000 iu	K		\$2.90		\$0.52
1683	Basiximab	K		\$1,624.44		\$324.89
1684	Corticorelin ovine triflutal	K		\$0.85		\$0.17
1685	Darbecoplin alfa, non-esrd	K		\$2.76		\$0.56
1686	Epoetin alfa, non-esrd	K		\$9.40		\$1.88
1687	Digoxin immune fab (ovine)	K		\$474.73		\$94.95
1688	Ethanolamine oleate	K		\$147.14		\$29.43
1689	Fomepizole	K		\$7.99		\$1.60
1690	Hemin	K		\$7.73		\$1.55
1693	Lepruridin	K		\$174.51		\$34.91
1694	Ziconotide injection	K		\$6.65		\$1.33
1695	Nesiritide injection	K		\$36.07		\$7.22
1696	Palifermin injection	K		\$11.06		\$2.22
1697	Pegaptanib sodium injection	K		\$19.93		\$3.99
1700	inj secretin synthetic human	K		\$1,014.11		\$202.83
1701	Treprostinil injection	K		\$54.83		\$10.97
1704	Humate-P, inj	K		\$0.87		\$0.18
1705	Factor vlla	K		\$1.29		\$0.26
1709	Azacitidine injection	K		\$4.78		\$0.96
1710	Clofarabine injection	K		\$114.21		\$22.85
1711	Vantus implant	G		\$1,473.60		\$289.16
1712	Paclitaxel protein bound	K		\$9.09		\$1.82
1716	Brachytx, non-str, Gold-198	U	0.6357	\$42.85		\$8.57
1717	Brachytx, non-str, HDR Ir-192	U	3.4326	\$231.38		\$46.28
1719	Brachytx, NS, Non-HDR Ir-192	U	0.9497	\$64.02		\$12.81
1738	Oxaliplatin	K		\$9.55		\$1.91
1739	Pegademase bovine, 25 iu	K		\$242.67		\$48.54
1740	Diazoxide injection	K		\$112.16		\$22.44
1741	Urofollitropin, 75 iu	K		\$59.26		\$11.86
2210	Methyldopa hcl injection	U	234.0942	\$15,779.35		\$5.53
2616	Brachytx, non-str, Yttrium-90	U	0.5626	\$37.92		\$7.59
2632	Iodine I-125 sodium iodide	U	0.8871	\$59.80		\$11.96
2634	Brachytx, non-str, HA, I-125	U	0.4242	\$26.59		\$5.72
2635	Brachytx, non-str, HA, P-103	U	0.2873	\$19.37		\$3.88
2638	Brachytx, stranded, I-125	U	0.6302	\$42.48		\$8.50
2639	Brachytx, non-stranded, I-125	U	0.5368	\$36.18		\$7.24
2640	Brachytx, stranded, P-103	U	0.8955	\$60.36		\$12.08
2641	Brachytx, non-stranded, P-103	U	0.8474	\$57.12		\$11.43
2642	Brachytx, stranded, C-131	U	1.6295	\$109.84		\$21.97
2643	Brachytx, non-stranded, C-131	U	0.9805	\$66.09		\$13.22
2698	Brachytx, stranded, NOS	U	0.6302	\$42.48		\$8.50
2699	Brachytx, non-stranded, NOS	U	0.4242	\$26.59		\$5.72

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
9231	Decitabine injection	K		\$28.42		\$5.68
9232	Idursulfase injection	K		\$446.44		\$89.29
9233	Panbuzumab injection	K		\$398.11		\$79.63
9234	Alglucosidase alfa injection	K		\$124.69		\$24.94
9235	Panitumumab injection	K		\$85.21		\$17.05
9236	Eculizumab injection	K		\$177.57		\$35.52
9237	Inj, lanreotide acetate	K		\$28.65		\$5.73
9238	Inj, levitracetam	G		\$0.75		\$0.15
9240	Injection, azabipione	G		\$63.74		\$12.51
9241	Injection, doxiprenem	G		\$0.57		\$0.11
9242	Injection, fosaprepitant	G		\$1.58		\$0.31
9243	Bandamustine injection	G		\$18.53		\$3.64
9244	Regadenoson injection	G		\$50.78		\$9.96
9245	Homiplostin injection	G		\$43.75		\$8.58
9246	Gastroxetate disodium inj	G		\$13.50		\$0.00
9247	Inj, iobenguane, I-123, dx	G		\$2,329.83		\$465.97
9248	Inj, clevudipine butyrate	G		\$3.39		\$0.67
9249	Centozumab pegol inj	G		\$3.80		\$0.75
9250	Artiss fibrin sealant	G		\$138.20		\$27.12
9251	CT esterase inhibitor inj	G		\$41.34		\$8.11
9252	Plexiator injection	G		\$268.51		\$52.69
9253	Temozolomide injection	G		\$4.90		\$0.96
9254	Injection, lacosamide	K		\$0.18		\$0.04
9255	Paliperidone palmitate inj	G		\$6.71		\$1.35
9256	Dexamethasone intravitreal	G		\$198.10		\$38.48
9300	Omalizumab injection	K		\$18.86		\$3.78
9356	TenoGlide tendon prot, cm2	K		\$24.86		\$0.00
9358	SurgiMend, fetal	G		\$10.76		\$2.11
9359	Impinj bon void filler-putty	G		\$63.54		\$0.00
9360	SurgiMend, neonatal	G		\$10.67		\$2.09
9361	NeuroMend nerve wrap	G		\$247.29		\$0.00
9362	Impinj bon void filler-strip	G		\$63.60		\$0.00
9363	Integra Meshed Bil Wound Mat	G		\$25.62		\$5.03
9364	Porcine implant, Permacol	G		\$17.21		\$0.00
9500	Platelets, irradiated	R	2.2920	\$150.45		\$30.09
9501	Platelet pheres leukoreduced	R	7.5974	\$512.11		\$102.43
9502	Platelet pheres irradiated	R	5.3105	\$357.96		\$71.60
9503	Fr tiz plasma donor released	R	1.0664	\$71.88		\$14.38
9504	RBC deglycerolized	R	5.3988	\$363.91		\$72.79
9505	RBC irradiated	R	3.3499	\$225.80		\$45.16
9506	Granulocytes, pheresis unit	R	0.6664	\$44.92		\$8.99
9507	Platelets, pheresis	R	6.9594	\$469.11		\$93.83
9508	Plasma 1 donor frz w/in 8 hr	R	1.1278	\$76.02		\$15.21

ADDENDUM A.—FINAL OPFS APCs FOR CY 2010

APC	Group Title	SI	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
9022	IM inj interferon beta 1-a	K		\$187.24		\$37.45
9023	Rho d immune globulin	K		\$25.78		\$5.16
9024	Amphotericin b lipid complex	K		\$9.66		\$1.94
9032	Baclofen 10 MG injection	K		\$195.31		\$39.07
9033	Cidofovir injection	K		\$746.46		\$149.30
9038	Inj estrogen conjugate	K		\$83.21		\$16.65
9042	Glucagon hydrochloride	K		\$79.20		\$15.84
9044	lulizide fumarate injection	K		\$404.01		\$80.81
9046	Iron sucrose injection	K		\$0.37		\$0.08
9104	Antithymocyte globulin rabbit	K		\$414.44		\$82.89
9108	Thyrotropin injection	K		\$98.38		\$19.68
9110	Alemtuzumab injection	K		\$559.46		\$111.90
9115	Zoledronic acid	K		\$214.94		\$42.99
9119	Injection, pegnigrastim 6mg	K		\$2,222.07		\$444.42
9120	Injection, Fulvestrant	K		\$80.63		\$16.13
9121	Injection, argatroban	K		\$18.10		\$3.62
9122	Triptorelin pamoate	K		\$160.83		\$32.17
9124	Daptomycin injection	K		\$0.40		\$0.08
9125	Risperidone, long acting	K		\$4.93		\$0.99
9126	Natalizumab injection	K		\$63.32		\$12.67
9133	Rabies ig, im/sc	K		\$142.79		\$28.56
9134	Rabies ig, heat treated	K		\$130.16		\$26.04
9135	Varicella-zoster ig, im	K		\$130.49		\$26.10
9137	Bcg vaccine, percult	K		\$111.66		\$22.34
9139	Rabies vaccine, im	K		\$151.97		\$30.40
9140	Rabies vaccine, id	K		\$96.27		\$19.26
9143	Meningococcal vaccine, sc	K		\$96.66		\$19.34
9144	Encephalitis vaccine, sc	K		\$100.15		\$20.03
9145	Meningococcal vaccine, im	K		\$102.46		\$20.50
9207	Bortezomib injection	K		\$56.54		\$7.31
9208	Agalsidase beta injection	K		\$133.69		\$26.74
9209	Larondase injection	K		\$25.08		\$5.02
9210	Palonosetron hcl	K		\$17.19		\$3.44
9213	Pemretaxed injection	K		\$48.50		\$9.70
9214	Bevacizumab injection	K		\$56.39		\$11.28
9215	Celuximab injection	K		\$48.79		\$9.76
9217	Leuprolide acetate suspension	K		\$210.52		\$42.11
9224	Galsulfase injection	K		\$339.04		\$67.81
9225	Fluocinolone acetonide impjt	K		\$18,980.00		\$3,796.00
9227	Micafungin sodium injection	K		\$1.08		\$0.22
9228	Tigecycline injection	K		\$1.15		\$0.23
9229	Ibandronate sodium injection	K		\$139.22		\$27.85
9230	Abatacept injection	K		\$18.98		\$3.80

ADDENDUM AA.—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
10022	Fria w/image	Y		G2	4.4	\$184.24
10040	Acne surgery	Y		P2	0.8408	\$35.21
10060	Drainage of skin abscess	Y		P3		\$42.61
10080	Drainage of pilonidal cyst	Y		P2	1.3927	\$58.32
10081	Drainage of pilonidal cyst	Y		P3	1.3927	\$58.32
10120	Remove foreign body	Y		P3		\$59.08
10121	Remove foreign body	Y		A2	15.1023	\$62.38
10140	Drainage of hematoma/fluid	Y		P3		\$63.35
10160	Puncture drainage of lesion	Y	CH	P3		\$52.55
10180	Complex drainage, wound	Y		A2	16.472	\$689.73
11000	Debride infected skin	Y		P3		\$20.17
11001	Debride infected skin add-on	Y		P3		\$6.82
11010	Debride skin, fx	Y		A2	4.5669	\$191.23
11011	Debride skin/muscle, fx	Y		A2	4.5669	\$191.23
11012	Debride skin/muscle/bone, fx	Y		A2	4.5669	\$191.23
11040	Debride skin, partial	Y		P3		\$18.75
11041	Debride skin, full	Y		P3		\$20.45
11042	Debride skin/lesion	Y		A2	2.947	\$123.40
11043	Debride lissue/muscle	Y		A2	2.947	\$123.40
11044	Debride lissue/muscle/bone	Y		A2	8.3025	\$347.65
11055	Trim skin lesion	Y		P3		\$21.87
11056	Trim skin lesions, 2 to 4	Y		P3		\$23.86
11057	Trim skin lesions, over 4	Y	CH	P3		\$26.99
11100	Bloopy, skin lesion	Y	CH	P3		\$49.71
11101	Biopsy, skin add-on	Y		P3		\$11.65
11200	Removal of skin tags	Y		P2	0.8408	\$35.21
11201	Remove skin tags add-on	Y		P3		\$4.83
11300	Shave skin lesion	Y	CH	P3		\$33.24
11301	Shave skin lesion	Y		P2	0.8408	\$35.21

Note 1: The Medicare program payment is 80 percent of the total payment amount and beneficiary coinsurance is 20 percent of the total payment amount, except for screening flexible sigmoidoscopies and screening colonoscopies for which the program payment is 75 percent and the beneficiary coinsurance is 25 percent.

Note 2: Payment indicators for "office-based" procedures (P2, P3) are based on a comparison of the final rates according to the ASC standard ratesetting methodology and the MPFS. Under current law, MPFS payment rates will have a negative update for CY 2010. For a discussion of these rates, we refer readers to the CY 2010 MPFS final rule.

: Asterisked codes () indicate that the procedure's "office-based," designation is temporary because we have insufficient claims data. We will reconsider this designation when new claims data become available.

ADDENDUM AA.—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
0016T	Thermax choroid vasc. lesion	Y		R2	5.5965	\$234.34
0017T	Photocoagulat macular drusen	Y		R2	5.5965	\$234.34
0084T	Temp prostatic urethral stent	N	CH	D5		
0098T	L ventricle fill pressure	N	CH	D5		
0098T*	Implant corneal ring	Y		R2	15.5449	\$650.91
0100T	Prosth retina receive&gen	Y		G2	38.2338	\$1,600.96
0101T	Extracorp shockwv tx, hi. enrg	Y		G2	30.396	\$1,272.77
0102T	Extracorp shockwv tx, anessth	Y		G2	30.396	\$1,272.77
0123T	Scialar fistulization	Y		G2	23.3455	\$977.55
0124T*	Conjunctival drug placement	Y		R2	4.3122	\$180.56
0170T	Anorectal fistula plug pr	N	CH	D5		
0176T	Aqu canal dilat w/o relent	Y		A2	37.7011	\$1,578.66
0177T	Aqu canal dilat w relent	Y		A2	37.7011	\$1,578.66
0186T	Suprachoroidal drug delivery	Y		G2	19.8176	\$829.82
0190T	Place intraoc radiation src	Y		G2	19.8176	\$829.82
0191T	Insert ant segment drain int	Y		G2	23.3455	\$977.55
0192T	Insert ant segment drain ext	Y		G2	40.0704	\$1,677.87
0193T	Rf bladder neck microremodel	Y	CH	G2	21.0617	\$881.92
0200T	Perq sacral augmt unilat inj	Y		G2	30.396	\$1,272.77
0201T	Perq sacral augmt bilat inj	Y		G2	30.396	\$1,272.77
0213T	Us facet jt inj cervit 1 lev	Y	NI	G2	6.8884	\$288.44
0214T	Us facet jt inj cervit 2 lev	Y	NI	G2	2.4451	\$102.38
0215T	Us facet jt inj cervit 3 lev	Y	NI	G2	2.4451	\$102.38
0216T	Us facet jt inj ls 1 level	Y	NI	G2	6.8884	\$288.44
0217T	Us facet jt inj ls 2 level	Y	NI	G2	2.4451	\$102.38
0218T	Us facet jt inj ls 3 level	Y	NI	G2	2.4451	\$102.38
10021	Fria w/o image	Y		P2	1.4457	\$60.54

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
11462	Removal, sweat gland lesion	Y		A2	19.3292	\$609.37
11463	Removal, sweat gland lesion	Y		A2	19.3292	\$609.37
11470	Removal, sweat gland lesion	Y		A2	19.3292	\$609.37
11471	Removal, sweat gland lesion	Y		A2	19.3292	\$609.37
11600	Exc tr-ext mlg+marq 0.5 < cm	Y		P3		\$80.39
11601	Exc tr-ext mlg+marq 0.6-1 cm	Y		P3		\$96.30
11602	Exc tr-ext mlg+marq 1.1-2 cm	Y		P3		\$105.39
11603	Exc tr-ext mlg+marq 2.1-3 cm	Y		P3		\$112.77
11604	Exc tr-ext mlg+marq 3.1-4 cm	Y		A2	8.2762	\$346.55
11606	Exc tr-ext mlg+marq > 4 cm	Y		A2	15.1023	\$632.38
11620	Exc h-f-nk-sp mlg+marq 0.5 <	Y		P3		\$82.66
11621	Exc h-f-nk-sp mlg+marq 0.6-1	Y		P3		\$97.43
11622	Exc h-f-nk-sp mlg+marq 1.1-2	Y		P3		\$107.94
11623	Exc h-f-nk-sp mlg+marq 2.1-3	Y		P3		\$117.03
11624	Exc h-f-nk-sp mlg+marq 3.1-4	Y		A2	15.1023	\$632.38
11626	Exc h-f-nk-sp mlg+mar > 4 cm	Y		A2	19.3292	\$609.37
11640	Exc face-mm malig+marq 0.5 <	Y		P3		\$86.92
11641	Exc face-mm malig+marq 0.6-1	Y		P3		\$101.98
11642	Exc face-mm malig+marq 1.1-2	Y		P3		\$113.62
11643	Exc face-mm malig+marq 2.1-3	Y		P3		\$123.28
11644	Exc face-mm malig+marq 3.1-4	Y		A2	15.1023	\$632.38
11646	Exc face-mm malig+marq > 4 cm	Y		A2	19.3292	\$609.37
11719	Trim nail(s)	Y		P3		\$10.23
11720	Debride nail, 1-5	Y		P3		\$12.50
11721	Debride nail, 6 or more	Y		P3		\$15.06
11730	Removal of nail plate	Y		P2	0.8408	\$35.21
11732	Remove nail plate, add-on	Y		P3		\$15.06
11740	Drain blood from under nail	Y		P2	0.4244	\$17.77
11750	Removal of nail bed	Y		P3		\$81.24
11752	Remove nail bed/finger tip	Y		P3		\$113.34

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
11302	Shave skin lesion	Y		P2	0.8408	\$35.21
11303	Shave skin lesion	Y	CH	P3		\$55.96
11305	Shave skin lesion	Y	CH	P3		\$29.54
11306	Shave skin lesion	Y		P2	0.8408	\$35.21
11307	Shave skin lesion	Y		P2	0.8408	\$35.21
11308	Shave skin lesion	Y		P2	0.8408	\$35.21
11310	Shave skin lesion	Y		P2	0.8408	\$35.21
11311	Shave skin lesion	Y		P2	0.8408	\$35.21
11312	Shave skin lesion	Y		P2	0.8408	\$35.21
11313	Shave skin lesion	Y		P3		\$55.21
11400	Exc tr-ext b9+marq 0.5 < cm	Y		P3		\$56.53
11401	Exc tr-ext b9+marq 0.6-1 cm	Y		P3		\$63.63
11402	Exc tr-ext b9+marq 1.1-2 cm	Y		P3		\$69.88
11403	Exc tr-ext b9+marq 2.1-3 cm	Y		P3		\$74.99
11404	Exc tr-ext b9+marq 3.1-4 cm	Y		A2	14.457	\$605.36
11406	Exc tr-ext b9+marq > 4.0 cm	Y		A2	15.1023	\$632.38
11420	Exc h-f-nk-sp b9+marq 0.5 <	Y		P3		\$53.12
11421	Exc h-f-nk-sp b9+marq 0.6-1	Y		P3		\$64.20
11422	Exc h-f-nk-sp b9+marq 1.1-2	Y		P3		\$70.16
11423	Exc h-f-nk-sp b9+marq 2.1-3	Y		P3		\$78.40
11424	Exc h-f-nk-sp b9+marq 3.1-4	Y		A2	15.1023	\$632.38
11426	Exc h-f-nk-sp b9+marq > 4 cm	Y		A2	19.3292	\$609.37
11440	Exc face-mm b9+marq 0.5 < cm	Y		P3		\$60.50
11441	Exc face-mm b9+marq 0.6-1 cm	Y		P3		\$70.16
11442	Exc face-mm b9+marq 1.1-2 cm	Y		P3		\$77.26
11443	Exc face-mm b9+marq 2.1-3 cm	Y		P3		\$85.50
11444	Exc face-mm b9+marq 3.1-4 cm	Y		A2	7.7878	\$326.10
11446	Exc face-mm b9+marq > 4 cm	Y		A2	19.3292	\$609.37
11450	Removal, sweat gland lesion	Y		A2		\$809.37
11451	Removal, sweat gland lesion	Y		A2	19.3292	\$609.37

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
12011	Repair superficial wound(s)	Y		P2	1.2956	\$54.25
12013	Repair superficial wound(s)	Y		P2	1.2956	\$54.25
12014	Repair superficial wound(s)	Y		P2	1.2956	\$54.25
12015	Repair superficial wound(s)	Y		G2	1.2956	\$54.25
12016	Repair superficial wound(s)	Y		A2	1.4926	\$62.50
12017	Repair superficial wound(s)	Y		A2	1.4926	\$62.50
12018	Repair superficial wound(s)	Y		A2	1.4926	\$62.50
12020	Closure of split wound	Y		A2	3.706	\$155.18
12021	Closure of split wound	Y		A2	2.782	\$116.49
12031	Intmd wnd repair s/r/text	Y		P2	1.2956	\$54.25
12032	Intmd wnd repair s/r/text	Y		P2	3.0144	\$126.22
12034	Intmd wnd repair s/r/text	Y		A2	1.4926	\$62.50
12035	Intmd wnd repair s/r/text	Y		A2	1.4926	\$62.50
12036	Intmd wnd repair s/r/text	Y		A2	2.782	\$116.49
12037	Intmd wnd repair s/r/text	Y		A2	4.1074	\$171.99
12041	Intmd wnd repair n-hg/genit	Y		P2	1.2956	\$54.25
12042	Intmd wnd repair n-hg/genit	Y		P2	1.2956	\$54.25
12044	Intmd wnd repair n-hg/genit	Y		A2	1.4926	\$62.50
12045	Intmd wnd repair n-hg/genit	Y		A2	2.782	\$116.49
12046	Intmd wnd repair n-hg/genit	Y		A2	2.782	\$116.49
12047	Intmd wnd repair n-hg/genit	Y		A2	4.1074	\$171.99
12051	Intmd wnd repair face/mm	Y		P2	1.2956	\$54.25
12052	Intmd wnd repair face/mm	Y		P2	1.2956	\$54.25
12053	Intmd wnd repair face/mm	Y		P2	1.2956	\$54.25
12054	Intmd wnd repair face/mm	Y		A2	1.4926	\$62.50
12055	Intmd wnd repair face/mm	Y		A2	2.782	\$116.49
12056	Intmd wnd repair face/mm	Y		A2	2.782	\$116.49
12057	Intmd wnd repair face/mm	Y		A2	4.1074	\$171.99
13100	Repair of wound or lesion	Y		A2	5.0314	\$210.68
13101	Repair of wound or lesion	Y		A2	5.0314	\$210.68

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11755	Biopsy, nail unit	Y		P3		\$55.68
11760	Repair of nail bed	Y		G2	1.2956	\$54.25
11762	Reconstruction of nail bed	Y		P3		\$104.25
11765	Excision of nail fold, toe	Y		P2	0.8408	\$35.21
11770	Removal of plantar lesion	Y		A2	19.8948	\$824.68
11771	Removal of plantar lesion	Y		A2	19.8948	\$824.68
11772	Removal of plantar lesion	Y		A2	19.8948	\$824.68
11900	Injection into skin lesions	Y		P3		\$24.71
11901	Added skin lesions injection	Y	CH	P3	\$27.27	\$27.27
11920	Correct skin color defects	Y		P3		\$77.26
11921	Correct skin color defects	Y		P3		\$65.79
11922	Correct skin color defects	Y		P3		\$28.12
11950	Therapy for contour defects	Y		P3		\$28.69
11951	Therapy for contour defects	Y		P3		\$36.93
11952	Therapy for contour defects	Y	CH	P3		\$47.15
11964	Therapy for contour defects	Y		P2	1.2956	\$54.25
11960	Insert tissue expander(s)	Y		A2	19.7192	\$825.70
11970	Replace tissue expander	Y		A2	36.3344	\$1,521.43
11971	Remove tissue expander(s)	Y		A2	18.6836	\$782.34
11976	Removal of contraceptive cap	Y		P3		\$51.70
11980	Implant hormone pellet(s)	N		P2	0.8403	\$26.81
11981	Insert drug implant device	N		P2	0.8403	\$26.81
11982	Remove drug implant device	N		P2	0.8403	\$26.81
11983	Remove/insert drug implant	N		P2	0.8403	\$26.81
12001	Repair superficial wound(s)	Y		P2	1.2956	\$54.25
12002	Repair superficial wound(s)	Y		P2	1.2956	\$54.25
12004	Repair superficial wound(s)	Y		P2	1.2956	\$54.25
12005	Repair superficial wound(s)	Y		A2	1.4926	\$62.50
12006	Repair superficial wound(s)	Y		A2	1.4926	\$62.50
12007	Repair superficial wound(s)	Y		A2	1.4926	\$62.50

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15100	Skin spl. grft. trnk/arm/leg	Y		A2	19,7192	\$825.70
15101	Skin spl. grft. /a/l, add-on	Y		A2	20,0845	\$841.00
15110	Epidrm autograft trnk/arm/leg	Y		A2	5,7323	\$240.03
15111	Epidrm autograft /a/l, add-on	Y		A2	5,0868	\$213.00
15115	Epidrm a-grft face/nck/hf/g	Y		A2	5,7323	\$240.03
15116	Epidrm a-grft /n/hf/g add	Y		A2	5,0868	\$213.00
15120	Skin spl. a-grft face/nck/hf/g	Y		A2	19,7192	\$825.70
15121	Skin spl. a-grft /n/hf/g add	Y		A2	20,0845	\$841.00
15130	Derm autograft, trnk/arm/leg	Y		A2	14,0668	\$589.02
15131	Derm autograft /a/l, add-on	Y		A2	13,4213	\$561.99
15135	Derm autograft face/nck/hf/g	Y		A2	14,0668	\$589.02
15136	Derm autograft, /n/hf/g add	Y		A2	13,4213	\$561.99
15150	Cult epidrm grft /arm/leg	Y		A2	5,7323	\$240.03
15151	Cult epidrm grft /a/l, add-on	Y		A2	5,0868	\$213.00
15152	Cult epidrm grft /n/hf/g	Y		A2	5,7323	\$240.03
15155	Cult epidrm grft, /n/hf/g	Y		A2	5,0868	\$213.00
15156	Cult epidrm grft /n/hf/g add	Y		A2	5,0868	\$213.00
15157	Cult epidrm grft /n/hf/g +*	Y		A2	5,0868	\$213.00
15170	Accl graft trunk/arms/legs	Y		G2	4,2464	\$177.81
15171	Accl graft /arm/leg add-on	Y		G2	3,0144	\$126.22
15175	Acclular graft, /n/hf/g	Y		G2	4,2464	\$177.81
15176	Accl graft, /n/hf/g add-on	Y		G2	4,2464	\$177.81
15200	Skin full graft, trunk	Y		A2	14,4325	\$604.33
15201	Skin full graft trunk add-on	Y		A2	13,3659	\$559.67
15220	Skin full graft soj/arm/leg	Y		A2	14,0668	\$589.02
15221	Skin full graft add-on	Y		A2	5,0314	\$210.68
15240	Skin full graft face/geni/hf	Y		A2	14,4325	\$604.33
15241	Skin full graft add-on	Y		A2	5,0314	\$210.68
15260	Skin full graft een & lips	Y		A2	14,0668	\$589.02
15261	Skin full graft add-on	Y		A2	13,3659	\$559.67

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13102	Repair wound/lesion add-on	Y		A2	3,706	\$155.18
13120	Repair of wound or lesion	Y		A2	2,782	\$116.49
13121	Repair of wound or lesion	Y		A2	2,782	\$116.49
13122	Repair wound/lesion add-on	Y		A2	1,4926	\$62.50
13131	Repair of wound or lesion	Y		A2	2,782	\$116.49
13132	Repair of wound or lesion	Y		A2	3,706	\$155.18
13133	Repair wound/lesion add-on	Y		A2	2,782	\$116.49
13150	Repair of wound or lesion	Y		A2	5,0314	\$210.68
13151	Repair of wound or lesion	Y		A2	5,0314	\$210.68
13152	Repair of wound or lesion	Y		A2	5,0314	\$210.68
13153	Repair wound/lesion add-on	Y		A2	2,782	\$116.49
13160	Late closure of wound	Y		A2	19,7192	\$825.70
14000	Skin tissue rearrangement	Y		A2	14,0668	\$589.02
14001	Skin tissue rearrangement	Y		A2	14,4325	\$604.33
14020	Skin tissue rearrangement	Y		A2	14,4325	\$604.33
14021	Skin tissue rearrangement	Y		A2	14,4325	\$604.33
14040	Skin tissue rearrangement	Y		A2	14,0668	\$589.02
14061	Skin tissue rearrangement	Y		A2	14,4325	\$604.33
14060	Skin tissue rearrangement	Y		A2	14,4325	\$604.33
14061	Skin tissue rearrangement	Y		A2	14,4325	\$604.33
14300	Skin tissue rearrangement	N	CH	D5		
14301	Skin tissue rearrangement	Y		G2	22,8955	\$958.70
14302	Skin tissue rearrange add-on	Y		G2	22,8955	\$958.70
14350	Skin tissue rearrangement	Y		A2	20,0845	\$841.00
15002	Wound prep, trk/arm/leg	Y		A2	5,0314	\$210.68
15003	Wound prep, addl 100 cm	Y		A2	5,0314	\$210.68
15004	Wound prep, /n/hf/g	Y		A2	5,0314	\$210.68
15005	Wound prep, /n/hf/g, addl cm	Y		A2	5,0314	\$210.68
15040	Harvest cultured skin graft	Y		A2	2,782	\$116.49
15050	Skin pinch graft	Y		A2	5,0314	\$210.68

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15732	Muscle-skin graft, head/neck	Y		A2	20.0845	\$841.00
15734	Muscle-skin graft, trunk	Y		A2	20.0845	\$841.00
15736	Muscle-skin graft, arm	Y		A2	20.0845	\$841.00
15738	Muscle-skin graft, leg	Y		A2	20.0845	\$841.00
15740	Island pedicle flap graft	Y		A2	14.0668	\$589.02
15750	Neurovascular pedicle graft	Y		A2	19.7192	\$825.70
15760	Composite skin graft	Y		A2	19.7192	\$825.70
15770	Derma-fat fascial graft	Y		A2	20.0845	\$841.00
15775	Hair transplant punch grafts	Y		A2	2.818	\$118.00
15776	Hair transplant punch grafts	Y		A2	2.818	\$118.00
15780	Abrasion treatment of skin	Y		P2		\$331.50
15781	Abrasion treatment of skin	Y		P2	4.1736	\$174.76
15782	Abrasion treatment of skin	Y		P2	4.1736	\$174.76
15783	Abrasion treatment of skin	Y		P2	2.677	\$112.09
15786	Abrasion, lesion, single	Y		P2	0.8408	\$35.21
15787	Abrasion, lesions, add-on	Y		P2		\$25.00
15788	Chemical peel, face, epiderm	Y		P2	0.8408	\$35.21
15789	Chemical peel, face, dermal	Y		P2	1.4745	\$61.74
15792	Chemical peel, nonfacial	Y		P2	1.4745	\$61.74
15793	Chemical peel, nonfacial	Y		P2	0.8408	\$35.21
15819	Plastic surgery, neck	Y		G2	3.0144	\$126.22
15820	Revision of lower eyelid	Y		A2	20.0845	\$841.00
15821	Revision of lower eyelid	Y		A2	20.0845	\$841.00
15822	Revision of upper eyelid	Y		A2	20.0845	\$841.00
15823	Revision of upper eyelid	Y		A2	21.2669	\$890.51
15824	Removal of forehead wrinkles	Y		A2	20.0845	\$841.00
15825	Removal of neck wrinkles	Y		A2	20.0845	\$841.00
15826	Removal of brow wrinkles	Y		A2	20.0845	\$841.00
15828	Removal of face wrinkles	Y		A2	20.0845	\$841.00
15829	Removal of skin wrinkles	Y		A2	21.2669	\$890.51

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
15300	Apply skin allograft, 1/arm/1/g	Y		A2	5.0314	\$210.68
15301	Apply skin allograft 1/a/1 add	Y		A2	5.0314	\$210.68
15320	Apply skin allograft 1/m/1/g	Y		A2	5.0314	\$210.68
15321	Apply skin allograft 1/m/1/g add	Y		A2	5.0314	\$210.68
15330	Apply acell allograft 1/arm/1/g	Y		A2	5.0314	\$210.68
15331	Apply acell graft 1/a/1 add-on	Y		A2	5.0314	\$210.68
15335	Apply acell graft, 1/m/1/g	Y		A2	5.0314	\$210.68
15336	Apply acell graft 1/m/1/g add	Y		A2	5.0314	\$210.68
15340	Apply cult skin substitute	Y		G2	3.0144	\$126.22
15341	Apply cult skin sub add-on	Y		G2	3.0144	\$126.22
15360	Apply cult derm sub, 1/a/1	Y		G2	3.0144	\$126.22
15361	Apply cult derm sub 1/a/1 add	Y		G2	3.0144	\$126.22
15365	Apply cult derm sub 1/m/1/g	Y		G2	3.0144	\$126.22
15366	Apply cult derm 1/m/1/g add	Y		G2	3.0144	\$126.22
15400	Apply skin xenograft, 1/a/1	Y		A2	5.0314	\$210.68
15401	Apply skin xenograft 1/a/1 add	Y		A2	5.0314	\$210.68
15420	Apply skin xgrft, 1/m/1/g	Y		A2	5.0314	\$210.68
15421	Apply skin xgrft 1/m/1/g add	Y		A2	5.0314	\$210.68
15431	Apply acellular xgrft add	Y		A2	5.0314	\$210.68
15432	Apply acellular xgrft add	Y		A2	5.0314	\$210.68
15570	Form skin pedicle flap	Y		A2	20.0845	\$841.00
15572	Form skin pedicle flap	Y		A2	20.0845	\$841.00
15574	Form skin pedicle flap	Y		A2	20.0845	\$841.00
15576	Form skin pedicle flap	Y		A2	20.0845	\$841.00
15600	Skin graft	Y		A2	20.0845	\$841.00
15610	Skin graft	Y		A2	20.0845	\$841.00
15620	Skin graft	Y		A2	20.7702	\$869.71
15650	Transfer skin pedicle flap	Y		A2	21.2669	\$890.51
15731	Forehead flap w/vasc pedicle	Y		A2	20.0845	\$841.00

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15940	Remove hip pressure sore	Y	A2	19.6948	\$824.68
15941	Remove hip pressure sore	Y	A2	19.6948	\$824.68
15942	Remove hip pressure sore	Y	A2	20.0845	\$841.00
15943	Remove hip pressure sore	Y	A2	20.7702	\$869.71
15944	Remove hip pressure sore	Y	A2	20.7702	\$869.71
15945	Remove thigh pressure sore	Y	A2	19.6948	\$824.68
15946	Remove thigh pressure sore	Y	A2	20.3802	\$853.38
15947	Remove thigh pressure sore	Y	A2	14.4325	\$604.33
15948	Remove thigh pressure sore	Y	A2	15.1179	\$633.03
15949	Remove thigh pressure sore	Y	A2	14.4325	\$604.33
15950	Remove thigh pressure sore	Y	A2	15.1179	\$633.03
16000	Initial treatment of burn(s)	Y	P3		\$22.72
16020	Dress/debrid p-thick burn, s	Y	P3		\$34.37
16025	Dress/debrid p-thick burn, m	Y	A2	1.4893	\$62.36
16030	Dress/debrid p-thick burn, l	Y	A2	1.676	\$70.18
16035	Incision of burn scab, initi	Y	G2	1.4745	\$61.74
17000	Destruct premalign lesion	Y	P2	0.8408	\$35.21
17003	Destruct premalign les, 2-14	Y	P3		\$3.12
17004	Destruct premalign lesions 15+	Y	P3		\$69.88
17106	Destruction of skin lesions	Y	P2	2.677	\$112.09
17107	Destruction of skin lesions	Y	P2	2.677	\$112.09
17108	Destruction of skin lesions	Y	P2	2.677	\$112.09
17110	Destruct b9 lesion, 1-14	Y	P2	0.8408	\$35.21
17111	Destruct lesion, 15 or more	Y	P2	1.4745	\$61.74
17250	Chemical cautery, tissue	Y	P3		\$38.06
17260	Destruction of skin lesions	Y	P3		\$39.77
17261	Destruction of skin lesions	Y	P2	1.4745	\$61.74
17262	Destruction of skin lesions	Y	P2	1.4745	\$61.74
17263	Destruction of skin lesions	Y	P2	1.4745	\$61.74
17264	Destruction of skin lesions	Y	P2	1.4745	\$61.74

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15830	Exc skin abd	Y	A2	19.6948	\$824.68
15832	Excise excessive skin tissue	Y	A2	19.6948	\$824.68
15833	Excise excessive skin tissue	Y	A2	19.6948	\$824.68
15834	Excise excessive skin tissue	Y	A2	19.6948	\$824.68
15835	Excise excessive skin tissue	Y	A2	18.6282	\$780.02
15836	Excise excessive skin tissue	Y	A2	15.468	\$647.69
15837	Excise excessive skin tissue	Y	G2	16.7399	\$700.95
15838	Excise excessive skin tissue	Y	G2	16.7399	\$700.95
15839	Excise excessive skin tissue	Y	A2	15.468	\$647.69
15840	Graft for face nerve palsy	Y	A2	20.7702	\$869.71
15841	Graft for face nerve palsy	Y	A2	20.7702	\$869.71
15842	Flap for face nerve palsy	Y	G2	22.9855	\$958.70
15845	Skin and muscle repair, face	Y	A2	20.7702	\$869.71
15847	Exc skin abd add-on	Y	A2	19.6948	\$824.68
15850	Removal of sutures	Y	G2	2.677	\$112.09
15851	Removal of sutures	Y	P3		\$41.19
15852	Dressing change not for burn	N	CH		\$26.81
15860	Test for blood flow in graft	N	G2	0.6403	\$26.81
15875	Suction assisted lipectomy	Y	A2	20.0845	\$841.00
15877	Suction assisted lipectomy	Y	A2	20.0845	\$841.00
15878	Suction assisted lipectomy	Y	A2	20.0845	\$841.00
15879	Suction assisted lipectomy	Y	A2	20.0845	\$841.00
15920	Removal of tail bone ulcer	Y	A2	4.5669	\$191.23
15931	Remove sacrum pressure sore	Y	A2	19.6948	\$824.68
15933	Remove sacrum pressure sore	Y	A2	19.6948	\$824.68
15934	Remove sacrum pressure sore	Y	A2	20.0845	\$841.00
15935	Remove sacrum pressure sore	Y	A2	20.0845	\$841.00
15936	Remove sacrum pressure sore	Y	A2	15.1179	\$633.03
15937	Remove sacrum pressure sore	Y	A2	20.7702	\$869.71

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19110	Nipple exploration	Y		A2	20.3074	\$950.33
19112	Excise breast duct fistula	Y		A2	20.6727	\$665.63
19120	Removal of breast lesion	Y		A2	20.6727	\$665.63
19125	Excision, breast lesion	Y		A2	20.6727	\$665.63
19126	Excision, add breast lesion	Y		A2	20.6727	\$665.63
19290	Place needle wire, breast	N		N1		
19291	Place needle wire, breast	N		N1		
19295	Place breast clip, percut	N		N1		
19296	Place po breast cath for rad	Y		A2	49.4283	\$2,069.71
19297	Place breast cath for rad	Y		A2	49.4283	\$2,069.71
19298	Place breast rad tube/caths	Y		A2	49.4283	\$2,069.71
19300	Removal of breast tissue	Y		A2	21.3584	\$894.34
19301	Partial mastectomy	Y		A2	20.6727	\$865.63
19302	P-mastectomy with removal	Y		A2	35.819	\$1,499.85
19303	Mast, simple, complete	Y		A2	28.1131	\$1,177.18
19304	Mast, subot	Y		A2	28.1131	\$1,177.18
19316	Suspension of breast	Y		A2	28.1131	\$1,177.18
19318	Reduction of large breast	Y		A2	33.7944	\$1,412.56
19324	Enlarge breast	Y		A2	33.7944	\$1,412.56
19325	Enlarge breast with implant	Y		A2	49.4283	\$2,069.71
19328	Removal of breast implant	Y		A2	26.4165	\$1,106.14
19330	Removal of implant material	Y		A2	26.4165	\$1,106.14
19340	Immediate breast prosthesis	Y		A2	32.6834	\$1,368.55
19342	Delayed breast prosthesis	Y		A2	44.693	\$1,871.43
19350	Breast reconstruction	Y		A2	21.3584	\$894.34
19355	Correct inverted nipple(s)	Y		A2	28.1131	\$1,177.18
19357	Breast reconstruction	Y		A2	45.8754	\$1,920.94
19366	Breast reconstruction	Y		A2	28.6098	\$1,197.98
19370	Surgery of breast capsule	Y		A2	28.1131	\$1,177.18
19371	Removal of breast capsule	Y		A2	28.1131	\$1,177.18

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17266	Destruction of skin lesions	Y	CH	P3		\$94.88
17270	Destruction of skin lesions	Y		P2	1.4745	\$61.74
17271	Destruction of skin lesions	Y		P2	1.4745	\$61.74
17272	Destruction of skin lesions	Y		P2	1.4745	\$61.74
17273	Destruction of skin lesions	Y	CH	P3		\$86.35
17274	Destruction of skin lesions	Y	CH	P3		\$87.72
17276	Destruction of skin lesions	Y	CH	P3		\$107.09
17280	Destruction of skin lesions	Y		P2	1.4745	\$61.74
17281	Destruction of skin lesions	Y		P3		\$73.86
17282	Destruction of skin lesions	Y		P3		\$84.37
17283	Destruction of skin lesions	Y	CH	P3		\$96.86
17284	Destruction of skin lesions	Y	CH	P3		\$108.86
17286	Destruction of skin lesions	Y		P2	2.677	\$112.09
17311	Mohs, 1 stage, 1/n/h/f/g	Y		P2	4.7201	\$197.64
17312	Mohs, 1 stage, 1/n/h/f/g	Y	CH	P3		\$192.88
17313	Mohs, 1 stage, 1/n/h/f/g	Y	CH	P3		\$197.64
17314	Mohs, add stage, 1/n/h/f/g	Y	CH	P3		\$178.96
17340	Cryotherapy of skin	Y		P3		\$32.67
17360	Skin peel therapy	Y		P3		\$12.78
17900	Hair removal by electrolysis	Y		P2	0.8408	\$35.21
19000	Drainage of breast lesion	Y		P3		\$54.82
19001	Drain breast lesion add-on	Y		P3		\$7.39
19020	Incision of breast lesion	Y		P3		\$689.73
19030	Injection for breast x-ray	N		N1	16.472	
19100	Bx breast percut w/o image	Y		A2	4.6708	\$195.58
19101	Biopsy of breast, open	Y		A2	20.9074	\$850.33
19102	Bx breast percut w/image	Y		A2	6.978	\$292.19
19103	Bx breast percut wideview	Y		A2	13.3824	\$560.36
19105	Cryosurg ablate 1a, each	Y	CH	P2	32.686	\$1,368.66

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20650	Insert and remove bone pin	Y		A2	18.7092	\$783.41
20662	Application of pelvis brace	Y		R2	21.0617	\$881.92
20663	Application of thigh brace	Y		R2	21.0617	\$881.92
20665	Removal of fixation device	N		G2	0.6403	\$26.81
20670	Removal of support implant	Y		A2	14.457	\$605.36
20680	Removal of support implant	Y		A2	19.6948	\$824.68
20690	Apply bone fixation device	Y		A2	25.3445	\$1,061.25
20692	Apply bone fixation device	Y		A2	25.7101	\$1,076.56
20693	Adjust bone fixation device	Y		A2	18.7092	\$783.41
20694	Remove bone fixation device	Y		A2	17.6983	\$741.08
20696	Comp multipiane ext fixation	Y		G2	30.396	\$1,272.77
20697	Comp ext fixate strut change	Y		G2	17.5996	\$736.95
20822	Replantation digit, complete	Y		G2	27.0149	\$1,131.19
20900	Removal of bone for graft	Y		A2	25.7101	\$1,076.56
20902	Removal of bone for graft	Y		A2	26.3955	\$1,105.26
20910	Remove cartilage for graft	Y		A2	20.0845	\$841.00
20912	Remove cartilage for graft	Y		A2	20.0845	\$841.00
20920	Removal of fascia for graft	Y		A2	15.1779	\$633.03
20922	Removal of fascia for graft	Y		A2	14.4325	\$604.33
20924	Removal of tendon for graft	Y		A2	26.3955	\$1,105.26
20926	Removal of tissue for graft	Y		A2	6.7934	\$284.04
20950	Fluid pressure, muscle	Y		G2	1.3927	\$58.32
20972	Bone/skin graft, metatarsal	Y		G2	50.2514	\$2,104.18
20973	Bone/skin graft, great toe	Y		G2	50.2514	\$2,104.18
20975	Electrical bone stimulation	N		N1		
20979	Us bone stimulation	N		P3		
20982	Ablate, bone tumor(s) periq	Y		G2	44.5617	\$1,865.93
20985	Cptr-asst dir ms px	N		N1		
21010	Incision of jaw joint	Y		A2	20.4597	\$856.71
21011	Exc face les sc < 2 cm	Y		P3		\$147.14

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19380	Revise breast reconstruction	Y		A2	34.2311	\$1,453.36
19396	Design custom breast implant	Y		G2	32.686	\$1,368.66
20000	Incision of abscess	Y		P2	1.3927	\$58.32
20005	Incision of deep abscess	Y		A2	18.3436	\$768.10
20103	Explore wound, extremity	Y		G2	12.0752	\$505.62
20150	Excise epiphyseal bar	Y		A2	44.5617	\$1,865.93
20200	Muscle biopsy	Y		A2	15.1023	\$632.38
20205	Deep muscle biopsy	Y		A2	15.488	\$647.69
20206	Needle biopsy, muscle	Y		A2	6.978	\$292.19
20220	Bone biopsy, trocar/needle	Y		A2	7.3224	\$306.61
20225	Bone biopsy, trocar/needle	Y		A2	14.9452	\$625.80
20240	Bone biopsy, excisional	Y		A2	19.2392	\$809.37
20245	Bone biopsy, excisional	Y		A2	19.6948	\$824.68
20250	Open bone biopsy	Y		A2	18.7092	\$783.41
20251	Open bone biopsy	Y		A2	18.7092	\$783.41
20500	Injection of sinus tract	Y		P3		\$44.03
20501	Inject sinus tract for x-ray	N		N1		
20520	Removal of foreign body	Y		P3		\$79.82
20525	Removal of foreign body	Y		A2	19.6948	\$824.68
20526	Ther injection, carp tunnel	Y		P3		\$25.28
20550	Inj tendon sheath/ligament	Y		P3		\$19.32
20551	Inj tendon origin/insertion	Y		P3		\$19.60
20552	Inj trigger point, 1/2 muscul	Y		P3		\$18.46
20553	Inject trigger points, => 3	Y		P3		\$21.02
20555	Place ndl muscles for rt	Y	CH	P2	30.396	\$1,272.77
20600	Drain/inject, joint/bursa	Y		P3		\$19.60
20605	Drain/inject, joint/bursa	Y		P3		\$21.87
20610	Drain/inject, joint/bursa	Y		P3		\$31.25
20612	Aspirate/inj ganglion cyst	Y		P3		\$21.02
20615	Treatment of bone cyst	Y		P3		\$85.50

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21086	Prepare facel/oral prosthesis	Y		P3		\$490.29
21087	Prepare facel/oral prosthesis	Y		P3		\$489.44
21088	Prepare facel/oral prosthesis	Y		R2	41.1215	\$1,721.88
21100	Maxillofacial fixation	Y		A2	33.3886	\$1,396.08
21110	Interdental fixation	Y		P2	7.2897	\$905.24
21116	Injection, jaw joint x-ray	N		N1		
21120	Reconstruction of chin	Y		A2	23.5956	\$988.02
21121	Reconstruction of chin	Y		A2	23.5956	\$988.02
21122	Reconstruction of chin	Y		A2	23.5956	\$988.02
21123	Reconstruction of chin	Y		A2	23.5956	\$988.02
21125	Augmentation, lower jaw bone	Y		A2	36.4895	\$1,611.67
21127	Augmentation, lower jaw bone	Y		A2	36.4895	\$1,611.67
21137	Reduction of forehead	Y		G2	23.8828	\$988.02
21138	Reduction of forehead	Y		G2	41.1215	\$1,721.88
21139	Reduction of forehead	Y		G2	41.1215	\$1,721.88
21150	Reconstruct midface, isfort	Y		G2	41.1215	\$1,721.88
21181	Contour cranial bone lesion	Y		A2	23.5956	\$988.02
21198	Reconst lwr jaw segment	Y		G2	41.1215	\$1,721.88
21199	Reconst lwr jaw w/advance	Y		G2	41.1215	\$1,721.88
21206	Reconstruct upper jaw bone	Y		A2	34.9366	\$1,462.90
21208	Augmentation of facial bones	Y		A2	36.5245	\$1,529.39
21209	Reduction of facial bones	Y		A2	34.9366	\$1,462.90
21210	Face bone graft	Y		A2	36.5245	\$1,529.39
21215	Lower jaw bone graft	Y		A2	36.5245	\$1,529.39
21230	Rib cartilage graft	Y		A2	36.5245	\$1,529.39
21235	Ear cartilage graft	Y		A2	23.5956	\$988.02
21240	Reconstruction of jaw joint	Y		A2	34.4396	\$1,442.09
21242	Reconstruction of jaw joint	Y		A2	34.9366	\$1,462.90
21243	Reconstruction of jaw joint	Y		A2	34.9366	\$1,462.90
21244	Reconstruction of lower jaw	Y		A2	36.5245	\$1,529.39

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
21012	Exc face les sc = 2 cm	Y	N1	R2	7.8476	\$328.60
21013	Exc face tum deep > 2 cm	Y	N1	P3		\$203.96
21014	Exc face tum deep = 2 cm	Y	N1	R2	7.8476	\$328.60
21015*	Resect face tum < 2 cm	Y	N1	R2	16.7399	\$700.95
21016	Resect face tum = 2 cm	Y	N1	G2	22.3753	\$936.92
21025	Excision of bone, lower jaw	Y		A2	33.3866	\$1,398.08
21026	Excision of facial bone(s)	Y		A2	33.3866	\$1,398.08
21029	Contour of face bone lesion	Y		A2	33.3866	\$1,398.08
21030	Excise max/zygoma b9 tumor	Y		P3		\$208.78
21031	Remove exostosis, mandible	Y		P3		\$171.86
21032	Remove exostosis, maxilla	Y		P3		\$174.98
21034	Excise max/zygoma mlg tumor	Y		A2	33.7542	\$1,413.39
21040	Excise mandible lesion	Y		A2	20.4597	\$856.71
21044	Removal of jaw bone lesion	Y		A2	33.3886	\$1,398.08
21046	Remove mandible cyst complex	Y		A2	33.3886	\$1,398.08
21047	Excise lwr jaw cyst w/repair	Y		A2	33.3886	\$1,398.08
21048	Remove maxilla cyst complex	Y		R2	41.1215	\$1,721.88
21050	Removal of jaw joint	Y		A2	33.7542	\$1,413.39
21060	Remove jaw joint cartilage	Y		A2	33.3886	\$1,398.08
21070	Remove coronoid process	Y		A2	33.7542	\$1,413.39
21073	Mipi of tmj w/invest	Y		P3		\$160.21
21076	Prepare facel/oral prosthesis	Y		P3		\$287.19
21077	Prepare facel/oral prosthesis	Y		P3		\$691.12
21079	Prepare facel/oral prosthesis	Y		P3		\$495.40
21080	Prepare facel/oral prosthesis	Y		P3		\$566.42
21081	Prepare facel/oral prosthesis	Y		P3		\$522.10
21082	Prepare facel/oral prosthesis	Y		P3		\$500.80
21083	Prepare facel/oral prosthesis	Y		P3		\$491.99
21084	Prepare facel/oral prosthesis	Y		P3		\$563.01
21085	Prepare facel/oral prosthesis	Y		P3		\$224.98

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
21406	Treat eye socket fracture	Y		G2	41.1215	\$1,721.88
21407	Treat eye socket fracture	Y		G2	41.1215	\$1,721.88
21421	Treat mouth roof fracture	Y		A2	21.5108	\$900.72
21440	Treat dental ridge fracture	Y		P3		\$281.22
21445	Treat dental ridge fracture	Y		A2	21.5108	\$900.72
21450	Treat lower jaw fracture	Y		A2	3.3186	\$138.96
21451	Treat lower jaw fracture	Y		A2	8.1184	\$339.94
21452	Treat lower jaw fracture	Y		A2	14.8802	\$623.08
21453	Treat lower jaw fracture	Y		A2	33.7542	\$1,413.39
21454	Treat lower jaw fracture	Y		A2	22.0077	\$921.53
21461	Treat lower jaw fracture	Y		A2	34.4396	\$1,442.09
21462	Treat lower jaw fracture	Y		A2	34.9366	\$1,462.90
21465	Treat lower jaw fracture	Y		A2	34.4396	\$1,442.09
21480	Reset dislocated jaw	Y		A2	1.6877	\$70.67
21485	Reset dislocated jaw	Y		A2	14.8802	\$623.08
21490	Repair dislocated jaw	Y		A2	33.7542	\$1,413.39
21496	Treat hyoid bone fracture	Y		G2	16.4437	\$688.55
21497	Interdental wiring	Y		A2	14.8802	\$623.08
21501	Drain neck/chest lesion	Y		A2	16.472	\$689.73
21502	Drain chest lesion	Y		A2	18.3436	\$768.10
21550	Biopsy of neck/chest	Y		G2	16.7989	\$700.95
21552	Exc neck les sc = 3 cm	Y	NI	G2	22.3753	\$936.92
21554	Exc neck tum deep = 5 cm	Y	NI	G2	22.3753	\$936.92
21555*	Exc neck les sc < 3 cm	Y	NI	P3		\$169.58
21556	Exc neck tum deep < 5 cm	Y	NI	G2	22.3753	\$936.92
21557	Resect neck tum < 5 cm	Y	NI	G2	16.7989	\$700.95
21558	Resect neck tum = 5 cm	Y	NI	G2	22.3753	\$936.92
21600	Partial removal of rib	Y		A2	25.3445	\$1,061.25
21610	Partial removal of rib	Y		A2	25.3445	\$1,061.25
21685	Hyoid myotomy & suspension	Y		G2	7.2897	\$305.24

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
21245	Reconstruction of jaw	Y		A2	36.5245	\$1,529.39
21246	Reconstruction of jaw	Y		A2	36.5245	\$1,529.39
21248	Reconstruction of jaw	Y		A2	36.5245	\$1,529.39
21249	Reconstruction of jaw	Y		A2	36.5245	\$1,529.39
21260	Revise eye sockets	Y		G2	41.1215	\$1,721.88
21267	Revise eye sockets	Y		A2	36.5245	\$1,529.39
21270	Augmentation, cheek bone	Y		A2	34.9366	\$1,462.90
21275	Revision, orbitofacial bones	Y		A2	36.5245	\$1,529.39
21280	Revision of eyelid	Y		A2	34.9366	\$1,462.90
21282	Revision of eyelid	Y		A2	16.4282	\$667.90
21295	Revision of jaw muscle/bone	Y		A2	7.3694	\$308.58
21296	Revision of jaw muscle/bone	Y		A2	19.8142	\$829.68
21310	Treatment of nose fracture	Y		A2	1.6877	\$70.67
21315	Treatment of nose fracture	Y		A2	13.1937	\$552.46
21320	Treatment of nose fracture	Y		A2	14.8802	\$623.08
21325	Treatment of nose fracture	Y		A2	21.5108	\$900.72
21330	Treatment of nose fracture	Y		A2	22.0077	\$921.53
21335	Treatment of nose fracture	Y		A2	23.5956	\$988.02
21336	Treat nasal septal fracture	Y		A2	22.1427	\$927.18
21337	Treat nasal septal fracture	Y		A2	14.8802	\$623.08
21338	Treat nasoinferior fracture	Y		A2	21.5108	\$900.72
21339	Treat nasoinferior fracture	Y		A2	22.0077	\$921.53
21340	Treatment of nose fracture	Y		A2	34.4396	\$1,442.09
21345	Treat nose/jaw fracture	Y		A2	23.5956	\$988.02
21355	Treat cheek bone fracture	Y		A2	33.7542	\$1,413.39
21356	Treat cheek bone fracture	Y		A2	20.8254	\$872.02
21360	Treat cheek bone fracture	Y		G2	23.8828	\$1,000.04
21390	Treat eye socket fracture	Y		G2	41.1215	\$1,721.88
21400	Treat eye socket fracture	Y		A2	8.0147	\$335.60
21401	Treat eye socket fracture	Y		A2	15.2459	\$638.39

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22904	Resect abd tum < 5 cm	Y	NI	G2	16.7399	\$700.95
22905	Resect abd tum > 5 cm	Y	NI	G2	22.3753	\$936.92
23000	Removal of calcium deposits	Y	A2	A2	15.1023	\$632.38
23002	Release shoulder joint	Y	A2	A2	35.969	\$1,506.13
23030	Drain shoulder lesion	Y	A2	A2	15.8264	\$662.70
23031	Drain shoulder bursa	Y	A2	A2	16.8376	\$705.04
23035	Drain shoulder bone lesion	Y	A2	A2	18.7092	\$783.41
23044	Exploratory shoulder surgery	Y	A2	A2	25.7101	\$1,076.56
23065	Biopsy shoulder	Y	P3	P3	26.3955	\$1,105.26
23066	Biopsy shoulder fissures	Y	A2	A2	19.3292	\$696.37
23071	Exc shoulder les sc > 3 cm	Y	NI	G2	22.3753	\$936.92
23073	Exc shoulder tum deep > 5 cm	Y	NI	G2	22.3753	\$936.92
23075*	Exc shoulder les sc < 3 cm	Y	NI	P3	16.7399	\$700.95
23076	Exc shoulder tum deep < 5 cm	Y	NI	G2	16.7399	\$700.95
23077	Resect shoulder tum < 5 cm	Y	NI	G2	22.3753	\$936.92
23078	Resect shoulder tum > 5 cm	Y	NI	G2	22.3753	\$936.92
23100	Biopsy of shoulder joint	Y	A2	A2	18.3436	\$768.10
23101	Shoulder joint surgery	Y	A2	A2	28.4804	\$1,192.56
23106	Remove shoulder joint lining	Y	A2	A2	26.3955	\$1,105.26
23107	Incision of collarbone joint	Y	A2	A2	26.3955	\$1,105.26
23120	Partial removal, collar bone	Y	A2	A2	26.3955	\$1,105.26
23125	Explore treat shoulder joint	Y	A2	A2	26.8925	\$1,126.07
23130	Remove of collar bone	Y	A2	A2	26.8925	\$1,126.07
23140	Remove shoulder bone, part	Y	A2	A2	37.5168	\$1,570.94
23145	Removal of bone lesion	Y	A2	A2	19.3946	\$812.11
23146	Removal of bone lesion	Y	A2	A2	26.8925	\$1,126.07
23150	Removal of humerus lesion	Y	A2	A2	26.3955	\$1,105.26
23155	Removal of humerus lesion	Y	A2	A2	26.8925	\$1,126.07

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21700	Revision of neck muscle	Y	A2	A2	18.3436	\$768.10
21720	Revision of neck muscle	Y	A2	A2	18.7092	\$783.41
21725	Revision of neck muscle	Y	A2	A2	1.5497	\$64.89
21800	Treatment of rib fracture	Y	A2	A2	1.7811	\$74.58
21805	Treatment of rib fracture	Y	A2	A2	21.0916	\$883.17
21820	Treat sternum fracture	Y	A2	A2	1.7811	\$74.58
21820	Biopsy soft tissue of back	Y	P3	P3	119.59	\$4,809.37
21825	Biopsy soft tissue of back	Y	A2	A2	19.3292	\$768.10
21930*	Exc back les sc < 3 cm	Y	NI	P3	18.0797	\$719.97
21931	Exc back les sc = 3 cm	Y	NI	G2	22.3753	\$936.92
21932	Exc back tum deep < 5 cm	Y	NI	G2	16.7399	\$700.95
21933	Exc back tum deep = 5 cm	Y	NI	G2	22.3753	\$936.92
21935	Resect back tum < 5 cm	Y	NI	G2	16.7399	\$700.95
21936	Resect back tum = 5 cm	Y	NI	G2	22.3753	\$936.92
22102	Remove part, lumbar vertebra	Y	G2	G2	47.0941	\$1,971.97
22103	Remove extra spine segment	Y	G2	G2	47.0941	\$1,971.97
22905	Treat spine process fracture	Y	A2	A2	1.7811	\$74.58
22310	Treat spine fracture	Y	A2	A2	4.0322	\$168.84
22315	Treat spine fracture	Y	A2	A2	13.7917	\$577.50
22505	Manipulation of spine	Y	A2	A2	13.5199	\$566.12
22520	Percut vertebroplasty thor	Y	A2	A2	30.4452	\$1,274.83
22521	Percut vertebroplasty lumb	Y	A2	A2	30.4452	\$1,274.83
22522	Percut vertebroplasty addl	Y	A2	A2	30.4452	\$1,274.83
22523	Percut kyphoplasty, thor	Y	G2	G2	84.8135	\$3,551.40
22524	Percut kyphoplasty, lumbar	Y	G2	G2	84.8135	\$3,551.40
22525	Percut kyphoplasty, add-on	Y	G2	G2	84.8135	\$3,551.40
22900	Exc back tum deep < 5 cm	Y	NI	G2	22.3753	\$936.92
22901	Exc back tum deep = 5 cm	Y	NI	G2	22.3753	\$936.92
22902	Exc abd les sc < 3 cm	Y	NI	G2	16.7399	\$700.95
22903	Exc abd les sc > 3 cm	Y	NI	G2	22.3753	\$936.92

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23485	Revision of collar bone	Y		A2	69.2933	\$2,901.52
23490	Reinforce clavicle	Y		A2	36.3344	\$1,521.43
23491	Reinforce shoulder bones	Y		A2	66.5231	\$2,765.52
23500	Treat clavicle fracture	Y		A2	1.7811	\$74.58
23505	Treat clavicle fracture	Y		A2	13.7917	\$577.50
23515	Treat clavicle fracture	Y		A2	48.8932	\$2,089.18
23520	Treat clavicle dislocation	Y		A2	4.0322	\$168.84
23525	Treat clavicle dislocation	Y		A2	4.0322	\$168.84
23530	Treat clavicle dislocation	Y		A2	35.5372	\$1,486.05
23532	Treat clavicle dislocation	Y		A2	22.1427	\$927.18
23540	Treat clavicle dislocation	Y		A2	1.7811	\$74.58
23545	Treat clavicle dislocation	Y		A2	4.0322	\$168.84
23550	Treat clavicle dislocation	Y		A2	35.5372	\$1,486.05
23552	Treat clavicle dislocation	Y		A2	36.2226	\$1,516.75
23570	Treat shoulder blade fx	Y		A2	1.7811	\$74.58
23575	Treat shoulder blade fx	Y		A2	4.0322	\$168.84
23585	Treat scapula fracture	Y		A2	48.8932	\$2,089.18
23600	Treat humerus fracture	Y		P2	1.5858	\$66.40
23605	Treat humerus fracture	Y		A2	13.7917	\$577.50
23615	Treat humerus fracture	Y		A2	50.5787	\$2,117.88
23616	Treat humerus fracture	Y		A2	50.5787	\$2,117.88
23620	Treat humerus fracture	Y		P2	1.5858	\$66.40
23625	Treat humerus fracture	Y		A2	13.7917	\$577.50
23630	Treat humerus fracture	Y		A2	51.0756	\$2,136.69
23650	Treat shoulder dislocation	Y		A2	1.7811	\$74.58
23655	Treat shoulder dislocation	Y		A2	12.8746	\$539.10
23665	Treat shoulder dislocation	Y		A2	35.5372	\$1,486.05
23670	Treat dislocation/fracture	Y		A2	4.0322	\$168.84
23675	Treat dislocation/fracture	Y		A2	48.8932	\$2,089.18
23675	Treat dislocation/fracture	Y		A2	1.7811	\$74.58

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23156	Removal of humerus lesion	Y		A2	26.8925	\$1,126.07
23170	Remove collar bone lesion	Y		A2	25.3445	\$1,061.25
23172	Remove shoulder blade lesion	Y		A2	25.3445	\$1,061.25
23174	Remove humerus lesion	Y		A2	25.3445	\$1,061.25
23180	Remove collar bone lesion	Y		A2	26.3955	\$1,105.26
23182	Remove shoulder blade lesion	Y		A2	26.3955	\$1,105.26
23184	Remove humerus lesion	Y		A2	26.3955	\$1,105.26
23190	Partial removal of scapula	Y		A2	26.3955	\$1,105.26
23195	Removal of head of humerus	Y		A2	26.8925	\$1,126.07
23330	Remove shoulder, foreign body	Y		A2	7.7878	\$326.10
23331	Remove shoulder, foreign body	Y		A2	18.6836	\$782.34
23350	Injection for shoulder X-ray	N		N1		
23395	Muscle transfer, shoulder/arm	Y		A2	37.5168	\$1,570.94
23397	Muscle transfers	Y		A2	69.2933	\$2,901.52
23400	Fixation of shoulder blade	Y		A2	28.4804	\$1,192.56
23405	Incision of tendon & muscle	Y		A2	25.3445	\$1,061.25
23406	Incise tendon(s) & muscle(s)	Y		A2	25.3445	\$1,061.25
23410	Repair rotator cuff, acute	Y		A2	37.5168	\$1,570.94
23412	Repair rotator cuff, chronic	Y		A2	39.1047	\$1,637.43
23415	Release of shoulder ligament	Y		A2	37.5168	\$1,570.94
23420	Repair of shoulder	Y		A2	39.1047	\$1,637.43
23430	Repair biceps tendon	Y		A2	37.02	\$1,550.14
23440	Remove/transplant tendon	Y		A2	37.02	\$1,550.14
23450	Repair shoulder capsule	Y		A2	67.7054	\$2,835.03
23455	Repair shoulder capsule	Y		A2	69.2933	\$2,901.52
23460	Repair shoulder capsule	Y		A2	67.7054	\$2,835.03
23462	Repair shoulder capsule	Y		A2	67.7054	\$2,835.03
23466	Repair shoulder capsule	Y		A2	39.1047	\$1,637.43
23480	Revision of collar bone	Y		A2	37.02	\$1,550.14

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Note 2: Payment indicators for "office-based" procedures (P2, P3) are based on a comparison of the final rates according to the ASC standard ratesetting methodology and the MPFS. Under current law, MPFS payment rates will have a negative update for CY 2010. For a discussion of these rates, we refer readers to the CY 2010 MPFS final rule.

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ADDENDUM AA—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
24136	Remove radius bone lesion	Y		A2	25.3445	\$1,061.25
24138	Remove elbow bone lesion	Y		A2	25.3445	\$1,061.25
24140	Partial removal of arm bone	Y		A2	25.7101	\$1,076.56
24145	Partial removal of radius	Y		A2	25.7101	\$1,076.56
24147	Partial removal of elbow	Y		A2	25.3445	\$1,061.25
24149	Radical resection of elbow	Y		G2	30.396	\$1,272.77
24152	Resect radius tumor	Y		G2	44.5617	\$1,665.93
24153	Extensive radius surgery	N	CH	D5		
24155	Removal of elbow joint	Y		A2	36.3344	\$1,521.43
24160	Remove elbow joint implant	Y		A2	25.3445	\$1,061.25
24164	Remove radius head implant	Y		A2	25.7101	\$1,076.56
24200	Removal of arm foreign body	Y		P3		\$65.22
24201	Removal of arm foreign body	Y		A2	15.1023	\$632.38
24220	Injection for elbow x-ray	N		N1		
24300	Manipulate elbow w/aneath	Y		G2	14.63	\$612.60
24301	Muscle/tendon transfer	Y		A2	26.3955	\$1,105.26
24305	Arm tendon lengthening	Y		A2	26.3955	\$1,105.26
24310	Revision of arm tendon	Y		A2	16.7092	\$783.41
24320	Repair of arm tendon	Y		A2	36.3344	\$1,521.43
24330	Revision of arm muscles	Y		A2	66.5231	\$2,785.52
24331	Revision of arm muscles	Y		A2	36.3344	\$1,521.43
24332	Tenolysis, triceps	Y		G2	21.0617	\$881.92
24340	Repair of biceps tendon	Y		A2	36.3344	\$1,521.43
24341	Repair arm tendon/muscle	Y		A2	36.3344	\$1,521.43
24342	Repair of ruptured tendon	Y		A2	36.3344	\$1,521.43
24343	Repr elbow lat ligmnt w/issu	Y		G2	30.396	\$1,272.77
24344	Reconstruct elbow lat ligmnt	Y		G2	84.8135	\$3,551.40
24345	Repr elbow med ligmnt w/issu	Y		A2	25.3445	\$1,061.25
24346	Reconstruct elbow med ligmnt	Y		G2	44.5617	\$1,665.93
24357	Repair elbow, perc	Y		G2	30.396	\$1,272.77

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
23680	Treat dislocation/fracture	Y		A2	35.5372	\$1,488.05
23700	Fixation of shoulder	Y		A2	12.8746	\$539.10
23800	Fusion of shoulder joint	Y		A2	67.2085	\$2,814.22
23900	Fusion of shoulder joint	Y		A2	39.1047	\$1,637.43
23921	Amputation follow-up surgery	Y		A2	13.3659	\$559.67
23930	Drainage of arm lesion	Y		A2	15.8264	\$662.70
23931	Drainage of arm bursa	Y		A2	16.472	\$689.73
23935	Drain armbow bone lesion	Y		A2	18.3436	\$788.10
24000	Exploratory elbow surgery	Y		A2	26.3955	\$1,105.26
24006	Release elbow joint	Y		A2	26.3955	\$1,105.26
24065	Biopsy arm/elbow soft tissue	Y		P3		\$115.04
24066	Biopsy arm/elbow soft tissue	Y		A2	15.1023	\$632.38
24071	Exc armbow iels sc = 3 cm	Y	NI	G2	22.3753	\$936.92
24073	Exc armbow tum deep > 5 cm	Y	NI	G2	22.3753	\$936.92
24075*	Exc armbow les sc < 3 cm	Y	NI	P3		\$210.21
24076	Exc armbow tum deep < 5 cm	Y	NI	G2	16.7399	\$700.95
24077	Resect armbow tum < 5 cm	Y	NI	G2	16.7399	\$700.95
24079	Resect armbow tum > 5 cm	Y	NI	G2	22.3753	\$936.92
24100	Biopsy elbow joint lining	Y		A2	17.6983	\$741.08
24101	Explore/treat elbow joint	Y		A2	26.3955	\$1,105.26
24102	Remove elbow joint lining	Y		A2	26.3955	\$1,105.26
24105	Removal of elbow bursa	Y		A2	18.7082	\$783.41
24110	Remove humerus lesion	Y		A2	18.3436	\$788.10
24115	Remove/graft bone lesion	Y		A2	25.7101	\$1,076.56
24116	Remove/graft bone lesion	Y		A2	25.7101	\$1,076.56
24120	Remove elbow lesion	Y		A2	18.7082	\$783.41
24125	Remove/graft bone lesion	Y		A2	25.7101	\$1,076.56
24130	Removal of head of radius	Y		A2	25.7101	\$1,076.56
24134	Removal of arm bone lesion	Y		A2	25.3445	\$1,061.25

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Note 2: Payment indicators for "office-based" procedures (P2, P3) are based on a comparison of the final rates according to the ASC standard ratesetting methodology and the MPFS. Under current law, MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
24577	Treat humerus fracture	Y		A2	4.0322	\$168.84
24579	Treat humerus fracture	Y		A2	49.8932	\$2,089.18
24582	Treat humerus fracture	Y		A2	21.0916	\$883.17
24586	Treat elbow fracture	Y		A2	50.5787	\$2,117.88
24587	Treat elbow fracture	Y		A2	51.0756	\$2,138.69
24600	Treat elbow dislocation	Y		A2	1.7811	\$74.58
24605	Treat elbow dislocation	Y		A2	13.5199	\$666.12
24615	Treat elbow dislocation	Y		A2	49.8932	\$2,089.18
24620	Treat elbow fracture	Y		A2	13.7917	\$677.50
24635	Treat elbow fracture	Y		A2	49.8932	\$2,089.18
24640	Treat elbow dislocation	Y		P3		\$47.72
24650	Treat radius fracture	Y		P2	1.5858	\$66.40
24655	Treat radius fracture	Y		A2	4.0322	\$168.84
24665	Treat radius fracture	Y		A2	36.2226	\$1,516.75
24666	Treat radius fracture	Y		A2	50.5787	\$2,117.88
24670	Treat ulnar fracture	Y		A2	1.7811	\$74.58
24675	Treat ulnar fracture	Y		A2	1.7811	\$74.58
24685	Treat ulnar fracture	Y		A2	35.5372	\$1,488.05
24800	Fusion of elbow joint	Y		A2	37.02	\$1,550.14
24802	Fusion/graft of elbow joint	Y		A2	37.5168	\$1,570.94
24925	Amputation follow-up surgery	Y		A2	18.7092	\$783.41
25000	Incision of tendon sheath	Y		A2	18.7092	\$783.41
25001	Incise flexor carpi radialis	Y		G2	21.0617	\$881.92
25020	Decompress forearm 1 space	Y		A2	25.7101	\$1,076.56
25023	Decompress forearm 1 space	Y		A2	25.7101	\$1,076.56
25024	Decompress forearm 2 spaces	Y		A2	25.7101	\$1,076.56
25025	Decompress forearm 2 spaces	Y		A2	25.7101	\$1,076.56
25028	Drainage of forearm lesion	Y		A2	17.6983	\$741.08
25031	Drainage of forearm bursa	Y		A2	18.3436	\$765.10
25035	Treat forearm bone lesion	Y		A2	18.3436	\$765.10

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Note 2: Payment indicators for "office-based" procedures (P2, P3) are based on a comparison of the final rates according to the ASC standard ratesetting methodology and the MPPS. Under current law, MPPS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPPS final rule.

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
24358	Repair elbow w/deb. open	Y		G2	30.3396	\$1,272.77
24359	Repair elbow debr/atch open	Y		G2	30.3396	\$1,272.77
24360	Reconstruct elbow joint	Y		A2	32.7158	\$1,369.91
24361	Reconstruct elbow joint	Y		H8	150.2982	\$6,293.48
24362	Reconstruct elbow joint	Y		A2	45.772	\$1,916.61
24363	Replace elbow joint	Y		H8	151.8871	\$6,359.97
24365	Reconstruct head of radius	Y		A2	32.7158	\$1,369.91
24366	Reconstruct head of radius	Y		H8	150.2982	\$6,293.48
24400	Revision of humerus	Y		A2	37.02	\$1,550.14
24410	Revision of humerus	Y		A2	37.02	\$1,550.14
24420	Revision of humerus	Y		A2	36.3344	\$1,521.43
24430	Repair of humerus	Y		A2	66.8231	\$2,785.52
24435	Repair humerus with graft	Y		A2	67.2085	\$2,814.22
24470	Revision of elbow joint	Y		A2	36.3344	\$1,521.43
24495	Decompression of forearm	Y		A2	25.3445	\$1,061.25
24498	Reinforce humerus	Y		A2	66.8231	\$2,785.52
24500	Treat humerus fracture	Y		A2	1.7811	\$74.58
24505	Treat humerus fracture	Y		A2	1.7811	\$74.58
24515	Treat humerus fracture	Y		A2	50.5787	\$2,117.88
24516	Treat humerus fracture	Y		A2	50.5787	\$2,117.88
24530	Treat humerus fracture	Y		A2	1.7811	\$74.58
24535	Treat humerus fracture	Y		A2	4.0322	\$168.84
24538	Treat humerus fracture	Y		A2	21.0916	\$883.17
24545	Treat humerus fracture	Y		A2	50.5787	\$2,117.88
24546	Treat humerus fracture	Y		A2	51.0756	\$2,138.69
24560	Treat humerus fracture	Y		A2	1.7811	\$74.58
24565	Treat humerus fracture	Y		A2	1.7811	\$74.58
24566	Treat humerus fracture	Y		A2	21.0916	\$883.17
24575	Treat humerus fracture	Y		A2	49.8932	\$2,089.18
24576	Treat humerus fracture	Y		A2	1.7811	\$74.58

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25151	Partial removal of radius	Y		A2	25.3445	\$1,061.25
25210	Removal of wrist bone	Y		A2	25.7101	\$1,076.56
25215	Removal of wrist bones	Y		A2	26.3955	\$1,105.26
25230	Partial removal of radius	Y		A2	26.3955	\$1,105.26
25240	Partial removal of ulna	Y		A2	26.3955	\$1,105.26
25246	Injection for wrist x-ray	N		N1		
25248	Remove forearm foreign body	Y		A2	18.3436	\$768.10
25250	Removal of wrist prosthesis	Y		A2	24.699	\$1,034.22
25251	Removal of wrist prosthesis	Y		A2	24.699	\$1,034.22
25259	Manipulate wrist w/anesth	Y		G2	17.5996	\$736.95
25260	Repair forearm tendon/muscle	Y		A2	26.3955	\$1,105.26
25263	Repair forearm tendon/muscle	Y		A2	25.3445	\$1,061.25
25265	Repair forearm tendon/muscle	Y		A2	25.7101	\$1,076.56
25270	Repair forearm tendon/muscle	Y		A2	26.3955	\$1,105.26
25272	Repair forearm tendon/muscle	Y		A2	25.7101	\$1,076.56
25274	Repair forearm tendon/muscle	Y		A2	26.3955	\$1,105.26
25275	Repair forearm tendon sheath	Y		A2	26.3955	\$1,105.26
25280	Revise wrist/forearm tendon	Y		A2	25.7101	\$1,076.56
25290	Incise wrist/forearm tendon	Y		A2	18.7092	\$783.41
25295	Release wrist/forearm tendon	Y		A2	25.7101	\$1,076.56
25300	Fusion of tendons at wrist	Y		A2	25.7101	\$1,076.56
25301	Fusion of tendons at wrist	Y		A2	25.7101	\$1,076.56
25310	Transplant forearm tendon	Y		A2	36.3344	\$1,521.43
25312	Transplant forearm tendon	Y		A2	37.02	\$1,550.14
25315	Revise palsy hand tendon(s)	Y		A2	36.3344	\$1,521.43
25316	Revise palsy hand tendon(s)	Y		A2	66.5231	\$2,785.52
25320	Repair/revise wrist joint	Y		A2	36.3344	\$1,521.43
25335	Realignment of hand	Y		A2	36.3344	\$1,521.43
25337	Reconstruct ulna/radioulnar	Y		A2	37.5168	\$1,570.94

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
25040	Explore/treat wrist joint	Y		A2	26.8925	\$1,126.07
25065	Biopsy forearm soft tissues	Y		P3		\$116.75
25066	Biopsy forearm soft tissues	Y		A2	19.3292	\$809.37
25071	Exc forearm les sc > 3 cm	Y	NI	G2	22.3753	\$936.92
25073	Exc forearm tum deep = 3 cm	Y	NI	G2	22.3753	\$936.92
25075*	Exc forearm les sc < 3 cm	Y	NI	P3		\$140.89
25076	Exc forearm tum deep < 3 cm	Y	NI	G2	16.7399	\$700.95
25077	Resect forearm/wrist tum<3cm	Y	NI	G2	16.7399	\$700.95
25078	Resect forearm/wrist tum=3cm	Y	NI	G2	22.3753	\$936.92
25085	Incision of wrist capsule	Y		A2	18.7092	\$783.41
25100	Biopsy of wrist joint	Y		A2	18.3436	\$768.10
25101	Explore/treat wrist joint	Y		A2	25.7101	\$1,076.56
25105	Remove wrist joint lining	Y		A2	26.3955	\$1,105.26
25107	Remove wrist joint cartilage	Y		A2	25.7101	\$1,076.56
25109	Excise tendon forearm/wrist	Y		G2	21.0617	\$881.92
25110	Remove wrist tendon lesion	Y		A2	18.7092	\$783.41
25111	Remove wrist tendon lesion	Y		A2	18.7092	\$783.41
25112	Remove wrist tendon lesion	Y		A2	19.3946	\$812.11
25115	Remove wrist/forearm lesion	Y		A2	19.3946	\$812.11
25116	Remove wrist/forearm lesion	Y		A2	25.3445	\$1,061.25
25118	Excise wrist tendon sheath	Y		A2	25.7101	\$1,076.56
25120	Partial removal of ulna	Y		A2	25.7101	\$1,076.56
25125	Removal of forearm lesion	Y		A2	25.7101	\$1,076.56
25126	Remove/graft forearm lesion	Y		A2	25.7101	\$1,076.56
25130	Remove/graft forearm lesion	Y		A2	25.7101	\$1,076.56
25136	Removal of wrist lesion	Y		A2	25.7101	\$1,076.56
25136	Remove & graft wrist lesion	Y		A2	25.7101	\$1,076.56
25145	Remove forearm bone lesion	Y		A2	25.3445	\$1,061.25
25150	Partial removal of ulna	Y		A2	25.3445	\$1,061.25

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
25490	Reinforce radius	Y		A2	36.3344	\$1,521.43
25491	Reinforce ulna	Y		A2	36.3344	\$1,521.43
25492	Reinforce radius and ulna	Y		A2	36.3344	\$1,521.43
25500	Treat fracture of radius	Y		P2	1.9858	\$66.40
25505	Treat fracture of radius	Y		A2	4.0322	\$168.84
25515	Treat fracture of radius	Y		A2	35.5372	\$1,488.05
25520	Treat fracture of radius	Y		A2	4.0322	\$168.84
25523	Treat fracture of radius	Y		A2	36.2226	\$1,516.75
25526	Treat fracture of radius	Y		A2	36.7196	\$1,537.56
25530	Treat fracture of ulna	Y		P2	1.5858	\$66.40
25535	Treat fracture of ulna	Y		A2	1.7811	\$74.58
25545	Treat fracture of ulna	Y		A2	35.5372	\$1,488.05
25560	Treat fracture radius & ulna	Y		P2	1.5858	\$66.40
25565	Treat fracture radius & ulna	Y		A2	4.0322	\$168.84
25574	Treat fracture radius & ulna	Y		A2	49.8932	\$2,089.18
25575	Treat fracture radius/ulna	Y		A2	49.8932	\$2,089.18
25600	Treat fracture radius/ulna	Y		P2	1.8658	\$66.40
25605	Treat fracture radius/ulna	Y		A2	4.0322	\$168.84
25606	Treat fx distal radial	Y		A2	21.4573	\$898.48
25607	Treat fx rad extra-articul	Y		A2	51.0756	\$2,138.69
25608	Treat fx rad intra-articul	Y		A2	51.0756	\$2,138.69
25609	Treat fx radial 3+ frag	Y		A2	51.0756	\$2,138.69
25622	Treat wrist bone fracture	Y		P2	1.5858	\$66.40
25624	Treat wrist bone fracture	Y		A2	4.0322	\$168.84
25628	Treat wrist bone fracture	Y		A2	35.5372	\$1,488.05
25630	Treat wrist bone fracture	Y		P2	1.5858	\$66.40
25635	Treat wrist bone fracture	Y		A2	4.0322	\$168.84
25645	Treat wrist bone fracture	Y		A2	35.5372	\$1,488.05
25650	Treat wrist bone fracture	Y		P2	1.5858	\$66.40
25651	Pin ulnar styloid fracture	Y		G2	24.7255	\$1,035.33

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25350	Revision of radius	Y		A2	36.3344	\$1,521.43
25355	Revision of radius	Y		A2	36.3344	\$1,521.43
25360	Revision of ulna	Y		A2	36.3344	\$1,521.43
25365	Revise radius & ulna	Y		A2	36.3344	\$1,521.43
25370	Revise radius or ulna	Y		A2	37.02	\$1,550.14
25375	Revise radius & ulna	Y		A2	36.3344	\$1,521.43
25380	Shorten radius or ulna	Y		A2	37.02	\$1,550.14
25391	Lengthen radius or ulna	Y		A2	25.7101	\$1,076.56
25392	Shorten radius & ulna	Y		A2	37.02	\$1,550.14
25393	Lengthen radius & ulna	Y		G2	44.5617	\$1,865.93
25394	Repair carpal bone, shorten	Y		A2	36.3344	\$1,521.43
25400	Repair radius or ulna	Y		A2	67.2085	\$2,814.22
25405	Repair/graft radius or ulna	Y		A2	66.5231	\$2,785.52
25415	Repair radius & ulna	Y		A2	67.2085	\$2,814.22
25420	Repair/graft radius & ulna	Y		A2	36.3344	\$1,521.43
25425	Repair/graft radius or ulna	Y		A2	37.02	\$1,550.14
25426	Repair/graft radius & ulna	Y		G2	44.5617	\$1,865.93
25430	Vasc graft into carpal bone	Y		G2	44.5617	\$1,865.93
25440	Repair/graft wrist bone	Y		A2	67.2085	\$2,814.22
25441	Reconstruct wrist joint	Y		H8	150.2992	\$6,293.48
25442	Reconstruct wrist joint	Y		H8	150.2992	\$6,293.48
25443	Reconstruct wrist joint	Y		A2	45.772	\$1,916.61
25444	Reconstruct wrist joint	Y		A2	45.772	\$1,916.61
25445	Reconstruct wrist joint	Y		H8	151.8871	\$6,359.97
25446	Wrist replacement	Y		A2	32.7158	\$1,369.91
25447	Repair wrist joint(s)	Y		A2	32.7158	\$1,369.91
25449	Remove wrist joint implant	Y		A2	36.3344	\$1,521.43
25450	Revision of wrist joint	Y		A2	36.3344	\$1,521.43
25455	Revision of wrist joint	Y		A2	36.3344	\$1,521.43

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
26055	Incise finger tendon sheath	Y		A2	14.7756	\$618.70
26060	Incision of finger tendon	Y		A2	14.7756	\$618.70
26070	Explore/treat hand joint	Y		A2	14.7756	\$618.70
26075	Explore/treat finger joint	Y		A2	15.8267	\$662.71
26080	Explore/treat finger joint	Y		A2	15.8267	\$662.71
26100	Biopsy hand joint lining	Y		A2	14.7756	\$618.70
26105	Biopsy finger joint lining	Y		A2	14.1301	\$591.67
26110	Biopsy finger joint lining	Y		A2	14.1301	\$591.67
26111	Exc hand les so > 1.5 cm	Y	NI	G2	22.3753	\$936.92
26113	Exc hand tum deep > 1.5 cm	Y	NI	G2	22.3753	\$936.92
26115	Exc hand les < 1.5 cm	Y	NI	P3		\$287.19
26116	Exc hand tum deep < 1.5 cm	Y	NI	G2	16.7999	\$700.95
26117	Exc hand tum ra < 3 cm	Y	NI	G2	16.7999	\$700.95
26118	Exc hand tum ra > 3 cm	Y	NI	G2	22.3753	\$936.92
26121	Release palm contracture	Y		A2	23.8595	\$993.07
26123	Release palm contracture	Y		A2	23.8595	\$993.07
26125	Release palm contracture	Y		A2	15.8267	\$662.71
26130	Remove wrist joint lining	Y		A2	15.141	\$634.00
26135	Revise finger joint, each	Y		A2	23.8595	\$993.07
26140	Revise finger joint, each	Y		A2	14.7756	\$618.70
26145	Tendon excision, palm/finger	Y		A2	15.141	\$634.00
26160	Remove tendon sheath lesion	Y		A2	15.141	\$634.00
26170	Removal of palm tendon, each	Y		A2	15.141	\$634.00
26180	Removal of finger tendon	Y		A2	15.141	\$634.00
26185	Remove hand bone	Y		A2	15.8267	\$662.71
26200	Remove hand bone lesion	Y		A2	14.7756	\$618.70
26205	Remove/graft bone lesion	Y		A2	23.1741	\$970.37
26210	Removal of finger lesion	Y		A2	14.7756	\$618.70
26215	Remove/graft finger lesion	Y		A2	15.141	\$634.00
26230	Partial removal of hand bone	Y		A2	17.8996	\$748.51

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25652	Treat fracture ulnar styloid	Y		G2	43.499	\$1,921.43
25660	Treat wrist dislocation	Y		A2	1.7811	\$74.58
25670	Treat wrist dislocation	Y		A2	21.4573	\$988.48
25671	Pin radioulnar dislocation	Y		A2	20.4461	\$856.14
25675	Treat wrist dislocation	Y		A2	1.7811	\$74.58
25676	Treat wrist dislocation	Y		A2	21.0916	\$883.17
25680	Treat wrist fracture	Y		A2	1.7811	\$74.58
25685	Treat wrist fracture	Y		A2	21.4573	\$988.48
25690	Treat wrist dislocation	Y		A2	13.7917	\$577.50
25695	Treat wrist dislocation	Y		A2	21.0916	\$883.17
25800	Fusion of wrist joint	Y		A2	67.2085	\$2,814.22
25805	Fusion/graft of wrist joint	Y		A2	37.5168	\$1,570.94
25810	Fusion/graft of wrist joint	Y		A2	67.7054	\$2,835.03
25820	Fusion of hand bones	Y		A2	37.02	\$1,550.14
25825	Fuse hand bones with graft	Y		A2	67.7054	\$2,835.03
25830	Fusion, radioulnar jnt/ulna	Y		A2	67.7054	\$2,835.03
25807	Amputation follow-up surgery	Y		A2	18.7092	\$783.41
25922	Amputate hand at wrist	Y		A2	18.7092	\$783.41
25929	Amputation follow-up surgery	Y		A2	14.4325	\$604.33
25931	Amputation follow-up surgery	Y		G2	21.0617	\$881.92
26010	Drainage of finger abscess	Y		P2	1.9827	\$58.32
26011	Drainage of finger abscess	Y		A2	10.9586	\$458.87
26020	Drain hand tendon sheath	Y		A2	14.7756	\$618.70
26025	Drainage of palm bursa	Y		A2	14.1301	\$591.67
26030	Drainage of palm bursa(s)	Y		A2	14.7756	\$618.70
26034	Treat hand bone lesion	Y		A2	14.7756	\$618.70
26035	Decompress fingers/hand	Y		G2	16.3041	\$682.70
26037	Decompress fingers/hand	Y	CH	G2	16.3041	\$682.70
26040	Release palm contracture	Y		A2	23.8595	\$993.07
26045	Release palm contracture	Y		A2	23.1741	\$970.37

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26437	Reassignment of tendons	Y		A2	15.141	\$634.00
26440	Release palm/finger tendon	Y		A2	15.141	\$634.00
26442	Release palm & finger tendon	Y		A2	23.1741	\$970.37
26445	Release hand/finger tendon	Y		A2	15.141	\$634.00
26449	Release forearm/hand tendon	Y		A2	23.1741	\$970.37
26450	Incision of palm tendon	Y		A2	15.141	\$634.00
26455	Incision of finger tendon	Y		A2	15.141	\$634.00
26460	Incise hand/finger tendon	Y		A2	15.141	\$634.00
26471	Fusion of finger tendons	Y		A2	14.7756	\$618.70
26474	Fusion of finger tendons	Y		A2	14.7756	\$618.70
26476	Tendon lengthening	Y		A2	14.1301	\$591.67
26477	Tendon shortening	Y		A2	14.1301	\$591.67
26478	Lengthening of hand tendon	Y		A2	14.1301	\$591.67
26479	Shortening of hand tendon	Y		A2	14.1301	\$591.67
26480	Transplant hand tendon	Y		A2	23.1741	\$970.37
26483	Transplant/graft hand tendon	Y		A2	23.1741	\$970.37
26489	Transplant palm tendon	Y		A2	22.8085	\$955.06
26490	Transplant/graft palm tendon	Y		A2	23.1741	\$970.37
26492	Revise thumb tendon	Y		A2	23.1741	\$970.37
26494	Tendon transfer with graft	Y		A2	23.1741	\$970.37
26496	Hand tendon/muscle transfer	Y		A2	23.1741	\$970.37
26497	Revise thumb tendon	Y		A2	23.1741	\$970.37
26498	Finger tendon transfer	Y		A2	23.8595	\$995.07
26499	Finger tendon transfer	Y		A2	23.1741	\$970.37
26500	Revision of finger	Y		A2	15.8267	\$662.71
26502	Hand tendon reconstruction	Y		A2	23.8595	\$995.07
26508	Hand tendon reconstruction	Y		A2	15.141	\$634.00
26510	Thumb tendon transfer	Y		A2	23.1741	\$970.37
26516	Fusion of knuckle joint	Y		A2	22.1632	\$928.04

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26235	Partial removal, finger bone	Y		A2	15.141	\$634.00
26236	Partial removal, finger bone	Y		A2	15.141	\$634.00
26250	Extensive hand surgery	Y		A2	15.141	\$634.00
26255	Extensive hand surgery	N	CH	D5		
26260	Resect prox finger tumor	Y		D5	15.141	\$634.00
26261	Extensive finger surgery	N	CH	D5		
26262	Resect distal finger tumor	Y		A2	14.7756	\$618.70
26320	Removal of implant from hand	Y		A2	15.1023	\$632.38
26340	Manipulate finger w/arsneth	Y		G2	4.587	\$192.07
26350	Repair finger/hand tendon	Y		A2	22.1632	\$928.04
26352	Repair/graft hand tendon	Y		A2	23.8595	\$999.07
26356	Repair finger/hand tendon	Y		A2	23.8595	\$999.07
26357	Repair finger/hand tendon	Y		A2	23.8595	\$999.07
26358	Repair/graft hand tendon	Y		A2	23.8595	\$999.07
26372	Repair finger/hand tendon	Y		A2	23.8595	\$999.07
26373	Repair finger/hand tendon	Y		A2	23.1741	\$970.37
26390	Revise hand/finger tendon	Y		A2	23.8595	\$999.07
26392	Repair/graft hand tendon	Y		A2	23.1741	\$970.37
26410	Repair hand tendon	Y		A2	15.141	\$634.00
26412	Repair/graft hand tendon	Y		A2	23.1741	\$970.37
26415	Excision, hand/finger tendon	Y		A2	23.8595	\$999.07
26416	Graft hand or finger tendon	Y		A2	23.1741	\$970.37
26418	Repair finger tendon	Y		A2	15.8267	\$662.71
26420	Repair/graft finger tendon	Y		A2	23.8595	\$999.07
26426	Repair finger/hand tendon	Y		A2	23.1741	\$970.37
26428	Repair/graft finger tendon	Y		A2	23.1741	\$970.37
26432	Repair finger tendon	Y		A2	15.141	\$634.00
26433	Repair finger tendon	Y		A2	15.141	\$634.00
26434	Repair/graft finger tendon	Y		A2	23.1741	\$970.37

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26607	Treat metacarpal fracture	Y		A2	13.7917	\$577.50
26608	Treat metacarpal fracture	Y		A2	22.1427	\$927.18
26615	Treat metacarpal fracture	Y		A2	36.2226	\$1,516.75
26641	Treat thumb dislocation	Y		P2	1.5858	\$66.40
26645	Treat thumb fracture	Y		A2	4.0322	\$168.84
26650	Treat thumb fracture	Y		A2	21.0916	\$883.17
26665	Treat thumb fracture	Y		A2	36.2226	\$1,516.75
26670	Treat hand dislocation	Y		P2	1.5858	\$66.40
26675	Treat hand dislocation	Y		A2	4.0322	\$168.84
26676	Pin hand dislocation	Y		A2	21.0916	\$883.17
26685	Treat hand dislocation	Y		A2	21.4573	\$898.48
26686	Treat hand dislocation	Y		A2	49.8932	\$2,089.18
26700	Treat knuckle dislocation	Y		P2	1.5858	\$66.40
26705	Treat knuckle dislocation	Y		A2	1.7811	\$74.58
26706	Pin knuckle dislocation	Y		A2	13.7917	\$577.50
26715	Treat knuckle dislocation	Y		A2	22.1427	\$927.18
26720	Treat finger fracture, each	Y		P2	1.5858	\$66.40
26725	Treat finger fracture, each	Y		A2	1.5858	\$66.40
26727	Treat finger fracture, each	Y		A2	24.2275	\$1,014.48
26735	Treat finger fracture, each	Y		A2	22.1427	\$927.18
26740	Treat finger fracture, each	Y		P2	1.5858	\$66.40
26742	Treat finger fracture, each	Y		A2	1.7811	\$74.58
26746	Treat finger fracture, each	Y		A2	22.6396	\$947.99
26750	Treat finger fracture, each	Y		P2	1.5858	\$66.40
26755	Treat finger fracture, each	Y		G2	1.5858	\$66.40
26756	Pin finger fracture, each	Y		A2	21.0916	\$883.17
26765	Treat finger fracture, each	Y		A2	22.1427	\$927.18
26775	Treat finger dislocation	Y		G2	1.5858	\$66.40
26776	Pin finger dislocation	Y		P3	21.0916	\$883.17

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26517	Fusion of knuckle joints	Y		A2	23.1741	\$970.37
26518	Fusion of knuckle joints	Y		A2	23.1741	\$970.37
26520	Release knuckle contracture	Y		A2	15.141	\$634.00
26525	Release finger contracture	Y		A2	15.141	\$634.00
26530	Revise knuckle joint	Y		A2	31.5334	\$1,320.40
26531	Revise knuckle with implant	Y		A2	47.3599	\$1,963.10
26535	Revise finger joint	Y		A2	32.7158	\$1,369.91
26536	Revise/implant finger joint	Y		A2	45.772	\$1,916.61
26540	Repair hand joint	Y		A2	15.8287	\$662.71
26541	Repair hand joint with graft	Y		A2	25.9444	\$1,086.37
26542	Repair hand joint with graft	Y		A2	15.8287	\$662.71
26545	Reconstruct finger joint	Y		A2	23.8595	\$999.07
26546	Repair nonunion hand	Y		A2	23.8595	\$999.07
26548	Reconstruct finger joint	Y		A2	23.8595	\$999.07
26550	Constrict thumb replacement	Y		A2	22.8085	\$965.06
26555	Positional change of finger	Y		A2	23.1741	\$970.37
26560	Repair of web finger	Y		A2	14.7756	\$618.70
26561	Repair of web finger	Y		A2	23.1741	\$970.37
26562	Repair of web finger	Y		A2	23.8595	\$999.07
26565	Correct metacarpal flaw	Y		A2	24.3565	\$1,019.88
26567	Correct finger deformity	Y		A2	24.3565	\$1,019.88
26568	Lengthen metacarpal/finger	Y		A2	23.1741	\$970.37
26587	Repair hand deformity	Y		A2	16.3234	\$683.51
26588	Reconstruct extra finger	Y		A2	16.3234	\$683.51
26590	Repair finger deformity	Y		A2	16.3234	\$683.51
26591	Repair muscles of hand	Y		A2	23.1741	\$970.37
26593	Release muscles of hand	Y		A2	15.141	\$634.00
26596	Excision constricting tissue	Y		A2	14.7756	\$618.70
26600	Treat metacarpal fracture	Y		P2	1.5858	\$66.40
26605	Treat metacarpal fracture	Y		A2	1.7811	\$74.58

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27052	Biopsy of hip joint	Y		A2	18.7092	\$783.41
27059	Resect hip/pelv tum > 5 cm	Y	NI	G2	22.3753	\$936.92
27060	Removal of ischial bursa	Y		A2	19.8916	\$832.92
27062	Removal femur lesion/bursa	Y		A2	19.8916	\$832.92
27065	Removal of hip bone lesion	Y		A2	26.8925	\$1,126.07
27067	Removal/graft hip bone lesion	Y		A2	26.8925	\$1,126.07
27080	Removal of tail bone	Y		A2	25.3445	\$1,061.25
27086	Remove hip foreign body	Y		A2	7.7878	\$326.10
27087	Remove hip foreign body	Y		A2	18.7092	\$783.41
27089	Injection for hip x-ray	N		N1		
27095	Injection for hip x-ray	N		N1		
27097	Revision of hip tendon	Y		A2	25.7101	\$1,076.56
27098	Transfer tendon to pelvis	Y		A2	25.7101	\$1,076.56
27100	Transfer of abdominal muscle	Y		A2	37.02	\$1,550.14
27105	Transfer of spinal muscle	Y		A2	37.02	\$1,550.14
27110	Transfer of ilioosios muscle	Y		A2	37.02	\$1,550.14
27111	Transfer of ilioosios muscle	Y		A2	37.02	\$1,550.14
27193	Treat pelvic ring fracture	Y		A2	1.7811	\$74.58
27194	Treat pelvic ring fracture	Y		A2	13.5199	\$566.12
27200	Treat tail bone fracture	Y	CH	P3		\$63.63
27202	Treat tail bone fracture	Y		A2	95.1716	\$1,472.74
27220	Treat hip socket fracture	Y		G2	1.5858	\$66.40
27230	Treat thigh fracture	Y		A2	1.7811	\$74.58
27238	Treat thigh fracture	Y		A2	4.0322	\$168.84
27246	Treat thigh fracture	Y		A2	4.0322	\$168.84
27252	Treat hip dislocation	Y		A2	1.7811	\$74.58
27256	Treat hip dislocation	Y		A2	13.5199	\$566.12
27257	Treat hip dislocation	Y		A2	13.8856	\$581.43

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
26785	Treat finger dislocation	Y		A2	21.0916	\$883.17
26820	Thumb fusion with graft	Y		A2	24.3565	\$1,019.88
26841	Fusion of thumb	Y		A2	23.8595	\$999.07
26842	Thumb fusion with graft	Y		A2	23.8595	\$999.07
26843	Fusion of hand joint	Y		A2	23.1741	\$970.37
26844	Fusion/graft of hand joint	Y		A2	23.1741	\$970.37
26850	Fusion of knuckle	Y		A2	23.8595	\$999.07
26852	Fusion of knuckle with graft	Y		A2	23.8595	\$999.07
26860	Fusion of finger joint	Y		A2	23.1741	\$970.37
26861	Fusion of finger jnt, add-on	Y		A2	22.8085	\$955.06
26862	Fusion/graft of finger joint	Y		A2	23.8595	\$999.07
26863	Fuse/graft added joint	Y		A2	23.1741	\$970.37
26910	Anipulate metacarpal bone	Y		A2	23.1741	\$970.37
26951	Amputation of finger/thumb	Y		A2	14.7756	\$618.70
26952	Amputation of finger/thumb	Y		A2	15.8267	\$662.71
26990	Drainage of pelvis lesion	Y		A2	17.6983	\$741.08
26991	Drainage of pelvis bursa	Y		A2	17.6983	\$741.08
27000	Incision of hip tendon	Y		A2	18.3436	\$768.10
27001	Incision of hip tendon	Y		A2	25.7101	\$1,076.56
27003	Incision of hip tendon	Y		A2	25.7101	\$1,076.56
27033	Exploration of hip joint	Y		A2	36.3344	\$1,521.43
27035	Dernervation of hip joint	Y		A2	37.02	\$1,550.14
27040	Biopsy of soft tissues	Y		A2	7.7878	\$326.10
27041	Biopsy of soft tissues	Y		A2	8.2762	\$346.55
27043	Exc hip/pelvis les sc > 3 cm	Y	NI	G2	22.3753	\$936.92
27045	Exc hip/pelvis tum deep > 5 cm	Y	NI	G2	22.3753	\$936.92
27047*	Exc hip/pelvis les sc < 3 cm	Y	NI	P3		\$200.26
27048	Exc hip/pelvis tum deep < 5 cm	Y	NI	G2	16.7399	\$700.95
27049	Resect hip/pelv tum < 5 cm	Y	NI	G2	16.7399	\$700.95
27050	Biopsy of sacroiliac joint	Y		A2	18.7092	\$783.41

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27357	Remove femur lesion/graft	Y		A2	26.8925	\$1,126.07
27358	Remove femur lesion/fixation	Y		A2	26.8925	\$1,126.07
27360	Partial removal, leg bone(s)	Y		A2	26.8925	\$1,126.07
27364	Resect thigh/knee tum >5 cm	Y	NI	G2	22.3753	\$936.92
27370	Injection for knee x-ray	N		N1		
27372	Removal of foreign body	Y		A2	22.4651	\$940.68
27380	Repair of kneecap tendon	Y		A2	17.6983	\$741.08
27381	Repair/graft kneecap tendon	Y		A2	18.7092	\$783.41
27385	Repair of thigh muscle	Y		A2	18.7092	\$783.41
27386	Repair/graft of thigh muscle	Y		A2	18.7092	\$783.41
27390	Incision of thigh tendon	Y		A2	17.6983	\$741.08
27391	Incision of thigh tendons	Y		A2	18.3436	\$768.10
27392	Incision of thigh tendons	Y		A2	18.7092	\$783.41
27393	Lengthening of thigh tendon	Y		A2	25.3445	\$1,061.25
27394	Lengthening of thigh tendons	Y		A2	25.7101	\$1,076.56
27395	Lengthening of thigh tendons	Y		A2	36.3344	\$1,521.43
27396	Transplant of thigh tendon	Y		A2	25.7101	\$1,076.56
27397	Transplants of thigh tendons	Y		A2	36.3344	\$1,521.43
27400	Revise thigh muscles/tendons	Y		A2	26.3955	\$1,105.26
27403	Repair of knee cartilage	Y		A2	37.02	\$1,550.14
27405	Repair of knee ligament	Y		A2	67.2085	\$2,814.22
27407	Repair of knee ligament	Y		A2	37.02	\$1,550.14
27409	Repair of knee ligaments	Y		A2	44.5617	\$1,865.93
27416	Osteochondral knee autograft	Y		G2	36.3344	\$1,521.43
27418	Repair degenerated kneecap	Y		A2	36.3344	\$1,521.43
27420	Revision of unstable kneecap	Y		A2	39.1047	\$1,637.43
27422	Revision/removal of kneecap	Y		A2	36.3344	\$1,521.43
27424	Revision/removal of kneecap	Y		A2	28.4804	\$1,192.56
27425	Lat retinacular release open	Y		A2	36.3344	\$1,521.43
27427	Reconstruction, knee	Y		A2	36.3344	\$1,521.43

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27265	Treat hip dislocation	Y		A2	1.7811	\$74.58
27266	Treat hip dislocation	Y		A2	13.5199	\$566.12
27267	Clx thigh fx	Y		G2	1.5858	\$66.40
27275	Manipulation of hip joint	Y		A2	13.5199	\$566.12
27301	Drain thigh/knee lesion	Y		A2	16.8376	\$705.04
27305	Incise thigh tendon & fascia	Y		A2	18.3436	\$768.10
27306	Incision of thigh tendon	Y		A2	18.7092	\$783.41
27307	Incision of thigh tendons	Y		A2	18.7092	\$783.41
27310	Exploration of knee joint	Y		A2	26.3955	\$1,105.26
27323	Biopsy, thigh soft tissues	Y		A2	7.7878	\$326.10
27324	Biopsy, thigh soft tissues	Y		A2	18.6936	\$782.34
27325	Neurectomy, hamstring	Y		A2	15.9795	\$669.11
27326	Neurectomy, popliteal	Y		A2	15.9795	\$669.11
27327*	Exc thigh/knee les sc < 3 cm	Y	NI	P3	16.7399	\$700.95
27328	Exc thigh/knee tum deep <5cm	Y	NI	G2	16.7399	\$700.95
27329	Resect thigh/knee tum < 5 cm	Y	NI	G2	26.3955	\$1,105.26
27330	Biopsy, knee joint lining	Y		A2	26.3955	\$1,105.26
27331	Explore/treat knee joint	Y		A2	26.3955	\$1,105.26
27332	Removal of knee cartilage	Y		A2	26.3955	\$1,105.26
27334	Remove knee joint lining	Y		A2	26.3955	\$1,105.26
27335	Remove knee joint lining	Y		A2	26.3955	\$1,105.26
27337	Exc thigh/knee les sc > 3 cm	Y	NI	G2	22.3753	\$936.92
27339	Exc thigh/knee tum deep >5cm	Y	NI	G2	22.3753	\$936.92
27340	Removal of kneecap bursa	Y		A2	18.7092	\$783.41
27345	Removal of knee cyst	Y		A2	19.3946	\$812.11
27347	Remove knee cyst	Y		A2	19.3946	\$812.11
27350	Removal of Kneecap	Y		A2	26.3955	\$1,105.26
27355	Remove femur lesion	Y		A2	25.7101	\$1,076.56
27356	Remove femur lesion/graft	Y		A2	26.3955	\$1,105.26

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27550	Treat knee dislocation	Y	A2	1.7811	\$74.58
27552	Treat knee dislocation	Y	A2	12.8746	\$539.10
27560	Treat kneecap dislocation	Y	A2	1.7811	\$74.58
27562	Treat kneecap dislocation	Y	A2	12.8746	\$539.10
27566	Treat kneecap dislocation	Y	A2	35.1716	\$1,472.74
27570	Fixation of knee joint	Y	A2	12.8746	\$539.10
27594	Amputation follow-up surgery	Y	A2	16.7092	\$783.41
27600	Decompression of lower leg	Y	A2	18.7092	\$783.41
27601	Decompression of lower leg	Y	A2	18.7092	\$783.41
27602	Decompression of lower leg	Y	A2	18.7092	\$783.41
27603	Decompression of lower leg	Y	A2	16.472	\$689.73
27604	Drain lower leg bursa	Y	A2	18.3436	\$768.10
27605	Incision of achilles tendon	Y	A2	17.5755	\$735.94
27606	Incision of achilles tendon	Y	A2	17.6963	\$741.08
27607	Treat lower leg bone lesion	Y	A2	18.3436	\$768.10
27610	Explore/treat ankle joint	Y	A2	25.3445	\$1,061.25
27612	Exploration of ankle joint	Y	A2	25.7101	\$1,076.56
27613	Biopsy lower leg soft tissue	Y	P3		\$110.78
27614	Biopsy lower leg soft tissue	Y	A2	19.3292	\$809.37
27615	Resect leg/ankle tum < 5 cm	Y	G2	16.7399	\$700.95
27616	Resect leg/ankle tum > 5 cm	Y	G2	22.3753	\$936.92
27618*	Exc leg/ankle tum < 3 cm	Y	NI		\$188.05
27619	Exc leg/ankle tum deep < 5 cm	Y	NI	16.7399	\$700.95
27620	Explore/treat ankle joint	Y	A2	26.3955	\$1,105.26
27625	Remove ankle joint lining	Y	A2	26.3955	\$1,105.26
27626	Remove ankle joint lining	Y	A2	26.3955	\$1,105.26
27630	Removal of tendon lesion	Y	A2	18.7092	\$783.41
27632	Exc leg/ankle les sc > 3 cm	Y	NI	22.3753	\$936.92
27634	Exc leg/ankle tum deep > 5 cm	Y	NI	22.3753	\$936.92
27635	Remove lower leg bone lesion	Y	A2	25.7101	\$1,076.56

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27428	Reconstruction, knee	Y	A2	67.2085	\$2,814.22
27429	Reconstruction, knee	Y	A2	67.2085	\$2,814.22
27430	Revision of thigh muscles	Y	A2	37.02	\$1,550.14
27435	Incision of knee joint	Y	A2	37.02	\$1,550.14
27437	Revise kneecap	Y	A2	32.2191	\$1,349.11
27438	Revise kneecap with implant	Y	A2	45.772	\$1,916.61
27440	Revision of knee joint	Y	G2	38.1606	\$1,597.90
27441	Revision of knee joint	Y	A2	32.7158	\$1,369.91
27442	Revision of knee joint	Y	A2	32.7158	\$1,369.91
27443	Revision of knee joint	Y	A2	32.7158	\$1,369.91
27446	Revision of knee joint	Y	J8	158.2621	\$6,626.91
27475	Surgery to stop leg growth	Y	CH	30.396	\$1,272.77
27479	Surgery to stop leg growth	Y	CH	30.396	\$1,272.77
27496	Decompression of thigh/knee	Y	A2	26.8925	\$1,126.07
27497	Decompression of thigh/knee	Y	A2	18.7092	\$783.41
27498	Decompression of thigh/knee	Y	A2	25.7101	\$1,076.56
27499	Decompression of thigh/knee	Y	A2	25.7101	\$1,076.56
27500	Treatment of thigh fracture	Y	A2	4.0322	\$168.84
27501	Treatment of thigh fracture	Y	A2	1.7811	\$74.58
27502	Treatment of thigh fracture	Y	A2	13.7917	\$577.50
27503	Treatment of thigh fracture	Y	A2	1.7811	\$74.58
27508	Treatment of thigh fracture	Y	A2	1.7811	\$74.58
27509	Treatment of thigh fracture	Y	A2	21.4573	\$898.48
27510	Treatment of thigh fracture	Y	A2	4.0322	\$168.84
27516	Treat thigh fx growth plate	Y	A2	1.7811	\$74.58
27517	Treat thigh fx growth plate	Y	A2	1.7811	\$74.58
27520	Treat kneecap fracture	Y	A2	1.7811	\$74.58
27530	Treat knee fracture	Y	A2	1.7811	\$74.58
27532	Treat knee fracture	Y	A2	13.7917	\$577.50
27538	Treat knee fracture(s)	Y	A2	1.7811	\$74.58

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27707	Incision of fibula	Y		A2	18.3436	\$768.10
27709	Incision of tibia & fibula	Y		A2	25.3445	\$1,061.25
27720	Repair of fibula	Y	CH	G2	43.499	\$1,821.43
27726	Repair fibula nonunion	Y		G2	43.499	\$1,821.43
27730	Repair of tibia epiphysis	Y		A2	25.3445	\$1,061.25
27732	Repair of fibula epiphysis	Y		A2	25.3445	\$1,061.25
27734	Repair lower leg epiphyses	Y		A2	25.3445	\$1,061.25
27740	Repair of leg epiphyses	Y		A2	25.3445	\$1,061.25
27742	Repair of leg epiphyses	Y		A2	35.969	\$1,506.13
27745	Reinforce tibia	Y		A2	66.5231	\$2,785.52
27750	Treatment of tibia fracture	Y		A2	1.7811	\$74.58
27752	Treatment of fibula fracture	Y		A2	13.7917	\$577.50
27756	Treatment of tibia fracture	Y		A2	21.4573	\$898.48
27758	Treatment of fibula fracture	Y		A2	36.2226	\$1,516.75
27759	Treatment of tibia fracture	Y		A2	50.5787	\$2,117.88
27760	Clix medial ankle fx	Y		A2	1.7811	\$74.58
27762	Clix med ankle fx w/mppj	Y		A2	13.7917	\$577.50
27766	Optx medial ankle fx	Y		A2	35.5372	\$1,488.05
27767	Clix post ankle fx	Y		G2	1.5858	\$66.40
27768	Clix post ankle fx w/mppj	Y		G2	1.5858	\$66.40
27769	Optx post ankle fx	Y		G2	43.499	\$1,821.43
27780	Treatment of fibula fracture	Y		A2	1.7811	\$74.58
27781	Treatment of fibula fracture	Y		A2	13.7917	\$577.50
27784	Treatment of fibula fracture	Y		A2	35.5372	\$1,488.05
27786	Treatment of ankle fracture	Y		A2	1.7811	\$74.58
27788	Treatment of ankle fracture	Y		A2	1.7811	\$74.58
27792	Treatment of ankle fracture	Y		A2	35.5372	\$1,488.05
27808	Treatment of ankle fracture	Y		A2	1.7811	\$74.58
27810	Treatment of ankle fracture	Y		A2	4.0322	\$168.84
27814	Treatment of ankle fracture	Y		A2	35.5372	\$1,488.05

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27637	Remove/graft leg bone lesion	Y		A2	25.7101	\$1,076.56
27638	Remove/graft leg bone lesion	Y		A2	25.7101	\$1,076.56
27640	Partial removal of fibula	Y		A2	35.969	\$1,506.13
27641	Partial removal of fibula	Y		A2	25.3445	\$1,061.25
27647	Resect talus/calcaneus tum	Y		A2	36.3344	\$1,521.43
27648	Injection for ankle x-ray	N		N1		
27650	Repair achilles tendon	Y		A2	36.3344	\$1,521.43
27652	Repair/graft achilles tendon	Y		A2	66.5231	\$2,785.52
27654	Repair of achilles tendon	Y		A2	1.521.43	\$768.10
27656	Repair leg fascia defect	Y		A2	17.9983	\$741.08
27658	Repair of leg tendon, each	Y		A2	18.3436	\$768.10
27659	Repair of leg tendon, each	Y		A2	25.3445	\$1,061.25
27664	Repair of leg tendon, each	Y		A2	25.3445	\$1,061.25
27665	Repair of leg tendon, each	Y		A2	18.3436	\$768.10
27675	Repair lower leg tendons	Y		A2	25.7101	\$1,076.56
27676	Repair lower leg tendons	Y		A2	25.7101	\$1,076.56
27680	Release of lower leg tendon	Y		A2	25.7101	\$1,076.56
27681	Release of lower leg tendons	Y		A2	25.3445	\$1,061.25
27685	Revision of lower leg tendon	Y		A2	25.7101	\$1,076.56
27686	Revision lower leg tendons	Y		A2	25.7101	\$1,076.56
27687	Revision of calf tendon	Y		A2	25.7101	\$1,076.56
27690	Revision lower leg tendon	Y		A2	37.02	\$1,550.14
27691	Revision lower leg tendon	Y		A2	37.02	\$1,550.14
27692	Revise additional leg tendon	Y		A2	36.3344	\$1,521.43
27695	Repair of ankle ligament	Y		A2	25.3445	\$1,061.25
27696	Repair of ankle ligaments	Y		A2	25.3445	\$1,061.25
27698	Repair of ankle ligament	Y		A2	25.3445	\$1,061.25
27700	Revision of ankle joint	Y		A2	32.7158	\$1,369.91
27704	Removal of ankle implant	Y		A2	18.3436	\$768.10
27705	Incision of tibia	Y		A2	35.969	\$1,506.13

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28010	Incision of toe tendon	Y		P3		\$80.11
28011	Incision of toe tendons	Y		A2	18.5864	\$78.27
28020	Exploration of foot joint	Y		A2	18.221	\$762.97
28022	Exploration of foot joint	Y		A2	18.221	\$762.97
28024	Exploration of toe joint	Y		A2	18.221	\$762.97
28035	Decompression of tibia nerve	Y		A2	17.0305	\$713.12
28039*	Exc foot/toe lum sc > 1.5 cm	Y	NI	P3		\$199.69
28041*	Exc foot/toe lum deep > 1.5cm	Y	NI	R2	22.3753	\$936.92
28043*	Exc foot/toe lum sc < 1.5 cm	Y	NI	P3		\$142.03
28045*	Exc foot/toe lum deep < 1.5cm	Y	NI	P3		\$194.87
28046*	Resect foot/toe tumor < 3 cm	Y	NI	R2	16.7399	\$700.95
28047	Resect foot/toe tumor > 3 cm	Y	NI	G2	22.3753	\$936.92
28050	Biopsy of foot joint lining	Y		A2	18.221	\$762.97
28052	Biopsy of foot joint lining	Y		A2	18.221	\$762.97
28054	Biopsy of toe joint lining	Y		A2	18.221	\$762.97
28055	Neurectomy, foot	Y		A2	17.0305	\$713.12
28060	Partial removal, foot fascia	Y		A2	18.221	\$762.97
28062	Removal of foot fascia	Y		A2	18.5864	\$778.27
28070	Removal of foot joint lining	Y		A2	18.5864	\$778.27
28072	Removal of foot joint lining	Y		A2	18.5864	\$778.27
28080	Removal of foot lesion	Y		A2	18.5864	\$778.27
28086	Excise foot tendon sheath	Y		A2	18.221	\$762.97
28088	Excise foot tendon sheath	Y		A2	18.221	\$762.97
28090	Removal of foot lesion	Y		A2	18.5864	\$778.27
28092	Removal of toe lesions	Y		A2	18.5864	\$778.27
28100	Removal of ankle/heel lesion	Y		A2	18.221	\$762.97
28102	Removal/graft foot lesion	Y		A2	40.6016	\$1,700.11
28103	Removal/graft foot lesion	Y		A2	40.6016	\$1,700.11
28104	Removal of foot lesion	Y		A2	18.221	\$762.97
28106	Removal/graft foot lesion	Y		A2	40.6016	\$1,700.11

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ADDENDUM AA—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPFS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
27816	Treatment of ankle fracture	Y		A2	1.7811	\$74.58
27818	Treatment of ankle fracture	Y		A2	4.0322	\$168.84
27822	Treatment of ankle fracture	Y		A2	35.5372	\$1,488.05
27823	Treatment of ankle fracture	Y		A2	49.8932	\$2,089.18
27824	Treat lower leg fracture	Y		A2	1.7811	\$74.58
27825	Treat lower leg fracture	Y		A2	13.7917	\$577.50
27826	Treat lower leg fracture	Y		A2	35.5372	\$1,488.05
27827	Treat lower leg fracture	Y		A2	49.8932	\$2,089.18
27828	Treat lower leg fracture	Y		A2	50.5787	\$2,117.88
27829	Treat lower leg joint	Y		A2	35.1716	\$1,472.74
27830	Treat lower leg dislocation	Y		A2	1.7811	\$74.58
27831	Treat lower leg dislocation	Y		A2	13.7917	\$577.50
27832	Treat lower leg dislocation	Y		A2	35.1716	\$1,472.74
27840	Treat ankle dislocation	Y		A2	4.0322	\$168.84
27842	Treat ankle dislocation	Y		A2	12.8746	\$539.10
27846	Treat ankle dislocation	Y		A2	35.5372	\$1,488.05
27848	Treat ankle dislocation	Y		A2	35.5372	\$1,488.05
27860	Fixation of ankle joint	Y		A2	12.8746	\$539.10
27870	Fusion of ankle joint, open	Y		A2	67.2085	\$2,814.22
27871	Fusion of fibiotalar joint	Y		A2	67.2085	\$2,814.22
27884	Amputation follow-up surgery	Y		A2	18.7092	\$783.41
27889	Amputation of foot at ankle	Y		A2	25.7101	\$1,076.56
27892	Decompression of leg	Y		A2	25.7101	\$1,076.56
27893	Decompression of leg	Y		A2	25.7101	\$1,076.56
27894	Decompression of leg	Y		A2	25.7101	\$1,076.56
28001	Drainage of bursa of foot	Y		P3		\$109.65
28002	Treatment of foot infection	Y		A2	18.7092	\$783.41
28003	Treatment of foot infection	Y		A2	18.7092	\$783.41
28005	Treat foot bone lesion	Y		A2	18.5864	\$778.27
28008	Incision of foot fascia	Y		A2	18.5864	\$778.27

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
28222	Release of foot tendons	Y		A2	17.5755	\$735.94
28225	Release of foot tendon	Y		A2	17.5755	\$735.94
28226	Release of foot tendons	Y		A2	17.5755	\$735.94
28230	Incision of foot tendon(s)	Y		P3		\$171.57
28232	Incision of toe tendon	Y		P3		\$164.47
28234	Incision of foot tendon	Y		A2	18.221	\$762.97
28238	Revision of foot tendon	Y		A2	40.6016	\$1,700.11
28240	Release of big toe	Y		A2	18.221	\$762.97
28250	Revision of foot fascia	Y		A2	18.5864	\$778.27
28260	Release of midfoot joint	Y		A2	18.5864	\$778.27
28261	Revision of foot tendon	Y		A2	18.5864	\$778.27
28262	Revision of foot and ankle	Y		A2	19.2721	\$806.98
28264	Release of midfoot joint	Y		A2	39.5907	\$1,657.78
28270	Release of foot contracture	Y		A2	18.5864	\$778.27
28272	Release of toe joint, each	Y		P3		\$158.51
28280	Fusion of toes	Y		A2	18.221	\$762.97
28285	Repair of hammer toe	Y		A2	18.5864	\$778.27
28286	Repair of hammer toe	Y		A2	19.2721	\$806.98
28288	Partial removal of foot bone	Y		A2	18.5864	\$778.27
28289	Repair hallux rigidus	Y		A2	18.5864	\$778.27
28290	Correction of bunion	Y		A2	25.1878	\$1,054.69
28292	Correction of bunion	Y		A2	25.1878	\$1,054.69
28293	Correction of bunion	Y		A2	25.5535	\$1,070.00
28294	Correction of bunion	Y		A2	25.5535	\$1,070.00
28296	Correction of bunion	Y		A2	25.5535	\$1,070.00
28297	Correction of bunion	Y		A2	25.5535	\$1,070.00
28298	Correction of bunion	Y		A2	26.7358	\$1,119.51
28299	Correction of bunion	Y		A2	40.2362	\$1,684.81
28300	Incision of heel bone	Y		A2	18.221	\$762.97
28302	Incision of ankle bone	Y		A2	18.221	\$762.97

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
28107	Remove/graft foot lesion	Y		A2	40.6016	\$1,700.11
28108	Removal of toe lesions	Y		A2	18.221	\$762.97
28110	Part removal of metatarsal	Y		A2	18.5864	\$778.27
28111	Part removal of metatarsal	Y		A2	18.5864	\$778.27
28112	Part removal of metatarsal	Y		A2	18.5864	\$778.27
28113	Part removal of metatarsal	Y		A2	18.5864	\$778.27
28114	Removal of metatarsal heads	Y		A2	18.5864	\$778.27
28116	Revision of foot	Y		A2	18.5864	\$778.27
28118	Removal of heel bone	Y		A2	19.2721	\$806.98
28119	Removal of heel spur	Y		A2	18.5864	\$778.27
28120	Part removal of ankle/heel	Y		A2	21.9567	\$894.27
28122	Partial removal of foot bone	Y		A2	18.5864	\$778.27
28124	Partial removal of toe	Y		P3		\$185.78
28126	Partial removal of toe	Y		A2	18.5864	\$778.27
28130	Removal of ankle bone	Y		A2	18.5864	\$778.27
28140	Removal of metatarsal	Y		A2	18.5864	\$778.27
28150	Removal of toe	Y		A2	18.5864	\$778.27
28153	Partial removal of toe	Y		A2	18.5864	\$778.27
28160	Partial removal of toe	Y		A2	18.5864	\$778.27
28171	Resect tarsal tumor	Y		A2	18.5864	\$778.27
28173	Resect metatarsal tumor	Y		A2	18.5864	\$778.27
28175	Resect phalanx of toe tumor	Y		A2	18.5864	\$778.27
28190	Removal of foot foreign body	Y		P3		\$113.62
28192	Removal of foot foreign body	Y		A2	15.1023	\$632.38
28193	Removal of foot foreign body	Y		A2	8.2762	\$346.55
28200	Repair of foot tendon	Y		A2	18.5864	\$778.27
28202	Repair/graft of foot tendon	Y		A2	18.5864	\$778.27
28208	Repair of foot tendon	Y		A2	18.5864	\$778.27
28210	Repair/graft of foot tendon	Y		A2	40.6016	\$1,700.11
28220	Release of foot tendon	Y		P3		\$175.55

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Year Transition Payment Weight	CY 2010 Third Year Transition Payment
28470	Treat metatarsal fracture	Y		1.5858	\$66.40
28475	Treat metatarsal fracture	Y		1.5858	\$66.40
28476	Treat metatarsal fracture	Y		21.0916	\$883.17
28485	Treat metatarsal fracture	Y		36.2226	\$1,516.75
28490	Treat big toe fracture	Y	CH		\$61.64
28495	Treat big toe fracture	Y		1.5858	\$66.40
28496	Treat big toe fracture	Y		21.0916	\$883.17
28505	Treat big toe fracture	Y		21.4573	\$898.48
28510	Treatment of toe fracture	Y			\$48.86
28515	Treatment of toe fracture	Y	CH		\$63.35
28525	Treat toe fracture	Y		21.4573	\$898.48
28530	Treat sesamoid bone fracture	Y			\$46.87
28531	Treat sesamoid bone fracture	Y		21.4573	\$898.48
28540	Treat foot dislocation	Y		20.4461	\$856.14
28545	Treat foot dislocation	Y		21.0916	\$883.17
28546	Treat foot dislocation	Y		35.1716	\$1,472.74
28555	Repair foot dislocation	Y			\$67.61
28570	Treat foot dislocation	Y		13.7917	\$577.50
28575	Treat foot dislocation	Y		21.4573	\$898.48
28576	Treat foot dislocation	Y		21.4573	\$898.48
28585	Repair foot dislocation	Y		1.5858	\$66.40
28600	Treat foot dislocation	Y		1.7811	\$74.58
28605	Treat foot dislocation	Y		21.0916	\$883.17
28615	Repair foot dislocation	Y		35.5372	\$1,488.05
28630	Treat toe dislocation	Y	CH		\$53.12
28635	Treat toe dislocation	Y		12.8746	\$539.10
28636	Treat toe dislocation	Y		21.4573	\$898.48
28645	Repair toe dislocation	Y		21.4573	\$898.48
28660	Treat toe dislocation	Y			\$38.92

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28304	Incision of midfoot bones	Y		40.2362	\$1,684.81
28305	Incise/graft midfoot bones	Y		40.8016	\$1,700.11
28306	Incision of metatarsal	Y		19.2721	\$806.98
28307	Incision of metatarsal	Y		18.221	\$762.97
28308	Incision of metatarsals	Y		41.2872	\$1,728.82
28310	Revision of big toe	Y		18.5864	\$778.27
28312	Revision of toe	Y		18.5864	\$778.27
28313	Repair deformity of toe	Y		18.221	\$762.97
28315	Removal of sesamoid bone	Y		19.2721	\$806.98
28320	Repair of foot bones	Y		41.2872	\$1,728.82
28322	Repair of metatarsals	Y		41.2872	\$1,728.82
28340	Resect enlarged toe tissue	Y		19.2721	\$806.98
28341	Resect enlarged toe	Y		19.2721	\$806.98
28344	Repair extra toe(s)	Y		19.2721	\$806.98
28345	Repair webbed toe(s)	Y		1.7811	\$74.58
28400	Treatment of heel fracture	Y		13.7917	\$577.50
28405	Treatment of heel fracture	Y		49.8932	\$2,069.18
28415	Treat heel fracture	Y		36.2226	\$1,516.75
28420	Treat/graft heel fracture	Y		1.5858	\$66.40
28430	Treatment of ankle fracture	Y		1.7811	\$74.58
28435	Treatment of ankle fracture	Y		21.0916	\$883.17
28436	Treatment of ankle fracture	Y		35.5372	\$1,488.05
28445	Treat ankle fracture	Y		50.2514	\$2,104.18
28446	Osteochondral talus autograft	Y			\$66.40
28450	Treat midfoot fracture, each	Y		1.5858	\$66.40
28455	Treat midfoot fracture, each	Y		21.0916	\$883.17
28456	Treat midfoot fracture, each	Y		35.5372	\$1,488.05

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
29075	Application of forearm cast	N		P3		\$37.21
29085	Apply hand/wrist cast	N	CH	P3		\$38.06
29086	Apply finger cast	N		P3		\$52.10
29105	Apply long arm splint	N		P3		\$33.52
29126	Apply forearm splint	N		P3		\$29.26
29128	Apply forearm splint	N		P3		\$30.96
29130	Application of finger splint	N		P3		\$13.35
29131	Application of finger splint	N		P3		\$19.03
29200	Strapping of chest	N		P3		\$18.75
29220	Strapping of low back	N	CH	D5		\$20.45
29240	Strapping of shoulder	N		P3		\$19.88
29260	Strapping of elbow or wrist	N		P3		\$20.17
29280	Strapping of hand or finger	N		P2	2.2441	\$93.97
29305	Application of hip cast	N		P2	2.2441	\$93.97
29325	Application of hip casts	N		P2	2.2441	\$93.97
29345	Application of long leg cast	N		P3		\$50.56
29355	Application of long leg cast	N		P3		\$49.99
29358	Apply long leg cast brace	N		P3		\$62.49
29365	Application of long leg cast	N		P3		\$47.72
29405	Apply short leg cast	N		P3		\$35.79
29425	Apply short leg cast	N		P3		\$36.08
29435	Apply short leg cast	N		P3		\$45.73
29440	Addition of walker to cast	N		P3		\$19.60
29445	Apply rigid leg cast	N		P3		\$47.72
29450	Application of leg cast	N		P2	1.0081	\$42.21
29505	Application, long leg splint	N		P3		\$32.67
29520	Application lower leg splint	N		P3		\$28.12
29530	Strapping of hip	N		P3		\$19.32
29540	Strapping of knee	N		P3		\$19.88
29540	Strapping of ankle and/or ft	N		P3		\$15.06

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28665	Treat toe dislocation	Y		A2	12.8746	\$539.10
28666	Treat toe dislocation	Y		A2	21.4573	\$898.48
28675	Repair of toe dislocation	Y		A2	21.4573	\$898.48
28705	Fusion of foot bones	Y		A2	41.2872	\$1,728.82
28715	Fusion of foot bones	Y		A2	67.2085	\$2,814.22
28725	Fusion of foot bones	Y		A2	41.2872	\$1,728.82
28730	Fusion of foot bones	Y		A2	41.2872	\$1,728.82
28735	Fusion of foot bones	Y		A2	41.2872	\$1,728.82
28737	Revision of foot bones	Y		A2	41.784	\$1,749.82
28740	Fusion of foot bones	Y		A2	41.2872	\$1,728.82
28750	Fusion of big toe joint	Y		A2	41.2872	\$1,728.82
28755	Fusion of big toe joint	Y		A2	19.2721	\$806.88
28760	Fusion of big toe joint	Y		A2	41.2872	\$1,728.82
28810	Amputation toe & metatarsal	Y		A2	18.221	\$762.97
28820	Amputation of toe	Y		A2	18.221	\$762.97
28825	Partial amputation of toe	Y		A2	18.221	\$762.97
28860	High energy swt. plantar f	Y		P3		\$141.46
29000	Application of body cast	N		G2	1.0081	\$42.21
29010	Application of body cast	N		P2	2.2441	\$93.97
29015	Application of body cast	N		P2	2.2441	\$93.97
29020	Application of body cast	N		G2	1.0081	\$42.21
29025	Application of body cast	N		P2	1.0081	\$42.21
29035	Application of body cast	N		P2	2.2441	\$93.97
29040	Application of body cast	N		G2	1.0081	\$42.21
29044	Application of body cast	N		P2	2.2441	\$93.97
29046	Application of body cast	N		G2	2.2441	\$93.97
29049	Application of figure eight	N		P3		\$34.66
29055	Application of shoulder cast	N	CH	P3		\$90.90
29058	Application of shoulder cast	N	CH	P3		\$38.06
29065	Application of long arm cast	N		P3		\$38.63

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29836	Elbow arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29837	Elbow arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29838	Elbow arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29840	Wrist arthroscopy	Y		A2	24,381.1	\$1,020.91
29843	Wrist arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29844	Wrist arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29845	Wrist arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29846	Wrist arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29847	Wrist arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29848	Wrist endoscopy/surgery	Y		A2	29,176.4	\$1,219.19
29850	Knee arthroscopy/surgery	Y		A2	25,066.7	\$1,049.62
29851	Knee arthroscopy/surgery	Y		A2	38,626.3	\$1,617.40
29855	Tibial arthroscopy/surgery	Y		A2	38,626.3	\$1,617.40
29856	Tibial arthroscopy/surgery	Y		A2	38,626.3	\$1,617.40
29860	Hip arthroscopy, dx	Y		A2	38,626.3	\$1,617.40
29861	Hip arthroscopy/surgery	Y		A2	38,626.3	\$1,617.40
29863	Hip arthroscopy/surgery	Y		A2	42,676.2	\$1,786.98
29863	Hip arthroscopy/surgery	Y		A2	38,626.3	\$1,617.40
29866	Augrit implant, knee w/scope	Y		G2	46,703.8	\$1,955.63
29870	Knee arthroscopy, dx	Y		A2	24,381.1	\$1,020.91
29871	Knee arthroscopy/drainage	Y		A2	24,381.1	\$1,020.91
29873	Knee arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29874	Knee arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29875	Knee arthroscopy/surgery	Y		A2	25,066.7	\$1,049.62
29876	Knee arthroscopy/surgery	Y		A2	25,066.7	\$1,049.62
29877	Knee arthroscopy/surgery	Y		A2	25,066.7	\$1,049.62
29879	Knee arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29880	Knee arthroscopy/surgery	Y		A2	25,066.7	\$1,049.62
29881	Knee arthroscopy/surgery	Y		A2	25,066.7	\$1,049.62
29882	Knee arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
29550	Strapping of toes	N		P3	15.34	\$15.34
29580	Application of paste boot	N		P3		\$20.74
29581	Apply military compres lwr leg	N	NI	P2	1,008.1	\$42.21
29590	Application of foot splint	N		P3		\$16.76
29700	Removal/revision of cast	N		P3	\$27.55	\$27.55
29705	Removal/revision of cast	N		P3	\$42.33	\$42.33
29710	Removal/revision of cast	N		P3	\$33.24	\$33.24
29715	Removal/revision of cast	N	CH	P3	\$34.94	\$34.94
29720	Repair of body cast	N		P3	\$22.72	\$22.72
29730	Windowing of cast	N		P3	\$30.68	\$30.68
29740	Wedging of cast	N		P3	\$33.52	\$33.52
29750	Wedging of clubfoot cast	N		P3	24,381.1	\$1,020.91
29800	Jaw arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29804	Jaw arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29805	Shoulder arthroscopy, dx	Y		A2	37,940.9	\$1,588.70
29806	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29807	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29819	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29820	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29821	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29822	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29823	Shoulder arthroscopy/surgery	Y		A2	25,563.5	\$1,070.42
29824	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29825	Shoulder arthroscopy/surgery	Y		A2	37,940.9	\$1,588.70
29826	Shoulder arthroscopy/surgery	Y		A2	38,123.3	\$1,638.21
29827	Arthroscopy rotator cuff repr	Y		A2	46,703.8	\$1,955.63
29828	Arthroscopy biceps tenodesis	Y		G2	24,381.1	\$1,020.91
29830	Elbow arthroscopy	Y		A2	24,381.1	\$1,020.91
29834	Elbow arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91
29835	Elbow arthroscopy/surgery	Y		A2	24,381.1	\$1,020.91

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HCPSC Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
30124	Removal of nose lesion	Y		R2	7.2897	\$305.24
30125	Removal of nose lesion	Y		A2	33.3886	\$1,398.08
30130	Excise inferior turbinate	Y		A2	15.2459	\$638.39
30140	Resect inferior turbinate	Y		A2	20.4597	\$858.71
30150	Partial removal of nose	Y		A2	33.7542	\$1,413.39
30160	Removal of nose	Y		A2	34.4396	\$1,442.09
30200	Injection treatment of nose	Y		P3		\$56.24
30210	Nasal sinus therapy	Y		P3		\$71.58
30220	Insert nasal septal button	Y		A2	8.1184	\$339.94
30300	Remove nasal foreign body	N		P2	0.6403	\$26.81
30310	Remove nasal foreign body	Y		A2	14.2349	\$596.06
30320	Remove nasal foreign body	Y		A2	14.8802	\$623.08
30400	Reconstruction of nose	Y		A2	34.4396	\$1,442.09
30420	Reconstruction of nose	Y		A2	34.9366	\$1,462.90
30430	Revision of nose	Y		A2	20.8254	\$872.02
30435	Revision of nose	Y		A2	34.9366	\$1,462.90
30450	Revision of nose	Y		A2	96.5245	\$1,529.39
30460	Revision of nose	Y		A2	36.5245	\$1,529.39
30465	Repair nasal stenosis	Y		A2	38.4895	\$1,611.67
30520	Repair of nasal septum	Y		A2	21.5108	\$900.72
30540	Repair nasal defect	Y		A2	34.9366	\$1,462.90
30545	Repair nasal defect	Y		A2	34.9366	\$1,462.90
30560	Release of nasal adhesions	Y		A2	3.3186	\$136.96
30580	Repair upper jaw fistula	Y		A2	34.4396	\$1,442.09
30600	Repair mouth/nose fistula	Y		A2	34.4396	\$1,442.09
30620	Intranasal reconstruction	Y		A2	36.5245	\$1,529.39
30630	Repair nasal septum defect	Y		A2	23.5956	\$988.02
30801	Ablate inf turbinate, superi	Y		A2	7.3694	\$308.58

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29883	Knee arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29884	Knee arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29885	Knee arthroscopy/surgery	Y		A2	37.9409	\$1,588.70
29886	Knee arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29887	Knee arthroscopy/surgery	Y		A2	66.5231	\$2,765.52
29888	Knee arthroscopy/surgery	Y		A2	66.5231	\$2,765.52
29891	Ankle arthroscopy/surgery	Y		A2	37.9409	\$1,588.70
29892	Ankle arthroscopy/surgery	Y		A2	66.5231	\$2,765.52
29893	Scope, plantar fasciotomy	Y		A2	22.845	\$956.59
29894	Ankle arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29895	Ankle arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29897	Ankle arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29898	Ankle arthroscopy/surgery	Y		A2	24.3811	\$1,020.91
29899	Ankle arthroscopy/surgery	Y		A2	37.9409	\$1,588.70
29900	Mcp joint arthroscopy, dx	Y		A2	24.3811	\$1,020.91
29901	Mcp joint arthroscopy, surg	Y		A2	24.3811	\$1,020.91
29902	Mcp joint arthroscopy, surg	Y		A2	24.3811	\$1,020.91
29904	Subtalar arthro w/ib rmlv	Y		G2	28.6243	\$1,198.59
29905	Subtalar arthro w/ib	Y		G2	28.6243	\$1,198.59
29906	Subtalar arthro w/ib	Y		G2	28.6243	\$1,198.59
29907	Subtalar arthro w/fusion	Y		G2	46.7038	\$1,955.63
30000	Drainage of nose lesion	Y		P3		\$119.02
30020	Drainage of nose lesion	Y		CH		\$115.33
30100	Intranasal biopsy	Y		P3		\$71.30
30110	Removal of nose poly(p/s)	Y		P3		\$110.50
30115	Removal of nose poly(p/s)	Y		A2	14.8602	\$623.08
30117	Removal of intranasal lesion	Y		A2	15.2459	\$638.39
30118	Removal of intranasal lesion	Y		A2	20.8254	\$872.02
30120	Revision of nose	Y		A2	19.8142	\$829.68

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31235	Nasal/sinus endoscopy, dx	Y	A2	17.4415	\$730.33
31237	Nasal/sinus endoscopy, surg	Y	A2	18.0871	\$757.36
31238	Nasal/sinus endoscopy, surg	Y	A2	17.4415	\$730.33
31239	Nasal/sinus endoscopy, surg	Y	A2	24.6018	\$1,030.15
31240	Nasal/sinus endoscopy, surg	Y	A2	18.0871	\$757.36
31254	Revision of ethmoid sinus	Y	A2	23.9164	\$1,001.45
31255	Removal of ethmoid sinus	Y	A2	25.0988	\$1,050.96
31256	Exploration maxillary sinus	Y	A2	23.9164	\$1,001.45
31267	Endoscopy, maxillary sinus	Y	A2	23.9164	\$1,001.45
31276	Sinus endoscopy, surgical	Y	A2	23.9164	\$1,001.45
31288	Nasal/sinus endoscopy, surg	Y	A2	23.9164	\$1,001.45
31300	Removal of larynx lesion	Y	A2	22.0077	\$921.53
31400	Revision of larynx	Y	A2	33.3886	\$1,398.08
31420	Removal of epiglottis	Y	A2	33.3886	\$1,398.08
31500	Insert emergency airway	N	G2	2.349	\$98.36
31502	Change of windpipe airway	N	G2	1.353	\$56.65
31505	Diagnostic laryngoscopy	Y	P2	0.766	\$32.07
31510	Laryngoscopy with biopsy	Y	A2	18.0871	\$757.36
31511	Remove foreign body, larynx	Y	A2	1.8155	\$76.02
31512	Removal of larynx lesion	Y	A2	18.0871	\$757.36
31513	Injection into vocal cord	Y	A2	1.8155	\$76.02
31515	Laryngoscopy for aspiration	Y	A2	17.4415	\$730.33
31520	Dx laryngoscopy, newborn	Y	G2	1.7827	\$73.81
31525	Dx laryngoscopy excl nb	Y	A2	17.4415	\$730.33
31526	Dx laryngoscopy wiper scope	Y	A2	18.0871	\$757.36
31527	Laryngoscopy for treatment	Y	A2	22.9055	\$959.12
31528	Laryngoscopy and dilation	Y	A2	18.0871	\$757.36
31529	Laryngoscopy and dilation	Y	A2	18.0871	\$757.36

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30902	Ablate inf turbinate submuc	Y	A2	14.2349	\$586.06
30901	Control of nosebleed	Y	P3		\$38.63
30903	Control of nosebleed	Y	CH	1.2409	\$51.96
30905	Control of nosebleed	Y	A2	1.2409	\$51.96
30906	Repeat control of nosebleed	Y	A2	1.2409	\$51.96
30915	Ligation, nasal sinus artery	Y	A2	21.613	\$905.00
30920	Ligation, upper jaw artery	Y	A2	21.9786	\$920.31
30930	Thier fx, nasal inf turbinate	Y	A2	15.9313	\$67.09
31000	Irrigation, maxillary sinus	Y	P3	\$91.18	
31002	Exploration, maxillary sinus	Y	R2	20.4597	\$305.24
31030	Exploration, maxillary sinus	Y	A2	33.7542	\$1,413.39
31032	Explore sinus, remove polyps	Y	A2	34.4396	\$1,442.09
31040	Exploration behind upper jaw	Y	R2	23.8828	\$1,000.04
31050	Exploration, sphenoid sinus	Y	A2	33.3886	\$1,398.08
31051	Sphenoid sinus surgery	Y	A2	34.4396	\$1,442.09
31070	Exploration of frontal sinus	Y	A2	20.4597	\$856.71
31075	Exploration of frontal sinus	Y	A2	34.4396	\$1,442.09
31080	Removal of frontal sinus	Y	A2	34.4396	\$1,442.09
31081	Removal of frontal sinus	Y	A2	34.4396	\$1,442.09
31084	Removal of frontal sinus	Y	A2	34.4396	\$1,442.09
31085	Removal of frontal sinus	Y	A2	34.4396	\$1,442.09
31086	Removal of frontal sinus	Y	A2	34.4396	\$1,442.09
31087	Removal of frontal sinus	Y	A2	34.4396	\$1,442.09
31090	Exploration of sinuses	Y	A2	34.4396	\$1,462.90
31200	Removal of ethmoid sinus	Y	A2	33.3886	\$1,398.08
31201	Removal of ethmoid sinus	Y	A2	34.9366	\$1,462.90
31205	Removal of ethmoid sinus	Y	A2	33.7542	\$1,413.39
31231	Nasal endoscopy, dx	Y	P2	1.7627	\$73.81
31233	Nasal/sinus endoscopy, dx	Y	A2	1.8155	\$76.02

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31622	Dx bronchoscope/wash	Y		A2	9,3433	\$391,23
31623	Dx bronchoscope/brush	Y		A2	9,9888	\$418,26
31624	Dx bronchoscope/lavage	Y		A2	9,9888	\$418,26
31625	Bronchoscopy w/biopsy(s)	Y		A2	9,9888	\$418,26
31626	Bronchoscopy w/markers	Y	NI	G2	9,9216	\$415,45
31627	Navigationl bronchoscopy	Y		N1		
31628	Bronchoscopy/lung bx, each	Y		A2	9,9888	\$418,26
31629	Bronchoscopy/needle bx, each	Y		A2	9,9888	\$418,26
31630	Bronchoscopy/dilate/fix repr	Y		A2	21,1332	\$884,91
31631	Bronchoscopy, dilate w/stent	Y		A2	21,1332	\$884,91
31632	Bronchoscopy/lung bx, addl	Y		G2	9,9216	\$415,45
31633	Bronchoscopy/needle bx addl	Y		G2	9,9216	\$415,45
31635	Bronchoscopy w/lf removal	Y		A2	9,9888	\$418,26
31636	Bronchoscopy, bronch stents	Y		A2	21,1332	\$884,91
31637	Bronchoscopy, stent add-on	Y		A2	9,3433	\$391,23
31638	Bronchoscopy, revise stent	Y		A2	21,1332	\$884,91
31640	Bronchoscopy w/tumor excise	Y		A2	21,1332	\$884,91
31641	Bronchoscopy, treat blockage	Y		A2	21,1332	\$884,91
31643	Diag bronchoscope/catheter	Y		A2	9,9888	\$418,26
31645	Bronchoscopy, clear airways	Y		A2	9,3433	\$391,23
31646	Bronchoscopy, reclear airway	Y		A2	9,3433	\$391,23
31656	Bronchoscopy, inj for x-ray	Y		A2	9,3433	\$391,23
31715	Injection for bronchus x-ray	N		N1		
31717	Bronchial brush biopsy	Y		A2	4,516	\$189,10
31720	Clearance of airways	Y		A2	0,5615	\$23,51
31730	Intra, windpipe wire/tube	Y		A2	4,516	\$189,10
31750	Repair of windpipe	Y		A2	34,9866	\$1,462,90
31755	Repair of windpipe	Y		A2	33,3886	\$1,398,08
31820	Closure of windpipe lesion	Y		A2	19,8142	\$829,68
31825	Repair of windpipe defect	Y		A2	20,4597	\$856,71

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31530	Laryngoscopy w/lf removal	Y		A2	18,0871	\$757,36
31531	Laryngoscopy w/lf & op scope	Y		A2	18,4525	\$772,66
31535	Laryngoscopy w/biopsy	Y		A2	18,0871	\$757,36
31536	Laryngoscopy w/bx & op scope	Y		A2	18,4525	\$772,66
31540	Laryngoscopy w/exc of tumor	Y		A2	18,4525	\$772,66
31541	Laryngoscopy w/tum exc + scope	Y		A2	19,1381	\$801,37
31545	Remove vc lesion w/scope	Y		A2	24,6018	\$1,030,15
31546	Remove vc lesion scope/graft	Y		A2	24,6018	\$1,030,15
31560	Laryngoscopy w/arytenoidectomy	Y		A2	25,0988	\$1,050,96
31561	Laryngoscopy, remove cart + scop	Y		A2	25,0988	\$1,050,96
31570	Laryngoscopy w/vc inj	Y		A2	18,0871	\$757,36
31571	Laryngoscopy w/vc inj + scope	Y		A2	23,5507	\$966,14
31575	Diagnostic laryngoscopy	Y		P3	\$51,13	
31576	Laryngoscopy with biopsy	Y		A2	18,0871	\$757,36
31577	Remove foreign body, larynx	Y		A2	4,516	\$189,10
31578	Removal of larynx lesion	Y		A2	23,5507	\$966,14
31579	Diagnostic laryngoscopy	Y		P3	\$90,33	
31580	Revision of larynx	Y		A2	34,9366	\$1,462,90
31582	Revision of larynx	Y		A2	34,9366	\$1,462,90
31588	Revision of larynx	Y		A2	34,9366	\$1,462,90
31590	Reinervate larynx	Y		A2	34,9366	\$1,462,90
31595	Larynx nerve surgery	Y		A2	33,3886	\$1,398,08
31603	Incision of windpipe	Y		A2	7,3694	\$308,58
31605	Incision of windpipe	Y		G2	7,2897	\$305,24
31611	Surgery/speech prosthesis	Y		A2	20,8254	\$872,02
31612	Puncture/clear windpipe	Y		A2	19,8142	\$829,68
31613	Repair windpipe opening	Y		A2	20,4597	\$856,71
31614	Repair windpipe opening	Y		A2	33,3886	\$1,398,08
31615	Visualization of windpipe	Y		A2	7,3694	\$308,58
31620	Endobronchial us add-on	N		N1		

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
33226	Reposition I ventric lead	Y		G2	21.9478	\$919.02
33233	Removal of pacemaker system	Y		A2	19.0084	\$795.94
33234	Removal of pacemaker system	Y		G2	21.9478	\$919.02
33235	Removal pacemaker electrode	Y		G2	21.9478	\$919.02
33241	Remove pulse generator	Y		J8	500.9758	\$20,977.36
33242	Remove pulse generator	Y		G2	21.9478	\$919.02
33249	Eltr/insert pace-defib	Y		J8	632.2356	\$26,473.60
33282	Implant pat-active ht record	Y		J8	111.8317	\$4,662.73
33284	Remove pat-active ht record	N		G2	7.8476	\$326.60
33500	Endoscopic vein harvest	N		N1		
34480	Removal of vein clot	Y		G2	39.1293	\$1,638.46
35188	Repair blood vessel lesion	Y		A2	32.9456	\$1,379.53
35207	Repair blood vessel lesion	Y		A2	32.9456	\$1,379.53
35460	Repair venous blockage	Y	CH	G2	48.4864	\$2,030.27
35473	Repair arterial blockage	Y		G2	48.4864	\$2,030.27
35475	Repair arterial blockage	Y	CH	G2	48.4864	\$2,030.27
35476	Repair venous blockage	Y		G2	48.4864	\$2,030.27
35492	Atherectomy, percutaneous	Y		G2	89.2835	\$3,738.57
35572	Harvest femoropopliteal vein	N		N1		
35761	Exploration of artery/vein	Y		G2	34.0556	\$1,426.01
35875	Removal of clot in graft	Y		A2	36.9952	\$1,549.10
35876	Removal of clot in graft	Y		A2	36.9952	\$1,549.10
36000	Place needle in vein	N		N1		
36002	Pseudoaneurysm injection rt	N		G2	2.2009	\$92.16
36005	Injection ext venography	N		N1		
36010	Place catheter in vein	N		N1		
36011	Place catheter in vein	N		N1		
36012	Place catheter in vein	N		N1		
36013	Place catheter in artery	N		N1		
36014	Place catheter in artery	N		N1		

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31830	Revise windpipe scar	Y		A2	20.4597	\$856.71
32400	Needle biopsy chest lining	Y		A2	8.9382	\$370.08
32405	Biopsy, lung or mediastinum	Y		A2	8.9382	\$370.08
32420	Puncture/clear lung	Y		A2	5.2561	\$220.09
32421	Thoracentesis for aspiration	Y		A2	5.2561	\$220.09
32422	Thoracentesis w/ultra insert	Y		G2	5.3117	\$222.42
32550	Insert pleural cath	Y		G2	29.1413	\$1,220.23
32552	Remove lung catheter	N	NI	G2	1.353	\$56.65
32553	Ins mark thor for rt perq	N	NI	G2	13.1619	\$551.13
32960	Therapeutic pneumothorax	Y		G2	5.3117	\$222.42
32998	Perq rf ablate tx, pul tumor	Y		G2	49.1378	\$2,057.55
33010	Drainage of heart sac	Y		A2	5.2561	\$220.09
33011	Repeat drainage of heart sac	Y		A2	5.2561	\$220.09
33206	Insertion of heart pacemaker	Y		J8	169.4488	\$7,095.33
33207	Insertion of heart pacemaker	Y		J8	169.4488	\$7,095.33
33208	Insertion of heart pacemaker	Y		J8	205.4713	\$8,603.70
33210	Insertion of heart electrode	Y		G2	46.8172	\$1,960.38
33211	Insertion of heart electrode	Y		G2	46.8172	\$1,960.38
33212	Insertion of pulse generator	Y		H8	138.5387	\$5,801.03
33213	Insertion of pulse generator	Y		H8	152.6905	\$6,393.61
33214	Upgrade of pacemaker system	Y		J8	205.4713	\$8,603.70
33215	Reposition pacing-defib lead	Y		G2	21.9478	\$919.02
33216	Insert 1 electrode pm-defib	Y		G2	46.8172	\$1,960.38
33217	Insert 2 electrode pm-defib	Y		G2	46.8172	\$1,960.38
33218	Repair lead pace-defib, one	Y		G2	21.9478	\$919.02
33220	Repair lead pace-defib, dual	Y		G2	21.9478	\$919.02
33222	Revis pocket, pacemaker	Y		A2	14.0668	\$589.02
33223	Revis pocket for defib	Y		A2	14.0668	\$589.02
33224	Insert pacing lead & connect	Y		J8	303.507	\$12,708.75
33225	L ventric pacing lead add-on	Y		J8	303.507	\$12,708.75

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36455	Bl exchange/transfuse non-nb	N		G2	3.2345	\$135.44
36468	Injection(s), spider veins	Y		R2	0.8408	\$35.21
36469	Injection(s), spider veins	Y		R2	0.8408	\$35.21
36470	Injection therapy of vein	Y		P2	0.8408	\$35.21
36471	Injection therapy of vein	Y		P2	0.8408	\$35.21
36475	Endovenous rf, 1st vein	Y		A2	39.958	\$1,673.16
36476	Endovenous rf, vein add-on	Y		A2	26.7139	\$1,118.59
36479	Endovenous laser, 1st vein	Y		A2	26.7139	\$1,118.59
36481	Endovenous laser, vein add-on	Y		A2	26.7139	\$1,118.59
36481	Insertion of catheter, vein	N		N1		
36500	Insertion of catheter, vein	N		N1		
36510	Insertion of catheter, vein	N		N1		
36511	Apheresis wbc	N		G2	11.4253	\$478.41
36512	Apheresis rbc	N		G2	11.4253	\$478.41
36513	Apheresis platelets	N		G2	11.4253	\$478.41
36514	Apheresis plasma	N		G2	11.4253	\$478.41
36515	Apheresis, adsorp/reinfuse	N		P2	31.8778	\$1,334.82
36522	Apheresis, selective	N		P2	31.8778	\$1,334.82
36555	Photopheresis	N		A2	9.914	\$415.13
36556	Insert non-tunneled cv cath	Y		A2	9.914	\$415.13
36557	Insert tunneled cv cath	Y		A2	20.7258	\$867.85
36558	Insert tunneled cv cath	Y		A2	20.7258	\$867.85
36560	Insert tunneled cv cath	Y		A2	24.7544	\$1,036.54
36561	Insert tunneled cv cath	Y		A2	24.7544	\$1,036.54
36563	Insert tunneled cv cath	Y		A2	24.7544	\$1,036.54
36565	Insert tunneled cv cath	Y		A2	24.7544	\$1,036.54
36566	Insert tunneled cv cath	Y		A2	24.7544	\$1,036.54
36568	Insert picc cath	Y		A2	9.914	\$415.13
36569	Insert picc cath	Y		A2	9.914	\$415.13

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36015	Place catheter in artery	N		N1		
36100	Establish access to artery	N		N1		
36120	Establish access to artery	N		N1		
36140	Establish access to artery	N		N1		
36145	Artery to vein shunt	N	CH	D5		
36147	Access av dial grft for eval	Y	NH	P2	2.2917	\$95.96
36148	Access av dial grft for proc	N		N1		
36160	Establish access to aorta	N		N1		
36200	Place catheter in aorta	N		N1		
36215	Place catheter in artery	N		N1		
36216	Place catheter in artery	N		N1		
36217	Place catheter in artery	N		N1		
36218	Place catheter in artery	N		N1		
36245	Place catheter in artery	N		N1		
36246	Place catheter in artery	N		N1		
36247	Place catheter in artery	N		N1		
36248	Place catheter in artery	N		N1		
36260	Insertion of infusion pump	N		N1		
36261	Revision of infusion pump	Y		A2	24.7544	\$1,036.54
36262	Removal of infusion pump	Y		A2	19.0084	\$795.94
36400	Bl draw < 3 yrs fem/jugular	N		N1	18.3631	\$768.92
36405	Bl draw < 3 yrs scalp vein	N		N1		
36406	Bl draw < 3 yrs other vein	N		N1		
36410	Non-routine bl draw > 3 yrs	N		N1		
36416	Capillary blood draw	N		N1		
36420	Vein access cutdown < 1 yr	N		R2	0.222	\$9.30
36425	Vein access cutdown > 1 yr	N	CH	R2	0.222	\$9.30
36430	Blood transfusion service	N		P3		\$25.85
36440	Bl push transfuse, 2 yr or <	N		R2	3.2345	\$135.44
36450	Bl exchange/transfuse, nb	N		R2	3.2345	\$135.44

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36820	AV fusion/forearm vein	Y		A2	32.2599	\$1,350.82
36821	AV fusion direct any site	Y		A2	32.2599	\$1,350.82
36825	Artery-vein autograft	Y		A2	32.9456	\$1,379.53
36830	Artery-vein nonautograft	Y		A2	32.9456	\$1,379.53
36831	Open thrombect av fistula	Y		A2	36.9952	\$1,549.10
36832	Av fistula revision, open	Y		A2	32.9456	\$1,379.53
36833	Av fistula revision	Y		A2	32.9456	\$1,379.53
36834	Repair A-V aneurysm	N	CH	D5		
36835	Artery to vein shunt	Y		A2	25.8262	\$1,081.42
36860	External cannula declotting	Y		A2	2.4464	\$102.44
36861	Cannula declotting	Y		A2	25.1408	\$1,052.72
36870	Percut thrombect av fistula	Y		A2	41.1482	\$1,723.00
37184	Prim art mech thrombectomy	Y		G2	39.1293	\$1,638.46
37185	Prim art m-thrombect add-on	Y		G2	39.1293	\$1,638.46
37186	Sec art m-thrombect add-on	Y		G2	39.1293	\$1,638.46
37187	Venous mech thrombectomy	Y		G2	39.1293	\$1,638.46
37188	Venous m-thrombectomy add-on	Y		G2	39.1293	\$1,638.46
37200	Transcatheter biopsy	Y		G2	29.1216	\$1,219.41
37203	Transcatheter retrieval	Y		G2	29.1216	\$1,219.41
37250	Iv us first vessel add-on	N		N1		
37251	Iv us each add vessel add-on	N		N1		
37500	Endoscopy ligate perit veins	Y		A2	35.2227	\$1,474.88
37607	Ligation of a-v fistula	Y		A2	21.9786	\$920.31
37609	Temporal artery procedure	Y		A2	15.1023	\$632.38
37650	Revision of major vein	Y		A2	21.613	\$905.00
37700	Revises leg vein	Y		A2	21.613	\$905.00
37718	Ligate/strip short leg vein	Y		A2	21.9786	\$920.31
37722	Ligate/strip long leg vein	Y		A2	35.2227	\$1,474.88
37735	Removal of leg veins/lesion	Y		A2	35.2227	\$1,474.88
37760	Ligate leg veins radical	Y		A2	21.9786	\$920.31

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36570	Insert picvad cath	Y		A2	21.0912	\$883.15
36571	Insert picvad cath	Y		A2	21.0912	\$883.15
36575	Repair tunneled cv cath	Y		A2	7.1211	\$298.18
36576	Repair tunneled cv cath	Y		A2	10.5593	\$442.15
36578	Replace tunneled cv cath	Y		A2	20.7258	\$867.85
36580	Replace tunneled cv cath	Y		A2	9.914	\$415.13
36581	Replace tunneled cv cath	Y		A2	20.7258	\$867.85
36582	Replace tunneled cv cath	Y		A2	24.7544	\$1,036.54
36583	Replace tunneled cv cath	Y		A2	24.7544	\$1,036.54
36584	Replace picc cath	Y		A2	9.914	\$415.13
36585	Replace picvad cath	Y		A2	21.0912	\$883.15
36589	Removal tunneled cv cath	Y		A2	6.4758	\$271.16
36590	Removal tunneled cv cath	Y		A2	9.914	\$415.13
36591	Draw blood off venous device	N		N1		
36592	Collect blood from picc	N		N1		
36593	Declot vascular device	Y		P3		\$20.45
36595	Mech remov tunneled cv cath	Y		G2	24.2374	\$1,014.89
36596	Mech remov tunneled cv cath	Y		G2	10.6825	\$447.31
36597	Reposition venous catheter	Y		G2	10.6825	\$447.31
36598	Inj w/fluor, eval cv device	Y		P3		\$63.63
36600	Withdrawal of arterial blood	N		N1		
36620	Insertion catheter, artery	N		N1		
36625	Insertion catheter, artery	N		N1		
36640	Insertion catheter, artery	Y		A2	23.7432	\$994.20
36680	Insert needle, bone cavity	Y		G2	1.4457	\$60.54
36800	Insertion of cannula	Y		A2	25.1408	\$1,052.72
36810	Insertion of cannula	Y		A2	25.1408	\$1,052.72
36818	Av fuse, uppr arm, cephalic	Y		A2	32.2599	\$1,350.82
36819	Av fuse, uppr arm, basilic	Y		A2	32.2599	\$1,350.82

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38740	Remove armpit lymph nodes	Y		A2	37.5311	\$1,571.54
38745	Remove armpit lymph nodes	Y		A2	38.5821	\$1,615.55
38760	Remove groin lymph nodes	Y		A2	20.2092	\$846.22
38790	Inject for lymphatic x-ray	N		N1		
38792	Identify sentinel node	N		N1		
38794	Access thoracic lymph duct	N		N1		
40490	Biopsy of lip	Y		P3		\$57.10
40500	Partial excision of lip	Y		A2	14.8802	\$623.08
40510	Partial excision of lip	Y		A2	20.4597	\$856.71
40520	Partial excision of lip	Y		A2	14.8802	\$623.08
40525	Reconstruct lip with flap	Y		A2	20.4597	\$856.71
40527	Reconstruct lip with flap	Y		A2	20.4597	\$856.71
40650	Partial removal of lip	Y		A2	20.4597	\$856.71
40650	Repair lip	Y		A2	8.1184	\$339.94
40654	Repair lip	Y		A2	8.1184	\$339.94
40654	Repair lip	Y		A2	8.1184	\$339.94
40700	Repair cleft lip/nasal	Y		A2	36.5245	\$1,529.39
40701	Repair cleft lip/nasal	Y		A2	41.1215	\$1,721.88
40702	Repair cleft lip/nasal	Y		R2	36.5245	\$1,529.39
40720	Repair cleft lip/nasal	Y		A2	33.7542	\$1,413.39
40761	Repair cleft lip/nasal	Y		A2	1.3927	\$56.32
40800	Drainage of mouth lesion	Y		P2	8.0147	\$335.60
40801	Drainage of mouth lesion	Y		A2	0.6403	\$26.81
40804	Removal, foreign body, mouth	N		P2		\$144.59
40805	Removal, foreign body, mouth	Y		P3		\$64.48
40806	Incision of lip fold	Y		P3		\$99.14
40810	Excision of mouth lesion	Y		P3		\$102.83
40812	Excise/repair mouth lesion	Y		P3		\$129.82
40814	Excise/repair mouth lesion	Y		A2	14.8802	\$623.08

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37761*	Ligate leg veins open	Y	N1	R2	25.4208	\$1,064.45
37765	Phleb veins - extrem - to 20	Y		R2	25.4208	\$1,064.45
37766	Phleb veins - extrem 20+	Y		R2	25.4208	\$1,064.45
37780	Revision of leg vein	Y		A2	21.9786	\$920.31
37785	Ligate/divide/excise vein	Y		A2	21.9786	\$920.31
37790	Penile venous occlusion	Y		A2	27.9431	\$1,170.06
38200	Injection for spleen x-ray	N		N1		
38204	Bi donor search management	N		N1		
38206	Harvest auto stem cells	N		G2	11.4253	\$478.41
38220	Bone marrow aspiration	Y		P3		\$64.65
38221	Bone marrow biopsy	Y		P3		\$64.65
38230	Bone marrow collection	N		G2	31.8778	\$1,334.82
38241	Bone marrow/stem transplant	N		G2	31.8778	\$1,334.82
38242	Lymphocyte infuse transplant	N		R2	11.4253	\$478.41
38300	Drainage, lymph node lesion	Y		A2	10.9586	\$458.87
38305	Incision of lymph channels	Y		A2	16.472	\$689.73
38500	Biopsy/removal, lymph nodes	Y		A2	20.2092	\$846.22
38505	Needle biopsy, lymph nodes	Y		A2	20.2092	\$846.22
38510	Biopsy/removal, lymph nodes	Y		A2	20.2092	\$846.22
38520	Biopsy/removal, lymph nodes	Y		A2	20.2092	\$846.22
38525	Biopsy/removal, lymph nodes	Y		A2	20.2092	\$846.22
38530	Biopsy/removal, lymph nodes	Y		A2	20.2092	\$846.22
38542	Explore deep node(s), neck	Y		A2	37.5311	\$1,571.54
38550	Removal, neck/arm/pit lesion	Y		A2	20.5746	\$861.52
38555	Removal, neck/arm/pit lesion	Y		A2	21.2602	\$890.23
38570	Laparoscopy, lymph node biop	Y		A2	41.2571	\$1,727.56
38572	Laparoscopy, lymphadenectomy	Y		A2	59.9976	\$2,512.28
38574	Laparoscopy, lymphadenectomy	Y		A2	41.2571	\$1,727.56
38700	Removal of lymph nodes, neck	Y		G2	23.5488	\$966.06

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
41115	Excision of tongue fold	Y		P3	14.2349	\$119.31
41116	Excision of mouth lesion	Y		A2	22.0077	\$596.06
41120	Partial removal of tongue	Y		A2	1.6877	\$921.53
41250	Repair tongue laceration	Y		A2	3.3186	\$70.67
41251	Repair tongue laceration	Y		A2	8.0147	\$138.96
41252	Repair tongue laceration	Y		A2	19.8142	\$335.60
41500	Fixation of tongue	Y		A2	14.2349	\$629.68
41510	Tongue to lip surgery	Y		A2	7.2897	\$596.06
41512	Tongue suspension	Y	CH	G2	8.0147	\$305.24
41520	Reconstruction, tongue fold	Y		G2	23.8828	\$335.60
41530	Tongue base vol reduction	Y		A2	1.5497	\$1,000.04
41800	Drainage of gum lesion	Y		A2		\$64.89
41805	Removal of gum lesion	Y		P3		\$132.37
41806	Removal foreign body, gum	Y		P3		\$159.07
41807	Removal foreign body, jawbone	Y		P3		\$305.24
41820	Excision, gum, each quadrant	Y		R2	7.2897	\$305.24
41821	Excision of gum flap	Y		G2	7.2897	\$305.24
41822	Excision of gum lesion	Y		P3		\$130.95
41823	Excision of gum lesion	Y		P3		\$190.04
41825	Excision of gum lesion	Y		P3		\$104.53
41826	Excision of gum lesion	Y		P3		\$135.78
41827	Excision of gum lesion	Y		A2	20.4597	\$856.71
41828	Excision of gum lesion	Y		P3		\$120.73
41830	Removal of gum tissue	Y		P3		\$170.15
41850	Treatment of gum lesion	Y		R2	16.4437	\$688.55
41870	Gum graft	Y		G2	23.8828	\$1,000.04
41872	Repair gum	Y		P3		\$168.16
41874	Repair tooth socket	Y		P3		\$163.90
42000	Drainage mouth roof lesion	Y		A2	3.3186	\$138.96
42100	Biopsy roof of mouth	Y		P3		\$66.19
42104	Excision lesion, mouth roof	Y		P3		\$99.14

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40816	Excision of mouth lesion	Y		A2	20.4597	\$856.71
40818	Excise oral mucosa for graft	Y		A2	3.3186	\$138.96
40819	Excise lip or cheek fold	Y		A2	7.3694	\$308.58
40820	Treatment of mouth lesion	Y		P3		\$146.29
40830	Repair mouth laceration	Y		G2	3.2767	\$137.21
40831	Repair mouth laceration	Y		A2	7.3694	\$308.58
40840	Reconstruction of mouth	Y		A2	20.4597	\$856.71
40842	Reconstruction of mouth	Y		A2	20.8254	\$872.02
40843	Reconstruction of mouth	Y		A2	20.8254	\$872.02
40844	Reconstruction of mouth	Y		A2	34.9366	\$1,462.90
40845	Reconstruction of mouth	Y		A2	34.9366	\$1,462.90
41000	Drainage of mouth lesion	Y		P3		\$73.86
41005	Drainage of mouth lesion	Y		A2	3.3186	\$138.96
41006	Drainage of mouth lesion	Y		A2	19.8142	\$829.68
41007	Drainage of mouth lesion	Y		A2	14.2349	\$596.06
41008	Drainage of mouth lesion	Y		A2	14.2349	\$596.06
41009	Drainage of mouth lesion	Y		A2	3.3186	\$138.96
41010	Incision of tongue fold	Y		A2	7.3694	\$308.58
41015	Drainage of mouth lesion	Y		A2	3.3186	\$138.96
41016	Drainage of mouth lesion	Y		A2	7.3694	\$308.58
41017	Drainage of mouth lesion	Y		A2	7.3694	\$308.58
41018	Drainage of mouth lesion	Y		G2	23.8828	\$1,000.04
41019	Place needles h&n for r	Y		P3		\$77.55
41105	Biopsy of tongue	Y		P3		\$76.98
41108	Biopsy of floor of mouth	Y		P3		\$71.30
41110	Excision of tongue lesion	Y		P3		\$103.11
41112	Excision of tongue lesion	Y		A2	14.8802	\$623.08
41113	Excision of tongue lesion	Y		A2	14.8802	\$623.08
41114	Excision of tongue lesion	Y		A2	20.4597	\$856.71

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42409	Drainage of salivary cyst	Y		A2	15,2459	\$638.39
42410	Excise parotid gland/flesion	Y		A2	33,7542	\$1,413.39
42415	Excise parotid gland/flesion	Y		A2	36,5245	\$1,529.39
42420	Excise parotid gland/flesion	Y		A2	36,5245	\$1,529.39
42425	Excise parotid gland/flesion	Y		A2	36,5245	\$1,529.39
42440	Excise submaxillary gland	Y		A2	33,7542	\$1,413.39
42450	Excise sublingual gland	Y		A2	20,4597	\$856.71
42500	Repair salivary duct	Y		A2	20,8254	\$872.02
42505	Repair salivary duct	Y		A2	34,4396	\$1,442.09
42507	Parotid duct diversion	Y		A2	33,7542	\$1,413.39
42508	Parotid duct diversion	Y		A2	34,4396	\$1,442.09
42510	Parotid duct diversion	Y		A2	34,4396	\$1,442.09
42550	Injection for salivary x-ray	N		N1		
42600	Closure of salivary fistula	Y		A2	14,2349	\$596.06
42650	Dilation of salivary duct	Y		P3		\$36.64
42660	Dilation of salivary duct	Y		P3		\$43.18
42665	Ligation of salivary duct	Y		A2	23,5966	\$988.02
42700	Drainage of tonsil abscess	Y		A2	3,3186	\$138.96
42720	Drainage of throat abscess	Y		A2	14,2349	\$596.06
42725	Drainage of throat abscess	Y		A2	33,8886	\$1,398.08
42800	Biopsy of throat	Y		P3		\$71.02
42802	Biopsy of throat	Y		A2	14,2349	\$596.06
42804	Biopsy of upper nose/throat	Y		A2	14,2349	\$596.06
42806	Biopsy of upper nose/throat	Y		A2	20,4597	\$856.71
42809	Excise pharynx/flesion	Y		A2	20,4597	\$856.71
42810	Remove pharynx foreign body	N		G2	0,6403	\$26.81
42815	Excision of neck cyst	Y		A2	20,8254	\$872.02
42820	Excision of neck cyst	Y		A2	34,9366	\$1,462.90
42825	Remove tonsils and adenoids	Y		A2	20,8254	\$872.02

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
42106	Excision lesion, mouth roof	Y		P3		\$124.13
42107	Excision lesion, mouth roof	Y		A2	20,4597	\$856.71
42120	Remove palate/lesion	Y		A2	34,4396	\$1,442.09
42140	Excision of uvula	Y		A2	8,0147	\$335.60
42145	Repair palate, pharynx/uvula	Y		A2	22,0077	\$921.53
42160	Treatment mouth roof lesion	Y		P3		\$114.76
42180	Repair palate	Y		A2	3,3186	\$138.96
42182	Repair palate	Y		A2	33,9886	\$1,398.08
42200	Reconstruct cleft palate	Y		A2	34,9366	\$1,462.90
42205	Reconstruct cleft palate	Y		A2	34,9366	\$1,462.90
42210	Reconstruct cleft palate	Y		A2	34,9366	\$1,462.90
42215	Reconstruct cleft palate	Y		A2	36,5245	\$1,529.39
42220	Reconstruct cleft palate	Y	CH	G2	34,9366	\$1,462.90
42225	Reconstruct cleft palate	Y		A2	41,1215	\$1,721.88
42226	Lengthening of palate	Y	CH	G2	34,9366	\$1,462.90
42227	Lengthening of palate	Y		A2	41,1215	\$1,721.88
42235	Repair palate	Y		A2	16,4282	\$687.90
42260	Repair nose to lip fistula	Y		A2	21,5108	\$900.72
42280	Preparation, palate mold	Y		P3		\$65.33
42281	Insertion, palate prosthesis	Y		G2	16,4437	\$688.55
42300	Drainage of salivary gland	Y		A2	14,2349	\$596.06
42305	Drainage of salivary gland	Y		A2	14,8802	\$623.08
42310	Drainage of salivary gland	Y		A2	3,3186	\$138.96
42320	Drainage of salivary gland	Y		A2	3,3186	\$138.96
42330	Removal of salivary stone	Y		P3		\$99.71
42335	Removal of salivary stone	Y		P3		\$165.61
42340	Removal of salivary stone	Y		A2	14,8802	\$623.08
42400	Biopsy of salivary gland	Y		P3		\$55.11
42405	Biopsy of salivary gland	Y		A2	20,4597	\$856.71
42408	Excision of salivary cyst	Y		A2	15,2459	\$638.39

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43226	Esoph endoscopy, dilation	Y	A2	8.1776	\$342.42
43227	Esoph endoscopy, repair	Y	A2	8.8231	\$369.45
43228	Esoph endoscopy, ablation	Y	A2	19.962	\$635.67
43231	Esoph endoscopy w/us exam	Y	A2	8.8231	\$369.45
43234	Esoph endoscopy w/us fn bx	Y	A2	8.8231	\$369.45
43234	Upper GI endoscopy, exam	Y	A2	8.1776	\$342.42
43235	Upper gi endoscopy, diagnosis	Y	A2	8.1776	\$342.42
43236	Upper gi scope w/submuc inj	Y	A2	8.8231	\$369.45
43237	Endoscopic us exam, esoph	Y	A2	8.8231	\$369.45
43238	Upper gi endoscopy w/us fn bx	Y	A2	8.8231	\$369.45
43239	Upper GI endoscopy, biopsy	Y	A2	8.8231	\$369.45
43240	Esoph endoscopy w/drain cyst	Y	A2	8.8231	\$369.45
43241	Upper GI endoscopy with tube	Y	A2	8.8231	\$369.45
43242	Upper gi endoscopy w/us fn bx	Y	A2	8.8231	\$369.45
43243	Upper GI endoscopy & inject	Y	A2	8.8231	\$369.45
43244	Upper GI endoscopy/ligation	Y	A2	8.8231	\$369.45
43245	Upper gi scope dilate strict	Y	A2	8.8231	\$369.45
43246	Place gastrostomy tube	Y	A2	8.8231	\$369.45
43247	Operative upper GI endoscopy	Y	A2	8.8231	\$369.45
43248	Upper gi endoscopy/guide wire	Y	A2	8.8231	\$369.45
43249	Esoph endoscopy, dilation	Y	A2	8.8231	\$369.45
43250	Upper GI endoscopy/tumor	Y	A2	8.8231	\$369.45
43251	Operative upper GI endoscopy	Y	A2	8.8231	\$369.45
43255	Operative upper GI endoscopy	Y	A2	8.8231	\$369.45
43256	Upper gi endoscopy w/sient	Y	A2	21.9213	\$917.91
43257	Upper gi scope w/intrn bxmt	Y	A2	20.3277	\$851.18
43258	Operative upper GI endoscopy	Y	A2	9.1887	\$384.76
43259	Endoscopic ultrasound exam	Y	A2	9.1887	\$384.76
43260	Endo cholangiopancreatograph	Y	A2	18.7713	\$786.01
43261	Endo cholangiopancreatograph	Y	A2	18.7713	\$786.01

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42821	Remove tonsils and adenoids	Y	A2	22.0077	\$921.53
42825	Removal of tonsils	Y	A2	21.5108	\$900.72
42826	Removal of tonsils	Y	A2	21.5108	\$900.72
42831	Removal of adenoids	Y	A2	21.5108	\$900.72
42835	Removal of adenoids	Y	A2	21.5108	\$900.72
42836	Removal of adenoids	Y	A2	21.5108	\$900.72
42860	Excision of tonsil tags	Y	A2	20.8254	\$872.02
42870	Excision of lingual tonsil	Y	A2	20.8254	\$872.02
42890	Partial removal of pharynx	Y	A2	36.5245	\$1,529.39
42892	Revision of pharyngeal walls	Y	A2	36.5245	\$1,529.39
42900	Repair throat wound	Y	A2	7.3694	\$308.58
42950	Reconstruction of throat	Y	A2	20.4597	\$856.71
42955	Surgical opening of throat	Y	A2	20.4597	\$856.71
42960	Control throat bleeding	Y	A2	33.3886	\$1,398.08
42970	Control nose/throat bleeding	Y	P2	1.1023	\$46.16
42972	Control nose/throat bleeding	Y	A2	15.2459	\$638.39
43030	Throat muscle surgery	Y	G2	16.4437	\$688.55
43130	Removal of esophagus pouch	Y	G2	41.1215	\$1,721.88
43200	Esophagus endoscopy	Y	A2	8.1776	\$342.42
43201	Esoph scope w/submucous inj	Y	A2	8.1776	\$342.42
43202	Esophagus endoscopy, biopsy	Y	A2	8.1776	\$342.42
43204	Esoph scope w/sclerosis inj	Y	A2	8.1776	\$342.42
43205	Esophagus endoscopy/ligation	Y	A2	8.1776	\$342.42
43215	Esophagus endoscopy	Y	A2	8.1776	\$342.42
43216	Esophagus endoscopy/lesion	Y	A2	8.1776	\$342.42
43217	Esophagus endoscopy	Y	A2	20.9104	\$875.58
43219	Esophagus endoscopy	Y	A2	20.9104	\$875.58
43220	Esoph endoscopy, dilation	Y	A2	8.1776	\$342.42

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44365	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44366	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44369	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44370	Small bowel endoscopy/stent	Y	A2	A2	26.6566	\$1,116.19
44372	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44373	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44376	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44377	Small bowel endoscopy/biopsy	Y	A2	A2	9.6153	\$402.62
44378	Small bowel endoscopy	Y	A2	A2	9.6153	\$402.62
44379	S bowel endoscopy w/stent	Y	A2	A2	26.6566	\$1,116.19
44380	Small bowel endoscopy	Y	A2	A2	8.97	\$375.60
44381	Small bowel endoscopy	Y	A2	A2	8.97	\$375.60
44383	ileoscopy w/stent	Y	A2	A2	26.6566	\$1,116.19
44385	Endoscopy of bowel pouch	Y	A2	A2	8.4353	\$353.21
44386	Endoscopy, bowel pouch/biops	Y	A2	A2	8.4353	\$353.21
44388	Colonoscopy	Y	A2	A2	8.4353	\$353.21
44389	Colonoscopy with biopsy	Y	A2	A2	8.4353	\$353.21
44390	Colonoscopy for foreign body	Y	A2	A2	8.4353	\$353.21
44391	Colonoscopy for bleeding	Y	A2	A2	8.4353	\$353.21
44392	Colonoscopy & polypectomy	Y	A2	A2	8.4353	\$353.21
44393	Colonoscopy, lesion removal	Y	A2	A2	8.4353	\$353.21
44394	Colonoscopy w/stnare	Y	A2	A2	8.4353	\$353.21
44397	Colonoscopy w/stent	Y	A2	A2	20.9104	\$875.58
44500	Intraop. gastrointestinal tube	Y	G2	G2	6.0982	\$255.35
44701	Intraop colon lavage add-on	N		N1		
45000	Drainage of pelvic abscess	Y		A2	11.9098	\$498.70
45005	Drainage of rectal abscess	Y		A2	12.6748	\$530.73
45020	Drainage of rectal abscess	Y		A2	12.6748	\$530.73
45100	Biopsy of rectum	Y		A2	19.1013	\$799.63
45108	Removal of anorectal lesion	Y		A2	19.7469	\$826.86

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43262	Endo cholangiopancreatograph	Y	A2	A2	18.7713	\$786.01
43263	Endo cholangiopancreatograph	Y	A2	A2	18.7713	\$786.01
43264	Endo cholangiopancreatograph	Y	A2	A2	18.7713	\$786.01
43265	Endo cholangiopancreatograph	Y	A2	A2	18.7713	\$786.01
43268	Endo cholangiopancreatograph	Y	A2	A2	21.5557	\$902.60
43269	Endo cholangiopancreatograph	Y	A2	A2	21.5557	\$902.60
43271	Endo cholangiopancreatograph	Y	A2	A2	18.7713	\$786.01
43272	Endo cholangiopancreatograph	Y	A2	A2	18.7713	\$786.01
43273	Endoscopic pancreatotomy	Y	G2	G2	21.632	\$905.80
43450	Dilate esophagus	Y	A2	A2	6.1801	\$258.78
43453	Dilate esophagus	Y	A2	A2	6.1801	\$258.78
43456	Dilate esophagus	Y	A2	A2	6.194	\$259.36
43458	Dilate esophagus	Y	A2	A2	8.1914	\$343.00
43600	Biopsy of stomach	Y	A2	A2	8.342.42	\$342.42
43653	Laparoscopy, gastrostomy	Y			41.2571	\$1,727.56
43752	Nasal/orogastric w/stent	N	CH	G2	1.2143	\$50.85
43760	Change gastrostomy tube	Y		A2	2.547	\$106.65
43761	Reposition gastrostomy tube	Y		A2	8.1776	\$342.42
43870	Repair stomach opening	Y		A2	22.8955	\$958.70
43886	Revis gastric port, open	Y		G2	4.2464	\$177.81
43887	Remove gastric port, open	Y		G2	22.8955	\$958.70
44100	Biopsy of bowel	Y		A2	8.1776	\$342.42
44312	Revision of ileostomy	Y		A2	19.0736	\$798.67
44340	Revision of colostomy	Y		A2	20.0845	\$841.00
44360	Small bowel endoscopy	Y		A2	9.6153	\$402.62
44361	Small bowel endoscopy/biopsy	Y		A2	9.6153	\$402.62
44363	Small bowel endoscopy	Y		A2	9.6153	\$402.62
44364	Small bowel endoscopy	Y		A2	9.6153	\$402.62

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45365	Surgical colonoscopy	Y		A2	8.4353	\$353.21
45378	Diagnostic colonoscopy	Y		A2	9.0806	\$380.23
45379	Colonoscopy w/ib removal	Y		A2	9.0806	\$380.23
45380	Colonoscopy and biopsy	Y		A2	9.0806	\$380.23
45381	Colonoscopy, submucosal inj	Y		A2	9.0806	\$380.23
45382	Colonoscopy/control bleeding	Y		A2	9.0806	\$380.23
45383	Lesion removal colonoscopy	Y		A2	9.0806	\$380.23
45384	Lesion remove colonoscopy	Y		A2	9.0806	\$380.23
45385	Lesion removal colonoscopy	Y		A2	9.0806	\$380.23
45386	Colonoscopy dilate stricture	Y		A2	9.0806	\$380.23
45387	Colonoscopy w/stent	Y		A2	20.9104	\$875.58
45391	Colonoscopy w/ndoscope us	Y		A2	9.0806	\$380.23
45392	Colonoscopy w/ndoscopic fmb	Y		A2	9.0806	\$380.23
45500	Repair of rectum	Y		A2	19.7469	\$826.86
45505	Repair of rectum	Y		A2	25.6382	\$1,073.55
45520	Treatment of rectal prolapse	Y		P2	0.8408	\$36.21
45541	Correct rectal prolapse	Y	CH	G2	30.7878	\$1,289.18
45560	Repair of rectocele	Y		A2	25.6382	\$1,073.55
45900	Reduction of rectal prolapse	Y		A2	5.632	\$235.83
45905	Dilation of anal sphincter	Y		A2	19.1013	\$799.83
45910	Dilation of rectal narrowing	Y		A2	11.9098	\$498.70
45915	Remove rectal obstruction	Y		A2	18.9819	\$794.83
45990	Surg dx exam, anorectal	Y		A2	20.1125	\$842.17
46020	Placement of seton	Y		A2	20.1125	\$842.17
46030	Removal of rectal marker	Y		A2	5.632	\$235.83
46040	Incision of rectal abscess	Y		A2	20.1125	\$842.17
46045	Incision of rectal abscess	Y		A2	19.7469	\$826.86
46050	Incision of anal abscess	Y		A2	11.9098	\$498.70
46060	Incision of rectal abscess	Y		A2	19.7469	\$826.86
46070	Incision of anal septum	Y		G2	13.5029	\$565.41

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45150	Excision of rectal stricture	Y		A2	19.7469	\$826.86
45160	Excision of rectal lesion	Y		A2	19.7469	\$826.86
45170	Excision of rectal lesion	N	CH	D5		
45171	Exc rect tum transanal part	Y	NI	G2	13.5029	\$565.41
45172	Exc rect tum transanal full	Y	NI	G2	22.9324	\$960.25
45190	Deslucation, rectal tumor	Y		A2	24.8475	\$1,040.44
45300	Proctosigmoidoscopy dx	Y		P3		\$55.96
45303	Proctosigmoidoscopy dilate	Y		P2	8.8447	\$370.35
45305	Proctosigmoidoscopy w/ib	Y		A2	8.5358	\$357.42
45307	Proctosigmoidoscopy fb	Y		A2	18.2767	\$765.30
45308	Proctosigmoidoscopy removal	Y		A2	8.5358	\$357.42
45309	Proctosigmoidoscopy removal	Y		A2	8.5358	\$357.42
45315	Proctosigmoidoscopy removal	Y		A2	8.5358	\$357.42
45320	Proctosigmoidoscopy ablate	Y		A2	18.2767	\$765.30
45321	Proctosigmoidoscopy volvul	Y		A2	18.2767	\$765.30
45327	Proctosigmoidoscopy w/stent	Y		A2	20.9104	\$875.58
45330	Diagnostic sigmoidoscopy	Y		P3		\$70.16
45331	Sigmoidoscopy and biopsy	Y		A2	5.8527	\$245.07
45332	Sigmoidoscopy w/ib removal	Y		A2	5.8527	\$245.07
45333	Sigmoidoscopy & polypectomy	Y		A2	8.5358	\$357.42
45334	Sigmoidoscopy for bleeding	Y		A2	8.5358	\$357.42
45335	Sigmoidoscopy w/submuc inj	Y		A2	5.8527	\$245.07
45337	Sigmoidoscopy & decompress	Y		A2	5.8527	\$245.07
45338	Sigmoidoscopy w/tumr remove	Y		A2	8.5358	\$357.42
45339	Sigmoidoscopy w/labiate tumr	Y		A2	8.5358	\$357.42
45340	Sig w/balloon dilatation	Y		A2	8.5358	\$357.42
45341	Sigmoidoscopy w/ultrasound	Y		A2	8.5358	\$357.42
45342	Sigmoidoscopy w/us guide bx	Y		A2	8.5358	\$357.42
45345	Sigmoidoscopy w/stent	Y		A2	20.9104	\$875.58

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46614	Anoscopy, control bleeding	Y		P3	18.922	\$593.08
46615	Anoscopy	Y		A2	18.922	\$792.32
46700	Repair of anal stricture	Y		A2	20.1125	\$842.17
46706	Repr of anal fistula w/ glue	Y		A2	24.9927	\$1,046.52
46707	Repair anorectal fist w/plug	Y	NI	G2	30.7878	\$1,289.18
46750	Repair of anal sphincter	Y		A2	26.0039	\$1,086.86
46753	Reconstruction of anus	Y		A2	20.1125	\$842.17
46754	Removal of suture from anus	Y		A2	19.7469	\$826.86
46760	Repair of anal sphincter	Y		A2	25.6382	\$1,073.55
46761	Repair of anal sphincter	Y		A2	26.0039	\$1,086.86
46762	Implant artificial sphincter	Y		A2	28.7742	\$1,204.86
46900	Destruction, anal lesion(s)	Y	CH	P3		\$102.83
46910	Destruction, anal lesion(s)	Y		P3		\$110.50
46916	Cryosurgery, anal lesion(s)	Y		P2	1.4745	\$61.74
46917	Laser surgery, anal lesions	Y		A2	17.1605	\$718.56
46922	Excision of anal lesion(s)	Y		A2	17.1605	\$718.56
46924	Destruction, anal lesion(s)	Y		A2	17.1605	\$718.56
46930*	Destroy internal hemorrhoids	Y		P3		\$100.56
46937	Cryotherapy of rectal lesion	N	CH	D5		\$80.96
46938	Cryotherapy of rectal lesion	N	CH	D5		\$80.96
46940	Treatment of anal fissure	Y		P3		\$79.25
46942	Treatment of anal fissure	Y		P3		\$79.25
46945	Remove by ligat int hem grps	Y		P3		\$132.09
46946	Remove by ligat int hem grps	Y		A2	12.0292	\$503.70
46947	Hemorrhoidectomy by stapling	Y		A2	26.7742	\$1,204.86
47000	Needle biopsy, liver	Y		A2	8.8362	\$370.08
47001	Needle biopsy, liver add-on	N		N1		
47382	Percut ablate liver f	Y		G2		\$2,057.55
47500	Injection for liver x-rays	N		N1	49.1378	\$2,057.55
47505	Injection for liver x-rays	N		N1		

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46080	Incision of anal sphincter	Y		A2	20.1125	\$842.17
46083	Incise external hemorrhoid	Y		P2	1.932	\$60.90
46200	Removal of anal fissure	Y		A2	19.7469	\$826.86
46210	Removal of anal crypt	N	CH	D5		
46211	Removal of anal crypts	N	CH	D5		
46220	Excise anal ext tag/papilla	Y		A2	12.0292	\$503.70
46221	Ligation of hemorrhoid(s)	Y		P3		\$105.39
46230	Removal of anal tags	Y		A2	19.1013	\$799.83
46250	Remove ext hem groups = 2	Y		A2	20.1125	\$842.17
46255	Remove in/ext hem 1 group	Y		A2	20.1125	\$842.17
46257	Remove in/ext hem grps & fiss	Y		A2	20.1125	\$842.17
46258	Remove in/ext hem grps w/fistu	Y		A2	20.1125	\$842.17
46260	Remove in/ext hem groups = 2	Y		A2	20.1125	\$842.17
46261	Remove in/ext hem grps & fiss	Y		A2	20.1125	\$842.17
46270	Remove anal fist subq	Y		A2	20.1125	\$842.17
46275	Remove anal fist inter	Y		A2	20.1125	\$842.17
46280	Remove anal fist complex	Y		A2	20.7979	\$870.87
46285	Remove anal fist 2 stage	Y		A2	20.7979	\$870.87
46288	Repair anal fistula	Y		A2	20.7979	\$870.87
46320	Removal of hemorrhoid clot	Y		P3		\$71.02
46500	Injection into hemorrhoid(s)	Y		P3		\$99.42
46505	Chemodestruction anal musc	Y		G2	22.9324	\$960.25
46600	Diagnostic anoscopy	N		P2	0.8403	\$26.81
46604	Anoscopy and dilation	Y	CH	P3		\$343.71
46606	Anoscopy and biopsy	Y		P3		\$113.91
46608	Anoscopy, remove for body	Y		A2	8.5358	\$357.42
46610	Anoscopy, remove lesion	Y		A2	18.2767	\$765.30
46611	Anoscopy	Y		A2	8.5358	\$357.42
46612	Anoscopy, remove lesions	Y		A2	18.2767	\$765.30

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49420	Insert abdom drain, temp	Y		A2	23.758	\$994.82
49421	Insert abdom drain, perm	Y		A2	23.758	\$994.82
49422	Remove perm catheter/catheter	Y		A2	18.3631	\$769.92
49423	Exchange drainage catheter	Y		G2	14.6474	\$613.33
49424	Assess cyst, contrast inject	N		N1		
49426	Revise abdom-venous shunt	Y		A2	22.0538	\$923.46
49427	Injection, abdominal shunt	N		N1		
49429	Removal of shunt	Y		G2	21.9478	\$919.02
49435	Insert subq exten to ip cath	Y	CH	G2	14.6474	\$613.33
49436	Embedded ip cath exit-site	Y	CH	G2	14.6474	\$613.33
49440	Place gastrostomy tube perc	Y		G2	8.3675	\$350.37
49441	Place duod/jej tube perc	Y		G2	8.3675	\$350.37
49442	Place cecostomy tube perc	Y	CH	G2	13.5029	\$565.41
49446	Change g-tube to g-j perc	Y		G2	8.3675	\$350.37
49451	Replace g/c tube perc	Y		G2	6.0982	\$255.35
49452	Replace duod/jej tube perc	Y		G2	6.0982	\$255.35
49462	Replace g-j tube perc	Y		G2	6.0982	\$255.35
49463	Fluoro exam of g/colon tube	Y		N1		
49465	Fluoro exam of g/colon tube	N		N1		
49495	Rpr ing hernia baby, reduc	Y		A2	26.5364	\$1,111.16
49496	Rpr ing hernia, init, reduce	Y		A2	26.5364	\$1,111.16
49500	Rpr ing hernia, init, reduce	Y		A2	26.5364	\$1,111.16
49501	Rpr ing hernia, init blocked	Y		A2	30.5863	\$1,280.74
49505	Rpr /hern init reduc >5 yr	Y		A2	26.5364	\$1,111.16
49507	Rpr /hern init block >5 yr	Y		A2	30.5863	\$1,280.74
49520	Repair ing hernia, reduce	Y		A2	26.6213	\$1,198.46
49521	Repair ing hernia, blocked	Y		A2	30.5863	\$1,280.74
49525	Repair ing hernia, sliding	Y		A2	26.5364	\$1,111.16
49540	Repair lumbar hernia	Y		A2	25.4854	\$1,067.15
49550	Rpr tem hernia, init, reduce	Y		A2	27.0334	\$1,131.97

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47510	Insert catheter, bile duct	Y		A2	24.6063	\$1,030.34
47511	Insert bile duct drain	Y		A2	29.1751	\$1,221.65
47525	Change bile duct catheter	Y		A2	12.8875	\$539.84
47530	Revis/reinsert bile tube	Y		A2	12.8875	\$539.84
47552	Biliary endoscopy thru skin	Y		A2	24.6063	\$1,030.34
47553	Biliary endoscopy thru skin	Y		A2	24.9719	\$1,045.65
47554	Biliary endoscopy thru skin	Y		A2	24.9719	\$1,045.65
47555	Biliary endoscopy thru skin	Y		A2	24.9719	\$1,045.65
47556	Biliary endoscopy thru skin	Y		A2	29.1751	\$1,221.65
47560	Laparoscopy w/cholangio	Y		A2	30.2178	\$1,265.31
47561	Laparoscopic cholecystectomy	Y		G2	44.8118	\$1,876.40
47562	Laparoscopic cholecystectomy	Y		G2	44.8118	\$1,876.40
47564	Laparo cholecystectomy/expl	Y		G2	44.8118	\$1,876.40
47630	Remove bile duct stone	Y		A2	24.9719	\$1,045.65
48102	Needle biopsy, pancreas	Y		A2	8.8382	\$370.08
49080	Puncture, peritoneal cavity	Y		A2	5.2561	\$220.09
49081	Removal of abdominal fluid	Y		A2	5.2561	\$220.09
49180	Biopsy, abdominal mass	Y		A2	8.8382	\$370.08
49250	Excision of umbilicus	Y		A2	23.1049	\$967.47
49320	Diag laparo separate proc	Y		A2	30.2178	\$1,265.31
49321	Laparoscopy, biopsy	Y		A2	30.9032	\$1,294.01
49322	Laparoscopy, aspiration	Y		A2	30.9032	\$1,294.01
49324	Lap insertion perm ip cath	Y		G2	36.4063	\$1,524.44
49325	Lap revision perm ip cath	Y		G2	36.4063	\$1,524.44
49326	Lap w/menotopy add-on	Y		G2	36.4063	\$1,524.44
49400	Air injection into abdomen	N		N1		
49402	Remove foreign body, abdomen	Y		A2	22.0538	\$923.46
49411	Ins mark abd/peel for rt perq	N	NI	P3		\$281.50
49419	Insert abdom cath for chemox	Y		A2	24.1299	\$1,010.39

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50386	Remove stent via transureth	Y	CH	P2	6.8253	\$285.80
50387	Change ext/int ureter stent	Y		G2	14.6474	\$613.33
50389	Remove renal tube w/fluoro	Y		G2	6.8253	\$285.80
50390	Drainage of kidney lesion	Y		A2	8.8382	\$370.08
50391	Instill vx agnt into mal tub	Y	CH	P3	14.1246	\$42.33
50392	Insert kidney drain	Y		A2	20.2152	\$691.44
50393	Insert ureteral tube	Y		A2	20.2152	\$646.47
50394	Injection for kidney x-ray	N		N1		
50395	Create passage to kidney	Y		A2	20.2152	\$846.47
50396	Measure kidney pressure	Y		A2	2.2002	\$92.13
50398	Change kidney tube	Y		A2	12.8875	\$539.64
50551	Kidney endoscopy	Y		A2	7.021	\$293.99
50553	Kidney endoscopy	Y		A2	20.2152	\$846.47
50555	Kidney endoscopy & biopsy	Y		A2	7.021	\$293.99
50557	Kidney endoscopy & treatment	Y		A2	20.2152	\$846.47
50561	Kidney endoscopy & treatment	Y		A2	20.2152	\$846.47
50562	Renal scope w/tumor resect	Y		G2	6.8253	\$285.80
50570	Kidney endoscopy	Y		G2	6.8253	\$285.80
50572	Kidney endoscopy	Y		G2	6.8253	\$285.80
50574	Kidney endoscopy & biopsy	Y		G2	6.8253	\$285.80
50575	Kidney endoscopy	Y		G2	34.6334	\$1,450.20
50576	Kidney endoscopy & treatment	Y		G2	16.2968	\$682.40
50580	Kidney endoscopy & treatment	Y		G2	16.2968	\$682.40
50590	Fragmenting of kidney stone	Y		G2	39.5716	\$1,656.98
50592	Perc r/ ablate renal tumor	Y		G2	49.1378	\$2,057.55
50684	Injection for ureter x-ray	N		N1		
50686	Measure ureter pressure	Y	CH	P3		\$37.78
50688	Change of ureter tube/stent	Y		A2	12.8875	\$539.64
50690	Injection for ureter x-ray	N		N1		
50727	Revise ureter	Y	CH	G2	19.1572	\$602.17

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
49553	Rpr fem hernia, int blocked	Y		A2	30.5863	\$1,280.74
49555	Rerepair fem hernia, reduce	Y		A2	27.0334	\$1,131.97
49557	Rerepair fem hernia, blocked	Y		A2	30.5863	\$1,280.74
49560	Rpr ventral hern int, reduce	Y		A2	26.5364	\$1,111.16
49561	Rpr ventral hern int, block	Y		A2	30.5863	\$1,280.74
49565	Rerepair ventrl hern, reduce	Y		A2	26.5364	\$1,111.16
49566	Rerepair ventrl hern, block	Y		A2	30.5863	\$1,280.74
49568	Hernia repair w/mesh	Y		A2	28.6213	\$1,198.46
49570	Rpr epigastric hern, reduce	Y		A2	26.5364	\$1,111.16
49572	Rpr epigastric hern, blocked	Y		A2	30.5863	\$1,280.74
49580	Rpr umbil hern, reduc < 5 yr	Y		A2	26.5364	\$1,111.16
49582	Rpr umbil hern, block < 5 yr	Y		A2	30.5863	\$1,280.74
49585	Rpr umbil hern, reduc > 5 yr	Y		A2	26.5364	\$1,111.16
49587	Rpr umbil hern, block > 5 yr	Y		A2	30.5863	\$1,280.74
49590	Repair spigelian hernia	Y		A2	25.851	\$1,082.46
49600	Repair umbilical lesion	Y		A2	26.5364	\$1,111.16
49650	Lap lng hernia repair int	Y		A2	37.2073	\$1,557.98
49651	Lap lng hernia repair recur	Y		A2	39.2921	\$1,645.28
49652	Lap vent/abd hernia repair	Y		G2	69.7991	\$2,922.70
49653	Lap vent/abd hern proe comp	Y		G2	69.7991	\$2,922.70
49654	Lap inc hernia repair	Y		G2	69.7991	\$2,922.70
49655	Lap inc hern repair comp	Y		G2	69.7991	\$2,922.70
49656	Lap inc hernia repair recur	Y		G2	69.7991	\$2,922.70
49657	Lap inc hern recur comp	Y		G2	69.7991	\$2,922.70
50080	Removal of kidney stone	Y	CH	G2	44.6588	\$1,870.00
50081	Removal of kidney stone	Y	CH	G2	44.6588	\$1,870.00
50200	Renal biopsy perq	Y		A2	8.8382	\$370.08
50382	Change ureter stent, percut	Y		G2	24.4172	\$1,022.42
50384	Remove ureter stent, percut	Y		G2	16.2968	\$682.40
50385	Change stent via transureth	Y		G2	24.4172	\$1,022.42

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51702	Insert temp bladder cath	N		P2	0.6403	\$26.81
51703	Insert bladder cath, complex	Y		P2	1.0484	\$43.90
51705	Change of bladder tube	Y	CH	P3		\$58.81
51710	Change of bladder tube	Y		A2	6.4758	\$271.16
51715	Endoscopic injection/implant	Y		A2	25.4269	\$1,064.70
51720	Treatment of bladder lesion	Y		P3		\$46.45
51725	Simple cystometrogram	Y	CH	P3		\$106.52
51726	Complex cystometrogram	Y		A2	3.3645	\$140.88
51727	Cystometrogram w/ulp	Y	NI	P2	2.8906	\$121.04
51728	Cystometrogram w/vp&ulp	Y	NI	P2	2.8906	\$121.04
51729	Cystometrogram w/vp&ulp	Y	NI	P2	2.8906	\$121.04
51736	Urine flow measurement	Y		P3		\$17.04
51741	Electro-uroflowmetry, first	Y	CH	P3		\$19.88
51772	Urethra pressure profile	Y		D5		\$43.90
51784	Ana/urinary muscle study	Y		A2	1.8313	\$76.68
51785	Ana/urinary muscle study	Y		P2	1.0484	\$43.90
51792	Urinary reflex study	Y		D5		\$74.71
51795	Urine voiding pressure study	Y	CH	P3		\$14.49
51797	Intraabdominal pressure test	Y	CH	P3		\$946.47
51798	Us urine capacity measure	Y		A2	20.2152	\$1,578.79
51880	Repair of bladder opening	Y		A2	37.7042	\$293.99
51992	Laparo sling operation	Y		A2	7.021	\$607.28
52000	Cystoscopy	Y		A2	14.5029	\$607.28
52001	Cystoscopy, removal of clots	Y		A2	20.8605	\$873.49
52005	Cystoscopy & ureter catheter	Y		A2	20.8605	\$873.49
52007	Cystoscopy and biopsy	Y		A2	7.3993	\$309.83
52010	Cystoscopy & duct catheter	Y		A2	20.8605	\$873.49
52204	Cystoscopy w/biopsy(s)	Y		A2	20.8605	\$873.49
52214	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52224	Cystoscopy and treatment	Y		A2	20.8605	\$873.49

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
50947	Laparo new ureter/bladder	Y		A2	41.2571	\$1,727.56
50948	Laparo new ureter/bladder	Y		A2	41.2571	\$1,727.56
50951	Endoscopy of ureter	Y		A2	7.021	\$293.99
50953	Endoscopy of ureter	Y		A2	7.021	\$293.99
50955	Ureter endoscopy & biopsy	Y		A2	20.2152	\$846.47
50957	Ureter endoscopy & treatment	Y		A2	20.2152	\$846.47
50961	Ureter endoscopy & treatment	Y		A2	20.2152	\$846.47
50970	Ureter endoscopy	Y		A2	7.021	\$293.99
50972	Ureter endoscopy & catheter	Y		A2	7.021	\$293.99
50974	Ureter endoscopy & biopsy	Y		A2	14.1246	\$591.44
50976	Ureter endoscopy & treatment	Y		A2	14.1246	\$591.44
50980	Ureter endoscopy & treatment	Y		A2	20.2152	\$846.47
51020	Incise & treat bladder	Y		A2	21.9115	\$917.50
51030	Incise & treat bladder	Y		A2	21.9115	\$917.50
51040	Incise & drain bladder	Y		A2	21.9115	\$917.50
51045	Incise bladder/obtain ureter	Y		A2	7.3993	\$309.83
51055	Removal of bladder stone	Y		A2	21.9115	\$917.50
51065	Remove ureter calculus	Y		A2	21.9115	\$917.50
51080	Drainage of bladder abscess	Y		A2	15.8264	\$662.70
51100	Drain bladder by needle	Y		P3		\$25.00
51101	Drain bladder by trocar/cath	Y		P2	1.0484	\$43.90
51102	Drain bl w/cath insertion	Y		A2	16.2699	\$681.27
51500	Removal of bladder cyst	Y		A2	26.5364	\$1,111.16
51520	Removal of bladder lesion	Y		A2	21.9115	\$917.50
51535	Repair of ureter lesion	Y	CH	G2	24.4172	\$1,022.42
51600	Injection for bladder x-ray	N		N1		
51605	Preparation for bladder xray	N		N1		
51610	Injection for bladder x-ray	N		N1		
51700	Irrigation of bladder	Y		P3		\$41.47
51701	Insert bladder catheter	N		P2	0.6403	\$26.81

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52343	Cysto wifrenal stricture bx	Y		A2	21.2261	\$888.80
52344	Cystouretero, stricture bx	Y		A2	21.2261	\$888.80
52345	Cystouretero w/np stricture	Y		A2	21.2261	\$888.80
52346	Cystouretero wifrenal strict	Y		A2	21.2261	\$888.80
52351	Cystouretero & or Pyeloscope	Y		A2	21.9115	\$917.50
52352	Cystouretero wistone remove	Y		A2	29.5735	\$1,238.33
52353	Cystouretero wifnithiopsy	Y		A2	21.9115	\$917.50
52354	Cystouretero wibiopsy	Y		A2	21.9115	\$917.50
52355	Cystouretero wilexice tumor	Y		A2	21.2261	\$888.80
52400	Cystouretero wicongen repr	Y		A2	21.2261	\$888.80
52402	Cystourethro cut ejacul duct	Y		A2	21.2261	\$888.80
52450	Incision of prostate	Y		A2	21.2261	\$888.80
52500	Revision of bladder neck	Y		A2	21.2261	\$888.80
52601	Prostatectomy (TURP)	Y		A2	29.5735	\$1,238.33
52630	Remove prostate regrowth	Y		A2	28.5224	\$1,194.32
52640	Relieve bladder contracture	Y		A2	20.8605	\$873.49
52647	Laser surgery of prostate	Y		A2	41.1423	\$1,722.75
52648	Laser surgery of prostate	Y		A2	41.1423	\$1,722.75
52700	Drainage of prostate abscess	Y		A2	20.8605	\$873.49
53000	Incision of urethra	Y		A2	16.4949	\$690.69
53010	Incision of urethra	Y		A2	16.4949	\$690.69
53020	Incision of urethra	Y		R2	19.4568	\$814.71
53040	Drainage of urethra abscess	Y		A2	17.1402	\$717.71
53060	Drainage of urethra abscess	Y		P3		\$58.52
53085	Drainage of urinary leakage	Y		G2	17.5058	\$735.02
53200	Biopsy of urethra	Y		A2	19.4568	\$814.71
53210	Removal of urethra	Y		A2	16.4949	\$690.69
53215	Removal of urethra	Y		A2	26.6093	\$1,114.21
53215	Removal of urethra	Y		A2	18.6882	\$782.53

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52234	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52235	Cystoscopy and treatment	Y		A2	21.2261	\$888.80
52240	Cystoscopy and treatment	Y		A2	21.2261	\$888.80
52250	Cystoscopy and radiotracer	Y		A2	21.9115	\$917.50
52260	Cystoscopy and treatment	Y		A2	14.7699	\$618.46
52265	Cystoscopy and treatment	Y	CH	P3	\$235.77	
52270	Cystoscopy & revise urethra	Y		A2	14.7699	\$618.46
52275	Cystoscopy & revise urethra	Y		A2	20.8605	\$873.49
52276	Cystoscopy and treatment	Y		A2	21.2261	\$888.80
52277	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52281	Cystoscopy and treatment	Y		A2	14.7699	\$618.46
52282	Cystoscopy, implant stent	Y		A2	33.6233	\$1,407.91
52283	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52285	Cystoscopy and treatment	Y		A2	14.7699	\$618.46
52290	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52300	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52301	Cystoscopy and treatment	Y		A2	21.2261	\$888.80
52305	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52310	Cystoscopy and treatment	Y		A2	14.5029	\$607.28
52315	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52317	Cystoscopy and treatment	Y		A2	20.2152	\$846.47
52318	Remove bladder stone	Y		A2	20.8605	\$873.49
52320	Cystoscopy and treatment	Y		A2	22.4085	\$938.31
52325	Cystoscopy, stone removal	Y		A2	21.9115	\$917.50
52327	Cystoscopy, inject material	Y		A2	28.5224	\$1,194.32
52330	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52332	Cystoscopy and treatment	Y		A2	20.8605	\$873.49
52334	Create passage to kidney	Y		A2	21.2261	\$888.80
52341	Cysto w/ureter stricture bx	Y		A2	21.2261	\$888.80
52342	Cysto w/np stricture bx	Y		A2	21.2261	\$888.80

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53600	Dilate urethra stricture	Y		P3		\$31.81
53601	Dilate urethra stricture	Y	CH	A2		\$36.93
53605	Dilate urethra stricture	Y		A2	14.7689	\$618.46
53620	Dilate urethra stricture	Y		P3		\$48.57
53621	Dilate urethra stricture	Y		P3		\$51.13
53680	Dilation of urethra	Y	CH	P3		\$36.08
53681	Dilation of urethra	Y	CH	P3		\$35.22
53685	Dilation of urethra	Y		A2	16.4949	\$940.69
53850	Prostatic microwave thermox	Y	CH	P3		\$1,568.87
53852	Prostatic rf thermox	Y	CH	P3		\$1,479.96
53855	Insert prost urethral stent	Y	NI	P2	1.932	\$60.90
54000	Slitting of prepuce	Y		A2	17.1402	\$717.71
54001	Slitting of prepuce	Y		A2	17.1402	\$717.71
54015	Drain penis lesion	Y		A2	17.523	\$733.74
54050	Destruction, penis lesion(s)	Y		P2	0.8408	\$35.21
54055	Destruction, penis lesion(s)	Y		P2		\$53.40
54056	Cryosurgery, penis lesion(s)	Y		P2	0.8408	\$35.21
54057	Laser surg, penis lesion(s)	Y		A2	17.1605	\$718.56
54060	Excision of penis lesion(s)	Y		A2	17.1605	\$718.56
54065	Destruction, penis lesion(s)	Y		A2	17.1605	\$718.56
54100	Biopsy of penis	Y		A2	14.457	\$605.36
54105	Biopsy of penis	Y		A2	16.6836	\$782.94
54110	Treatment of penis lesion	Y		A2	27.5774	\$1,154.75
54111	Treat penis lesion, graft	Y		A2	27.5774	\$1,154.75
54112	Treat penis lesion, graft	Y		A2	27.5774	\$1,154.75
54115	Treatment of penis lesion	Y		A2	15.8264	\$662.70
54120	Partial removal of penis	Y		A2	27.5774	\$1,154.75
54150	Circumcision w/regional block	Y		A2	18.6086	\$779.20
54160	Circumcision, neonate	Y		A2	19.2542	\$806.23
54161	Circum 28 days or older	Y		A2	19.2542	\$806.23

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53220	Treatment of urethra lesion	Y		A2	25.0613	\$1,049.39
53230	Removal of urethra lesion	Y		A2	25.0613	\$1,049.39
53235	Removal of urethra lesion	Y		A2	17.5058	\$733.02
53240	Surgery for urethra pouch	Y		A2	25.0613	\$1,049.39
53250	Removal of urethra gland	Y		A2	17.1402	\$717.71
53260	Treatment of urethra lesion	Y		A2	17.1402	\$717.71
53265	Treatment of urethra lesion	Y		A2	17.1402	\$717.71
53270	Removal of urethra gland	Y		A2	17.1402	\$717.71
53275	Repair of urethra defect	Y		A2	25.4269	\$1,064.70
53400	Revis urethra, stage 1	Y		A2	25.0613	\$1,049.39
53405	Revis urethra, stage 2	Y		A2	25.0613	\$1,049.39
53410	Reconstitution of urethra	Y		A2	25.0613	\$1,049.39
53420	Reconstruct urethra, stage 1	Y		A2	25.4289	\$1,064.70
53425	Reconstruct urethra, stage 2	Y		A2	25.0613	\$1,049.39
53430	Reconstitution of urethra	Y		A2	25.0613	\$1,049.39
53431	Reconstruct urethra/bladder	Y		A2	25.0613	\$1,049.39
53440	Male sling procedure	N		H8	124.7551	\$5,223.87
53442	Remove/revise male sling	Y		A2	24.416	\$1,022.37
53444	Insert tandem cuff	N		H8	124.7551	\$5,223.87
53445	Insert urolves nck sphincter	N		H8	223.2126	\$9,346.58
53446	Remove uro sphincter	Y		A2	24.416	\$1,022.37
53447	Remove/replace ur sphincter	N		H8	223.2126	\$9,346.58
53449	Repair uro sphincter	Y		A2	24.416	\$1,022.37
53450	Revision of urethra	Y		A2	24.416	\$1,022.37
53460	Revision of urethra	Y		A2	16.4949	\$690.69
53502	Repair of urethra injury	Y		A2	17.1402	\$717.71
53505	Repair of urethra injury	Y		A2	25.0613	\$1,049.39
53510	Repair of urethra injury	Y		A2	17.1402	\$717.71
53515	Repair of urethra injury	Y		A2	25.0613	\$1,049.39
53520	Repair of urethra defect	Y		A2	25.0613	\$1,049.39

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54405	Insert multi-comp penis pros	N		H8	224.2235	\$9,388.91
54406	Remove multi-comp penis pros	Y		A2	27.9431	\$1,170.06
54408	Repair multi-comp penis pros	Y		A2	27.9431	\$1,170.06
54410	Remove/replace penis prosth	N		H8	224.2235	\$9,388.91
54415	Remove self-contd penis pros	Y		A2	27.9431	\$1,170.06
54416	Remw/rep penis contain pros	Y		H8	224.2235	\$9,388.91
54420	Revision of penis	Y		A2	26.6285	\$1,198.76
54436	Revision of penis	Y		A2	26.6285	\$1,198.76
54440	Repair of penis	Y		A2	28.6285	\$1,198.76
54450	Preputial stretching	Y		A2	3.3645	\$140.88
54500	Blopsy of testis	Y		A2	13.0239	\$545.35
54505	Blopsy of testis	Y		A2	18.6086	\$779.20
54512	Excise lesion testis	Y		A2	19.2542	\$806.23
54520	Removal of testis	Y		A2	19.6196	\$821.53
54522	Orchiectomy, partial	Y		A2	19.6196	\$821.53
54530	Removal of testis	Y		A2	26.5364	\$1,111.16
54560	Exploration for testis	Y		A2	26.5364	\$1,111.16
54560	Exploration for testis	Y		G2	22.2756	\$932.75
54620	Reduce testis torsion	Y		A2	20.3052	\$850.24
54620	Suspension of testis	Y		A2	19.6196	\$821.53
54640	Suspension of testis	Y		A2	26.5364	\$1,111.16
54660	Revision of testis	Y		A2	19.2542	\$806.23
54670	Repair testis injury	Y		A2	19.6196	\$821.53
54680	Relocation of testis(es)	Y		A2	19.6196	\$821.53
54690	Laparoscopy, orchiectomy	Y		A2	41.2571	\$1,727.56
54692	Laparoscopy, orchiopexy	Y		G2	69.7991	\$2,922.70
54700	Drainage of scrotum	Y		A2	19.2542	\$806.23
54800	Blopsy of epididymis	Y		A2	4.0262	\$168.59
54830	Remove epididymis lesion	Y		A2	19.6196	\$821.53
54840	Remove epididymis lesion	Y		A2	20.3052	\$850.24

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HCPSC Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
54162	Lysis penit circumcison	Y		A2	19.2542	\$806.23
54163	Repair of circumcision	Y		A2	19.2542	\$806.23
54164	Frenulotomy of penis	Y		A2	19.2542	\$806.23
54200	Treatment of penis lesion	Y		P3		\$53.97
54205	Treatment of penis lesion	Y		A2	28.6285	\$1,198.76
54220	Treatment of penis lesion	Y		A2	2.2002	\$92.13
54230	Prepare penis study	N		N1		
54231	Dynamic cavernosometry	Y		P3		\$51.13
54235	Penile injection	Y		P3		\$36.36
54240	Penis study	Y		P3		\$25.85
54250	Penis study	Y		P3		\$9.09
54300	Revision of penis	Y		A2	27.9431	\$1,170.06
54304	Revision of penis	Y		A2	27.9431	\$1,170.06
54308	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54312	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54316	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54318	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54322	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54324	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54326	Reconstruction of urethra	Y		A2	27.9431	\$1,170.06
54328	Revisse penis/urethra	Y		A2	27.9431	\$1,170.06
54340	Secondary urethral surgery	Y		A2	27.9431	\$1,170.06
54344	Secondary urethral surgery	Y		A2	27.9431	\$1,170.06
54348	Secondary urethral surgery	Y		A2	27.9431	\$1,170.06
54352	Reconstruct urethra/penis	Y		A2	27.9431	\$1,170.06
54360	Penis plastic surgery	Y		A2	27.9431	\$1,170.06
54380	Repair penis	Y		A2	27.9431	\$1,170.06
54385	Repair penis	Y		A2	27.9431	\$1,170.06
54400	Insert semi-rigid prosthesis	N		H8	125.1205	\$5,239.17
54401	Insert self-contd prosthesis	N		H8	224.2235	\$9,388.91

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55706	Prostate saturation sampling	Y	G2	11.893	\$498.00
55720	Drainage of prostate abscess	Y	A2	20.2152	\$848.47
55725	Drainage of prostate abscess	Y	A2	20.8605	\$873.49
55860	Surgical exposure, prostate	Y	G2	19.1572	\$802.17
55870	Electrocoagulation	Y	P3		\$63.06
55873	Cryoblate prostate	Y	H8	146.626	\$6,139.67
55875	Transperineal needle biopsy, pros	N	A2	33.6293	\$1,407.91
55876	Place rt device/marker, pros	N	P3		\$57.66
55920	Place needles pelvic for rt	Y	G2	26.0084	\$1,089.05
56405	I & D of vulva/perineum	Y	P3		\$35.51
56420	Drainage of gland abscess	Y	P3		\$48.57
56440	Surgery for vulva lesion	Y	A2	16.9305	\$706.93
56441	Lysis of labial lesion(s)	Y	A2	16.285	\$681.90
56442	Hymenotomy	Y	A2	16.285	\$681.90
56501	Destroy, vulva lesions, sim	Y	P3		\$48.57
56515	Destroy, vulva lesions, compl	Y	A2	18.1714	\$760.89
56605	Biopsy of vulva/perineum	Y	P3		\$26.12
56606	Biopsy of vulva/perineum	Y	P3		\$11.65
56620	Partial removal of vulva	Y	A2	18.4785	\$773.75
56625	Complete removal of vulva	Y	A2	20.0664	\$840.24
56700	Partial removal of hymen	Y	A2	16.285	\$681.90
56740	Remove vagina gland lesion	Y	A2	17.2961	\$724.24
56800	Repair of vagina	Y	A2	17.2961	\$724.24
56805	Repair clitoris	Y	G2	19.1772	\$803.01
56810	Repair of perineum	Y	A2	18.4785	\$773.75
56820	Exam/biopsy of vulva w/scope	Y	P3		\$36.08
57000	Exploration of vagina	Y	CH		\$46.59
57010	Drainage of pelvic abscess	Y	A2	16.285	\$681.90
57020	Drainage of pelvic fluid	Y	A2	16.9305	\$706.93
57020	Drainage of pelvic fluid	Y	A2	7.2135	\$302.05

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54860	Removal of epididymis	Y	A2	19.6196	\$821.53
54861	Removal of epididymis	Y	A2	20.3052	\$850.24
54865	Explore epididymis	Y	A2	18.6086	\$779.20
54900	Fusion of spermatic ducts	Y	A2	20.3052	\$850.24
54901	Fusion of spermatic ducts	Y	A2	20.3052	\$850.24
55000	Drainage of hydrocele	Y	P3		\$52.55
55040	Removal of hydrocele	Y	A2	25.851	\$1,082.46
55041	Removal of hydroceles	Y	A2	27.0394	\$1,131.97
55060	Repair of hydrocele	Y	A2	20.3052	\$850.24
55100	Drainage of scrotum abscess	Y	A2	10.9586	\$458.87
55110	Explore scrotum	Y	A2	19.2542	\$806.23
55120	Removal of scrotum lesion	Y	A2	19.2542	\$806.23
55150	Removal of scrotum	Y	A2	18.6086	\$779.20
55175	Revision of scrotum	Y	A2	18.6086	\$779.20
55180	Revision of scrotum	Y	A2	19.2542	\$806.23
55200	Revision of sperm duct	Y	A2	19.2542	\$806.23
55250	Removal of sperm duct(s)	Y	A2	19.2542	\$806.23
55300	Prepare, sperm duct x-ray	N	N1		
55400	Repair of sperm duct	Y	A2	18.6086	\$779.20
55450	Ligation of sperm duct	Y	P3		\$166.74
55500	Removal of hydrocele	Y	A2	19.6196	\$821.53
55520	Removal of sperm cord lesion	Y	A2	20.3052	\$850.24
55530	Revise spermatic cord veins	Y	A2	20.3052	\$850.24
55535	Revise spermatic cord veins	Y	A2	26.5364	\$1,111.16
55540	Revise hernia & sperm veins	Y	A2	27.0394	\$1,131.97
55550	Laparoscopic spermatic vein	Y	A2	41.2571	\$1,727.56
55600	Incise sperm duct pouch	Y	P2	22.2756	\$932.75
55680	Remove sperm duct lesion	Y	A2	18.6086	\$779.20
55700	Biopsy of prostate	Y	A2	10.8951	\$456.21
55705	Biopsy of prostate	Y	A2	10.8951	\$456.21

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57400	Dilation of vagina	Y		A2	16.9305	\$708.93
57410	Pubic examination	Y		A2	16.9305	\$708.93
57415	Remove vaginal foreign body	Y		A2	16.9305	\$708.93
57420	Exam of vagina w/scope	Y		P3		\$37.21
57421	Exam/biopsy of vag w/scope	Y		P3		\$48.57
57426	Revise prosth vag graft lap	Y	NI	G2	19.1772	\$803.01
57462	Exam of cervix w/scope	Y		P3		\$35.22
57464	Bx/curett of cervix w/scope	Y		P3		\$43.46
57455	Biopsy of cervix w/scope	Y		P3		\$45.45
57456	Endocerv curettage w/scope	Y		P3		\$44.03
57460	Bx of cervix w/scope, loop	Y		P3		\$130.95
57461	Conz of cervix w/scope, loop	Y		P3		\$138.76
57500	Biopsy of cervix	Y		P3		\$60.22
57505	Endocervical curettage	Y		P3		\$39.20
57510	Cauterization of cervix	Y		P3		\$40.05
57511	Cryocautery of cervix	Y	CH	P3		\$48.57
57513	Laser surgery of cervix	Y		A2	16.9305	\$708.93
57522	Conization of cervix	Y		A2	16.9305	\$708.93
57530	Removal of residual cervix	Y		A2	28.2177	\$1,181.56
57550	Removal of residual cervix	Y		A2	28.2177	\$1,181.56
57556	Remove cervix, repair bowel	Y		A2	36.3764	\$1,523.19
57558	D&C of cervical stump	Y		A2	17.2961	\$724.24
57700	Revision of cervix	Y		A2	16.285	\$681.90
57720	Revision of cervix	Y		A2	17.2961	\$724.24
57800	Dilation of cervical canal	Y		P3		\$21.02
58100	Bx done w/colposcopy add-on	Y		P3		\$34.94
58120	Dilation and curettage	Y		N1		
58145	Myomectomy vag method	Y		A2	29.4001	\$1,231.07

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57022	I & d vaginal hematoma, pp	Y	CH	R2	12.0732	\$505.62
57023	I & d vag hematoma, non-ob	Y		A2	15.8264	\$662.70
57061	Destroy vag lesions, simple	Y		P3		\$44.60
57065	Destroy vag lesions, complex	Y		A2	16.285	\$681.90
57100	Biopsy of vagina	Y		P3		\$28.69
57105	Biopsy of vagina	Y		A2	16.9305	\$708.93
57130	Remove vagina lesion	Y		A2	16.9305	\$708.93
57135	Remove vagina lesion	Y		A2	16.9305	\$708.93
57150	Treat vagina infection	Y		P3		\$19.88
57155	Insert uteri tandem/s/ovoids	Y		A2	7.2135	\$302.05
57160	Insert pessary/other device	Y		P3		\$29.83
57170	Filing of diaphragm/cap	Y		P2	0.1263	\$5.29
57180	Treat vaginal bleeding	Y		A2	2.1133	\$88.49
57200	Repair of vagina	Y		A2	16.285	\$681.90
57210	Repair vagina/perineum	Y		A2	16.9305	\$708.93
57220	Revision of urethra	Y		A2	35.194	\$1,473.68
57230	Repair of urethral lesion	Y		A2	28.2177	\$1,181.56
57240	Repair bladder & vagina	Y		A2	29.4001	\$1,231.07
57250	Repair rectum & vagina	Y		A2	29.4001	\$1,231.07
57260	Repair of vagina	Y		A2	29.4001	\$1,231.07
57265	Extensive repair of vagina	Y		A2	37.9643	\$1,589.68
57267	Insert mesh/pelvic fir addon	Y		A2	30.988	\$1,297.56
57268	Repair of bowel bulge	Y		A2	28.2177	\$1,181.56
57287	Revise/remove sling repair	Y		G2	33.7396	\$1,412.78
57288	Repair bladder defect	Y		A2	36.3764	\$1,523.19
57289	Repair bladder & vagina	Y		A2	29.4001	\$1,231.07
57291	Construction of vagina	Y		A2	29.4001	\$1,231.07
57295	Revise vag graft via vagina	Y	CH	G2	19.1772	\$803.01
57300	Repair rectum-vagina fistula	Y		A2	28.2177	\$1,181.56
57320	Repair bladder-vagina lesion	Y		G2	33.7396	\$1,412.78

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58673	Laparoscopy, salpingostomy	Y		A2	37.7042	\$1,578.79
58800	Drainage of ovarian cyst(s)	Y		A2	17.2961	\$724.24
58805	Drainage of ovarian cyst(s)	Y		G2	33.7396	\$1,412.78
58820	Drain ovary abscess, open	Y		A2	28.2177	\$1,181.56
58900	Blopsy of ovar(s)	Y		A2	17.2961	\$724.24
58970	Retrieval of oocyte	Y		A2	3.8502	\$161.22
58974	Transfer of embryo	Y		A2	3.8502	\$161.22
58976	Transfer of embryo	Y		A2	3.8502	\$161.22
59001	Amniocentesis, diagnostic	Y		P3		\$51.98
59001	Amniocentesis, therapeutic	Y		R2	6.5007	\$272.20
59012	Fetal cord puncture, prenatal	Y		G2	3.2609	\$136.54
59016	Chorion biopsy	Y		P3		\$43.18
59020	Fetal contract stress test	Y		P3		\$22.44
59025	Fetal non-stress test	Y		P3		\$11.93
59070	Transabdom amniocentesis w/ius	Y		G2	1.4616	\$61.20
59072	Umbilical cord occlud w/ius	Y		G2	3.2609	\$136.54
59076	Fetal shunt placement, w/ius	Y		G2		\$136.54
59100	Remove uterus lesion	Y		R2	33.7396	\$1,412.78
59150	Treat ectopic pregnancy	Y		G2	44.8118	\$1,876.40
59151	Treat ectopic pregnancy	Y		G2	44.8118	\$1,876.40
59160	D & c after delivery	Y		A2	17.2961	\$724.24
59200	Insert cervical dilator	Y		P3		\$28.41
59300	Episiotomy or vaginal repair	Y		P3		\$62.78
59320	Revision of cervix	Y		A2	16.285	\$681.90
59412	Antepartum manipulation	Y		G2	19.1772	\$803.01
59414	Deliver placenta	Y		G2	19.1772	\$803.01
59820	Care of miscarriage	Y		A2	18.4785	\$773.75
59821	Treatment of miscarriage	Y		A2	18.4785	\$773.75
59840	Abortion	Y		A2	19.4785	\$773.75

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58301	Remove intrauterine device	Y		P3		\$32.38
58321	Artificial insemination	Y		P3		\$30.39
58322	Artificial insemination	Y		P3		\$30.96
58323	Sperm washing	Y		P3		\$6.53
58340	Catheter for hysteroscopy	N		N1		
58345	Reopen fallopian tube	Y		R2	19.1772	\$803.01
58346	Insert heyman uteri capsule	Y		A2	16.9305	\$708.93
58350	Reopen fallopian tube	Y		A2	28.2177	\$1,181.56
58353	Endometrial ablate, thermal	Y		A2	30.998	\$1,297.56
58356	Endometrial cryoablation	Y	CH	P3		\$1,302.70
58545	Laparoscopic myomectomy	Y		A2	34.9531	\$1,463.59
58546	Laparo-myomectomy, complex	Y		A2	41.2571	\$1,727.66
58550	Laparo-assist vag hysterectomy	Y		A2	59.9976	\$2,512.28
58552	Laparo-vag hyst incl t/o	Y		G2	44.8118	\$1,876.40
58555	Hysteroscopy, dx, sep proc.	Y		A2	18.1238	\$758.44
58558	Hysteroscopy, biopsy	Y		A2	19.1238	\$800.77
58559	Hysteroscopy, lysis	Y		A2	18.7584	\$785.47
58560	Hysteroscopy, resect septum	Y		A2	29.6193	\$1,240.25
58561	Hysteroscopy, remove myoma	Y		A2	29.6193	\$1,240.25
58562	Hysteroscopy, remove fb	Y		A2	19.1238	\$800.77
58563	Hysteroscopy, ablation	Y		A2	34.3546	\$1,438.53
58565	Hysteroscopy, sterilization	Y		A2	39.9293	\$1,671.96
58800	Division of fallopian tube	Y		G2	33.7396	\$1,412.78
58615	Occlude fallopian tube(s)	Y		G2	19.1772	\$803.01
58660	Laparoscopy, lysis	Y		A2	37.7042	\$1,578.79
58661	Laparoscopy, remove adnexa	Y		A2	37.7042	\$1,578.79
58662	Laparoscopy, excise lesions	Y		A2	36.5219	\$1,529.28
58670	Laparoscopy, tubal cautery	Y		A2	36.5219	\$1,529.28
58671	Laparoscopy, tubal block	Y		A2	36.5219	\$1,529.28
58672	Laparoscopy, fibrioplasty	Y		A2	37.7042	\$1,578.79

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61886	Implant neurostim arrays	N		H8	416.231	\$17,428.84
61888	Revise/remove neuroceiver	Y		A2	22.4689	\$940.84
62160	Neuroendoscopy add-on	N		N1		
62194	Replace/irrigate catheter	Y		A2	7.0685	\$295.98
62225	Replace/irrigate catheter	Y		A2	12.8875	\$539.64
62230	Replace/revise brain shunt	Y		A2	31.9867	\$1,339.38
62252	Cst shunt reprogram	N		P3		\$39.48
62263	Epidural lysis mult sessions	Y		A2	7.0685	\$295.98
62264	Epidural lysis on single day	Y		A2	11.405	\$477.56
62267	Interdiscal perq aspir, dx	Y		G2	4.4	\$184.24
62268	Drain spinal cord cyst	Y		A2	6.2164	\$260.30
62269	Needle biopsy, spinal cord	Y		A2	8.982	\$370.08
62270	Spinal fluid tap, diagnostic	Y		A2	3.4648	\$145.08
62272	Drain cerebro spinal fluid	Y		A2	4.5729	\$191.48
62280	Inject epidural patch	Y		A2	7.0685	\$295.98
62281	Treat spinal cord lesion	Y		A2	7.0685	\$295.98
62282	Treat spinal canal lesion	Y		A2	7.0685	\$295.98
62284	Injection for myelogram	N		N1		
62287	Percutaneous disectomy	Y		A2	34.3981	\$1,440.35
62290	Inject for spine disk x-ray	N		N1		
62291	Inject for spine disk x-ray	N		N1		
62292	Injection into disk lesion	Y		R2	6.8884	\$288.44
62294	Injection into spinal artery	Y		A2	6.2164	\$260.30
62310	Inject spine c/t	Y		A2	7.0685	\$295.98
62311	Inject spine l/s (cd)	Y		A2	7.0685	\$295.98
62319	Inject spine w/cath l/s (cd)	Y		A2	7.0685	\$295.98
62350	Implant spinal canal cath	Y		A2	31.9867	\$1,339.38
62355	Remove spinal canal catheter	Y		A2	12.0502	\$504.58

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59841	Abortion	Y		A2	18.4785	\$773.75
59866	Abortion (mtp)	Y		G2	3.2609	\$136.54
59870	Evacuate mole of uterus	Y		A2	18.4785	\$773.75
59871	Remove cerclage suture	Y		A2	18.4785	\$773.75
60000	Drain thyroid/tongue cyst	Y		A2	7.3634	\$308.58
60100	Biopsy of thyroid	Y		P3		\$38.63
60200	Remove thyroid lesion	Y		A2	37.5311	\$1,571.54
60210	Partial thyroid excision	Y	CH	G2	46.645	\$1,953.17
60212	Partial thyroid excision	Y	CH	G2	46.645	\$1,953.17
60220	Partial removal of thyroid	Y	CH	G2	46.645	\$1,953.17
60225	Partial removal of thyroid	Y	CH	G2	46.645	\$1,953.17
60280	Remove thyroid duct lesion	Y		A2	38.5821	\$1,615.55
60281	Remove thyroid duct lesion	Y		A2	38.5821	\$1,615.55
60300	Aspir/inj thyroid cyst	Y		P3		\$51.98
61000	Remove cranial cavity fluid	Y		R2	6.8884	\$288.44
61001	Remove cranial cavity fluid	Y		R2	6.8884	\$288.44
61020	Remove brain cavity fluid	Y		A2	6.2164	\$260.30
61050	Injection into brain canal	Y		A2	6.2164	\$260.30
61055	Injection into brain canal	Y		A2	6.2164	\$260.30
61070	Brain canal shunt procedure	Y		A2	5.6237	\$235.48
61215	Insert brain-fluid device	Y		A2	32.3524	\$1,354.89
61330	Decompress eye socket	Y		G2	41.1215	\$1,721.88
61334	Explore orbit/remove object	Y		G2	41.1215	\$1,721.88
61770	Hoist skull for treatment	Y	CH	G2	35.6664	\$1,493.46
61790	Treat trigeminal nerve	Y		A2	16.3451	\$684.42
61791	Treat trigeminal tract	Y		A2	11.5129	\$482.08
61795	Brain surgery using computer	N		N1		
61880	Revise/remove neuroelectrode	Y		G2	18.7878	\$786.70
61885	Instr/reco neurostim 1 array	N		H8	307.5302	\$12,877.21

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64418	N block inj, suprascapular	Y		P3	3.4648	\$59,94
64420	N block inj, intercost, stng	Y		A2	7.0685	\$145.08
64421	N block inj, intercost, mit	Y		A2	7.0685	\$295.98
64425	N block inj, ilio-ingu/hypogl	Y		P3	5.9604	\$42.61
64430	N block inj, pudendal	Y		A2	5.9604	\$249.58
64435	N block inj, paracervical	Y		P3		\$59.65
64445	N block inj, sciatic, stng	Y		P3		\$54.26
64446	N blk inj, sciatic, cont inf	Y		G2	6.8884	\$286.44
64447	N block inj fem, single	Y		R2	3.5609	\$149.11
64448	N block inj fem, cont inf	Y		G2	6.8884	\$286.44
64449	N block inj, lumbar plexus	Y		G2	6.8884	\$286.44
64450	N block, other peripheral	Y		P3		\$37.50
64455*	N block inj, plantar digit	Y		P3		\$15.62
64470	Inj paravertebral c/t	N	CH	D5		
64472	Inj paravertebral c/t add-on	N	CH	D5		
64475	Inj paravertebral l/s	N	CH	D5		
64476	Inj paravertebral l/s add-on	N	CH	D5		
64479	Inj foramen epidural c/t	Y		A2	7.0685	\$295.98
64480	Inj foramen epidural add-on	Y		A2	4.5729	\$191.48
64484	Inj foramen epidural l/s	Y		A2	7.0685	\$295.98
64484	Inj foramen epidural add-on	Y		A2	4.5729	\$191.48
64490	Inj paravert f/jnt c/t 1 lev	Y	NI	G2	6.8884	\$286.44
64491	Inj paravert f/jnt c/t 2 lev	Y	NI	G2	2.4451	\$102.38
64492	Inj paravert f/jnt c/t 3 lev	Y	NI	G2	2.4451	\$102.38
64493	Inj paravert f/jnt l/s 1 lev	Y	NI	G2	6.8884	\$286.44
64494	Inj paravert f/jnt l/s 2 lev	Y	NI	G2	2.4451	\$102.38
64495	Inj paravert f/jnt l/s 3 lev	Y	NI	G2	2.4451	\$102.38
64508	N block, sphenopalatine gangl	Y		P3		\$35.95
64510	N block, carotid sinus s/p	Y		P3		\$74.71
64510	N block, stellate ganglion	Y		A2	7.0685	\$295.98

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62360	Insert spine infusion device	Y		A2	31.9867	\$1,339.38
62361	Implant spine infusion pump	Y		H8	291.6404	\$12,211.86
62362	Implant spine infusion pump	Y		H8	291.6404	\$12,211.86
62365	Remove spine infusion device	Y		A2	29.2974	\$1,226.77
62367	Analyze spine infusion pump	N		P3		\$14.77
62368	Analyze spine infusion pump	N		P3		\$19.32
63600	Remove spinal cord lesion	Y		A2	15.9795	\$669.11
63610	Stimulation of spinal cord	Y		A2	15.3342	\$642.09
63615	Remove lesion of spinal cord	Y		R2	17.9094	\$749.92
63650	Implant neuroelectrodes	N		H8	83.4896	\$3,495.96
63655	Implant neuroelectrodes	N		J8	118.6891	\$4,969.87
63660	Reviser/remove neuroelectrode	N	CH	D5		
63661	Remove spine eltrd perq array	Y	NI	G2	18.7878	\$786.70
63662	Remove spine eltrd plate	Y	NI	G2	18.7878	\$786.70
63663	Reviser spine eltrd perq array	Y	NI	G2	18.7878	\$786.70
63664	Reviser spine eltrd plate	Y	NI	G2	18.7878	\$786.70
63685	Instr/redc spine n generator	N		H8	307.5302	\$12,877.21
63688	Reviser/remove neurorecoiler	Y		A2	22.4689	\$940.84
63744	Revision of spinal shunt	Y		A2	32.3524	\$1,354.69
63746	Removal of spinal shunt	Y		A2	12.0502	\$504.58
64400	N block inj, trigeminal	Y		P3		\$46.59
64402	N block inj, facial	Y		P3		\$44.03
64405	N block inj, occipital	Y		P3		\$38.35
64408	N block inj, vagus	Y		P3		\$46.02
64410	N block inj, phrenic	Y		A2	7.0685	\$295.98
64412	N block inj, spinal accessor	Y		P3		\$69.03
64413	N block inj, cervical plexus	Y		P3		\$42.89
64415	N block inj, brachial plexus	Y		A2	3.4648	\$145.08
64416	N block cont infuse, b plex	Y		G2	6.8884	\$286.44
64417	N block inj, axillary	Y		A2	3.4648	\$145.08

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64650	Chemodermiv eccrine glands	Y		P3		\$28.69
64653	Chemodermiv eccrine glands	Y		P3		\$31.53
64680	Injection treatment of nerve	Y		A2	7.3995	\$309.84
64681	Injection treatment of nerve	Y		A2	12.0502	\$304.98
64702	Revise finger/toe nerve	Y		A2	15.3342	\$642.09
64704	Revise hand/foot nerve	Y		A2	15.3342	\$642.09
64708	Revise arm/leg nerve	Y		A2	15.9795	\$669.11
64712	Revision of sciatic nerve	Y		A2	15.9795	\$669.11
64713	Revision of arm nerve(s)	Y		A2	15.9795	\$669.11
64714	Revise low back nerve(s)	Y		A2	15.9795	\$669.11
64716	Revision of cranial nerve	Y		A2	16.3451	\$684.42
64718	Revise ulnar nerve at elbow	Y		A2	15.9795	\$669.11
64719	Revise ulnar nerve at wrist	Y		A2	15.9795	\$669.11
64721	Carpal tunnel surgery	Y		A2	15.9795	\$669.11
64722	Relieve pressure on nerve(s)	Y		A2	15.3342	\$642.09
64726	Release foot/toe nerve	Y		A2	15.3342	\$642.09
64727	Internal nerve revision	Y		A2	15.9795	\$669.11
64732	Incision of brow nerve	Y		A2	15.9795	\$669.11
64734	Incision of cheek nerve	Y		A2	15.9795	\$669.11
64736	Incision of chin nerve	Y		A2	15.9795	\$669.11
64738	Incision of jaw nerve	Y		A2	15.9795	\$669.11
64740	Incision of tongue nerve	Y		A2	15.9795	\$669.11
64742	Incision of facial nerve	Y		A2	15.9795	\$669.11
64744	Incise nerve, back of head	Y		A2	15.9795	\$669.11
64746	Incise diaphragm nerve	Y		A2	15.9795	\$669.11
64761	Incision of pelvis nerve	Y		G2	17.9094	\$749.92
64763	Incise hip/high nerve	Y		G2	17.9094	\$749.92
64766	Incise hip/high nerve	Y		G2	35.8684	\$1,493.46
64771	Sever cranial nerve	Y		A2	15.9795	\$669.11
64772	Incision of spinal nerve	Y		A2	15.9795	\$669.11

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64517	N block inj, hypogas plex	Y		A2	5.9604	\$249.58
64520	N block, lumbar/thoracic	Y		A2	7.0685	\$295.98
64530	N block inj, celiac plexus	Y		A2	7.0685	\$295.98
64553	Implant neuroelectrodes	N		H8	82.8441	\$3,468.93
64555	Implant neuroelectrodes	N		J8	87.5994	\$3,668.05
64561	Implant neuroelectrodes	N		H8	87.5994	\$3,668.05
64565	Implant neuroelectrodes	N		H8	83.8562	\$3,511.27
64566	Implant neuroelectrodes	N		J8	87.5994	\$3,668.05
64573	Implant neuroelectrodes	N		H8	218.1981	\$9,136.61
64575	Implant neuroelectrodes	N		H8	113.0528	\$4,733.86
64577	Implant neuroelectrodes	N		H8	113.0528	\$4,733.86
64580	Implant neuroelectrodes	N		H8	113.0528	\$4,733.86
64581	Implant neuroelectrodes	N		H8	114.064	\$4,776.20
64585	Revise/remove neuroelectrode	Y		A2	15.9929	\$669.67
64590	Instr/reco pr/gastr stimul	N		H8	307.5302	\$12,877.21
64595	Revise/rmv pr/gastr stimul	Y		A2	22.4689	\$940.84
64600	Injection treatment of nerve	Y		A2	11.405	\$477.56
64605	Injection treatment of nerve	Y		A2	15.3342	\$642.09
64610	Injection treatment of nerve	Y		A2	15.3342	\$642.09
64612	Destroy nerve, face muscle	Y		P3		\$54.82
64613	Destroy nerve, neck muscle	Y		P3		\$51.98
64614	Destroy nerve, extrem muscle	Y		P3		\$59.08
64620	Injection treatment of nerve	Y		A2	7.0685	\$295.98
64622	Destr paravertebral nerve l/s	Y		A2	11.405	\$477.56
64623	Destr paravertebral n add-on	Y		A2	7.0685	\$295.98
64626	Destr paravertebral nerve c/t	Y		A2	7.0685	\$295.98
64627	Destr paravertebral n add-on	Y		A2	3.7361	\$156.44
64630	Injection treatment of nerve	Y		A2	7.1765	\$300.50
64632*	N block inj, common digit	Y		P3		\$28.41
64640	Injection treatment of nerve	Y		P3		\$60.67

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64864	Repair of facial nerve	Y		A2	29.6628	\$1,242.07
64865	Repair of facial nerve	Y		A2	30.3484	\$1,270.78
64870	Fusion of facial/other nerve	Y		A2	30.3484	\$1,270.78
64872	Subsequent repair of nerve	Y		A2	29.2974	\$1,226.77
64874	Repair & revise nerve add-on	Y		A2	29.6628	\$1,242.07
64876	Repair nerve/shorten bone	Y		A2	29.2974	\$1,242.07
64885	Nerve graft, head or neck	Y		A2	29.2974	\$1,226.77
64886	Nerve graft, head or neck	Y		A2	29.2974	\$1,226.77
64890	Nerve graft, hand or foot	Y		A2	29.2974	\$1,226.77
64891	Nerve graft, hand or foot	Y		A2	29.2974	\$1,226.77
64892	Nerve graft, arm or leg	Y		A2	29.2974	\$1,226.77
64893	Nerve graft, arm or leg	Y		A2	29.2974	\$1,226.77
64895	Nerve graft, hand or foot	Y		A2	29.6628	\$1,242.07
64896	Nerve graft, hand or foot	Y		A2	29.6628	\$1,242.07
64897	Nerve graft, arm or leg	Y		A2	29.6628	\$1,242.07
64898	Nerve graft, arm or leg	Y		A2	29.6628	\$1,242.07
64901	Nerve graft add-on	Y		A2	29.2974	\$1,226.77
64902	Nerve graft add-on	Y		A2	29.2974	\$1,226.77
64905	Nerve pedicle transfer	Y		A2	29.2974	\$1,226.77
64907	Nerve pedicle transfer	Y		A2	26.6519	\$1,199.74
64910	Nerve repair w/allograft	Y		G2	35.6664	\$1,483.46
65091	Revise eye	Y		A2	30.9049	\$1,294.08
65093	Revise eye with implant	Y		A2	30.9049	\$1,294.08
65101	Removal of eye	Y		A2	30.9049	\$1,294.08
65103	Remove eye/insert implant	Y		A2	30.9049	\$1,294.08
65105	Remove eyelid/attach implant	Y		A2	31.5903	\$1,322.78
65110	Removal of eye	Y		A2	32.0873	\$1,343.59
65112	Remove eye/revise socket	Y		A2	33.6752	\$1,410.08
65114	Remove eye/revise socket	Y		A2	33.6752	\$1,410.08
65125	Revise ocular implant	Y		G2	25.6774	\$1,075.19

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64774	Remove skin nerve lesion	Y		A2	15.9795	\$669.11
64776	Remove digit nerve lesion	Y		A2	16.3451	\$684.42
64778	Digit nerve surgery add-on	Y		A2	15.9795	\$669.11
64782	Remove limb nerve lesion	Y		A2	16.3451	\$684.42
64783	Limb nerve surgery add-on	Y		A2	15.9795	\$669.11
64784	Remove nerve lesion	Y		A2	16.3451	\$684.42
64786	Remove sciatic nerve lesion	Y		A2	29.6628	\$1,242.07
64787	Implant nerve end	Y		A2	15.9795	\$669.11
64788	Remove skin nerve lesion	Y		A2	16.3451	\$684.42
64790	Removal of nerve lesion	Y		A2	16.3451	\$684.42
64792	Removal of nerve lesion	Y		A2	29.6628	\$1,242.07
64795	Biopsy of nerve	Y		A2	15.9795	\$669.11
64802	Remove sympathetic nerves	Y		A2	15.9795	\$669.11
64820	Remove sympathetic nerves	Y		G2	17.9094	\$749.92
64821	Remove sympathetic nerves	Y		A2	23.6595	\$959.07
64822	Remove sympathetic nerves	Y		G2	27.0149	\$1,131.19
64823	Remove sympathetic nerves	Y		G2	27.0149	\$1,131.19
64831	Repair of digit nerve	Y		A2	30.3484	\$1,270.78
64832	Repair nerve add-on	Y		A2	28.6519	\$1,199.74
64834	Repair of hand or foot nerve	Y		A2	29.2974	\$1,226.77
64835	Repair of hand or foot nerve	Y		A2	29.6628	\$1,242.07
64836	Repair of hand or foot nerve	Y		A2	29.6628	\$1,242.07
64837	Repair nerve add-on	Y		A2	28.6519	\$1,199.74
64840	Repair of leg nerve	Y		A2	29.2974	\$1,226.77
64856	Repair/transpose nerve	Y		A2	29.2974	\$1,226.77
64857	Repair arm/leg nerve	Y		A2	29.2974	\$1,226.77
64858	Repair sciatic nerve	Y		A2	29.2974	\$1,226.77
64859	Nerve surgery	Y		A2	28.6519	\$1,199.74
64861	Repair of arm nerves	Y		A2	29.6628	\$1,242.07
64862	Repair of low back nerves	Y		A2	29.6628	\$1,242.07

Note 1: The Medicare program payment is 80 percent of the total payment amount and beneficiary coinsurance is 20 percent of the total payment amount, except for screening flexible sigmoidoscopies and screening colonoscopies for which the program payment is 75 percent and the beneficiary coinsurance is 25 percent.

Note 2: Payment indicators for "office-based" procedures (P2, P3) are based on a comparison of the final rates according to the ASC standard ratesetting methodology and the MPFS. Under current law, MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
65730	Corneal transplant	Y		A2	32.6184	\$1,365.83
65750	Corneal transplant	Y		A2	32.6184	\$1,365.83
65755	Corneal transplant	Y		A2	32.6184	\$1,365.83
65756	Corneal trnspl, endothelial	Y		G2	35.9134	\$1,503.80
65757	Prep corneal endo allograft	N		H1		
65770	Revise cornea with implant	Y		H8	134.3348	\$5,625.00
65772	Correction of astigmatism	Y		A2	15.2571	\$638.86
65773	Correction of astigmatism	Y		A2	15.2571	\$638.86
65780	Ocular reconst, transplant	Y		A2	31.0305	\$1,299.34
65781	Ocular reconst, transplant	Y		A2	31.0305	\$1,299.34
65782	Ocular reconst, transplant	Y		A2	31.0305	\$1,299.34
65800	Drainage of eye	Y		A2	13.5608	\$567.83
65805	Drainage of eye	Y		A2	13.5608	\$567.83
65810	Drainage of eye	Y		A2	20.4222	\$855.14
65815	Drainage of eye	Y		A2	20.0566	\$839.83
65820	Relieve inner eye pressure	Y		A2	5.1362	\$215.07
65850	Incision of eye	Y		A2	21.1076	\$883.84
65855	Laser surgery of eye	Y		P3		\$115.33
65860	Incise inner eye adhesions	Y		P3		\$107.38
65865	Incise inner eye adhesions	Y		A2	13.5608	\$567.83
65870	Incise inner eye adhesions	Y		A2	21.1076	\$883.84
65875	Incise inner eye adhesions	Y		A2	21.1076	\$883.84
65880	Incise inner eye adhesions	Y		A2	15.2571	\$638.86
65900	Remove eye lesion	Y		A2	15.7541	\$659.67
65920	Remove implant of eye	Y		A2	23.1925	\$971.14
65930	Remove blood clot from eye	Y		A2	21.6046	\$904.65
66020	Injection treatment of eye	Y		A2	13.5608	\$567.83
66130	Remove eye lesion	Y		A2	5.1362	\$215.07
66150	Glaucoma surgery	Y		A2	21.1076	\$883.84

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
65130	Insert ocular implant	Y		A2	22.1711	\$928.37
65135	Insert ocular implant	Y		A2	21.8055	\$913.06
65140	Attach ocular implant	Y		A2	30.9049	\$1,294.08
65150	Revise ocular implant	Y		A2	21.8055	\$913.06
65155	Reinsert ocular implant	Y		A2	30.9049	\$1,294.08
65175	Removal of ocular implant	Y		A2	14.9309	\$625.20
65205	Remove foreign body from eye	N		P3		\$17.90
65210	Remove foreign body from eye	N		P3		\$23.01
65220	Remove foreign body from eye	N		G2	0.9139	\$38.27
65225	Remove foreign body from eye	N		P3		\$25.00
65235	Remove foreign body from eye	Y		A2	14.2061	\$594.85
65260	Remove foreign body from eye	Y		A2	7.1103	\$297.73
65265	Remove foreign body from eye	Y		A2	18.4618	\$773.05
65270	Repair of eye wound	Y		A2	15.5764	\$652.23
65275	Repair of eye wound	Y		A2	20.0566	\$839.83
65275	Repair of eye wound	Y		A2	21.1076	\$883.84
65280	Repair of eye wound	Y		A2	18.4618	\$773.05
65285	Repair of eye wound	Y		A2	32.274	\$1,351.41
65285	Repair of eye wound	Y		P2	4.3122	\$180.56
65290	Repair of eye socket wound	Y		A2	19.7528	\$827.11
65400	Removal of eye lesion	Y		A2	13.5608	\$567.83
65410	Biopsy of cornea	Y		A2	14.2061	\$594.85
65420	Removal of eye lesion	Y		A2	14.2061	\$594.85
65425	Removal of eye lesion	Y		A2	21.6046	\$904.65
65430	Corneal smear	N		P3		\$35.79
65435	Curette/treat cornea	Y		P3		\$27.55
65436	Curette/treat cornea	Y		P3		\$122.43
65450	Treatment of corneal lesion	N		G2	1.8727	\$78.42
65600	Revision of cornea	Y		P3		\$142.31
65710	Corneal transplant	Y		A2	32.6184	\$1,365.83

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66830	Removal of lens lesion	Y		A2	5.3646	\$224.63
66840	Removal of lens material	Y		A2	14.9416	\$625.65
66850	Removal of lens material	Y		A2	27.4454	\$1,149.22
66852	Removal of lens material	Y		A2	25.3605	\$1,061.92
66920	Extraction of lens	Y		A2	25.3605	\$1,061.92
66930	Extraction of lens	Y		A2	25.8575	\$1,082.73
66940	Extraction of lens	Y		A2	15.4386	\$646.46
66982	Cataract surgery, complex	Y		A2	22.9847	\$962.44
66983	Cataract surg w/ol, 1 stage	Y		A2	22.9847	\$962.44
66984	Cataract surg w/ol, 1 stage	Y		A2	22.9847	\$962.44
66985	Insert lens prosthesis	Y		A2	22.1453	\$927.29
66986	Exchange lens prosthesis	Y		A2	22.1453	\$927.29
66990	Ophthalmic endoscope add-on	N		N1	18.4618	\$773.05
67005	Partial removal of eye fluid	Y		A2	32.274	\$1,351.41
67010	Partial removal of eye fluid	Y		A2	30.5775	\$1,280.37
67015	Release of eye fluid	Y		A2	16.7652	\$702.01
67025	Replace eye fluid	Y		A2	32.274	\$1,351.41
67027	Implant eye drug system	Y		P3	16.7652	\$71.58
67030	Injection eye drug	Y		A2	5.589	\$234.03
67031	Intraocular eye strands	Y		A2	32.274	\$1,351.41
67036	Removal of inner eye fluid	Y		A2	34.3589	\$1,438.71
67039	Laser treatment of retina	Y		A2	38.2338	\$1,600.96
67040	Laser treatment of retina	Y		G2	38.2338	\$1,600.96
67041	Vit for macular pucker	Y		G2	38.2338	\$1,600.96
67042	Vit for macular hole	Y		G2	38.2338	\$1,600.96
67043	Vit for membrane dissect	Y		G2	38.2338	\$1,600.96
67101	Repair detached retina	Y	CH	P2	5.0718	\$212.37
67105	Repair detached retina	Y		P2	5.0718	\$212.37
67107	Repair detached retina	Y		A2	32.771	\$1,372.22

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66155	Glaucoma surgery	Y		A2	21.1076	\$883.84
66160	Glaucoma surgery	Y		A2	20.0566	\$839.83
66165	Glaucoma surgery	Y		A2	21.1076	\$883.84
66170	Glaucoma surgery	Y		A2	21.1076	\$883.84
66172	Implant eye shunt	Y		A2	34.1483	\$1,429.89
66185	Revis eye shunt	Y		A2	20.0566	\$839.83
66220	Repair eye lesion	Y		A2	31.5886	\$1,322.71
66225	Repair/graft eye lesion	Y		A2	33.6513	\$1,409.08
66250	Follow-up surgery of eye	Y		A2	14.2061	\$594.85
66500	Incision of iris	Y		A2	5.1362	\$215.07
66505	Incision of iris	Y		A2	5.1362	\$215.07
66600	Remove iris and lesion	Y		A2	20.4222	\$855.14
66605	Removal of iris	Y		A2	20.4222	\$855.14
66625	Removal of iris	Y		A2	13.7888	\$577.36
66630	Removal of iris	Y		A2	20.4222	\$855.14
66635	Removal of iris	Y		A2	20.4222	\$855.14
66680	Repair iris & ciliary body	Y		A2	20.4222	\$855.14
66682	Repair iris & ciliary body	Y		A2	20.0566	\$839.83
66700	Destruction, ciliary body	Y		A2	14.2061	\$594.85
66710	Ciliary transscleral therapy	Y		A2	14.2061	\$594.85
66720	Destruction, ciliary body	Y		A2	14.2061	\$594.85
66740	Destruction, ciliary body	Y		A2	20.0566	\$839.83
66761	Revision of iris	Y		P3	16.1106	\$661.06
66762	Revision of iris	Y		P3	16.1106	\$661.06
66770	Removal of inner eye lesion	Y		P3	4.3122	\$178.39
66820	Incision, secondary cataract	Y	CH	G2	5.589	\$234.03
66821	After cataract laser surgery	Y		A2	5.589	\$234.03
66825	Reposition intraocular lens	Y		A2	21.1076	\$883.84

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HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
67340	Revise eye muscle add-on	Y		A2	20.4385	\$855.82
67343	Release eye lissue	Y		A2	22.5231	\$943.11
67345	Destroy nerve of eye muscle	Y		P3		\$71.98
67346	Biopsy, eye muscle	Y		A2	13.053	\$546.57
67400	Explore/biopsy eye socket	Y		A2	15.942	\$667.54
67405	Explore/drain eye socket	Y		A2	22.8665	\$957.07
67412	Explore/drain eye socket	Y		A2	17.1244	\$717.05
67413	Explore/treat eye socket	Y		A2	23.3535	\$977.88
67414	Expli/decompress eye socket	Y		G2	37.3223	\$1,562.80
67415	Aspiration, orbital contents	Y		A2	14.9309	\$625.20
67420	Explore/treat eye socket	Y		A2	32.0873	\$1,343.59
67440	Explore/drain eye socket	Y		A2	32.0873	\$1,343.59
67445	Expli/decompress eye socket	Y		A2	32.0873	\$1,343.59
67450	Explore/biopsy eye socket	Y		A2	32.0873	\$1,343.59
67500	Inject/treat eye socket	N		G2	1.8727	\$78.42
67505	Inject/treat eye socket	Y		P3		\$23.86
67515	Inject/treat eye socket	Y		P3		\$24.43
67550	Insert eye socket implant	Y		A2	31.5903	\$1,322.78
67560	Revise eye socket implant	Y		A2	21.8055	\$913.06
67570	Drainage of eyelid abscess	Y		A2	31.5903	\$1,322.78
67700	Decompress optic nerve	Y		P2	3.067	\$129.26
67710	Incision of eyelid	Y		P3		\$119.87
67715	Incision of eyelid fold	Y		A2	14.9309	\$625.20
67800	Remove eyelid lesion	Y		P3		\$45.17
67805	Remove eyelid lesions	Y		P3		\$54.82
67808	Remove eyelid lesion(s)	Y		P3		\$70.73
67810	Biopsy of eyelid	Y	CH	A2	15.5764	\$652.23
67820	Revise eyelidties	N		P3		\$108.80
						\$14.77

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67108	Repair detached retina	Y		A2	34.3589	\$1,438.71
67110	Repair detached retina	Y		P3		\$230.88
67112	Repair detached retina	Y		A2	34.3589	\$1,438.71
67113	Repair retinal detach, opix	Y		G2	38.2338	\$1,600.96
67115	Release encircling material	Y		A2	17.4107	\$729.04
67120	Remove eye implant material	Y		A2	17.4107	\$729.04
67121	Remove eye implant material	Y		A2	17.4107	\$729.04
67141	Treatment of retina	Y		A2	5.5783	\$233.58
67145	Treatment of retina	Y		P3		\$172.14
67208	Treatment of retinal lesion	Y		P3		\$184.64
67210	Treatment of retinal lesion	Y	CH	P3		\$197.14
67218	Treatment of retinal lesion	Y		A2	18.9585	\$793.85
67220	Treatment of choroid lesion	Y		P2	5.5965	\$234.34
67221	Ocular photodynamic ther	Y		P3		\$100.56
67225	Eye photodynamic ther add-on	Y		P3	\$7.39	\$702.01
67227	Treatment of retinal lesion	Y		A2	16.7652	\$721.37
67229*	Tr retinal les preterm inf	Y		R2	5.0718	\$212.37
67250	Reinforce eye wall	Y		A2	15.942	\$667.54
67255	Reinforce/graft eye wall	Y		A2	17.7761	\$744.34
67311	Revise eye muscle	Y		A2	19.7528	\$827.11
67312	Revise two eye muscles	Y		A2	20.4385	\$855.82
67314	Revise eye muscle	Y		A2	20.4385	\$855.82
67316	Revise two eye muscles	Y		A2	20.4385	\$855.82
67318	Revise eye muscle(s)	Y		A2	20.4385	\$855.82
67320	Revise eye muscle(s) add-on	Y		A2	20.4385	\$855.82
67331	Eye surgery follow-up add-on	Y		A2	20.4385	\$855.82
67332	Revise eye muscles add-on	Y		A2	20.4385	\$855.82
67334	Revise eye muscle w/suture	Y		A2	20.4385	\$855.82
67335	Eye suture during surgery	Y		A2	20.4385	\$855.82

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67961	Revision of eyelid	Y		A2	15.942	\$667.54
67966	Revision of eyelid	Y		A2	15.942	\$667.54
67971	Reconstruction of eyelid	Y		A2	15.942	\$667.54
67973	Reconstruction of eyelid	Y		A2	22.1711	\$928.37
67974	Reconstruction of eyelid	Y		A2	15.942	\$667.54
67975	Reconstruction of eyelid	Y		A2	15.942	\$667.54
68020	Incise/drain eyelid lining	Y		P3		\$40.05
68040	Treatment of eyelid lesions	N		P3		\$19.88
68100	Biopsy of eyelid lining	Y		P3		\$76.41
68110	Remove eyelid lining lesion	Y		P3		\$99.42
68115	Remove eyelid lining lesion	Y		A2	15.5764	\$652.23
68130	Remove eyelid lining lesion	Y		A2	14.2061	\$594.85
68135	Remove eyelid lining lesion	Y		P3		\$51.70
68200	Treat eyelid by injection	N		P3		\$14.49
68320	Revise/graft eyelid lining	Y		A2	22.8565	\$957.07
68325	Revise/graft eyelid lining	Y		A2	22.8565	\$957.07
68326	Revise/graft eyelid lining	Y		A2	16.6274	\$696.24
68328	Revise/graft eyelid lining	Y		A2	22.8565	\$957.07
68330	Revise eyelid lining	Y		A2	21.1076	\$883.84
68335	Revise/graft eyelid lining	Y		A2	22.8565	\$957.07
68340	Separate eyelid adhesions	Y		A2	16.6274	\$696.24
68360	Revise eyelid lining	Y		A2	20.0566	\$839.83
68362	Revise eyelid lining	Y		A2	20.0566	\$839.83
68371	Harvest eye tissue, allograft	Y		A2	14.2061	\$594.85
68400	Incise/drain tear gland	Y		P2	3.087	\$129.26
68420	Incise/drain tear sac	Y		P3		\$46.28
68440	Incise tear duct opening	Y		P3		\$43.18
68500	Removal of tear gland	Y		A2	22.1711	\$928.37
68505	Partial removal, tear gland	Y		A2	22.1711	\$928.37
68510	Biopsy of tear gland	Y		A2	14.9309	\$625.20

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67825	Revise eyelashes	Y		P3		\$45.73
67830	Revise eyelashes	Y		A2	8.031	\$336.28
67835	Revise eyelashes	Y		A2	15.5764	\$652.23
67840	Remove eyelid lesion	Y		P3		\$126.98
67850	Treat eyelid lesion	Y		P3		\$99.99
67875	Closure of eyelid by suture	Y		G2	7.3112	\$306.14
67880	Revision of eyelid	Y		A2	14.5717	\$610.16
67882	Revision of eyelid	Y		A2	15.942	\$667.54
67900	Repair brow defect	Y		A2	22.8565	\$957.07
67901	Repair eyelid defect	Y		A2	17.1244	\$717.05
67902	Repair eyelid defect	Y		A2	23.3535	\$977.88
67903	Repair eyelid defect	Y		A2	16.6274	\$696.24
67904	Repair eyelid defect	Y		A2	16.6274	\$696.24
67906	Repair eyelid defect	Y		A2	17.1244	\$717.05
67908	Repair eyelid defect	Y		A2	16.6274	\$696.24
67909	Repair eyelid defect	Y		A2	16.6274	\$696.24
67911	Revise eyelid defect	Y		A2	15.942	\$667.54
67912	Correction eyelid w/implant	Y		A2	15.942	\$667.54
67914	Repair eyelid defect	Y		A2	15.942	\$667.54
67915	Repair eyelid defect	Y		P3		\$142.03
67916	Repair eyelid defect	Y		A2	16.6274	\$696.24
67917	Repair eyelid defect	Y		A2	16.6274	\$696.24
67921	Repair eyelid defect	Y		A2	15.942	\$667.54
67922	Repair eyelid defect	Y		P3		\$138.05
67923	Repair eyelid defect	Y		A2	16.6274	\$696.24
67924	Repair eyelid defect	Y		A2	16.6274	\$696.24
67930	Repair eyelid wound	Y		P3		\$143.17
67935	Repair eyelid wound	Y		A2	15.5764	\$652.23
67938	Remove eyelid foreign body	N		P2	1.8727	\$78.42
67950	Revision of eyelid	Y		A2	15.5764	\$652.23

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Note 2: Payment indicators for "office-based" procedures (P2, P3) are based on a comparison of the final rates according to the ASC standard ratesetting methodology and the MPPS. Under current law, MPPS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPPS final rule.

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ADDENDUM AA.—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPFS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
69200	Clear outer ear canal	N		P2	0.6403	\$26.61
69205	Clear outer ear canal	Y		A2	18.6836	\$782.34
69210	Remove impacted ear wax	N		P3		\$17.90
69220	Clean out mastoid cavity	Y		P2	0.8408	\$35.21
69222	Clean out mastoid cavity	Y		P3		\$114.76
69300	Revise external ear	Y		A2	20.8254	\$672.02
69310	Rebuild outer ear canal	Y		A2	33.7542	\$1,413.39
69320	Rebuild outer ear canal	Y		A2	36.5245	\$1,529.39
69400	Inflate middle ear canal	Y		P3		\$77.83
69401	Inflate middle ear canal	Y		P3		\$41.47
69405	Catheterize middle ear canal	Y		P3		\$108.23
69420	Incision of eardrum	Y		P3		\$96.01
69421	Incision of eardrum	Y		A2	15.2459	\$638.39
69424	Remove ventilating tube	Y		P3		\$67.04
69433	Create eardrum opening	Y		P3		\$96.01
69436	Create eardrum opening	Y		A2	15.2459	\$638.39
69440	Exploration of middle ear	Y		A2	20.8254	\$872.02
69450	Eardrum revision	Y		A2	32.7433	\$1,371.06
69501	Mastoidectomy	Y		A2	36.5245	\$1,529.39
69502	Mastoidectomy	Y		A2	23.5956	\$988.02
69505	Remove mastoid structures	Y		A2	36.5245	\$1,529.39
69511	Extensive mastoid surgery	Y		A2	36.5245	\$1,529.39
69530	Extensive mastoid surgery	Y		A2	36.5245	\$1,529.39
69540	Remove ear lesion	Y		P3		\$112.20
69550	Remove ear lesion	Y		A2	34.9366	\$1,462.90
69552	Remove ear lesion	Y		A2	36.5245	\$1,529.39
69601	Mastoid surgery revision	Y		A2	36.5245	\$1,529.39
69602	Mastoid surgery revision	Y		A2	36.5245	\$1,529.39
69603	Mastoid surgery revision	Y		A2	36.5245	\$1,529.39
69604	Mastoid surgery revision	Y		A2	36.5245	\$1,529.39

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ADDENDUM AA.—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPFS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
68520	Removal of tear sac	Y		A2	22.1711	\$928.37
68525	Biopsy of tear sac	Y		A2	14.9309	\$625.20
68530	Clearance of tear duct	Y		P2	3.087	\$129.26
68540	Remove tear gland lesion	Y		A2	15.942	\$667.54
68550	Remove tear gland lesion	Y		A2	22.1711	\$928.37
68700	Repair tear ducts	Y		A2	15.5764	\$652.23
68705	Revise tear duct opening	Y		P3		\$99.42
68720	Create tear sac drain	Y		A2	22.8565	\$957.07
68745	Create tear duct drain	Y		A2	22.8565	\$957.07
68750	Create tear duct drain	Y		A2	22.8565	\$957.07
68760	Close tear duct opening	Y		P3		\$64.65
68761	Close tear duct opening	Y		P3		\$59.08
68770	Close tear system fistula	Y		A2	22.8565	\$957.07
68801	Dilate tear duct opening	N		P2	0.9139	\$38.27
68810	Probe nasolacrimal duct	Y		A2	3.0683	\$128.48
68811	Probe nasolacrimal duct	Y		A2	15.5764	\$652.23
68815	Probe nasolacrimal duct	Y		G2	15.5764	\$652.23
68840	Explore/irrigate tear ducts	Y		G2	17.3718	\$727.41
68850	Injection for tear sac x-ray	N		P3		\$47.44
69000	Drain external ear lesion	Y		N1		
69005	Drain external ear lesion	Y		P2	1.3927	\$58.32
69020	Drain outer ear canal lesion	Y		P3		\$88.91
69100	Biopsy of external ear	Y		P2	1.3927	\$58.32
69105	Biopsy of external ear canal	Y		P3		\$51.13
69110	Remove external ear, partial	Y		A2	14.457	\$74.99
69120	Removal of external ear	Y		A2	20.4597	\$856.71
69140	Remove ear canal lesion(s)	Y		A2	20.4597	\$856.71
69145	Remove ear canal lesion(s)	Y		A2	15.1023	\$632.38
69150	Extensive ear canal surgery	Y		A2	8.1184	\$339.94

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ADDENDUM AA—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
69740	Repair facial nerve	Y		A2	34.9366	\$1,462.90
69745	Repair facial nerve	Y		A2	34.9366	\$1,462.90
69801	Incise inner ear	Y		A2	22.0077	\$921.53
69802	Incise inner ear	Y		A2	23.5956	\$988.02
69805	Explore inner ear	Y		A2	36.5245	\$1,529.39
69806	Explore inner ear	Y		A2	36.5245	\$1,529.39
69820	Establish inner ear window	Y		A2	34.9366	\$1,462.90
69840	Remove inner ear window	Y		A2	34.9366	\$1,462.90
69905	Remove inner ear	Y		A2	36.5245	\$1,529.39
69910	Remove inner ear & mastoid	Y		A2	36.5245	\$1,529.39
69915	Incise inner ear nerve	Y		A2	36.5245	\$1,529.39
69930	Implant cochlear device	Y		H8	636.8197	\$26,665.55
C9716	Radiofrequency energy to anu	N		N1		
C9724	EPS gast cardia pic	Y		G2	30.7878	\$1,289.18
C9726	Place endorectal app	Y		G2	23.2194	\$972.27
C9727	Rxt breast appl place/remov	Y		G2	5.1327	\$214.92
C9728	Insert palate implants	Y		G2	23.6799	\$991.55
G0104	Place device/marker, non pro	N		R2	7.2897	\$305.24
G0105	Colorectal scrn; hi risk ind	N		P3	13.1619	\$551.13
G0121	Colon ca scrn not hi risk ind	Y		A2	8.3531	\$349.77
G0127	Trim nail(s)	Y		P3	8.3531	\$349.77
G0186	Dstry eye lesn,ldr,vssl tech	Y		R2	5.5965	\$234.34
G0247	Flouitne footcare pt w tops	Y		P3		\$18.75
G0259	Inject for sacroiliac joint	N		N1		
G0260	lnj for sacroiliac jt anesih	Y		A2	7.0685	\$295.98
G0268	Removal of impacted wax,md	N		N1		
G0289	Occlusive device in vein art	N		N1		
G0289	Arthro, loose body + chondro	N		N1		

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ADDENDUM AA—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
69605	Mastoid surgery revision	Y		A2	36.5245	\$1,529.39
69610	Repair of eardrum	Y		P3		\$150.84
69620	Repair of eardrum	Y		A2	20.4597	\$856.71
69631	Repair eardrum structures	Y		A2	34.9366	\$1,462.90
69632	Rebuild eardrum structures	Y		A2	34.9366	\$1,462.90
69633	Rebuild eardrum structures	Y		A2	34.9366	\$1,462.90
69635	Repair eardrum structures	Y		A2	36.5245	\$1,529.39
69636	Rebuild eardrum structures	Y		A2	36.5245	\$1,529.39
69645	Rebuild eardrum structures	Y		A2	36.5245	\$1,529.39
69641	Revise middle ear & mastoid	Y		A2	36.5245	\$1,529.39
69642	Revise middle ear & mastoid	Y		A2	36.5245	\$1,529.39
69643	Revise middle ear & mastoid	Y		A2	36.5245	\$1,529.39
69644	Revise middle ear & mastoid	Y		A2	36.5245	\$1,529.39
69645	Revise middle ear & mastoid	Y		A2	36.5245	\$1,529.39
69646	Revise middle ear & mastoid	Y		A2	36.5245	\$1,529.39
69650	Release middle ear bone	Y		A2	23.5956	\$988.02
69660	Release middle ear bone	Y		A2	34.9366	\$1,462.90
69661	Revise middle ear bone	Y		A2	34.9366	\$1,462.90
69662	Revise middle ear bone	Y		A2	34.9366	\$1,462.90
69666	Repair middle ear structures	Y		A2	34.9366	\$1,442.09
69667	Repair middle ear structures	Y		A2	34.9366	\$1,442.09
69670	Remove mastoid air cells	Y		A2	33.7542	\$1,413.39
69676	Remove middle ear nerve	Y		A2	33.7542	\$1,413.39
69700	Close mastoid fistula	Y		A2	33.7542	\$1,413.39
69711	Remove/repair hearing aid	Y		A2	32.7433	\$1,371.06
69714	Implant temple bone w/stimul	Y		H8	153.8521	\$6,442.25
69715	Temple bone implant w/stimul	Y		H8	153.8521	\$6,442.25
69717	Temple bone implant revision	Y		H8	153.8521	\$6,442.25
69718	Revise temple bone implant	Y		H8	153.8521	\$6,442.25
69720	Release facial nerve	Y		A2	34.9366	\$1,462.90

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ADDENDUM B.—FINAL OPFS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
00100	Anesth, salivary gland	N	N					
00102	Anesth, repair of cleft lip	N	N					
00103	Anesth, blepharoplasty	N	N					
00104	Anesth, electroshock	N	N					
00120	Anesth, ear surgery	N	N					
00124	Anesth, ear exam	N	N					
00126	Anesth, tympanotomy	N	N					
00140	Anesth, procedures on eye	N	N					
00142	Anesth, lens surgery	N	N					
00144	Anesth, corneal transplant	N	N					
00145	Anesth, vitreoretinal surg	N	N					
00147	Anesth, iridectomy	N	N					
00148	Anesth, eye exam	N	N					
00160	Anesth, nose/sinus surgery	N	N					
00162	Anesth, nose/sinus surgery	N	N					
00164	Anesth, biopsy of nose	N	N					
00170	Anesth, procedure on mouth	N	N					
00172	Anesth, cleft palate repair	N	N					
00174	Anesth, pharyngeal surgery	N	N					
00176	Anesth, pharyngeal surgery	C	N					
00190	Anesth, face/skull bone surg	N	N					
00192	Anesth, facial bone surgery	C	N					
00210	Anesth, cranial surg nos	N	N					
00211	Anesth, cran surg, hemotoma	C	N					
00212	Anesth, skull drainage	C	N					
00214	Anesth, skull drainage	C	N					
00215	Anesth, skull repair/fract	C	N					
00216	Anesth, head vessel surgery	N	N					
00218	Anesth, special head surgery	N	N					
00220	Anesth, intrcn nerve	N	N					
00222	Anesth, head nerve surgery	N	N					
00300	Anesth, head/neck/trunk	N	N					
00320	Anesth, neck organ, 1 & over	N	N					
00322	Anesth, biopsy of thyroid	N	N					
00326	Anesth, larynx/trach, < 1 yr	N	N					
00350	Anesth, neck vessel surgery	N	N					
00352	Anesth, neck vessel surgery	N	N					
00400	Anesth, skin, ext/peit/atunk	N	N					
00402	Anesth, surgery of breast	N	N					
00404	Anesth, surgery of breast	N	N					

ADDENDUM AA.—FINAL ASC COVERED SURGICAL PROCEDURES FOR CY 2010 (INCLUDING SURGICAL PROCEDURES FOR WHICH PAYMENT IS PACKAGED)

HCPCS Code	Short Descriptor	Subject To Multiple Procedure Discounting	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
G0364	Bone marrow aspirate & biopsy	N	CH	P3		\$4.54
G0392	AV fistula or graft arterial	N	CH	D5		
G0393	AV fistula or graft venous	N	CH	D5		

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ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
00670	Anesth, spine, cord surgery	C						
00700	Anesth, abdominal wall surg		N					
00702	Anesth, for liver biopsy		N					
00730	Anesth, abdominal wall surg		N					
00740	Anesth, upper gi visualize		N					
00750	Anesth, repair of hernia		N					
00762	Anesth, repair of hernia		N					
00754	Anesth, repair of hernia		N					
00756	Anesth, repair of hernia		N					
00770	Anesth, blood vessel repair		N					
00790	Anesth, surg upper abdomen		N					
00792	Anesth, hemorrhoidectomy		C					
00794	Anesth, pancreas removal		C					
00796	Anesth, for liver transplant		C					
00797	Anesth, surgery for obesity		N					
00800	Anesth, abdominal wall surg		N					
00802	Anesth, fat layer removal		C					
00810	Anesth, low intestine scope		N					
00820	Anesth, abdominal wall surg		N					
00830	Anesth, repair of hernia		N					
00832	Anesth, repair of hernia		N					
00834	Anesth, hernia repair< 1 yr		N					
00836	Anesth hernia repair preemie		N					
00840	Anesth, surg lower abdomen		N					
00842	Anesth, amniocentesis		N					
00844	Anesth, pelvis surgery		C					
00846	Anesth, hysterectomy		C					
00848	Anesth, pelvic organ surg		C					
00851	Anesth, tubal ligation		N					
00860	Anesth, surgery of abdomen		N					
00862	Anesth, kidney/ureter surg		C					
00864	Anesth, removal of bladder		C					
00865	Anesth, removal of prostate		C					
00866	Anesth, removal of adrenal		C					
00868	Anesth, kidney transplant		C					
00870	Anesth, bladder stone surg		N					
00872	Anesth kidney stone destruct		N					
00873	Anesth kidney stone destruct		N					
00880	Anesth, abdomen vessel surg		N					
00882	Anesth, major vein ligation		C					
00902	Anesth, anorectal surgery		N					
00904	Anesth, perineal surgery		C					
00906	Anesth, removal of vulva		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
00406	Anesth, surgery of breast		N					
00410	Anesth, correct heart rhythm		N					
00450	Anesth, surgery of shoulder		N					
00452	Anesth, surgery of shoulder		C					
00454	Anesth, collar bone biopsy		N					
00470	Anesth, removal of rib		N					
00472	Anesth, chest wall repair		N					
00474	Anesth, surgery of rib(s)		C					
00500	Anesth, esophageal surgery		N					
00520	Anesth, chest procedure		N					
00522	Anesth, chest lining biopsy		N					
00524	Anesth, chest drainage		C					
00528	Anesth, chest partition view		N					
00529	Anesth, chest partition view		N					
00530	Anesth, pacemaker insertion		N					
00532	Anesth, vascular access		N					
00534	Anesth, cardioverter/defib		N					
00537	Anesth, cardiac electrophys		N					
00539	Anesth, trach-bronch reconist		N					
00540	Anesth, chest surgery		C					
00541	Anesth, one lung ventilation		N					
00542	Anesth, release of lung		C					
00546	Anesth, lung,chest wall surg		C					
00548	Anesth, trachea,bronchi surg		N					
00550	Anesth, sternal debridement		N					
00560	Anesth, heart surg w/o pump		C					
00561	Anesth, heart surg < age 1		C					
00562	Anesth hrt surg w/pmp age 1+		C					
00563	Anesth, heart surg w/arest		N					
00566	Anesth, cabg w/o pump		N					
00567	Anesth, cabg w/pump		C					
00580	Anesth, heart/lung transplnt		C					
00600	Anesth, spine, cord surgery		N					
00604	Anesth, sitling procedure		C					
00620	Anesth, spine, cord surgery		N					
00622	Anesth, removal of nerves		C					
00625	Anes spine transthor w/o vent		N					
00626	Anes, spine transthor w/vent		N					
00630	Anesth, spine, cord surgery		N					
00632	Anesth, removal of nerves		C					
00634	Anesth for chemoradiolysis		N					
00635	Anesth, lumbar puncture		N					
00640	Anesth, spine manipulation		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
01250	Anesth, upper leg surg	N	N					
01260	Anesth, upper leg veins surg	N	N					
01270	Anesth, thigh arteries surg	N	N					
01272	Anesth, femoral artery surg	C	C					
01274	Anesth, femoral embolectomy	N	N					
01320	Anesth, knee area surgery	N	N					
01340	Anesth, knee area procedure	N	N					
01360	Anesth, knee area surgery	N	N					
01380	Anesth, knee joint procedure	N	N					
01382	Anesth, dx knee arthroscopy	N	N					
01390	Anesth, knee area procedure	N	N					
01392	Anesth, knee area surgery	N	N					
01400	Anesth, knee joint surgery	N	N					
01402	Anesth, knee arthroplasty	C	C					
01404	Anesth, amputation at knee	N	N					
01420	Anesth, knee joint casting	N	N					
01430	Anesth, knee veins surgery	N	N					
01432	Anesth, knee vessel surg	N	N					
01440	Anesth, knee arteries surg	N	N					
01442	Anesth, knee artery surg	C	C					
01444	Anesth, knee artery repair	N	N					
01462	Anesth, lower leg procedure	N	N					
01464	Anesth, ankle/ft arthroscopy	N	N					
01470	Anesth, lower leg surgery	N	N					
01472	Anesth, achilles tendon surg	N	N					
01474	Anesth, lower leg surgery	N	N					
01480	Anesth, lower leg bone surg	N	N					
01482	Anesth, radical leg surgery	N	N					
01484	Anesth, lower leg revision	N	N					
01486	Anesth, ankle replacement	C	C					
01490	Anesth, lower leg casting	N	N					
01500	Anesth, leg arteries surg	N	N					
01502	Anesth, lwr leg embolectomy	C	C					
01520	Anesth, lower leg vein surg	N	N					
01522	Anesth, lower leg vein surg	N	N					
01610	Anesth, surgery of shoulder	N	N					
01620	Anesth, shoulder procedure	N	N					
01622	Anes dx shoulder arthroscopy	N	N					
01630	Anesth, surgery of shoulder	N	N					
01632	Anesth, surgery of shoulder	CH	D					
01634	Anesth, shoulder joint amput	N	C					
01636	Anesth, forequarter amput	N	C					
01638	Anesth, shoulder replacement	N	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
00908	Anesth, removal of prostate	N	C					
00910	Anesth, bladder surgery	N	N					
00912	Anesth, bladder tumor surg	N	N					
00914	Anesth, removal of prostate	N	N					
00916	Anesth, bleeding control	N	N					
00918	Anesth, stone removal	N	N					
00920	Anesth, genitalia surgery	N	N					
00921	Anesth, vasectomy	N	N					
00922	Anesth, sperm duct surgery	N	N					
00924	Anesth, testis exploration	N	N					
00926	Anesth, removal of testis	N	N					
00928	Anesth, removal of testis	N	N					
00930	Anesth, testis suspension	N	N					
00932	Anesth, amputation of penis	C	C					
00934	Anesth, penis, nodes removal	N	N					
00936	Anesth, penis, nodes removal	C	C					
00938	Anesth, insert penis device	N	N					
00940	Anesth, vaginal procedures	N	N					
00942	Anesth, surg on vag/urethral	N	N					
00944	Anesth, vaginal hysterectomy	C	C					
00948	Anesth, repair of cervix	N	N					
00950	Anesth, vaginal endoscopy	N	N					
00952	Anesth, hysteroscope/graph	N	N					
01112	Anesth, bone aspirate/bx	N	N					
01120	Anesth, pelvis surgery	N	N					
01130	Anesth, body cast procedure	N	N					
01140	Anesth, amputation at pelvis	C	C					
01150	Anesth, pelvic tumor surgery	C	C					
01160	Anesth, pelvis procedure	N	N					
01170	Anesth, pelvis surgery	N	N					
01173	Anesth, tx repair, pelvis	N	N					
01180	Anesth, pelvis nerve removal	N	N					
01190	Anesth, pelvis nerve removal	N	N					
01200	Anesth, hip joint procedure	N	N					
01202	Anesth, arthroscopy of hip	N	N					
01210	Anesth, hip joint surgery	N	N					
01212	Anesth, hip disarticulation	C	C					
01214	Anesth, hip arthroplasty	C	C					
01215	Anesth, revise hip repair	N	N					
01220	Anesth, procedure on femur	N	N					
01230	Anesth, surgery of femur	N	N					
01232	Anesth, amputation of femur	C	C					
01234	Anesth, radical femur surg	N	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
01933	Anes, tx interv rad, cran v	N						
01935	Anesth, peric ring dx sp proc	N						
01936	Anesth, peric ring tx sp proc	N						
01951	Anesth, burn, less 4 percent	N						
01952	Anesth, burn, 4-9 percent	N						
01953	Anesth, burn, each 9 percent	N						
01958	Anesth, antepartum manipul	N						
01960	Anesth, vaginal delivery	N						
01961	Anesth, cs delivery	N						
01962	Anesth, emer hysterectomy	N						
01963	Anesth, cs hysterectomy	N						
01965	Anesth, inc/misssed ab proc	N						
01966	Anesth, induced ab procedure	N						
01967	Anesth/analg, vag delivery	N						
01968	Anes/analg cs deliver add-on	N						
01969	Anesth/analg cs hyst add-on	N						
01990	Support for organ donor	C						
01991	Anesth, nerve block(in)	N						
01992	Anesth, n block(in), prone	N						
01996	Hosp manage cont drug admin	N						
01999	Unlisted anesth procedure	N						
10021	Fna w/o image	T	0002		1.5111	\$101.86		\$20.38
10022	Fna w/image	T	0004		4.5991	\$310.01		\$62.01
10040	Acne surgery	T	0013		0.8789	\$59.24		\$11.85
10060	Drainage of skin abscess	T	0006		1.4557	\$98.12		\$19.63
10061	Drainage of skin abscess	T	0006		1.4557	\$98.12		\$19.63
10080	Drainage of pilonidal cyst	T	0006		1.4557	\$98.12		\$19.63
10081	Drainage of pilonidal cyst	T	0007		12.6217	\$850.78		\$170.16
10120	Remove foreign body	T	0016		2.7982	\$188.62		\$37.73
10121	Remove foreign body	T	0021		17.4975	\$1,179.44		\$235.89
10140	Drainage of hematoma/fluid	T	0007		12.6217	\$850.78		\$170.16
10160	Puncture drainage of lesion	T	0006		1.4557	\$98.12		\$19.63
10180	Complex drainage, wound	T	0008		19.4063	\$1,308.10		\$261.62
11000	Debride infected skin	T	0015		1.5412	\$103.89		\$20.78
11001	Debride infected skin add-on	T	0013		0.8789	\$59.24		\$11.85
11004	Debride genitalia & perineum	C						
11006	Debride abdomen wall	C						
11008	Remove mesh from abd wall	C						
11010	Debride skin, fx	T	0019		4.3625	\$294.06	\$64.51	\$58.82
11011	Debride skin/muscle, fx	T	0019		4.3625	\$294.06	\$64.51	\$58.82
11012	Debride skin/muscle/bone, fx	T	0019		4.3625	\$294.06	\$64.51	\$58.82
11040	Debride skin, partial	T	0015		1.5412	\$103.89		\$20.78

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
01650	Anesth, shoulder artery surg	N						
01652	Anesth, shoulder vessel surg	C						
01654	Anesth, shoulder vessel surg	C						
01656	Anesth, arm-leg vessel surg	C						
01670	Anesth, shoulder vein surg	N						
01680	Anesth, shoulder casting	N						
01682	Anesth, airplane cast	N						
01710	Anesth, elbow area surgery	N						
01712	Anesth, uppr arm tendon surg	N						
01714	Anesth, uppr arm tendon surg	N						
01716	Anesth, biceps tendon repair	N						
01730	Anesth, uppr arm procedure	N						
01732	Anesth, dx elbow arthroscopy	N						
01740	Anesth, upper arm surgery	N						
01742	Anesth, humerus surgery	N						
01744	Anesth, humerus repair	N						
01756	Anesth, radical humerus surg	N						
01758	Anesth, humeral lesion surg	N						
01760	Anesth, elbow replacement	N						
01770	Anesth, uppr arm artery surg	N						
01772	Anesth, uppr arm embolectomy	N						
01780	Anesth, upper arm vein surg	N						
01782	Anesth, uppr arm vein repair	N						
01810	Anesth, lower arm surgery	N						
01820	Anesth, lower arm procedure	N						
01829	Anesth, dx wrist arthroscopy	N						
01830	Anesth, lower arm surgery	N						
01832	Anesth, wrist replacement	N						
01840	Anesth, lwr arm artery surg	N						
01842	Anesth, lwr arm embolectomy	N						
01844	Anesth, vascular shunt surg	N						
01850	Anesth, lower arm vein surg	N						
01852	Anesth, lwr arm vein repair	N						
01860	Anesth, lower arm casting	N						
01916	Anesth, dx arteriography	N						
01920	Anesth, catheterize heart	N						
01922	Anesth, cat or MRI scan	N						
01924	Anes, ther interven rad, art	N						
01925	Anes, ther interven rad, car	N						
01926	Anes, tx interv rad hrt/cran	N						
01930	Anes, ther interven rad, vei	N						
01931	Anes, ther interven rad, lip	N						
01932	Anes, tx interv rad, th vein	N						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
11446	Exc face-mm b9+marq > 4 cm	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11450	Removal, sweat gland lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11451	Removal, sweat gland lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11462	Removal, sweat gland lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11463	Removal, sweat gland lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11470	Removal, sweat gland lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11471	Removal, sweat gland lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11600	Exc tr-ext mlg+marq 0.5 < cm	CH		0020	8.2028	\$552.92		\$110.59
11601	Exc tr-ext mlg+marq 0.6-1 cm	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11602	Exc tr-ext mlg+marq 1.1-2 cm	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11603	Exc tr-ext mlg+marq 2.1-3 cm	T		0020	8.2028	\$552.92		\$110.59
11604	Exc tr-ext mlg+marq 3.1-4 cm	T		0020	8.2028	\$552.92		\$110.59
11606	Exc tr-ext mlg+marq > 4 cm	T		0021	17.4975	\$1,179.44		\$235.89
11620	Exc h-f-nk-sp mlg+marq 0.5 <	T		0020	8.2028	\$552.92	\$64.51	\$110.59
11621	Exc h-f-nk-sp mlg+marq 0.6-1	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11622	Exc h-f-nk-sp mlg+marq 1.1-2	T		0020	8.2028	\$552.92		\$110.59
11623	Exc h-f-nk-sp mlg+marq 2.1-3	T		0021	17.4975	\$1,179.44		\$235.89
11624	Exc h-f-nk-sp mlg+marq 3.1-4	T		0021	17.4975	\$1,179.44		\$235.89
11626	Exc h-f-nk-sp mlg+marq > 4 cm	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11640	Exc face-mm malig+marq 0.6-1	T		0020	8.2028	\$552.92		\$110.59
11641	Exc face-mm malig+marq 0.6-1	T		0020	8.2028	\$552.92		\$110.59
11642	Exc face-mm malig+marq 1.1-2	T		0020	8.2028	\$552.92		\$110.59
11643	Exc face-mm malig+marq 2.1-3	T		0020	8.2028	\$552.92		\$110.59
11644	Exc face-mm malig+marq 3.1-4	T		0021	17.4975	\$1,179.44		\$235.89
11646	Exc face-mm mlg+marq > 4 cm	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11719	Trim nail(s)	CH		0012	0.4436	\$29.90		\$5.98
11720	Debride nail, 1-5	T		0013	0.8789	\$59.24		\$11.85
11721	Debride nail, 6 or more	T		0013	0.8789	\$59.24		\$11.85
11730	Removal of nail plate	T		0013	0.8789	\$59.24		\$11.85
11732	Remove nail plate, add-on	T		0013	0.8789	\$59.24		\$11.85
11740	Drain blood from under nail	T		0012	0.4436	\$29.90		\$5.98
11750	Removal of nail bed	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11752	Remove nail bed/finger tip	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11755	Biopsy, nail unit	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11760	Repair of nail bed	CH		0133	1.3542	\$91.28	\$25.67	\$18.26
11762	Reconstruction of nail bed	T		0136	16.0542	\$1,082.15		\$216.43
11765	Excision of nail fold, toe	T		0013	0.8789	\$59.24		\$11.85
11770	Removal of pilonidal lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11771	Removal of pilonidal lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11772	Removal of pilonidal lesion	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11900	Injection into skin lesions	T		0013	0.8789	\$59.24		\$11.85
11901	Added skin lesions injection	T		0013	0.8789	\$59.24		\$11.85

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
11041	Debride skin, full	T		0015	1.5412	\$103.89		\$20.78
11042	Debride skin/fleshy	T		0016	2.7982	\$188.62		\$37.73
11043	Debride tissue/muscle	T		0016	2.7982	\$188.62		\$37.73
11044	Debride tissue/muscle/bone	CH		0020	8.2028	\$552.92		\$110.59
11055	Trim skin lesion	T		0013	0.8789	\$59.24		\$11.85
11056	Trim skin lesions, 2 to 4	T		0013	0.8789	\$59.24		\$11.85
11057	Trim skin lesions, over 4	T		0013	0.8789	\$59.24		\$11.85
11100	Biopsy, skin lesion	T		0015	1.5412	\$103.89		\$20.78
11101	Biopsy, skin add-on	T		0013	0.8789	\$59.24		\$11.85
11200	Removal of skin tags	T		0013	0.8789	\$59.24		\$11.85
11201	Remove skin tags add-on	T		0013	0.8789	\$59.24		\$11.85
11300	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11301	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11302	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11303	Shave skin lesion	T		0015	1.5412	\$103.89		\$20.78
11305	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11306	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11307	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11308	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11310	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11311	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11312	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11313	Shave skin lesion	T		0013	0.8789	\$59.24		\$11.85
11400	Exc tr-ext b9+marq 0.5 < cm	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11401	Exc tr-ext b9+marq 0.6-1 cm	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11402	Exc tr-ext b9+marq 1.1-2 cm	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11403	Exc tr-ext b9+marq 2.1-3 cm	T		0020	8.2028	\$552.92		\$110.59
11404	Exc tr-ext b9+marq 3.1-4 cm	T		0021	17.4975	\$1,179.44		\$235.89
11406	Exc tr-ext b9+marq > 4.0 cm	T		0021	17.4975	\$1,179.44		\$235.89
11420	Exc h-f-nk-sp b9+marq 0.5 <	T		0020	8.2028	\$552.92		\$110.59
11421	Exc h-f-nk-sp b9+marq 0.6-1	T		0020	8.2028	\$552.92		\$110.59
11422	Exc h-f-nk-sp b9+marq 1.1-2	T		0020	8.2028	\$552.92		\$110.59
11423	Exc h-f-nk-sp b9+marq 2.1-3	T		0021	17.4975	\$1,179.44		\$235.89
11424	Exc h-f-nk-sp b9+marq 3.1-4	T		0021	17.4975	\$1,179.44		\$235.89
11426	Exc h-f-nk-sp b9+marq > 4 cm	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
11440	Exc face-mm b9+marq 0.5 <	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11441	Exc face-mm b9+marq 0.6-1	T		0019	4.3625	\$294.06	\$64.51	\$58.82
11442	Exc face-mm b9+marq 1.1-2	T		0020	8.2028	\$552.92		\$110.59
11443	Exc face-mm b9+marq 2.1-3	T		0020	8.2028	\$552.92		\$110.59
11444	Exc face-mm b9+marq 3.1-4	T		0020	8.2028	\$552.92		\$110.59

ADDENDUM B.—FINAL OPFS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
12047	Intmd wnd repair n-hg/genit	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12051	Intmd wnd repair face/mm	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12052	Intmd wnd repair face/mm	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12053	Intmd wnd repair face/mm	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12054	Intmd wnd repair face/mm	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12055	Intmd wnd repair face/mm	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12056	Intmd wnd repair face/mm	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12057	Intmd wnd repair face/mm	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
13100	Repair of wound or lesion	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13101	Repair of wound or lesion	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13102	Repair wound/lesion add-on	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13120	Repair of wound or lesion	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
13121	Repair of wound or lesion	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
13122	Repair wound/lesion add-on	CH	0133	1.3542	\$91.28	\$25.67	\$18.26	
13131	Repair of wound or lesion	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
13132	Repair of wound or lesion	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13133	Repair wound/lesion add-on	CH	0134	3.1508	\$212.38	\$42.48	\$42.48	
13150	Repair of wound or lesion	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13151	Repair of wound or lesion	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13152	Repair of wound or lesion	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
13153	Repair wound/lesion add-on	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
13160	Late closure of wound	T	0137	23.9317	\$1,613.14	\$322.63	\$322.63	
14000	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14001	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14020	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14021	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14040	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14060	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14061	Skin tissue rearrangement	T	0136	16.0542	\$1,082.15	\$216.43	\$216.43	
14300	Skin tissue rearrangement	CH	D					
14301	Skin tissue rearrangement	NI	T	0137	23.9317	\$1,613.14	\$322.63	
14302	Skin tissue rearrangement add-on	NI	T	0137	23.9317	\$1,613.14	\$322.63	
14350	Skin tissue rearrangement	T	0137	23.9317	\$1,613.14	\$322.63	\$322.63	
15002	Wound prep, trn/arm/leg	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
15003	Wound prep, addl 100 cm	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
15004	Wound prep, l/n/h/g	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
15005	Wound prep, l/n/h/g, addl cm	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
15040	Harvest cultured skin graft	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
15050	Skin pinch graft	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
15100	Skin split grft, trnk/arm/leg	T	0137	23.9317	\$1,613.14	\$322.63	\$322.63	
15101	Skin split grft 1/air, add-on	T	0137	23.9317	\$1,613.14	\$322.63	\$322.63	
15110	Epidrm autogrtf trnk/arm/leg	T	0135	4.4386	\$299.19	\$59.84	\$59.84	

ADDENDUM B.—FINAL OPFS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
11920	Correct skin color defects	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
11921	Correct skin color defects	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
11922	Correct skin color defects	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
11950	Therapy for contour defects	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
11951	Therapy for contour defects	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
11952	Therapy for contour defects	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
11954	Therapy for contour defects	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
11960	Insert tissue expander(s)	T	0137	23.9317	\$1,613.14	\$322.63	\$322.63	
11970	Replace tissue expander	T	0051	46.5786	\$3,139.68	\$627.94	\$627.94	
11971	Remove tissue expander(s)	T	0022	23.368	\$1,576.49	\$354.45	\$315.30	
11975	Insert contraceptive cap	E						
11976	Removal of contraceptive cap	T	0019	4.3625	\$294.06	\$64.51	\$58.82	
11977	Removal/reinsert contra cap	E						
11980	Implant hormone pellet(s)	X	0340	0.6693	\$45.11	\$9.03	\$9.03	
11981	Insert drug implant device	X	0340	0.6693	\$45.11	\$9.03	\$9.03	
11982	Remove drug implant device	X	0340	0.6693	\$45.11	\$9.03	\$9.03	
11983	Remove/insert drug implant	X	0340	0.6693	\$45.11	\$9.03	\$9.03	
12001	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12002	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12004	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12005	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12006	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12007	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12011	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12013	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12014	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12015	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12016	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12017	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12018	Repair superficial wound(s)	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12020	Closure of split wound	T	0135	4.4386	\$299.19	\$59.84	\$59.84	
12021	Closure of split wound	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12031	Intmd wnd repair s/n/ext	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12032	Intmd wnd repair s/n/ext	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12034	Intmd wnd repair s/n/ext	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12035	Intmd wnd repair s/n/ext	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12036	Intmd wnd repair s/n/ext	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12037	Intmd wnd repair s/n/ext	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12041	Intmd wnd repair n-hg/genit	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12042	Intmd wnd repair n-hg/genit	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12044	Intmd wnd repair n-hg/genit	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
12045	Intmd wnd repair n-hg/genit	T	0134	3.1508	\$212.38	\$42.48	\$42.48	
12046	Intmd wnd repair n-hg/genit	T	0134	3.1508	\$212.38	\$42.48	\$42.48	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
15420	Apply skin xgrft, f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15421	Apply skin xgrft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15430	Apply acellular xenograft	T	0135	4.4386	\$299.19		\$59.84	
15431	Apply acellular xgrft add	T	0135	4.4386	\$299.19		\$59.84	
15570	Form skin pedicle flap	T	0137	23.9317	\$1,613.14		\$322.63	
15572	Form skin pedicle flap	T	0137	23.9317	\$1,613.14		\$322.63	
15574	Form skin pedicle flap	T	0137	23.9317	\$1,613.14		\$322.63	
15576	Form skin pedicle flap	T	0137	23.9317	\$1,613.14		\$322.63	
15610	Skin graft	T	0137	23.9317	\$1,613.14		\$322.63	
15620	Skin graft	T	0137	23.9317	\$1,613.14		\$322.63	
15630	Skin graft	T	0137	23.9317	\$1,613.14		\$322.63	
15650	Transfer skin pedicle flap	T	0137	23.9317	\$1,613.14		\$322.63	
15731	Forehead flap w/vasc pedicle	T	0137	23.9317	\$1,613.14		\$322.63	
15732	Muscle-skin graft, head/neck	T	0137	23.9317	\$1,613.14		\$322.63	
15734	Muscle-skin graft, trunk	T	0137	23.9317	\$1,613.14		\$322.63	
15736	Muscle-skin graft, arm	T	0137	23.9317	\$1,613.14		\$322.63	
15738	Muscle-skin graft, leg	T	0137	23.9317	\$1,613.14		\$322.63	
15740	Island pedicle flap graft	T	0136	16.0542	\$1,082.15		\$216.43	
15750	Free myo/skin flap microvasc	T	0137	23.9317	\$1,613.14		\$322.63	
15756	Free fascial flap, microvasc	C						
15757	Free skin flap, microvasc	C						
15758	Composite skin graft	T	0137	23.9317	\$1,613.14		\$322.63	
15760	Derma-fat-fascia graft	T	0137	23.9317	\$1,613.14		\$322.63	
15770	Hair transplant punch grafts	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
15775	Hair transplant punch grafts	T	0133	1.3542	\$91.28	\$25.67	\$18.26	
15776	Abrasion treatment of skin	T	0022	23.388	\$1,576.49	\$354.45	\$315.30	
15780	Abrasion treatment of skin	T	0019	4.3625	\$294.06	\$64.51	\$58.82	
15782	Abrasion treatment of skin	T	0016	2.7982	\$188.62	\$44.51	\$37.73	
15783	Abrasion, lesion, single	T	0013	0.8789	\$59.24		\$11.85	
15787	Abrasion, lesions, add-on	T	0013	0.8789	\$59.24		\$11.85	
15788	Chemical peel, face, epiderm	T	0015	1.5412	\$103.89		\$20.78	
15789	Chemical peel, face, dermal	T	0015	1.5412	\$103.89		\$20.78	
15792	Chemical peel, nonfacial	T	0013	0.8789	\$59.24		\$11.85	
15793	Chemical peel, nonfacial	T	0013	0.8789	\$59.24		\$11.85	
15819	Plastic surgery, neck	T	0134	3.1508	\$212.38		\$42.48	
15820	Revision of lower eyelid	T	0137	23.9317	\$1,613.14		\$322.63	
15821	Revision of lower eyelid	T	0137	23.9317	\$1,613.14		\$322.63	
15822	Revision of upper eyelid	T	0137	23.9317	\$1,613.14		\$322.63	
15823	Revision of upper eyelid	T	0137	23.9317	\$1,613.14		\$322.63	
15824	Removal of forehead wrinkles	T	0137	23.9317	\$1,613.14		\$322.63	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
15111	Epiderm a-grft face/nck/h/g	T	0135	4.4386	\$299.19		\$59.84	
15115	Epiderm a-grft face/nck/h/g	T	0135	4.4386	\$299.19		\$59.84	
15116	Epiderm a-grft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15120	Skin split a-grft face/nck/h/g	T	0137	23.9317	\$1,613.14		\$322.63	
15121	Skin split a-grft f/n/h/g add	T	0137	23.9317	\$1,613.14		\$322.63	
15130	Derm autograft, trnk/arm/leg	T	0136	16.0542	\$1,082.15		\$216.43	
15131	Derm autograft, f/n/h/g add-on	T	0136	16.0542	\$1,082.15		\$216.43	
15135	Derm autograft face/nck/h/g	T	0136	16.0542	\$1,082.15		\$216.43	
15136	Derm autograft, f/n/h/g add	T	0136	16.0542	\$1,082.15		\$216.43	
15150	Cult epiderm, grft f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15151	Cult epiderm, grft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15152	Cult epiderm, grft f/n/h/g +%	T	0135	4.4386	\$299.19		\$59.84	
15155	Cult epiderm, grft, f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15156	Cult epiderm, grft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15157	Cult epiderm, grft f/n/h/g +%	CH						
15171	Acclular, grft, f/n/h/g	T	0134	3.1508	\$212.38		\$42.48	
15175	Acclular, grft, f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15200	Skin full graft, trunk	T	0136	16.0542	\$1,082.15		\$216.43	
15201	Skin full graft trunk add-on	T	0136	16.0542	\$1,082.15		\$216.43	
15220	Skin full graft, sep/arm/leg	T	0136	16.0542	\$1,082.15		\$216.43	
15221	Skin full graft add-on	T	0135	4.4386	\$299.19		\$59.84	
15240	Skin full graft face/genit/hf	T	0136	16.0542	\$1,082.15		\$216.43	
15241	Skin full graft add-on	T	0135	4.4386	\$299.19		\$59.84	
15260	Skin full graft een & lips	T	0136	16.0542	\$1,082.15		\$216.43	
15261	Skin full graft add-on	T	0135	4.4386	\$299.19		\$59.84	
15300	Apply sknallogrt, t/arm/leg	T	0135	4.4386	\$299.19		\$59.84	
15301	Apply sknallogrt, f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15320	Apply skin allogrt f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15321	Apply skin allogrt f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15330	Apply acell grft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15331	Apply acell grft f/n/h/g add-on	T	0135	4.4386	\$299.19		\$59.84	
15335	Apply acell grft, f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15336	Apply acell grft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	
15340	Apply cut skin substitute	T	0134	3.1508	\$212.38		\$42.48	
15341	Apply cut skin sub add-on	T	0134	3.1508	\$212.38		\$42.48	
15360	Apply cult derm sub, f/n/h/g	T	0134	3.1508	\$212.38		\$42.48	
15361	Apply cult derm sub f/n/h/g add	T	0134	3.1508	\$212.38		\$42.48	
15365	Apply cult derm sub f/n/h/g	T	0134	3.1508	\$212.38		\$42.48	
15366	Apply cult derm f/n/h/g add	T	0134	3.1508	\$212.38		\$42.48	
15400	Apply skin xenograft, f/n/h/g	T	0135	4.4386	\$299.19		\$59.84	
15401	Apply skin xenograft f/n/h/g add	T	0135	4.4386	\$299.19		\$59.84	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
15956	Remove thigh pressure sore	T	T	0136	16.0542	\$1,082.15		\$216.43
15958	Remove thigh pressure sore	T	T	0136	16.0542	\$1,082.15		\$216.43
15989	Removal of pressure sore	T	T	0019	4.3625	\$294.06	\$64.51	\$58.82
16000	Initial treatment of burn(s)	T	T	0013	0.8789	\$59.24		\$11.85
16020	Dress/debrid p-thick burn, s	T	T	0015	1.5412	\$103.89		\$20.78
16025	Dress/debrid p-thick burn, m	T	T	0015	1.5412	\$103.89		\$20.78
16030	Dress/debrid p-thick burn, l	T	T	0015	1.5412	\$103.89		\$20.78
16035	Incision of burn scab, infl	T	T	0015	1.5412	\$103.89		\$20.78
16036	Escharotomy, addl incision	C						
17000	Deconstruct premalign lesion	T	T	0013	0.8789	\$59.24		\$11.85
17003	Deconstruct premalign les, 2-14	T	T	0012	0.4436	\$29.90		\$5.98
17004	Destroy premig lesions 15+	T	T	0016	2.7982	\$188.62		\$37.73
17106	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17107	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17108	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17110	Destruct b9 lesion, 1-14	T	T	0013	0.8789	\$59.24		\$11.85
17111	Destruct lesion, 15 or more	T	T	0015	1.5412	\$103.89		\$20.78
17250	Chemical cautery, tissue	T	T	0015	1.5412	\$103.89		\$20.78
17260	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17261	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17262	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17263	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17264	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17266	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17270	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17271	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17272	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17273	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17274	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17276	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17280	Destruction of skin lesions	T	T	0015	1.5412	\$103.89		\$20.78
17281	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17282	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17283	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17284	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17296	Destruction of skin lesions	T	T	0016	2.7982	\$188.62		\$37.73
17311	Mohs, 1 stage, h/m/h/g	T	T	0694	4.9337	\$332.56	\$91.69	\$66.52
17312	Mohs addl stage	T	T	0694	4.9337	\$332.56	\$91.69	\$66.52
17313	Mohs, 1 stage, V/a/l	T	T	0694	4.9337	\$332.56	\$91.69	\$66.52
17314	Mohs, addl stage, t/a/l	T	T	0694	4.9337	\$332.56	\$91.69	\$66.52
17315	Mohs surg, addl block	T	T	0694	4.9337	\$332.56	\$91.69	\$66.52
17340	Cryotherapy of skin	T	T	0013	0.8789	\$59.24		\$11.85
17360	Skin peel therapy	T	T	0013	0.8789	\$59.24		\$11.85

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
15825	Removal of neck wrinkles	T	T	0137	23.9317	\$1,613.14		\$322.63
15826	Removal of brow wrinkles	T	T	0137	23.9317	\$1,613.14		\$322.63
15828	Removal of face wrinkles	T	T	0137	23.9317	\$1,613.14		\$322.63
15829	Removal of skin wrinkles	T	T	0137	23.9317	\$1,613.14		\$322.63
15830	Exc skin abd	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15832	Excise excessive skin tissue	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15833	Excise excessive skin tissue	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15834	Excise excessive skin tissue	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15835	Excise excessive skin tissue	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15836	Excise excessive skin tissue	T	T	0021	17.4975	\$1,179.44		\$235.89
15837	Excise excessive skin tissue	T	T	0021	17.4975	\$1,179.44		\$235.89
15838	Excise excessive skin tissue	T	T	0021	17.4975	\$1,179.44		\$235.89
15839	Excise excessive skin tissue	T	T	0021	17.4975	\$1,179.44		\$235.89
15840	Graft for face nerve palsy	T	T	0137	23.9317	\$1,613.14		\$322.63
15841	Graft for face nerve palsy	T	T	0137	23.9317	\$1,613.14		\$322.63
15845	Skin and muscle repair, face	T	T	0137	23.9317	\$1,613.14		\$322.63
15847	Exc skin abd add-on	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15850	Removal of sutures	T	T	0016	2.7982	\$188.62		\$37.73
15851	Removal of sutures	T	T	0016	2.7982	\$188.62		\$37.73
15852	Dressing change not for burn	X	X	0340	0.6693	\$45.11		\$9.03
15860	Test for blood flow in graft	X	X	0340	0.6693	\$45.11		\$9.03
15876	Suction assisted lipectomy	T	T	0137	23.9317	\$1,613.14		\$322.63
15877	Suction assisted lipectomy	T	T	0137	23.9317	\$1,613.14		\$322.63
15878	Suction assisted lipectomy	T	T	0137	23.9317	\$1,613.14		\$322.63
15879	Suction assisted lipectomy	T	T	0137	23.9317	\$1,613.14		\$322.63
15920	Removal of fat bone ulcer	T	T	0019	4.3625	\$294.06	\$64.51	\$58.82
15922	Removal of fat bone ulcer	T	T	0137	23.9317	\$1,613.14		\$322.63
15931	Remove sacrum pressure sore	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15933	Remove sacrum pressure sore	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15934	Remove sacrum pressure sore	T	T	0137	23.9317	\$1,613.14		\$322.63
15935	Remove sacrum pressure sore	T	T	0137	23.9317	\$1,613.14		\$322.63
15936	Remove sacrum pressure sore	T	T	0136	16.0542	\$1,082.15		\$216.43
15937	Remove sacrum pressure sore	T	T	0137	23.9317	\$1,613.14		\$322.63
15940	Remove hip pressure sore	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15941	Remove hip pressure sore	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15944	Remove hip pressure sore	T	T	0137	23.9317	\$1,613.14		\$322.63
15945	Remove hip pressure sore	T	T	0137	23.9317	\$1,613.14		\$322.63
15950	Remove thigh pressure sore	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15951	Remove thigh pressure sore	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
15952	Remove thigh pressure sore	T	T	0136	16.0542	\$1,082.15		\$216.43
15953	Remove thigh pressure sore	T	T	0136	16.0542	\$1,082.15		\$216.43

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
19357	Breast reconstruction	T	T	0648	58.2278	\$3,924.90		\$784.98
19361	Breast reconstruction w/lat flap	C	C					
19364	Breast reconstruction	C	C					
19366	Breast reconstruction	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19367	Breast reconstruction	C	C					
19368	Breast reconstruction	C	C					
19369	Breast reconstruction	C	C					
19370	Surgery of breast capsule	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19371	Removal of breast capsule	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19380	Revise breast reconstruction	T	T	0030	41.9997	\$2,831.03	\$747.07	\$566.21
19386	Design custom breast implant	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19396	Breast surgery procedure	T	T	0028	24.7516	\$1,668.41		\$333.69
19499	Incision of abscess	T	T	0006	1.4557	\$98.12		\$19.63
20000	Incision of deep abscess	T	T	0049	22.0149	\$1,483.94		\$296.79
20005	Explore wound, neck	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
20100	Explore wound, chest	T	T	0137	23.9317	\$1,613.14		\$322.63
20102	Explore wound, abdomen	T	T	0137	23.9317	\$1,613.14		\$322.63
20103	Explore wound, extremity	CH	T	0007	12.6217	\$860.78		\$170.16
20150	Excise epiphyseal bar	T	T	0051	46.5786	\$3,139.68		\$627.94
20200	Muscle biopsy	T	T	0021	17.4975	\$1,179.44		\$235.89
20205	Deep muscle biopsy	T	T	0021	17.4975	\$1,179.44		\$235.89
20206	Needle biopsy, muscle	T	T	0005	7.8145	\$526.74		\$105.35
20220	Bone biopsy, trocar/needle	T	T	0020	8.2028	\$552.92		\$110.59
20225	Bone biopsy, trocar/needle	T	T	0021	17.4975	\$1,179.44		\$235.89
20240	Bone biopsy, excisional	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
20245	Bone biopsy, excisional	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
20250	Open bone biopsy	T	T	0049	22.0149	\$1,483.94		\$296.79
20251	Open bone biopsy	T	T	0049	22.0149	\$1,483.94		\$296.79
20500	Injection of sinus tract	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
20501	Inject sinus tract for x-ray	N	N					
20520	Removal of foreign body	T	T	0019	4.3625	\$294.06	\$64.51	\$58.82
20525	Removal of foreign body	T	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
20526	Thor injection, carp tunnel	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20550	Inf tendon sheath/ligament	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20551	Inf tendon origin/insertion	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20552	Inf trigger point, 1/2 muscl	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20553	Inject trigger points, => 3	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20555	Place ind muscils for rt	T	T	0050	31.7717	\$2,141.60	\$428.32	\$387.46
20600	Drain/inject, joint/bursa	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20605	Drain/inject, joint/bursa	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20610	Drain/inject, joint/bursa	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20612	Aspirate/inf joint/bursa cyst	T	T	0204	2.5558	\$172.28	\$40.13	\$34.46
20615	Treatment of bone cyst	T	T	0004	4.5991	\$310.01		\$62.01

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
17380	Hair removal by electrolysis	T	T	0013	0.8789	\$59.24		\$11.85
17999	Skin tissue procedure	T	T	0012	0.4436	\$29.90		\$5.98
19000	Drainage of breast lesion	T	T	0004	4.5991	\$310.01		\$62.01
19001	Drain breast lesion add-on	T	T	0002	1.5111	\$101.86		\$20.38
19020	Incision of breast lesion	T	T	0008	19.4063	\$1,308.10		\$261.62
19030	Injection for breast x-ray	N	N					
19100	Bx breast percut w/o image	T	T	0004	4.5991	\$310.01		\$62.01
19101	Biopsy of breast, open	T	T	0028	24.7516	\$1,668.41		\$333.69
19102	Bx breast percut w/image	T	T	0005	7.8145	\$526.74		\$105.35
19103	Bx breast percut w/device	T	T	0037	15.0003	\$1,044.81	\$228.76	\$208.87
19105	Cryosurg ablate fa, each	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19110	Nipple exploration	T	T	0028	24.7516	\$1,668.41		\$333.69
19112	Excise breast duct fistula	T	T	0028	24.7516	\$1,668.41		\$333.69
19120	Removal of breast lesion	T	T	0028	24.7516	\$1,668.41		\$333.69
19125	Excision, breast lesion	T	T	0028	24.7516	\$1,668.41		\$333.69
19126	Excision, addl breast lesion	T	T	0028	24.7516	\$1,668.41		\$333.69
19260	Removal of chest wall lesion	T	T	0021	17.4975	\$1,179.44		\$235.89
19271	Revision of chest wall	C	C					
19272	Extensive chest wall surgery	C	C					
19290	Place needle wire, breast	N	N					
19291	Place needle wire, breast	N	N					
19295	Place breast clip, percut	N	N					
19296	Place po breast cath for rad	T	T	0648	58.2278	\$3,924.90		\$784.98
19297	Place breast cath for rad	T	T	0648	58.2278	\$3,924.90		\$784.98
19298	Place breast rad tube/caths	T	T	0648	58.2278	\$3,924.90		\$784.98
19300	Removal of breast tissue	T	T	0028	24.7516	\$1,668.41		\$333.69
19301	Partial mastectomy	T	T	0028	24.7516	\$1,668.41		\$333.69
19302	P-mastectomy w/in removal	T	T	0030	41.9997	\$2,831.03	\$747.07	\$566.21
19303	Mast, simple, complete	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19304	Mast, subq	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19305	Mast, radical	C	C					
19306	Mast, rad, urban type	C	C					
19307	Mast, mod rad	T	T	0030	41.9997	\$2,831.03	\$747.07	\$566.21
19316	Suspension of breast	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19318	Reduction of large breast	T	T	0030	41.9997	\$2,831.03	\$747.07	\$566.21
19324	Enlarge breast	T	T	0030	41.9997	\$2,831.03	\$747.07	\$566.21
19325	Enlarge breast with implant	T	T	0648	58.2278	\$3,924.90		\$784.98
19328	Removal of breast implant	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19330	Removal of implant material	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59
19340	Immediate breast prosthesis	T	T	0030	41.9997	\$2,831.03	\$747.07	\$566.21
19342	Delayed breast prosthesis	T	T	0648	58.2278	\$3,924.90		\$784.98
19350	Breast reconstruction	T	T	0028	24.7516	\$1,668.41		\$333.69
19355	Correct inverted nipple(s)	T	T	0029	34.1654	\$2,302.95	\$581.52	\$460.59

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
20972	Bone/skin graft, metatarsal	T	T	0056	52.5258	\$3,540.55		\$708.11
20973	Bone/skin graft, great toe	T	T	0056	52.5258	\$3,540.55		\$708.11
20974	Electrical bone stimulation	A						
20975	Electrical bone stimulation	X	N	0340	0.6693	\$45.11		\$9.03
20979	Us bone stimulation	X	N	0051	46.5786	\$3,139.68		\$627.94
20982	Ablate, bone tumor(s) petq	T	N					
20985	Cptr-assst dir ms px	T	N					
20989	Musculoskeletal surgery	T	T	0049	22.0149	\$1,483.94		\$296.79
21010	Incision of jaw joint	T	T	0254	24.9637	\$1,682.70		\$336.54
21011	Exc face les sc < 2 cm	NI	T	0020	8.2028	\$552.92		\$110.59
21012	Exc face les sc = 2 cm	NI	T	0020	8.2028	\$552.92		\$110.59
21013	Exc face turn deep < 2 cm	NI	T	0020	8.2028	\$552.92		\$110.59
21014	Exc face turn deep = 2 cm	NI	T	0020	8.2028	\$552.92		\$110.59
21015	Resect face turn < 2 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
21016	Resect face turn = 2 cm	NI	T	0022	23.988	\$1,576.49	\$354.45	\$315.30
21025	Excision of bone, lower jaw	T	T	0256	42.9827	\$2,897.29		\$579.46
21026	Excision of facial bone(s)	T	T	0256	42.9827	\$2,897.29		\$579.46
21029	Contour of face bone lesion	T	T	0256	42.9827	\$2,897.29		\$579.46
21030	Excise max/zygoma b9 tumor	T	T	0254	24.9637	\$1,682.70		\$336.54
21031	Remove exostosis, mandible	T	T	0254	24.9637	\$1,682.70		\$336.54
21032	Remove exostosis, maxilla	T	T	0254	24.9637	\$1,682.70		\$336.54
21034	Excise max/zygoma mlg tumor	T	T	0256	42.9827	\$2,897.29		\$579.46
21040	Excise mandible lesion	T	T	0254	24.9637	\$1,682.70		\$336.54
21044	Removal of jaw bone lesion	T	T	0256	42.9827	\$2,897.29		\$579.46
21045	Extensive jaw surgery	C						
21046	Remove mandible cyst complex	T	T	0256	42.9827	\$2,897.29		\$579.46
21047	Excise lwr jaw cyst w/repair	T	T	0256	42.9827	\$2,897.29		\$579.46
21048	Remove maxilla cyst complex	T	T	0256	42.9827	\$2,897.29		\$579.46
21049	Excis uppr jaw cyst w/repair	T	T	0256	42.9827	\$2,897.29		\$579.46
21050	Removal of jaw joint	T	T	0256	42.9827	\$2,897.29		\$579.46
21050	Remove jaw joint cartilage	T	T	0256	42.9827	\$2,897.29		\$579.46
21070	Remove coronoid process	T	T	0256	42.9827	\$2,897.29		\$579.46
21073	Mingl of tmj w/anessth	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
21076	Prepare face/oral prosthesis	T	T	0254	24.9637	\$1,682.70		\$336.54
21077	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21079	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21080	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21081	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21082	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21083	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21084	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46
21085	Prepare face/oral prosthesis	T	T	0253	17.1879	\$1,168.57	\$282.29	\$231.72
21086	Prepare face/oral prosthesis	T	T	0256	42.9827	\$2,897.29		\$579.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
20650	Insert and remove bone pin	T	T	0049	22.0149	\$1,483.94		\$296.79
20660	Apply, rem fixation device	T	T	0138	4.7946	\$323.18		\$64.64
20661	Application of head brace	C						
20662	Application of pelvis brace	T	T	0049	22.0149	\$1,483.94		\$296.79
20663	Application of thigh brace	T	T	0049	22.0149	\$1,483.94		\$296.79
20664	Halo brace application	C						
20665	Removal of fixation device	X	N	0340	0.6693	\$45.11		\$9.03
20670	Removal of support implant	T	T	0021	17.4975	\$1,179.44		\$235.89
20680	Removal of support implant	T	T	0022	23.988	\$1,576.49	\$354.45	\$315.30
20690	Apply bone fixation device	T	T	0050	31.7717	\$2,141.60		\$428.32
20692	Apply bone fixation device	T	T	0050	31.7717	\$2,141.60		\$428.32
20693	Adjust bone fixation device	T	T	0049	22.0149	\$1,483.94		\$296.79
20694	Remove bone fixation device	T	T	0049	22.0149	\$1,483.94		\$296.79
20696	Comp multiplane ext fixation	T	T	0050	31.7717	\$2,141.60		\$428.32
20697	Comp ext fixate strut change	T	T	0139	18.9982	\$1,240.01		\$248.01
20802	Replantation, arm, complete	C						
20805	Replant forearm, complete	C						
20816	Replantation hand, complete	C						
20822	Replantation digit, complete	C						
20824	Replantation thumb, complete	C						
20827	Replantation thumb, complete	C						
20838	Replantation foot, complete	C						
20900	Removal of bone for graft	T	T	0050	31.7717	\$2,141.60		\$428.32
20910	Remove cartilage for graft	T	T	0050	31.7717	\$2,141.60		\$428.32
20912	Remove cartilage for graft	T	T	0137	23.9317	\$1,613.14		\$322.63
20920	Removal of fascia for graft	T	T	0136	16.0542	\$1,082.15		\$216.43
20922	Removal of fascia for graft	T	T	0136	16.0542	\$1,082.15		\$216.43
20924	Removal of tendon for graft	T	T	0050	31.7717	\$2,141.60		\$428.32
20926	Removal of tissue for graft	T	T	0135	4.4386	\$299.19		\$59.84
20930	Sp bone algrft morsel add-on	C						
20931	Sp bone algrft local add-on	C						
20936	Sp bone agrft local add-on	C						
20937	Sp bone agrft morsel add-on	C						
20938	Sp bone agrft struct add-on	C						
20950	Fluid pressure, muscle	T	T	0006	1.4557	\$98.12		\$19.63
20955	Fibula bone graft, microvasc	C						
20956	Iliac bone graft, microvasc	C						
20957	Mt bone graft, microvasc	C						
20962	Other bone graft, microvasc	C						
20969	Bone/skin graft, microvasc	C						
20970	Bone/skin graft, iliac crest	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
21208	Augmentation of facial bones	T	0256	42.9827	\$2,897.29			\$579.46
21209	Reduction of facial bones	T	0256	42.9827	\$2,897.29			\$579.46
21210	Face bone graft	T	0256	42.9827	\$2,897.29			\$579.46
21215	Lower jaw bone graft	T	0256	42.9827	\$2,897.29			\$579.46
21230	Rib cartilage graft	T	0256	42.9827	\$2,897.29			\$579.46
21235	Ear cartilage graft	T	0254	24.9637	\$1,682.70			\$336.54
21240	Reconstruction of jaw joint	T	0256	42.9827	\$2,897.29			\$579.46
21242	Reconstruction of jaw joint	T	0256	42.9827	\$2,897.29			\$579.46
21243	Reconstruction of jaw joint	T	0256	42.9827	\$2,897.29			\$579.46
21244	Reconstruction of lower jaw	T	0256	42.9827	\$2,897.29			\$579.46
21245	Reconstruction of jaw	T	0256	42.9827	\$2,897.29			\$579.46
21246	Reconstruction of jaw	T	0256	42.9827	\$2,897.29			\$579.46
21247	Reconstruct lower jaw bone	C						
21248	Reconstruction of jaw	T	0256	42.9827	\$2,897.29			\$579.46
21249	Reconstruction of jaw	T	0256	42.9827	\$2,897.29			\$579.46
21255	Reconstruct lower jaw bone	C						
21256	Reconstruction of orbit	CH	0256	42.9827	\$2,897.29			\$579.46
21260	Revise eye sockets	T	0256	42.9827	\$2,897.29			\$579.46
21261	Revise eye sockets	T	0256	42.9827	\$2,897.29			\$579.46
21263	Revise eye sockets	T	0256	42.9827	\$2,897.29			\$579.46
21267	Revise eye sockets	T	0256	42.9827	\$2,897.29			\$579.46
21268	Revise eye sockets	C						
21270	Augmentation, cheek bone	T	0256	42.9827	\$2,897.29			\$579.46
21275	Revision, orbitofacial bones	T	0256	42.9827	\$2,897.29			\$579.46
21280	Revision of eyelid	T	0256	42.9827	\$2,897.29			\$579.46
21282	Revision of eyelid	T	0253	17.1879	\$1,158.57	\$282.29		\$231.72
21295	Revision of jaw muscle/bone	T	0252	7.6196	\$513.61	\$109.16		\$102.73
21296	Revision of jaw muscle/bone	T	0254	24.9637	\$1,682.70			\$336.54
21299	Cranio/maxillofacial surgery	T	0250	1.1522	\$77.67	\$25.10		\$15.54
21310	Treatment of nose fracture	T	0250	1.1522	\$77.67	\$25.10		\$15.54
21325	Treatment of nose fracture	T	0253	17.1879	\$1,158.57	\$282.29		\$231.72
21326	Treatment of nose fracture	T	0254	24.9637	\$1,682.70	\$282.29		\$231.72
21330	Treatment of nose fracture	T	0254	24.9637	\$1,682.70			\$336.54
21335	Treatment of nose fracture	T	0254	24.9637	\$1,682.70			\$336.54
21336	Treat nasal septal fracture	T	0062	25.8446	\$1,742.08	\$372.87		\$348.42
21337	Treat nasal septal fracture	T	0253	17.1879	\$1,158.57	\$282.29		\$231.72
21338	Treat nasosethmoid fracture	T	0254	24.9637	\$1,682.70			\$336.54
21339	Treat nasosethmoid fracture	T	0254	24.9637	\$1,682.70			\$336.54
21340	Treatment of nose fracture	T	0256	42.9827	\$2,897.29			\$579.46
21343	Treatment of sinus fracture	C						
21344	Treatment of sinus fracture	C						
21345	Treat nose/jaw fracture	T	0254	24.9637	\$1,682.70			\$336.54

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
21087	Prepare face/oral prosthesis	T	0256	42.9827	\$2,897.29			\$579.46
21088	Prepare face/oral prosthesis	T	0256	42.9827	\$2,897.29			\$579.46
21089	Prepare face/oral prosthesis	T	0250	1.1522	\$77.67	\$25.10		\$15.54
21100	Maxillofacial fixation	T	0256	42.9827	\$2,897.29			\$579.46
21110	Interdental fixation	T	0252	7.6196	\$513.61	\$109.16		\$102.73
21116	Injection, jaw joint x-ray	N						
21120	Reconstruction of chin	T	0254	24.9637	\$1,682.70			\$336.54
21121	Reconstruction of chin	T	0254	24.9637	\$1,682.70			\$336.54
21122	Reconstruction of chin	T	0254	24.9637	\$1,682.70			\$336.54
21123	Reconstruction of chin	T	0254	24.9637	\$1,682.70			\$336.54
21125	Augmentation, lower jaw bone	T	0254	24.9637	\$1,682.70			\$336.54
21127	Augmentation, lower jaw bone	T	0256	42.9827	\$2,897.29			\$579.46
21137	Reduction of forehead	T	0254	24.9637	\$1,682.70			\$336.54
21138	Reduction of forehead	T	0256	42.9827	\$2,897.29			\$579.46
21139	Reduction of forehead	T	0256	42.9827	\$2,897.29			\$579.46
21141	Reconstruct midface, left	C						
21142	Reconstruct midface, left	C						
21143	Reconstruct midface, left	C						
21145	Reconstruct midface, left	C						
21146	Reconstruct midface, left	C						
21147	Reconstruct midface, left	C						
21150	Reconstruct midface, left	C						\$579.46
21151	Reconstruct midface, left	C						
21154	Reconstruct midface, left	C						
21155	Reconstruct midface, left	C						
21159	Reconstruct midface, left	C						
21160	Reconstruct midface, left	C						
21172	Reconstruct orbit/forehead	T	0256	42.9827	\$2,897.29			\$579.46
21175	Reconstruct orbit/forehead	T	0256	42.9827	\$2,897.29			\$579.46
21179	Reconstruct entire forehead	C						
21180	Reconstruct entire forehead	C						
21181	Contour cranial bone lesion	T	0254	24.9637	\$1,682.70			\$336.54
21182	Reconstruct cranial bone	C						
21183	Reconstruct cranial bone	C						
21184	Reconstruct cranial bone	C						
21188	Reconstruction of midface	C						
21193	Reconst lwr jaw w/o graft	C						
21194	Reconst lwr jaw w/graft	T	0256	42.9827	\$2,897.29			\$579.46
21195	Reconst lwr jaw w/o fixation	T	0256	42.9827	\$2,897.29			\$579.46
21196	Reconst lwr jaw w/fixation	C						
21198	Reconst lwr jaw segment	T	0256	42.9827	\$2,897.29			\$579.46
21199	Reconst lwr jaw w/advance	T	0256	42.9827	\$2,897.29			\$579.46
21206	Reconstruct upper jaw bone	T	0256	42.9827	\$2,897.29			\$579.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
21501	Drain neck/chest lesion	T	0008	0049	19.4063	\$1,308.10		\$261.62
21502	Drain chest lesion	T	0049	0049	22.0149	\$1,483.94		\$296.79
21510	Drainage of bone lesion	C						
21550	Biopsy of neck/chest	T	0021	0021	17.4975	\$1,179.44		\$235.89
21552	Exc neck les sc = 3 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
21554	Exc neck tum deep = 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
21555	Exc neck les sc < 3 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
21556	Exc neck tum deep < 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
21557	Resect neck tum < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
21558	Resect neck tum = 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
21600	Partial removal of rib	T	0050	0050	31.7717	\$2,141.60		\$428.32
21610	Partial removal of rib	T	0050	0050	31.7717	\$2,141.60		\$428.32
21615	Removal of rib	C						
21616	Removal of rib and nerves	C						
21620	Partial removal of sternum	C						
21627	Sternal debridement	C						
21630	Extensive sternum surgery	C						
21632	Extensive sternum surgery	C						
21685	Hyoid myotomy & suspension	T	0252	0252	7.6196	\$513.61	\$109.16	\$102.73
21700	Revision of neck muscle	T	0049	0049	22.0149	\$1,483.94		\$296.79
21705	Revision of neck muscle/rib	C						
21720	Revision of neck muscle	T	0049	0049	22.0149	\$1,483.94		\$296.79
21725	Revision of neck muscle	T	0006	0006	1.4657	\$98.12		\$19.63
21740	Reconstruction of sternum	C						
21742	Repair sternum/muss w/ scope	T	0051	0051	46.5786	\$3,139.68		\$627.94
21743	Repair sternum/muss w/ scope	T	0051	0051	46.5786	\$3,139.68		\$627.94
21750	Repair of sternum separation	C						
21800	Treatment of rib fracture	T	0129	0129	1.6576	\$111.73		\$22.35
21805	Treatment of rib fracture	T	0062	0062	25.8446	\$1,742.08	\$372.87	\$348.42
21810	Treatment of rib fracture(s)	C						
21820	Treat sternum fracture	T	0129	0129	1.6576	\$111.73		\$22.35
21825	Treat sternum fracture	C						
21899	Neck/chest surgery procedure	T	0250	0250	1.1522	\$77.67	\$25.10	\$15.54
21920	Biopsy soft tissue of back	T	0020	0020	8.2028	\$552.92		\$10.59
21925	Biopsy soft tissue of back	T	0022	0022	23.388	\$1,576.49	\$354.45	\$315.30
21930	Exc back les sc < 3 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
21931	Exc back les sc = 3 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
21932	Exc back tum deep < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
21933	Exc back tum deep = 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
21935	Resect back tum < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
21936	Resect back tum = 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
22010	l&d, p-spine, c/c/cerv-thor	C						
22015	l&d, p-spine, l/s/l/s	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
21346	Treat nose/jaw fracture	C						
21347	Treat nose/jaw fracture	C						
21348	Treat nose/jaw fracture	C						
21355	Treat cheek bone fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21356	Treat cheek bone fracture	T	0254	0254	24.9637	\$1,682.70		\$336.54
21360	Treat cheek bone fracture	T	0254	0254	24.9637	\$1,682.70		\$336.54
21365	Treat cheek bone fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21366	Treat cheek bone fracture	C						
21385	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21386	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21387	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21390	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21395	Treat eye socket fracture	C						
21400	Treat eye socket fracture	T	0252	0252	7.6196	\$513.61	\$109.16	\$102.73
21401	Treat eye socket fracture	T	0253	0253	17.1879	\$1,158.57	\$282.29	\$231.72
21406	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21407	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21408	Treat eye socket fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21421	Treat mouth roof fracture	T	0254	0254	24.9637	\$1,682.70		\$336.54
21422	Treat mouth roof fracture	C						
21423	Treat mouth roof fracture	C						
21431	Treat craniofacial fracture	C						
21432	Treat craniofacial fracture	C						
21433	Treat craniofacial fracture	C						
21435	Treat craniofacial fracture	C						
21436	Treat craniofacial fracture	C						
21440	Treat dental ridge fracture	T	0254	0254	24.9637	\$1,682.70		\$336.54
21445	Treat dental ridge fracture	T	0254	0254	24.9637	\$1,682.70		\$336.54
21450	Treat lower jaw fracture	T	0251	0251	3.425	\$230.87	\$46.18	\$46.18
21451	Treat lower jaw fracture	T	0252	0252	7.6196	\$513.61	\$109.16	\$102.73
21452	Treat lower jaw fracture	T	0253	0253	17.1879	\$1,158.57	\$282.29	\$231.72
21453	Treat lower jaw fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21454	Treat lower jaw fracture	T	0254	0254	24.9637	\$1,682.70		\$336.54
21461	Treat lower jaw fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21462	Treat lower jaw fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21465	Treat lower jaw fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21470	Treat lower jaw fracture	T	0256	0256	42.9827	\$2,897.29		\$579.46
21480	Reset dislocated jaw	T	0250	0250	1.1522	\$77.67	\$25.10	\$15.54
21485	Reset dislocated jaw	T	0253	0253	17.1879	\$1,158.57	\$282.29	\$231.72
21490	Repair dislocated jaw	T	0256	0256	42.9827	\$2,897.29		\$579.46
21495	Treat hyoid bone fracture	T	0253	0253	17.1879	\$1,158.57	\$282.29	\$231.72
21497	Interdental wiring	T	0253	0253	17.1879	\$1,158.57	\$282.29	\$231.72
21499	Head surgery procedure	T	0250	0250	1.1522	\$77.67	\$25.10	\$15.54

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
22568	Lumbar spine fusion		C					
22585	Additional spinal fusion		C					
22590	Spine & skull spinal fusion		C					
22595	Neck spinal fusion		C					
22600	Neck spine fusion		C					
22610	Thorax spine fusion		C					
22612	Lumbar spine fusion		T	0208	49.2256	\$3,318.10		\$663.62
22614	Spine fusion, extra segment		T	0208	49.2256	\$3,318.10		\$663.62
22630	Lumbar spine fusion		C					
22632	Spine fusion, extra segment		C					
22800	Fusion of spine		C					
22802	Fusion of spine		C					
22804	Fusion of spine		C					
22808	Fusion of spine		C					
22810	Fusion of spine		C					
22818	Kyphectomy, 1-2 segments		C					
22819	Kyphectomy, 3 or more		C					
22830	Exploration of spinal fusion		C					
22840	Insert spine fixation device		C					
22841	Insert spine fixation device		C					
22842	Insert spine fixation device		C					
22843	Insert spine fixation device		C					
22844	Insert spine fixation device		C					
22845	Insert spine fixation device		C					
22846	Insert spine fixation device		C					
22847	Insert spine fixation device		C					
22848	Insert pelv fixation device		C					
22849	Reinsert spinal fixation		C					
22850	Remove spine fixation device		T	0049	22.0149	\$1,483.94		\$296.79
22851	Apply spine prosth device		C					
22852	Remove spine fixation device		C					
22855	Remove spine fixation device		C					
22856	Cerv artilc. diskectomy		C					
22857	Lumbar artilc diskectomy		C					
22861	Revise cerv artilc disc		C					
22862	Revise lumbar artilc disc		C					
22864	Remove cerv artilc disc		C					
22865	Remove lumbar artilc disc		C					
22899	Spine surgery procedure		T	0049	22.0149	\$1,483.94		\$296.79
22900	Exc back tum deep < 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
22901	Exc back tum deep = 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
22902	Exc abd les sc < 3 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
22100	Remove part of neck vertebra		T	0208	49.2256	\$3,318.10		\$663.62
22101	Remove part, thorax vertebra		T	0208	49.2256	\$3,318.10		\$663.62
22102	Remove part, lumbar vertebra		T	0208	49.2256	\$3,318.10		\$663.62
22103	Remove extra spine segment		T	0208	49.2256	\$3,318.10		\$663.62
22110	Remove part of neck vertebra		C					
22112	Remove part, thorax vertebra		C					
22114	Remove part, lumbar vertebra		C					
22116	Remove extra spine segment		C					
22206	Cut spine 3 col, thor		C					
22207	Cut spine 3 col, lumb		C					
22208	Cut spine 3 col, addl seg		C					
22210	Revision of neck spine		C					
22212	Revision of thorax spine		C					
22214	Revision of lumbar spine		C					
22216	Revise, extra spine segment		C					
22220	Revision of neck spine		C					
22222	Revision of thorax spine		C	0208	49.2256	\$3,318.10		\$663.62
22224	Revision of lumbar spine		C					
22226	Revise, extra spine segment		C					
22305	Treat spine process fracture		T	0129	1.6576	\$111.73		\$22.35
22310	Treat spine fracture		T	0138	4.7946	\$323.18		\$64.64
22315	Treat spine fracture		T	0139	18.3962	\$1,240.01		\$248.01
22318	Treat odontoid fx w/o graft		C					
22319	Treat odontoid fx w/graft		C					
22325	Treat spine fracture		C					
22326	Treat neck spine fracture		C					
22327	Treat thorax spine fracture		C					
22328	Treat each add spine fx		C					
22505	Manipulation of spine		T	0045	15.2922	\$1,030.79	\$268.47	\$206.16
22520	Percut vertebroplasty thor		T	0050	31.7717	\$2,141.60		\$428.32
22521	Percut vertebroplasty lumb		T	0050	31.7717	\$2,141.60		\$428.32
22522	Percut vertebroplasty addl		T	0050	31.7717	\$2,141.60		\$428.32
22523	Percut kyphoplasty, thor		T	0052	88.6521	\$5,975.68		\$1,195.14
22524	Percut kyphoplasty, lumbar		T	0052	88.6521	\$5,975.68		\$1,195.14
22525	Percut kyphoplasty, add-on		T	0052	88.6521	\$5,975.68		\$1,195.14
22526	Idet, single level		E					
22527	Idet, 1 or more levels		E					
22532	Lat thorax spine fusion		C					
22533	Lat lumbar spine fusion		C					
22534	Lat thor/lumb, addl seg		C					
22548	Neck spine fusion		C					
22554	Neck spine fusion		C					
22556	Thorax spine fusion		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
23220	Resect prox humerus tumor	C	D					
23221	Partial removal of humerus	CH	D					
23222	Partial removal of humerus	CH	D					
23330	Remove shoulder foreign body	T	D	0020	8.2028	\$552.92		\$110.59
23331	Remove shoulder foreign body	T	D	0022	23.388	\$1,576.49	\$354.45	\$315.30
23332	Remove shoulder foreign body	T	C					
23350	Injection for shoulder x-ray	N						
23395	Muscle transfer, shoulder/arm	T		0051	46.5786	\$3,139.68		\$627.94
23397	Muscle transfers	T		0052	88.6521	\$5,975.68		\$1,195.14
23400	Fixation of shoulder blade	T		0050	31.7717	\$2,141.60		\$428.32
23405	Incision of tendon & muscle	T		0050	31.7717	\$2,141.60		\$428.32
23406	Incise tendon(s) & muscle(s)	T		0050	31.7717	\$2,141.60		\$428.32
23410	Repair rotator cuff, acute	T		0051	46.5786	\$3,139.68		\$627.94
23412	Repair rotator cuff, chronic	T		0051	46.5786	\$3,139.68		\$627.94
23415	Release of shoulder ligament	T		0051	46.5786	\$3,139.68		\$627.94
23420	Repair of shoulder	T		0051	46.5786	\$3,139.68		\$627.94
23430	Repair/transplant tendon	T		0051	46.5786	\$3,139.68		\$627.94
23450	Repair shoulder capsule	T		0052	88.6521	\$5,975.68		\$1,195.14
23455	Repair shoulder capsule	T		0052	88.6521	\$5,975.68		\$1,195.14
23460	Repair shoulder capsule	T		0052	88.6521	\$5,975.68		\$1,195.14
23462	Repair shoulder capsule	T		0051	46.5786	\$3,139.68		\$627.94
23465	Repair shoulder capsule	T		0052	88.6521	\$5,975.68		\$1,195.14
23466	Repair shoulder capsule	T		0051	46.5786	\$3,139.68		\$627.94
23470	Reconstruct shoulder joint	T		0425	118.767	\$8,005.61		\$1,601.13
23472	Reconstruct shoulder joint	C						
23480	Revision of collar bone	T		0051	46.5786	\$3,139.68		\$627.94
23485	Revision of collar bone	T		0052	88.6521	\$5,975.68		\$1,195.14
23490	Reinforce clavicle	T		0051	46.5786	\$3,139.68		\$627.94
23491	Reinforce shoulder bones	T		0052	88.6521	\$5,975.68		\$1,195.14
23500	Treat clavicle fracture	T		0129	1.6576	\$111.73		\$22.35
23505	Treat clavicle fracture	T		0139	18.3962	\$1,240.01		\$248.01
23515	Treat clavicle fracture	T		0064	65.4752	\$4,413.42		\$882.69
23520	Treat clavicle dislocation	T		0138	4.7946	\$323.18		\$64.64
23525	Treat clavicle dislocation	T		0138	4.7946	\$323.18		\$64.64
23530	Treat clavicle dislocation	T		0063	45.4678	\$3,064.80		\$612.96
23532	Treat clavicle dislocation	T		0062	25.8446	\$1,742.08	\$372.87	\$348.42
23540	Treat clavicle dislocation	T		0129	1.6576	\$111.73		\$22.35
23545	Treat clavicle dislocation	T		0138	4.7946	\$323.18		\$64.64
23550	Treat clavicle dislocation	T		0063	45.4678	\$3,064.80		\$612.96
23552	Treat clavicle dislocation	T		0063	45.4678	\$3,064.80		\$612.96
23570	Treat shoulder blade fx	T		0129	1.6576	\$111.73		\$22.35
23575	Treat shoulder blade fx	T		0138	4.7946	\$323.18		\$64.64

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
22903	Exc. abd les sc > 3 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
22904	Resect abd tum < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
22905	Resect abd tum > 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
22999	Abdomen surgery procedure	T		0049	22.0149	\$1,483.94		\$296.79
23000	Removal of calcium deposits	T		0021	17.4975	\$1,179.44		\$235.89
23020	Release shoulder joint	T		0051	46.5786	\$3,139.68		\$627.94
23030	Drain shoulder lesion	T		0008	19.4063	\$1,308.10		\$261.62
23031	Drain shoulder bursa	T		0008	19.4063	\$1,308.10		\$261.62
23035	Drain shoulder bone lesion	T		0049	22.0149	\$1,483.94		\$296.79
23040	Exploratory shoulder surgery	T		0050	31.7717	\$2,141.60		\$428.32
23044	Exploratory shoulder surgery	T		0050	31.7717	\$2,141.60		\$428.32
23065	Biopsy shoulder tissues	T		0020	8.2028	\$552.92		\$110.59
23066	Biopsy shoulder tissues	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
23071	Exc. shoulder, les sc > 3 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
23073	Exc. shoulder, tum deep > 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
23075	Exc. shoulder, les sc < 3 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
23076	Exc. shoulder, tum deep < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
23077	Resect shoulder tum < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
23078	Resect shoulder tum > 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
23100	Biopsy of shoulder joint	T		0049	22.0149	\$1,483.94		\$296.79
23101	Shoulder joint surgery	T		0050	31.7717	\$2,141.60		\$428.32
23105	Remove shoulder joint lining	T		0050	31.7717	\$2,141.60		\$428.32
23106	Incision of collarbone joint	T		0050	31.7717	\$2,141.60		\$428.32
23107	Explore treat shoulder joint	T		0050	31.7717	\$2,141.60		\$428.32
23120	Partial removal, collar bone	T		0050	31.7717	\$2,141.60		\$428.32
23125	Removal of collar bone	T		0050	31.7717	\$2,141.60		\$428.32
23130	Remove shoulder bone, part	T		0051	46.5786	\$3,139.68		\$627.94
23140	Removal of bone lesion	T		0049	22.0149	\$1,483.94		\$296.79
23145	Removal of bone lesion	T		0050	31.7717	\$2,141.60		\$428.32
23146	Removal of bone lesion	T		0050	31.7717	\$2,141.60		\$428.32
23155	Removal of humerus lesion	T		0050	31.7717	\$2,141.60		\$428.32
23156	Removal of humerus lesion	T		0050	31.7717	\$2,141.60		\$428.32
23170	Remove collar bone lesion	T		0050	31.7717	\$2,141.60		\$428.32
23172	Remove shoulder blade lesion	T		0050	31.7717	\$2,141.60		\$428.32
23174	Remove humerus lesion	T		0050	31.7717	\$2,141.60		\$428.32
23180	Remove collar bone lesion	T		0050	31.7717	\$2,141.60		\$428.32
23182	Remove shoulder blade lesion	T		0050	31.7717	\$2,141.60		\$428.32
23184	Remove humerus lesion	T		0050	31.7717	\$2,141.60		\$428.32
23190	Partial removal of scapula	T		0050	31.7717	\$2,141.60		\$428.32
23195	Removal of head of humerus	T		0050	31.7717	\$2,141.60		\$428.32
23200	Resect clavicle tumor	C						
23210	Resect scapula tumor	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
24125	Remove/graft bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24126	Remove/graft bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24130	Removal of head of radius	T	0050	31.7717	\$2,141.60			\$428.32
24134	Removal of arm bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24136	Remove radius bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24138	Remove elbow bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24140	Partial removal of arm bone	T	0050	31.7717	\$2,141.60			\$428.32
24145	Partial removal of radius	T	0050	31.7717	\$2,141.60			\$428.32
24147	Partial removal of elbow	T	0050	31.7717	\$2,141.60			\$428.32
24149	Radical resection of elbow	T	0050	31.7717	\$2,141.60			\$428.32
24150	Resect distal humerus tumor	T	0051	46.5786	\$3,139.68			\$627.94
24151	Extensive humerus surgery	CH	D					
24152	Resect radius tumor	CH	D					
24153	Extensive radius surgery	CH	D					
24155	Removal of elbow joint	T	0051	46.5786	\$3,139.68			\$627.94
24160	Remove elbow joint implant	T	0050	31.7717	\$2,141.60			\$428.32
24164	Remove radius head implant	T	0050	31.7717	\$2,141.60			\$428.32
24200	Removal of arm foreign body	T	0019	4.3625	\$294.06		\$64.51	\$58.82
24201	Removal of arm foreign body	T	0021	17.4975	\$1,179.44			\$236.89
24220	Injection for elbow x-ray	N						
24300	Manipulate elbow w/aresth	T	0045	15.2922	\$1,030.79		\$268.47	\$206.16
24301	Muscle/tendon transfer	T	0050	31.7717	\$2,141.60			\$428.32
24305	Arm tendon lengthening	T	0050	31.7717	\$2,141.60			\$428.32
24310	Revision of arm tendon	T	0049	22.0149	\$1,483.94			\$296.79
24320	Repair of arm tendon	T	0051	46.5786	\$3,139.68			\$627.94
24330	Revision of arm muscles	T	0052	88.6521	\$5,975.68			\$1,195.14
24331	Revision of arm muscles	T	0051	46.5786	\$3,139.68			\$627.94
24332	Tenolysis, triceps	T	0049	22.0149	\$1,483.94			\$296.79
24340	Repair of biceps tendon	T	0051	46.5786	\$3,139.68			\$627.94
24341	Repair arm tendon/muscle	T	0051	46.5786	\$3,139.68			\$627.94
24342	Repair of ruptured tendon	T	0051	46.5786	\$3,139.68			\$627.94
24343	Repr elbow lat ligmnt w/issr	T	0050	31.7717	\$2,141.60			\$428.32
24345	Repr elbow med ligmnt w/issr	T	0052	88.6521	\$5,975.68			\$1,195.14
24346	Reconstruct elbow med ligmnt	T	0051	46.5786	\$3,139.68			\$627.94
24357	Repair elbow, perc	T	0050	31.7717	\$2,141.60			\$428.32
24358	Repair elbow wideb, open	T	0050	31.7717	\$2,141.60			\$428.32
24359	Repair elbow deb/atch open	T	0050	31.7717	\$2,141.60			\$428.32
24360	Reconstruct elbow joint	T	0047	39.8877	\$2,688.67			\$537.74
24361	Reconstruct elbow joint	T	0425	118.767	\$8,005.61			\$1,601.13
24362	Reconstruct elbow joint	T	0048	58.0838	\$3,915.20			\$783.04
24363	Replace elbow joint	T	0425	118.767	\$8,005.61			\$1,601.13
24365	Reconstruct head of radius	T	0047	39.8877	\$2,688.67			\$537.74

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
23585	Treat scapula fracture	T	0064	65.4752	\$4,413.42			\$882.69
23600	Treat humerus fracture	T	0129	1.6576	\$111.73			\$22.35
23605	Treat humerus fracture	T	0139	18.3962	\$1,240.01			\$248.01
23615	Treat humerus fracture	T	0064	65.4752	\$4,413.42			\$882.69
23616	Treat humerus fracture	T	0064	65.4752	\$4,413.42			\$882.69
23620	Treat humerus fracture	T	0129	1.6576	\$111.73			\$22.35
23625	Treat humerus fracture	T	0139	18.3962	\$1,240.01			\$248.01
23630	Treat humerus fracture	T	0064	65.4752	\$4,413.42			\$882.69
23650	Treat shoulder dislocation	T	0129	1.6576	\$111.73			\$22.35
23655	Treat shoulder dislocation	T	0045	15.2922	\$1,030.79			\$206.16
23660	Treat shoulder dislocation	T	0063	45.4678	\$3,064.80		\$268.47	\$612.96
23665	Treat dislocation/fracture	T	0138	4.7946	\$323.18			\$64.64
23670	Treat dislocation/fracture	T	0064	65.4752	\$4,413.42			\$882.69
23675	Treat dislocation/fracture	T	0129	1.6576	\$111.73			\$22.35
23680	Treat dislocation/fracture	T	0063	45.4678	\$3,064.80			\$612.96
23700	Fixation of shoulder	T	0045	15.2922	\$1,030.79		\$268.47	\$206.16
23800	Fusion of shoulder joint	T	0052	88.6521	\$5,975.68			\$1,195.14
23802	Fusion of shoulder joint	T	0051	46.5786	\$3,139.68			\$627.94
23900	Amputation of arm & girdle	C						
23920	Amputation at shoulder joint	C						
23921	Amputation follow-up surgery	T	0136	16.0542	\$1,082.15			\$216.43
23929	Shoulder surgery procedure	T	0129	1.6576	\$111.73			\$22.35
23930	Drainage of arm lesion	T	0008	19.4063	\$1,308.10			\$261.62
23931	Drainage of arm bursa	T	0008	19.4063	\$1,308.10			\$261.62
23935	Drain arm/elbow bone lesion	T	0049	22.0149	\$1,483.94			\$296.79
24000	Exploratory elbow surgery	T	0050	31.7717	\$2,141.60			\$428.32
24006	Release elbow joint	T	0050	31.7717	\$2,141.60			\$428.32
24065	Biopsy arm/elbow soft tissue	T	0021	17.4975	\$1,179.44			\$236.89
24066	Biopsy arm/elbow soft tissue	T	0021	17.4975	\$1,179.44			\$236.89
24071	Exc arm/elbow les sc = 3 cm	NI	0022	23.388	\$1,576.49		\$354.45	\$315.30
24073	Exc arm/elbow tum deep > 5 cm	NI	0022	23.388	\$1,576.49		\$354.45	\$315.30
24075	Exc arm/elbow les sc < 3 cm	NI	0021	17.4975	\$1,179.44			\$236.89
24076	Exc arm/elbow tum deep < 5 cm	NI	0021	17.4975	\$1,179.44			\$236.89
24077	Resect arm/elbow tum < 5 cm	NI	0021	17.4975	\$1,179.44			\$236.89
24079	Resect arm/elbow tum > 5 cm	NI	0022	23.388	\$1,576.49		\$354.45	\$315.30
24100	Biopsy elbow joint lining	T	0049	22.0149	\$1,483.94			\$296.79
24101	Explore/treat elbow joint	T	0050	31.7717	\$2,141.60			\$428.32
24102	Remove elbow joint lining	T	0050	31.7717	\$2,141.60			\$428.32
24105	Removal of elbow bursa	T	0049	22.0149	\$1,483.94			\$296.79
24110	Remove humerus lesion	T	0049	22.0149	\$1,483.94			\$296.79
24115	Remove/graft bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24116	Remove/graft bone lesion	T	0050	31.7717	\$2,141.60			\$428.32
24120	Remove elbow lesion	T	0049	22.0149	\$1,483.94			\$296.79

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
24900	Amputation of upper arm	C						
24920	Amputation of upper arm	C						
24925	Amputation follow-up surgery	T		0049	22.0149	\$1,483.94		\$296.79
24930	Amputation follow-up surgery	C						
24931	Amputate upper arm & implant	C						
24940	Revision of amputation	T		0052	88.6521	\$5,975.68		\$1,195.14
24940	Revision of upper arm	C						
24999	Upper arm/wrist surgery	T		0129	1.6576	\$111.73		\$22.35
25001	Incision of tendon sheath	T		0049	22.0149	\$1,483.94		\$296.79
25001	Incise flexor carpi radialis	T		0049	22.0149	\$1,483.94		\$296.79
25020	Decompress forearm 1 space	CH		0050	31.7717	\$2,141.60		\$428.32
25023	Decompress forearm 1 space	T		0050	31.7717	\$2,141.60		\$428.32
25024	Decompress forearm 2 spaces	T		0050	31.7717	\$2,141.60		\$428.32
25025	Decompress forearm 2 spaces	T		0050	31.7717	\$2,141.60		\$428.32
25028	Drainage of forearm lesion	T		0049	22.0149	\$1,483.94		\$296.79
25031	Treatment of forearm bursa	T		0049	22.0149	\$1,483.94		\$296.79
25035	Treat forearm bone lesion	T		0049	22.0149	\$1,483.94		\$296.79
25040	Explore/treat wrist joint	T		0050	31.7717	\$2,141.60		\$428.32
25065	Biopsy forearm soft tissues	T		0020	8.2028	\$552.92		\$10.59
25066	Biopsy forearm soft tissues	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
25071	Exc forearm les > 3 cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
25073	Exc forearm lum deep = 3 cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
25075	Exc forearm les < 3 cm	NI		0021	17.4975	\$1,179.44		\$235.89
25076	Exc forearm lum deep < 3 cm	NI		0021	17.4975	\$1,179.44		\$235.89
25077	Resect forearm/wrist tum<3cm	NI		0021	17.4975	\$1,179.44		\$235.89
25078	Resect forearm/wrist tum=3cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
25085	Incision of wrist capsule	T		0049	22.0149	\$1,483.94		\$296.79
25100	Biopsy of wrist joint	T		0049	22.0149	\$1,483.94		\$296.79
25101	Explore/treat wrist joint	T		0050	31.7717	\$2,141.60		\$428.32
25105	Remove wrist joint lining	T		0050	31.7717	\$2,141.60		\$428.32
25107	Remove wrist joint cartilage	T		0050	31.7717	\$2,141.60		\$428.32
25109	Excise tendon forearm/wrist	T		0049	22.0149	\$1,483.94		\$296.79
25110	Remove wrist tendon lesion	T		0049	22.0149	\$1,483.94		\$296.79
25111	Remove wrist tendon lesion	T		0049	22.0149	\$1,483.94		\$296.79
25112	Remove wrist tendon lesion	T		0049	22.0149	\$1,483.94		\$296.79
25115	Remove wrist/forearm lesion	T		0049	22.0149	\$1,483.94		\$296.79
25116	Remove wrist/forearm lesion	T		0049	22.0149	\$1,483.94		\$296.79
25118	Excise wrist tendon sheath	T		0050	31.7717	\$2,141.60		\$428.32
25119	Partial removal of ulna	T		0050	31.7717	\$2,141.60		\$428.32
25120	Removal of forearm lesion	T		0050	31.7717	\$2,141.60		\$428.32
25125	Remove/graft forearm lesion	T		0050	31.7717	\$2,141.60		\$428.32
25126	Remove/graft forearm lesion	T		0050	31.7717	\$2,141.60		\$428.32
25130	Removal of wrist lesion	T		0050	31.7717	\$2,141.60		\$428.32

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
24366	Reconstruct head of radius	T		0425	118.767	\$8,005.61		\$1,601.13
24400	Revision of humerus	CH		0051	46.5786	\$3,139.68		\$627.94
24410	Revision of humerus	CH		0051	46.5786	\$3,139.68		\$627.94
24420	Revision of humerus	T		0051	46.5786	\$3,139.68		\$627.94
24430	Repair of humerus	T		0052	88.6521	\$5,975.68		\$1,195.14
24435	Repair humerus with graft	T		0052	88.6521	\$5,975.68		\$1,195.14
24470	Revision of elbow joint	T		0051	46.5786	\$3,139.68		\$627.94
24495	Decompression of forearm	T		0050	31.7717	\$2,141.60		\$428.32
24498	Reinforce humerus	T		0052	88.6521	\$5,975.68		\$1,195.14
24500	Treat humerus fracture	T		0129	1.6576	\$111.73		\$22.35
24505	Treat humerus fracture	T		0129	1.6576	\$111.73		\$22.35
24515	Treat humerus fracture	T		0064	65.4752	\$4,413.42		\$882.69
24516	Treat humerus fracture	T		0064	65.4752	\$4,413.42		\$882.69
24530	Treat humerus fracture	T		0129	1.6576	\$111.73		\$22.35
24535	Treat humerus fracture	T		0138	4.7946	\$323.18		\$64.64
24538	Treat humerus fracture	T		0062	25.8446	\$1,742.08	\$372.87	\$348.42
24545	Treat humerus fracture	T		0064	65.4752	\$4,413.42		\$882.69
24546	Treat humerus fracture	T		0064	65.4752	\$4,413.42		\$882.69
24560	Treat humerus fracture	T		0129	1.6576	\$111.73		\$22.35
24565	Treat humerus fracture	T		0129	1.6576	\$111.73		\$22.35
24566	Treat humerus fracture	T		0062	25.8446	\$1,742.08	\$372.87	\$348.42
24575	Treat humerus fracture	T		0064	65.4752	\$4,413.42		\$882.69
24576	Treat humerus fracture	T		0129	1.6576	\$111.73		\$22.35
24577	Treat humerus fracture	T		0138	4.7946	\$323.18		\$64.64
24579	Treat humerus fracture	T		0064	65.4752	\$4,413.42		\$882.69
24582	Treat humerus fracture	T		0062	25.8446	\$1,742.08	\$372.87	\$348.42
24586	Treat elbow fracture	T		0064	65.4752	\$4,413.42		\$882.69
24587	Treat elbow fracture	T		0064	65.4752	\$4,413.42		\$882.69
24600	Treat elbow dislocation	T		0129	1.6576	\$111.73	\$268.47	\$22.35
24605	Treat elbow dislocation	T		0045	15.2922	\$1,030.79		\$206.16
24615	Treat elbow dislocation	T		0064	65.4752	\$4,413.42		\$882.69
24620	Treat elbow fracture	T		0139	18.3962	\$1,240.01		\$248.01
24635	Treat elbow fracture	T		0064	65.4752	\$4,413.42		\$882.69
24640	Treat elbow dislocation	T		0129	1.6576	\$111.73		\$22.35
24650	Treat radius fracture	T		0129	1.6576	\$111.73		\$22.35
24655	Treat radius fracture	T		0138	4.7946	\$323.18		\$64.64
24665	Treat radius fracture	T		0063	45.4678	\$3,064.80		\$612.96
24666	Treat radius fracture	T		0064	65.4752	\$4,413.42		\$882.69
24670	Treat ulnar fracture	T		0129	1.6576	\$111.73		\$22.35
24675	Treat ulnar fracture	T		0129	1.6576	\$111.73		\$22.35
24685	Treat ulnar fracture	T		0063	45.4678	\$3,064.80		\$612.96
24800	Fusion of elbow joint	T		0051	46.5786	\$3,139.68		\$627.94
24802	Fusion/graft of elbow joint	T		0051	46.5786	\$3,139.68		\$627.94

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
25392	Shorten radius & ulna	T		0050	31.7717	\$2,141.60		\$428.32
25393	Lengthen radius & ulna	T		0051	46.5786	\$3,139.68		\$627.94
25394	Repair carpal bone, shorten	T		0051	46.5786	\$3,139.68		\$627.94
25400	Repair radius or ulna	T		0051	46.5786	\$3,139.68		\$627.94
25405	Repair/graft radius or ulna	T		0052	88.6521	\$5,975.68		\$1,195.14
25410	Repair radius & ulna	T		0052	88.6521	\$5,975.68		\$1,195.14
25420	Repair/graft radius & ulna	T		0052	88.6521	\$5,975.68		\$1,195.14
25425	Repair/graft radius or ulna	T		0051	46.5786	\$3,139.68		\$627.94
25426	Repair/graft radius & ulna	T		0051	46.5786	\$3,139.68		\$627.94
25430	Vasc graft into carpal bone	T		0051	46.5786	\$3,139.68		\$627.94
25431	Repair nonunion carpal bone	T		0051	46.5786	\$3,139.68		\$627.94
25440	Repair/graft wrist bone	T		0052	88.6521	\$5,975.68		\$1,195.14
25441	Reconstruct wrist joint	T		0045	118.767	\$8,005.61		\$1,601.13
25442	Reconstruct wrist joint	T		0425	118.767	\$8,005.61		\$1,601.13
25443	Reconstruct wrist joint	T		0048	58.0838	\$3,915.20		\$783.04
25444	Reconstruct wrist joint	T		0048	58.0838	\$3,915.20		\$783.04
25445	Reconstruct wrist joint	T		0048	58.0838	\$3,915.20		\$783.04
25446	Wrist replacement	T		0425	118.767	\$8,005.61		\$1,601.13
25447	Repair wrist joint(s)	T		0047	39.8877	\$2,688.67		\$537.74
25449	Remove wrist joint implant	T		0047	39.8877	\$2,688.67		\$537.74
25450	Revision of wrist joint	T		0051	46.5786	\$3,139.68		\$627.94
25455	Revision of wrist joint	T		0051	46.5786	\$3,139.68		\$627.94
25490	Reinforce radius	T		0051	46.5786	\$3,139.68		\$627.94
25491	Reinforce ulna	T		0051	46.5786	\$3,139.68		\$627.94
25492	Reinforce radius and ulna	T		0051	46.5786	\$3,139.68		\$627.94
25500	Treat fracture of radius	T		0129	1.6576	\$111.73		\$22.35
25505	Treat fracture of radius	T		0138	4.7946	\$323.18		\$64.64
25515	Treat fracture of radius	T		0063	45.4678	\$3,064.80		\$612.96
25520	Treat fracture of radius	T		0063	45.4678	\$3,064.80		\$612.96
25525	Treat fracture of radius	T		0063	45.4678	\$3,064.80		\$612.96
25528	Treat fracture of radius	T		0063	45.4678	\$3,064.80		\$612.96
25530	Treat fracture of ulna	T		0129	1.6576	\$111.73		\$22.35
25535	Treat fracture of ulna	T		0129	1.6576	\$111.73		\$22.35
25545	Treat fracture of ulna	T		0063	45.4678	\$3,064.80		\$612.96
25560	Treat fracture radius & ulna	T		0129	1.6576	\$111.73		\$22.35
25565	Treat fracture radius & ulna	T		0138	4.7946	\$323.18		\$64.64
25574	Treat fracture radius & ulna	T		0064	65.4752	\$4,413.42		\$882.69
25575	Treat fracture radius/ulna	T		0064	65.4752	\$4,413.42		\$882.69
25600	Treat fracture radius/ulna	T		0129	1.6576	\$111.73		\$22.35
25605	Treat fracture radius/ulna	T		0138	4.7946	\$323.18		\$64.64
25606	Treat fx distal radial	T		0062	25.8446	\$1,742.08		\$348.42
25607	Treat fx rad extra-articul	T		0064	65.4752	\$4,413.42		\$882.69
25608	Treat fx rad intra-articul	T		0064	65.4752	\$4,413.42		\$882.69

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
25135	Remove & graft wrist lesion	T		0050	31.7717	\$2,141.60		\$428.32
25136	Remove & graft wrist lesion	T		0050	31.7717	\$2,141.60		\$428.32
25145	Remove forearm bone lesion	T		0050	31.7717	\$2,141.60		\$428.32
25150	Partial removal of radius	T		0050	31.7717	\$2,141.60		\$428.32
25151	Partial removal of ulna	T		0051	46.5786	\$3,139.68		\$627.94
25170	Resect radius/ulnar tumor	T		0051	46.5786	\$3,139.68		\$627.94
25210	Remove of wrist bone	T		0050	31.7717	\$2,141.60		\$428.32
25215	Removal of wrist bones	T		0050	31.7717	\$2,141.60		\$428.32
25230	Partial removal of radius	T		0050	31.7717	\$2,141.60		\$428.32
25240	Partial removal of ulna	T		0050	31.7717	\$2,141.60		\$428.32
25246	Injection for wrist x-ray	N						
25248	Remove forearm foreign body	T		0049	22.0149	\$1,483.94		\$296.79
25250	Removal of wrist prosthesis	T		0050	31.7717	\$2,141.60		\$428.32
25251	Removal of wrist prosthesis	T		0050	31.7717	\$2,141.60		\$428.32
25259	Manipulate wrist w/anesthes	T		0139	18.9962	\$1,240.01		\$248.01
25260	Repair forearm tendon/muscle	T		0050	31.7717	\$2,141.60		\$428.32
25265	Repair forearm tendon/muscle	T		0050	31.7717	\$2,141.60		\$428.32
25265	Repair forearm tendon/muscle	T		0050	31.7717	\$2,141.60		\$428.32
25270	Repair forearm tendon/muscle	T		0050	31.7717	\$2,141.60		\$428.32
25272	Repair forearm tendon/muscle	T		0050	31.7717	\$2,141.60		\$428.32
25274	Repair forearm tendon sheath	T		0050	31.7717	\$2,141.60		\$428.32
25280	Revise wrist/forearm tendon	T		0050	31.7717	\$2,141.60		\$428.32
25290	Incise wrist/forearm tendon	T		0050	31.7717	\$2,141.60		\$428.32
25295	Release wrist/forearm tendon	T		0049	22.0149	\$1,483.94		\$296.79
25300	Fusion of tendons at wrist	T		0050	31.7717	\$2,141.60		\$428.32
25301	Fusion of tendons at wrist	T		0050	31.7717	\$2,141.60		\$428.32
25310	Transplant forearm tendon	T		0051	46.5786	\$3,139.68		\$627.94
25312	Transplant forearm tendon	T		0051	46.5786	\$3,139.68		\$627.94
25315	Revise palsy hand tendon(s)	T		0051	46.5786	\$3,139.68		\$627.94
25316	Revise palsy hand tendon(s)	T		0052	88.6521	\$5,975.68		\$1,195.14
25320	Repair/revise wrist joint	T		0051	46.5786	\$3,139.68		\$627.94
25332	Revise wrist joint	T		0047	39.8877	\$2,688.67		\$537.74
25335	Realignment of hand	T		0051	46.5786	\$3,139.68		\$627.94
25337	Reconstruct ulna/rad/oulnar	CH		0051	46.5786	\$3,139.68		\$627.94
25350	Revision of radius	T		0051	46.5786	\$3,139.68		\$627.94
25355	Revision of radius	T		0051	46.5786	\$3,139.68		\$627.94
25360	Revision of ulna	CH		0051	46.5786	\$3,139.68		\$627.94
25365	Revise radius & ulna	CH		0051	46.5786	\$3,139.68		\$627.94
25370	Revise radius or ulna	T		0051	46.5786	\$3,139.68		\$627.94
25375	Revise radius & ulna	T		0051	46.5786	\$3,139.68		\$627.94
25390	Shorten radius or ulna	CH		0051	46.5786	\$3,139.68		\$627.94
25391	Lengthen radius or ulna	T		0051	46.5786	\$3,139.68		\$627.94

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
26035	Decompress fingers/hand	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26037	Decompress fingers/hand	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26040	Release palm contracture	T	T	0054	28.2376	\$1,903.98		\$380.68
26045	Release palm contracture	T	T	0054	28.2376	\$1,903.98		\$380.68
26055	Incise finger tendon sheath	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26060	Incision of finger tendon	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26070	Explore/treat hand joint	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26075	Explore/treat hand joint	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26080	Explore/treat finger joint	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26100	Biopsy hand joint lining	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26105	Biopsy finger joint lining	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26110	Biopsy finger joint lining	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26111	Exc hand les > 1.5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
26113	Exc hand tum deep > 1.5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
26115	Exc hand tum deep < 1.5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
26116	Exc hand tum deep < 1.5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
26117	Exc hand tum ra < 3 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
26118	Exc hand tum ra > 3 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
26121	Release palm contracture	T	T	0054	28.2376	\$1,903.98		\$380.68
26125	Release palm contracture	T	T	0054	28.2376	\$1,903.98		\$380.68
26130	Remove wrist joint lining	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26135	Revise finger joint, each	T	T	0054	28.2376	\$1,903.98		\$380.68
26140	Revise finger joint, each	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26145	Tendon excision, palm/finger	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26160	Remove tendon sheath lesion	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26170	Removal of palm tendon, each	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26180	Removal of finger tendon	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26185	Remove finger bone	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26200	Remove hand bone lesion	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26205	Remove/graft bone lesion	T	T	0054	28.2376	\$1,903.98		\$380.68
26210	Removal of finger lesion	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26215	Remove/graft finger lesion	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26230	Partial removal of hand bone	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26235	Partial removal, finger bone	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26236	Partial removal, finger bone	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26250	Extensive hand surgery	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26255	Extensive hand surgery	CH	D					
26260	Resect prox finger tumor	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26261	Extensive finger surgery	CH	D					
26262	Resect distal finger tumor	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26320	Removal of implant from hand	T	T	0021	17.4975	\$1,179.44		\$235.89
26340	Manipulate finger w/anesth	T	T	0138	4.7946	\$323.18		\$64.64

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
25609	Treat tx radial 3+ frag	T	T	0064	65.4752	\$4,413.42		\$882.69
25622	Treat wrist bone fracture	T	T	0129	1.6576	\$111.73		\$22.35
25624	Treat wrist bone fracture	T	T	0138	4.7946	\$323.18		\$64.64
25628	Treat wrist bone fracture	T	T	0063	45.4678	\$3,064.80		\$612.96
25630	Treat wrist bone fracture	T	T	0129	1.6576	\$111.73		\$22.35
25635	Treat wrist bone fracture	T	T	0138	4.7946	\$323.18		\$64.64
25645	Treat wrist bone fracture	T	T	0063	45.4678	\$3,064.80		\$612.96
25650	Treat wrist bone fracture	T	T	0129	1.6576	\$111.73		\$22.35
25651	Pin ulnar styloid fracture	T	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
25652	Treat fracture ulnar styloid	T	T	0063	45.4678	\$3,064.80		\$612.96
25660	Treat wrist dislocation	T	T	0129	1.6576	\$111.73		\$22.35
25670	Treat wrist dislocation	T	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
25671	Pin radioulnar dislocation	T	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
25675	Treat wrist dislocation	T	T	0129	1.6576	\$111.73		\$22.35
25676	Treat wrist dislocation	T	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
25680	Treat wrist fracture	T	T	0129	1.6576	\$111.73		\$22.35
25685	Treat wrist fracture	T	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
25690	Treat wrist dislocation	T	T	0139	18.3962	\$1,240.01		\$248.01
25695	Treat wrist dislocation	T	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
25900	Fusion of wrist joint	T	T	0052	88.6521	\$5,975.68		\$1,195.14
25905	Fusion/graft of wrist joint	T	T	0051	46.5786	\$3,139.68		\$627.94
25810	Fusion/graft of wrist joint	T	T	0052	88.6521	\$5,975.68		\$1,195.14
25820	Fusion of hand bones	T	T	0051	46.5786	\$3,139.68		\$627.94
25825	Fuse hand bones with graft	T	T	0052	88.6521	\$5,975.68		\$1,195.14
25830	Fusion, radioulnar jnt/ulna	T	T	0052	88.6521	\$5,975.68		\$1,195.14
25900	Amputation of forearm	C						
25905	Amputation of forearm	C						
25907	Amputation follow-up surgery	T	T	0049	25.0149	\$1,483.94		\$296.79
25909	Amputation follow-up surgery	C						
25915	Amputation of forearm	C						
25920	Amputate hand at wrist	T	T	0049	25.0149	\$1,483.94		\$296.79
25924	Amputate hand at wrist	C						
25927	Amputation of hand	C						
25929	Amputation follow-up surgery	T	T	0136	16.0542	\$1,082.15		\$216.43
25931	Amputation follow-up surgery	T	T	0049	25.0149	\$1,483.94		\$296.79
25999	Forearm or wrist surgery	T	T	0129	1.6576	\$111.73		\$22.35
26010	Drainage of finger abscess	T	T	0006	1.4557	\$98.12		\$19.63
26011	Drainage of finger abscess	T	T	0007	12.6217	\$850.78		\$170.16
26020	Drain hand tendon sheath	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26025	Drainage of palm bursa	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26030	Drainage of palm bursa(s)	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75
26034	Treat hand bone lesion	T	T	0053	17.042	\$1,148.73	\$253.49	\$229.75

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
26952	Amputation of finger/thumb		T	0053	17.042	\$1,148.73	\$253.49	\$228.75
26989	Hand/finger surgery		T	0129	1.6576	\$111.73		\$22.35
26990	Drainage of pelvis lesion		T	0049	22.0149	\$1,483.94		\$296.79
26991	Drainage of bursae		T	0049	22.0149	\$1,483.94		\$296.79
26992	Drainage of bone lesion		C					
27000	Incision of hip tendon		T	0049	22.0149	\$1,483.94		\$296.79
27001	Incision of hip tendon		T	0050	31.7717	\$2,141.60		\$428.32
27003	Incision of hip tendon		T	0050	31.7717	\$2,141.60		\$428.32
27006	Incision of hip tendons		T	0050	31.7717	\$2,141.60		\$428.32
27025	Incision of high/igh fascia		C					
27027	Buttock fasciotomy		T	0049	22.0149	\$1,483.94		\$296.79
27030	Drainage of hip joint		C					
27033	Exploration of hip joint		T	0051	46.5786	\$3,139.68		\$627.94
27035	Denervation of hip joint		T	0051	46.5786	\$3,139.68		\$627.94
27036	Excision of hip joint/muscle		C					
27040	Biopsy of soft tissues		T	0020	8.2028	\$552.92		\$110.59
27041	Biopsy of soft tissues		T	0020	8.2028	\$552.92		\$110.59
27043	Exc hip/pelvis les sc > 3 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
27045	Exc hip/pelvis tum deep > 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
27047	Exc hip/pelvis les sc < 3 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
27048	Exc hip/pelvis tum deep < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
27049	Resect hip/pelvis tum < 5 cm	NI	T	0021	17.4975	\$1,179.44		\$235.89
27050	Biopsy of sacroiliac joint		T	0049	22.0149	\$1,483.94		\$296.79
27052	Biopsy of hip joint		T	0049	22.0149	\$1,483.94		\$296.79
27054	Removal of hip joint lining		C					
27057	Buttock fasciotomy w/dbrmt		T	0049	22.0149	\$1,483.94		\$296.79
27059	Resect hip/pelvis tum > 5 cm	NI	T	0022	23.388	\$1,576.49	\$354.45	\$315.30
27060	Removal of ischial bursa		T	0049	22.0149	\$1,483.94		\$296.79
27062	Remove femur lesion/bursa		T	0049	22.0149	\$1,483.94		\$296.79
27065	Removal of hip bone lesion		T	0049	22.0149	\$1,483.94		\$296.79
27066	Removal of hip bone lesion		T	0050	31.7717	\$2,141.60		\$428.32
27067	Remove/graft hip bone lesion		T	0050	31.7717	\$2,141.60		\$428.32
27070	Partial removal of hip bone		C					
27071	Partial removal of hip bone		C					
27075	Resect hip tumor		C					
27076	Resect hip tum incl acetalabul		C					
27077	Resect hip tum w/innom bone		C					
27078	Resect hip tum incl femur	CH	D					
27079	Extensive hip surgery		C					
27080	Removal of tail bone		T	0050	31.7717	\$2,141.60		\$428.32
27086	Remove hip foreign body		T	0020	8.2028	\$552.92		\$110.59
27087	Remove hip foreign body		T	0049	22.0149	\$1,483.94		\$296.79

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
26608	Treat metacarpal fracture		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26615	Treat metacarpal fracture		T	0063	45.4678	\$3,064.80		\$612.96
26641	Treat thumb dislocation		T	0129	1.6576	\$111.73		\$22.35
26645	Treat thumb fracture		T	0138	4.7946	\$323.18		\$64.84
26650	Treat thumb fracture		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26665	Treat thumb fracture		T	0063	45.4678	\$3,064.80		\$612.96
26670	Treat hand dislocation		T	0129	1.6576	\$111.73		\$22.35
26675	Treat hand dislocation		T	0138	4.7946	\$323.18		\$64.84
26676	Pin hand dislocation		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26685	Treat hand dislocation		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26686	Treat hand dislocation		T	0064	65.4752	\$4,413.42		\$882.69
26700	Treat knuckle dislocation		T	0129	1.6576	\$111.73		\$22.35
26705	Treat knuckle dislocation		T	0129	1.6576	\$111.73		\$22.35
26706	Pin knuckle dislocation		T	0139	18.3962	\$1,240.01		\$248.01
26715	Treat knuckle dislocation		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26720	Treat finger fracture, each		T	0129	1.6576	\$111.73		\$22.35
26725	Treat finger fracture, each		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26727	Treat finger fracture, each		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26735	Treat finger fracture, each		T	0129	1.6576	\$111.73		\$22.35
26740	Treat finger fracture, each		T	0129	1.6576	\$111.73		\$22.35
26742	Treat finger fracture, each		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26746	Treat finger fracture, each		T	0129	1.6576	\$111.73		\$22.35
26750	Treat finger fracture, each		T	0129	1.6576	\$111.73		\$22.35
26755	Treat finger fracture, each		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26756	Pin finger fracture, each		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26765	Treat finger fracture, each		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26770	Treat finger dislocation		T	0129	1.6576	\$111.73		\$22.35
26775	Treat finger dislocation		T	0045	15.2922	\$1,030.79	\$268.47	\$206.16
26776	Pin finger dislocation		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26785	Treat finger dislocation		T	0062	25.8446	\$1,742.08	\$372.87	\$348.42
26820	Thumb fusion with graft		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26841	Fusion of thumb		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26842	Thumb fusion with graft		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26843	Fusion of hand joint		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26844	Fusion/graft of hand joint		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26850	Fusion of knuckle		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26852	Fusion of knuckle with graft		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26860	Fusion of finger joint		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26861	Fusion of finger jnt, add-on		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26862	Fusion/graft of finger joint		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26863	Fuse/graft added joint		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26910	Amputate metacarpal bone		T	0054	28.2376	\$1,903.38	\$380.68	\$380.68
26951	Amputation of finger/thumb		T	0053	17.042	\$1,148.73	\$253.49	\$229.75

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27218	Treat pelvic ring fracture	E	T	0129	1.6576	\$111.73		\$22.35
27220	Treat hip socket fracture	C						
27222	Treat hip socket fracture	C						
27226	Treat hip wall fracture	C						
27227	Treat hip fracture(s)	C						
27228	Treat hip fracture(s)	C						
27230	Treat thigh fracture	T	0129		1.6576	\$111.73		\$22.35
27232	Treat thigh fracture	C						
27235	Treat thigh fracture	T	0050		31.7717	\$2,141.60		\$428.32
27236	Treat thigh fracture	C						
27238	Treat thigh fracture	T	0138		4.7946	\$323.18		\$64.64
27240	Treat thigh fracture	C						
27244	Treat thigh fracture	C						
27245	Treat thigh fracture	C						
27246	Treat thigh fracture	T	0138		4.7946	\$323.18		\$64.64
27248	Treat thigh fracture	C						
27250	Treat hip dislocation	T	0129		1.6576	\$111.73		\$22.35
27252	Treat hip dislocation	T	0045		15.2922	\$1,030.79	\$268.47	\$206.16
27253	Treat hip dislocation	C						
27254	Treat hip dislocation	C						
27256	Treat hip dislocation	T	0129		1.6576	\$111.73		\$22.35
27257	Treat hip dislocation	T	0045		15.2922	\$1,030.79	\$268.47	\$206.16
27258	Treat hip dislocation	C						
27259	Treat hip dislocation	C						
27265	Treat hip dislocation	T	0129		1.6576	\$111.73		\$22.35
27266	Treat hip dislocation	T	0045		15.2922	\$1,030.79	\$268.47	\$206.16
27267	Cfx thigh fx	T	0129		1.6576	\$111.73		\$22.35
27268	Cfx thigh fx w/mnpl	C						
27269	Optx thigh fx	C						
27275	Manipulation of hip joint	T	0045		15.2922	\$1,030.79	\$268.47	\$206.16
27280	Fusion of sacroiliac joint	C						
27282	Fusion of pubic bones	C						
27284	Fusion of hip joint	C						
27286	Fusion of hip joint	C						
27290	Amputation of leg at hip	C						
27295	Amputation of leg at hip	C						
27299	Penkt/hip joint surgery	T	0129		1.6576	\$111.73		\$22.35
27301	Drain thigh/knee lesion	T	0008		19.4063	\$1,308.10	\$261.62	\$206.16
27303	Drainage of bone lesion	C						
27305	Incise thigh tendon & fascia	T	0049		22.0149	\$1,483.94	\$296.79	\$206.16
27306	Incision of thigh tendon	T	0049		22.0149	\$1,483.94	\$296.79	\$206.16
27307	Incision of thigh tendons	T	0049		22.0149	\$1,483.94	\$296.79	\$206.16
27310	Exploration of knee joint	T	0050		31.7717	\$2,141.60	\$428.32	\$206.16

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27090	Removal of hip prosthesis	C						
27091	Removal of hip prosthesis	C						
27093	Injection for hip x-ray	N						
27095	Injection for hip x-ray	N						
27096	Inject sacroiliac joint	N						
27097	Revision of hip tendon	B						
27098	Revision of hip tendon	T	0050		31.7717	\$2,141.60	\$428.32	\$206.16
27100	Transfer of abdominal muscle	T	0050		31.7717	\$2,141.60	\$428.32	\$206.16
27105	Transfer of spinal muscle	T	0051		46.5786	\$3,139.68	\$627.94	\$206.16
27110	Transfer of iliopectus muscle	T	0051		46.5786	\$3,139.68	\$627.94	\$206.16
27111	Transfer of iliopectus muscle	T	0051		46.5786	\$3,139.68	\$627.94	\$206.16
27120	Reconstruction of hip socket	C						
27122	Reconstruction of hip socket	C						
27125	Partial hip replacement	C						
27130	Total hip arthroplasty	C						
27132	Total hip arthroplasty	C						
27134	Revise hip joint replacement	C						
27137	Revise hip joint replacement	C						
27138	Revise hip joint replacement	C						
27140	T transplant femur ridge	C						
27146	Incision of hip bone	C						
27147	Revision of hip bone	C						
27151	Incision of hip bones	C						
27156	Revision of hip bones	C						
27158	Revision of pelvis	C						
27161	Incision of neck of femur	C						
27165	Incision/fixation of femur	C						
27170	Repair/graft femur head/neck	C						
27175	Treat slipped epiphysis	C						
27176	Treat slipped epiphysis	C						
27177	Treat slipped epiphysis	C						
27178	Treat slipped epiphysis	C						
27179	Revise head/neck of femur	CH	T	0052	88.6521	\$5,975.68	\$1,195.14	\$206.16
27181	Treat slipped epiphysis	C						
27185	Revision of femur epiphysis	C						
27187	Reinforce hip bones	C						
27193	Treat pelvic ring fracture	T	0129		1.6576	\$111.73		\$22.35
27194	Treat pelvic ring fracture	T	0045		15.2922	\$1,030.79	\$268.47	\$206.16
27200	Treat tail bone fracture	T	0129		1.6576	\$111.73		\$22.35
27202	Treat tail bone fracture	T	0063		45.4678	\$3,064.80	\$612.96	\$206.16
27215	Treat pelvic fracture(s)	E						
27216	Treat pelvic ring fracture	E						
27217	Treat pelvic ring fracture	E						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27407	Repair of knee ligaments	T		0052	88.6521	\$5,975.68		\$1,195.14
27409	Repair of knee ligaments	T		0051	46.5786	\$3,139.68		\$627.94
27412	Autochondrocyte implant knee	T		0052	88.6521	\$5,975.68		\$1,195.14
27415	Osteochondral knee autograft	T		0051	46.5786	\$3,139.68		\$627.94
27416	Osteochondral knee autograft	T		0051	46.5786	\$3,139.68		\$627.94
27418	Repair degenerated kneecap	T		0051	46.5786	\$3,139.68		\$627.94
27420	Revision of unstable kneecap	T		0051	46.5786	\$3,139.68		\$627.94
27422	Revision of unstable kneecap	T		0051	46.5786	\$3,139.68		\$627.94
27424	Revision/removal of kneecap	T		0051	46.5786	\$3,139.68		\$627.94
27425	Lat retinacular release open	T		0050	31.7717	\$2,141.60		\$428.32
27427	Reconstruction, knee	T		0051	46.5786	\$3,139.68		\$627.94
27428	Reconstruction, knee	T		0052	88.6521	\$5,975.68		\$1,195.14
27429	Reconstruction, knee	T		0052	88.6521	\$5,975.68		\$1,195.14
27430	Revision of thigh muscles	T		0051	46.5786	\$3,139.68		\$627.94
27435	Incision of knee joint	T		0051	46.5786	\$3,139.68		\$627.94
27437	Revise kneecap	T		0047	39.8877	\$2,688.67		\$537.74
27438	Revise kneecap with implant	T		0048	58.0638	\$3,915.20		\$783.04
27440	Revision of knee joint	T		0047	39.8877	\$2,688.67		\$537.74
27441	Revision of knee joint	T		0047	39.8877	\$2,688.67		\$537.74
27442	Revision of knee joint	T		0047	39.8877	\$2,688.67		\$537.74
27443	Revision of knee joint	T		0047	39.8877	\$2,688.67		\$537.74
27445	Revision of knee joint	T		0047	39.8877	\$2,688.67		\$537.74
27446	Revision of knee joint	T		0047	39.8877	\$2,688.67		\$537.74
27447	Total knee arthroplasty	CH		0425	118.767	\$8,005.61		\$1,601.13
27448	Incision of thigh	C						
27450	Incision of thigh	C						
27454	Realignment of thigh bone	C						
27455	Realignment of knee	C						
27457	Realignment of knee	C						
27465	Shortening of thigh bone	C						
27466	Lengthening of thigh bone	C						
27468	Shorten/lengthen thighs	C						
27470	Repair of thigh	C						
27472	Repair/graft of thigh	C						
27475	Surgery to stop leg growth	T		0050	31.7717	\$2,141.60		\$428.32
27477	Surgery to stop leg growth	T		0050	31.7717	\$2,141.60		\$428.32
27479	Surgery to stop leg growth	T		0050	31.7717	\$2,141.60		\$428.32
27485	Surgery to stop leg growth	T		0050	31.7717	\$2,141.60		\$428.32
27486	Revise/replace knee joint	C						
27487	Revise/replace knee joint	C						
27488	Removal of knee prosthesis	C						
27495	Reinforce thigh	C						
27496	Decompression of thigh/knee	CH		0050	31.7717	\$2,141.60		\$428.32

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27323	Biopsy, thigh soft tissues	T		0020	8.2028	\$552.92		\$110.59
27324	Biopsy, thigh soft tissues	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
27325	Neurectomy, hamstring	T		0220	18.72	\$1,261.84		\$252.37
27326	Neurectomy, popliteal	T		0220	18.72	\$1,261.84		\$252.37
27327	Exc thigh/knee les sc < 3 cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
27328	Exc thigh/knee tum deep <5cm	NI		0021	17.4975	\$1,179.44		\$235.89
27329	Resect thigh/knee tum < 5 cm	NI		0021	17.4975	\$1,179.44		\$235.89
27330	Biopsy, knee joint lining	T		0050	31.7717	\$2,141.60		\$428.32
27331	Explore/treat knee joint	T		0050	31.7717	\$2,141.60		\$428.32
27332	Removal of knee cartilage	T		0050	31.7717	\$2,141.60		\$428.32
27333	Removal of knee cartilage	T		0050	31.7717	\$2,141.60		\$428.32
27334	Remove knee joint lining	T		0050	31.7717	\$2,141.60		\$428.32
27335	Remove knee joint lining	T		0050	31.7717	\$2,141.60		\$428.32
27337	Exc thigh/knee les sc > 3 cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
27339	Exc thigh/knee tum deep >5cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
27340	Removal of kneecap bursa	T		0049	22.0149	\$1,483.94		\$296.79
27345	Removal of knee cyst	T		0049	22.0149	\$1,483.94		\$296.79
27347	Remove knee cyst	T		0049	22.0149	\$1,483.94		\$296.79
27350	Removal of kneecap	T		0050	31.7717	\$2,141.60		\$428.32
27355	Remove femur lesion	T		0050	31.7717	\$2,141.60		\$428.32
27356	Remove femur lesion/graft	T		0050	31.7717	\$2,141.60		\$428.32
27357	Remove femur lesion/graft	T		0050	31.7717	\$2,141.60		\$428.32
27358	Remove femur lesion/fixation	T		0050	31.7717	\$2,141.60		\$428.32
27360	Partial removal, leg bone(s)	T		0050	31.7717	\$2,141.60		\$428.32
27364	Resect thigh/knee tum >5 cm	NI		0022	23.388	\$1,576.49	\$354.45	\$315.30
27365	Resect femur/knee tumor	C						
27370	Injection for knee x-ray	N						
27372	Removal of foreign body	T		0022	23.388	\$1,576.49	\$354.45	\$315.30
27380	Repair of kneecap tendon	T		0049	22.0149	\$1,483.94		\$296.79
27381	Repair/graft kneecap tendon	T		0049	22.0149	\$1,483.94		\$296.79
27385	Repair of thigh muscle	T		0049	22.0149	\$1,483.94		\$296.79
27386	Repair/graft of thigh muscle	T		0049	22.0149	\$1,483.94		\$296.79
27390	Incision of thigh tendon	T		0049	22.0149	\$1,483.94		\$296.79
27391	Incision of thigh tendons	T		0049	22.0149	\$1,483.94		\$296.79
27392	Incision of thigh tendons	T		0049	22.0149	\$1,483.94		\$296.79
27393	Lengthening of thigh tendon	T		0050	31.7717	\$2,141.60		\$428.32
27394	Lengthening of thigh tendons	T		0050	31.7717	\$2,141.60		\$428.32
27395	Lengthening of thigh tendons	T		0051	46.5786	\$3,139.68		\$627.94
27396	Transplant of thigh tendon	T		0050	31.7717	\$2,141.60		\$428.32
27397	Transplants of thigh tendons	T		0051	46.5786	\$3,139.68		\$627.94
27400	Revise thigh muscles/tendons	T		0051	46.5786	\$3,139.68		\$627.94
27403	Repair of knee cartilage	T		0050	31.7717	\$2,141.60		\$428.32
27405	Repair of knee ligament	T		0051	46.5786	\$3,139.68		\$627.94

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27600	Decompression of lower leg	T	0049	22.0149	\$1,483.94		\$296.79	
27601	Decompression of lower leg	T	0049	22.0149	\$1,483.94		\$296.79	
27602	Decompression of lower leg	T	0049	22.0149	\$1,483.94		\$296.79	
27603	Drain lower leg bursa	T	0049	22.0149	\$1,483.94		\$296.79	
27604	Drain lower leg bursa	T	0049	22.0149	\$1,483.94		\$296.79	
27605	Incision of achilles tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$296.79	
27606	Incision of achilles tendon	T	0049	22.0149	\$1,483.94		\$296.79	
27607	Treat lower leg bone lesion	T	0049	22.0149	\$1,483.94		\$296.79	
27610	Explore/treat ankle joint	T	0050	31.7717	\$2,141.60		\$428.32	
27612	Exploration of ankle joint	T	0050	31.7717	\$2,141.60		\$428.32	
27613	Biopsy lower leg soft tissue	T	0020	8.2028	\$552.92		\$110.59	
27614	Biopsy lower leg soft tissue	T	0022	23.388	\$1,576.49	\$354.45	\$315.30	
27615	Resect leg/ankle tum < 5 cm	NI	0021	17.4975	\$1,179.44		\$235.89	
27616	Resect leg/ankle tum > 5 cm	NI	0022	23.388	\$1,576.49	\$354.45	\$315.30	
27618	Exc leg/ankle tum < 3 cm	NI	0021	17.4975	\$1,179.44		\$235.89	
27619	Exc leg/ankle tum deep < 5 cm	NI	0021	17.4975	\$1,179.44		\$235.89	
27620	Explore/treat ankle joint	NI	0050	31.7717	\$2,141.60		\$428.32	
27625	Remove ankle joint lining	T	0050	31.7717	\$2,141.60		\$428.32	
27626	Remove ankle joint lining	T	0050	31.7717	\$2,141.60		\$428.32	
27630	Removal of tendon lesion	T	0049	22.0149	\$1,483.94		\$296.79	
27632	Exc leg/ankle les sc > 3 cm	NI	0022	23.388	\$1,576.49	\$354.45	\$315.30	
27634	Exc leg/ankle tum deep > 5 cm	NI	0022	23.388	\$1,576.49	\$354.45	\$315.30	
27635	Remove lower leg bone lesion	T	0050	31.7717	\$2,141.60		\$428.32	
27637	Remove/graft leg bone lesion	T	0050	31.7717	\$2,141.60		\$428.32	
27638	Remove/graft leg bone lesion	T	0050	31.7717	\$2,141.60		\$428.32	
27640	Partial removal of fibula	T	0051	46.5786	\$3,139.68		\$627.94	
27641	Partial removal of fibula	T	0050	31.7717	\$2,141.60		\$428.32	
27645	Resect fibula tumor	C						
27646	Resect fibula tumor	C						
27647	Resect talus/calcaneus tum	T	0051	46.5786	\$3,139.68		\$627.94	
27648	Injection for ankle x-ray	N						
27650	Repair achilles tendon	T	0051	46.5786	\$3,139.68		\$627.94	
27652	Repair/graft achilles tendon	T	0052	88.6521	\$5,975.68		\$1,195.14	
27654	Repair of achilles tendon	T	0051	46.5786	\$3,139.68		\$627.94	
27656	Repair leg fascia defect	T	0049	22.0149	\$1,483.94		\$296.79	
27658	Repair of leg tendon, each	T	0049	22.0149	\$1,483.94		\$296.79	
27659	Repair of leg tendon, each	T	0049	22.0149	\$1,483.94		\$296.79	
27664	Repair of leg tendon, each	CH	T	0050	31.7717	\$2,141.60	\$428.32	
27665	Repair of leg tendon, each	T	0050	31.7717	\$2,141.60		\$428.32	
27675	Repair lower leg tendons	T	0049	22.0149	\$1,483.94		\$296.79	
27676	Repair lower leg tendons	T	0050	31.7717	\$2,141.60		\$428.32	
27680	Release of lower leg tendon	T	0050	31.7717	\$2,141.60		\$428.32	
27681	Release of lower leg tendons	T	0050	31.7717	\$2,141.60		\$428.32	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27497	Decompression of thigh/knee	T	0049	22.0149	\$1,483.94		\$296.79	
27498	Decompression of thigh/knee	CH	T	0050	31.7717	\$2,141.60	\$428.32	
27499	Decompression of thigh/knee	CH	T	0050	31.7717	\$2,141.60	\$428.32	
27500	Treatment of thigh fracture	T	0138	4.7946	\$323.18		\$64.64	
27501	Treatment of thigh fracture	T	0129	1.6576	\$111.73		\$22.35	
27502	Treatment of thigh fracture	T	0139	18.3962	\$1,240.01		\$248.01	
27503	Treatment of thigh fracture	T	0129	1.6576	\$111.73		\$22.35	
27506	Treatment of thigh fracture	C						
27507	Treatment of thigh fracture	C						
27508	Treatment of thigh fracture	T	0129	1.6576	\$111.73		\$22.35	
27509	Treatment of thigh fracture	T	0062	25.8446	\$1,742.08	\$372.87	\$348.42	
27510	Treatment of thigh fracture	T	0138	4.7946	\$323.18		\$64.64	
27511	Treatment of thigh fracture	C						
27513	Treatment of thigh fracture	C						
27514	Treatment of thigh fracture	C						
27516	Treat thigh fx growth plate	T	0129	1.6576	\$111.73		\$22.35	
27517	Treat thigh fx growth plate	T	0129	1.6576	\$111.73		\$22.35	
27519	Treat thigh fx growth plate	C						
27520	Treat kneecap fracture	T	0129	1.6576	\$111.73		\$22.35	
27524	Treat kneecap fracture	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
27530	Treat knee fracture	T	0129	1.6576	\$111.73		\$22.35	
27532	Treat knee fracture	T	0139	18.3962	\$1,240.01		\$248.01	
27535	Treat knee fracture	C						
27536	Treat knee fracture	C						
27538	Treat knee fracture(s)	T	0129	1.6576	\$111.73		\$22.35	
27540	Treat knee fracture	C						
27550	Treat knee dislocation	T	0129	1.6576	\$111.73		\$22.35	
27552	Treat knee dislocation	T	0045	15.2922	\$1,030.79	\$268.47	\$206.16	
27556	Treat knee dislocation	C						
27557	Treat knee dislocation	C						
27558	Treat knee dislocation	C						
27560	Treat kneecap dislocation	T	0129	1.6576	\$111.73		\$22.35	
27562	Treat kneecap dislocation	T	0045	15.2922	\$1,030.79	\$268.47	\$206.16	
27566	Treat kneecap dislocation	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
27570	Fixation of knee joint	T	0045	15.2922	\$1,030.79	\$268.47	\$206.16	
27580	Fusion of knee	C						
27590	Amputate leg at thigh	C						
27591	Amputate leg at thigh	C						
27592	Amputate leg at thigh	C						
27594	Amputation follow-up surgery	T	0049	22.0149	\$1,483.94		\$296.79	
27596	Amputation follow-up surgery	C						
27598	Amputate lower leg at knee	C						
27599	Leg surgery procedure	T	0129	1.6576	\$111.73		\$22.35	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27784	Treatment of fibula fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27786	Treatment of ankle fracture	T	0129	1.6576	\$111.73	\$22.35		
27788	Treatment of ankle fracture	T	0129	1.6576	\$111.73	\$22.35		
27792	Treatment of ankle fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27810	Treatment of ankle fracture	T	0138	4.7946	\$323.18	\$64.64		
27814	Treatment of ankle fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27818	Treatment of ankle fracture	T	0138	4.7946	\$323.18	\$64.64		
27822	Treatment of ankle fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27826	Treatment of ankle fracture	T	0064	65.4752	\$4,413.42	\$882.69		
27828	Treat lower leg fracture	T	0139	18.3962	\$1,240.01	\$248.01		
27829	Treat lower leg fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27830	Treat lower leg fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27831	Treat lower leg fracture	T	0129	1.6576	\$111.73	\$22.35		
27832	Treat lower leg fracture	T	0139	18.3962	\$1,240.01	\$248.01		
27842	Treat ankle dislocation	T	0063	45.4678	\$3,064.80	\$612.96		
27844	Treat ankle dislocation	T	0063	45.4678	\$3,064.80	\$612.96		
27846	Treat ankle dislocation	T	0063	45.4678	\$3,064.80	\$612.96		
27860	Fixation of ankle joint, open	T	0045	15.2922	\$1,030.79	\$206.16		
27870	Fusion of ankle joint	T	0052	88.6521	\$5,975.68	\$1,195.14		
27871	Fusion of tibiofibular joint	T	0052	88.6521	\$5,975.68	\$1,195.14		
27880	Amputation of lower leg	C						
27881	Amputation of lower leg	C						
27882	Amputation of lower leg	C						
27884	Amputation follow-up surgery	T	0049	22.0149	\$1,483.94	\$296.79		
27886	Amputation follow-up surgery	C						
27888	Amputation of foot at ankle	C						
27889	Amputation of foot at ankle	C						
27892	Decompression of leg	CH	0050	31.7717	\$2,141.60	\$428.32		
27893	Decompression of leg	CH	0050	31.7717	\$2,141.60	\$428.32		
27894	Decompression of leg	CH	0050	31.7717	\$2,141.60	\$428.32		
27899	Leg/ankle surgery procedure	T	0129	1.6576	\$111.73	\$22.35		
28001	Drainage of bursa of foot	T	0007	12.6217	\$850.78	\$170.16		
28002	Treatment of foot infection	T	0049	22.0149	\$1,483.94	\$296.79		
28003	Treatment of foot infection	T	0049	22.0149	\$1,483.94	\$296.79		
28005	Treat foot bone lesion	T	0055	21.8439	\$1,472.41	\$294.49		
28008	Incision of foot fascia	T	0055	21.8439	\$1,472.41	\$294.49		

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
27685	Revision of lower leg tendon	T	0050	31.7717	\$2,141.60	\$428.32		
27686	Revise lower leg tendons	T	0050	31.7717	\$2,141.60	\$428.32		
27687	Revision of calf tendon	T	0050	31.7717	\$2,141.60	\$428.32		
27690	Revise lower leg tendon	T	0051	46.5786	\$3,139.68	\$627.94		
27691	Revise lower leg tendon	T	0051	46.5786	\$3,139.68	\$627.94		
27692	Revise additional leg tendon	T	0051	46.5786	\$3,139.68	\$627.94		
27695	Repair of ankle ligament	T	0050	31.7717	\$2,141.60	\$428.32		
27696	Repair of ankle ligaments	T	0050	31.7717	\$2,141.60	\$428.32		
27698	Repair of ankle ligament	T	0050	31.7717	\$2,141.60	\$428.32		
27700	Revision of ankle joint	T	0047	39.8877	\$2,688.67	\$537.74		
27702	Reconstruct ankle joint	C						
27703	Reconstruction, ankle joint	C						
27704	Removal of ankle implant	T	0049	22.0149	\$1,483.94	\$296.79		
27705	Incision of tibia	T	0051	46.5786	\$3,139.68	\$627.94		
27707	Incision of fibula	T	0049	22.0149	\$1,483.94	\$296.79		
27712	Incision of tibia & fibula	T	0050	31.7717	\$2,141.60	\$428.32		
27715	Realignment of lower leg	C						
27720	Repair of lower leg	T	0063	45.4678	\$3,064.80	\$612.96		
27722	Repair/graft of tibia	T	0064	65.4752	\$4,413.42	\$882.69		
27724	Repair/graft of tibia	C						
27725	Repair of lower leg	C						
27726	Repair fibula nonunion	C						
27727	Repair of lower leg	C						
27730	Repair of tibia epiphysis	T	0050	31.7717	\$2,141.60	\$428.32		
27732	Repair of fibula epiphysis	T	0050	31.7717	\$2,141.60	\$428.32		
27734	Repair lower leg epiphyses	T	0050	31.7717	\$2,141.60	\$428.32		
27740	Repair of leg epiphyses	T	0051	46.5786	\$3,139.68	\$627.94		
27742	Repair of leg epiphyses	T	0051	46.5786	\$3,139.68	\$627.94		
27745	Reinforce tibia	T	0052	88.6521	\$5,975.68	\$1,195.14		
27750	Treatment of tibia fracture	T	0129	1.6576	\$111.73	\$22.35		
27752	Treatment of tibia fracture	T	0139	18.3962	\$1,240.01	\$248.01		
27756	Treatment of tibia fracture	T	0062	25.8446	\$1,742.08	\$348.42		
27758	Treatment of tibia fracture	T	0063	45.4678	\$3,064.80	\$612.96		
27759	Treatment of tibia fracture	T	0064	65.4752	\$4,413.42	\$882.69		
27760	Cltx medial ankle fx	T	0129	1.6576	\$111.73	\$22.35		
27762	Cltx med ankle fx w/mmpj	T	0139	18.3962	\$1,240.01	\$248.01		
27766	Cltx medial ankle fx	T	0063	45.4678	\$3,064.80	\$612.96		
27767	Cltx post ankle fx	T	0129	1.6576	\$111.73	\$22.35		
27768	Cltx post ankle fx w/mmpj	T	0129	1.6576	\$111.73	\$22.35		
27769	Cltx post ankle fx	T	0063	45.4678	\$3,064.80	\$612.96		
27780	Treatment of fibula fracture	T	0129	1.6576	\$111.73	\$22.35		
27781	Treatment of fibula fracture	T	0139	18.3962	\$1,240.01	\$248.01		

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
28126	Partial removal of toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28130	Removal of ankle bone	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28140	Removal of metatarsal	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28150	Removal of toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28160	Partial removal of toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28171	Resect tarsal tumor	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28173	Resect metatarsal tumor	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28175	Resect phalanx of toe tumor	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28190	Removal of foot foreign body	T	0020	8.2028	\$552.92		\$110.59	
28192	Removal of foot foreign body	T	0021	17.4975	\$1,179.44		\$235.89	
28193	Removal of foot foreign body	T	0020	6.2028	\$552.92		\$110.59	
28200	Repair of foot tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28202	Repair/graft of foot tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28208	Repair of foot tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28210	Repair/graft of foot tendon	T	0056	52.5258	\$3,540.55		\$708.11	
28220	Release of foot tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28225	Release of foot tendons	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28226	Release of foot tendons	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28230	Revision of foot tendon(s)	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28232	Incision of toe tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28234	Incision of foot tendon	T	0056	52.5258	\$3,540.55		\$708.11	
28240	Revision of big toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28250	Revision of foot fascia	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28260	Release of midfoot joint	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28262	Revision of foot tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28264	Revision of foot and ankle	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28270	Release of midfoot joint	T	0056	52.5258	\$3,540.55		\$708.11	
28272	Release of toe joint, each	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28280	Fusion of toes	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28285	Repair of hammer toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28286	Repair of hammer toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28288	Partial removal of foot bone	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28289	Repair hallux rigidus	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28290	Correction of bunion	T	0057	31.5534	\$2,126.89	\$475.91	\$425.38	
28293	Correction of bunion	T	0057	31.5534	\$2,126.89	\$475.91	\$425.38	
28294	Correction of bunion	T	0057	31.5534	\$2,126.89	\$475.91	\$425.38	
28296	Correction of bunion	T	0057	31.5534	\$2,126.89	\$475.91	\$425.38	
28297	Correction of bunion	T	0057	31.5534	\$2,126.89	\$475.91	\$425.38	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
28010	Incision of toe tendon	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28011	Incision of toe tendons	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28020	Exploration of foot joint	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28022	Exploration of foot joint	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28024	Exploration of toe joint	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28035	Decompression of fibia nerve	T	0220	18.72	\$1,261.84	\$252.37		
28039	Exc foot/toe tum sc > 1.5 cm	NI	0022	23.988	\$1,576.49	\$354.45	\$315.30	
28041	Exc foot/toe tum deep > 1.5 cm	NI	0022	23.988	\$1,576.49	\$354.45	\$315.30	
28043	Exc foot/toe tum sc < 1.5 cm	NI	0021	17.4975	\$1,179.44	\$235.89	\$235.89	
28045	Exc foot/toe tum deep < 1.5 cm	NI	0021	17.4975	\$1,179.44	\$235.89	\$235.89	
28046	Resect foot/toe tumor < 3 cm	NI	0021	17.4975	\$1,179.44	\$235.89	\$235.89	
28047	Resect foot/toe tumor > 3 cm	NI	0022	23.988	\$1,576.49	\$354.45	\$315.30	
28050	Biospy of foot joint lining	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28052	Biospy of foot joint lining	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28054	Biospy of toe joint lining	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28055	Neurectomy, foot	T	0220	18.72	\$1,261.84	\$252.37		
28060	Partial removal, foot fascia	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28062	Removal of foot fascia	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28070	Removal of foot joint lining	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28072	Removal of foot joint lining	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28080	Removal of foot lesion	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28086	Excise foot tendon sheath	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28088	Excise foot tendon sheath	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28090	Removal of foot lesion	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28092	Removal of toe lesions	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28100	Removal of ankle/heel lesion	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28102	Remove/graft foot lesion	T	0056	52.5258	\$3,540.55		\$708.11	
28103	Remove/graft foot lesion	T	0056	52.5258	\$3,540.55		\$708.11	
28104	Removal of foot lesion	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28106	Remove/graft foot lesion	T	0056	52.5258	\$3,540.55		\$708.11	
28107	Remove/graft foot lesion	T	0056	52.5258	\$3,540.55		\$708.11	
28108	Removal of toe lesions	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28110	Part removal of metatarsal	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28111	Part removal of metatarsal	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28112	Part removal of metatarsal	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28113	Part removal of metatarsal heads	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28116	Revision of foot	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28118	Removal of heel bone	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28119	Removal of heel spur	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28120	Part removal of ankle/heel	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28122	Partial removal of foot bone	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	
28124	Partial removal of toe	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
28510	Treatment of toe fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28515	Treatment of toe fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28525	Treat toe fracture	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28530	Treat sesamoid bone fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28531	Treat sesamoid bone fracture	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28540	Treat foot dislocation	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28545	Treat foot dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28555	Treat foot dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28555	Repair foot dislocation	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
28570	Treat foot dislocation	T	0138	4.7946	\$323.16	\$64.64	\$64.64	
28575	Treat foot dislocation	T	0139	18.3962	\$1,240.01	\$248.01	\$248.01	
28585	Treat foot dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28585	Repair foot dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28600	Treat foot dislocation	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28605	Treat foot dislocation	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28605	Repair foot dislocation	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
28615	Treat toe dislocation	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28630	Treat toe dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28635	Treat toe dislocation	T	0045	15.2922	\$1,030.79	\$206.16	\$206.16	
28645	Repair toe dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28660	Treat toe dislocation	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28665	Treat toe dislocation	T	0045	15.2922	\$1,030.79	\$206.16	\$206.16	
28666	Treat toe dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28675	Repair of toe dislocation	T	0062	25.8446	\$1,742.08	\$372.87	\$372.87	
28705	Fusion of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28715	Fusion of foot bones	T	0052	88.6521	\$5,975.68	\$1,195.14	\$1,195.14	
28725	Fusion of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28730	Fusion of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28735	Fusion of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28737	Revision of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28740	Fusion of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28750	Fusion of big toe joint	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28755	Fusion of big toe joint	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28760	Fusion of big toe joint	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28800	Amputation of midfoot	C						
28805	Amputation thru metatarsal	CH	T	0055	21.8439	\$1,472.41	\$294.49	
28810	Amputation toe & metatarsal	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28820	Amputation of toe	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28825	Partial amputation of toe	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28890	High energy eswt, plantar	T	0050	31.7717	\$2,141.60	\$428.32	\$428.32	
28899	Foot/toes surgery procedure	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
29000	Application of body cast	S	0058	1.0537	\$71.03	\$14.21	\$14.21	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
28298	Correction of bunion	T	0057	31.5534	\$2,126.89	\$425.38	\$425.38	
28299	Correction of bunion	T	0057	31.5534	\$2,126.89	\$425.38	\$425.38	
28300	Incision of heel bone	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28302	Incision of ankle bone	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28304	Incision of ankle bone	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28305	Incise/graft midfoot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28306	Incision of metatarsal	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28307	Incision of metatarsal	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28308	Incision of metatarsals	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28309	Incision of metatarsals	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28310	Revision of big toe	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28312	Revision of toe	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28313	Repair deformity of toe	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28315	Removal of sesamoid bone	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28320	Repair of foot bones	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28322	Repair of metatarsals	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28340	Resect enlarged toe tissue	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28341	Resect enlarged toe	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28344	Repair extra toe(s)	T	0055	21.8439	\$1,472.41	\$294.49	\$294.49	
28360	Reconstruct cleft foot	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28400	Treatment of heel fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28405	Treatment of heel fracture	T	0139	18.3962	\$1,240.01	\$248.01	\$248.01	
28406	Treatment of heel fracture	T	0062	25.8446	\$1,742.08	\$348.42	\$348.42	
28415	Treat heel fracture	T	0064	65.4752	\$4,413.42	\$882.69	\$882.69	
28420	Treat/graft heel fracture	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
28430	Treatment of ankle fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28435	Treatment of ankle fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28436	Treatment of ankle fracture	T	0062	25.8446	\$1,742.08	\$348.42	\$348.42	
28445	Treat ankle fracture	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
28446	Osteochondral talus autograft	T	0056	52.5258	\$3,540.55	\$708.11	\$708.11	
28450	Treat midfoot fracture, each	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28455	Treat midfoot fracture, each	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28456	Treat midfoot fracture	T	0062	25.8446	\$1,742.08	\$348.42	\$348.42	
28465	Treat midfoot fracture, each	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
28470	Treat metatarsal fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28475	Treat metatarsal fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28476	Treat metatarsal fracture	T	0062	25.8446	\$1,742.08	\$348.42	\$348.42	
28485	Treat metatarsal fracture	T	0063	45.4678	\$3,064.80	\$612.96	\$612.96	
28490	Treat big toe fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28495	Treat big toe fracture	T	0129	1.6576	\$111.73	\$22.35	\$22.35	
28496	Treat big toe fracture	T	0062	25.8446	\$1,742.08	\$348.42	\$348.42	
28505	Treat big toe fracture	T	0062	25.8446	\$1,742.08	\$348.42	\$348.42	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
29581	Application of paste boot		S	0058	1.0537	\$71.03		\$14.21
29580	Apply mullay comps lwr leg	NI	S	0058	1.0537	\$71.03		\$14.21
29590	Application of foot splint		S	0058	1.0537	\$71.03		\$14.21
29705	Removal/revision of cast		S	0058	1.0537	\$71.03		\$14.21
29710	Removal/revision of cast		S	0058	2.3457	\$158.11		\$31.63
29715	Removal/revision of cast		S	0058	1.0537	\$71.03		\$14.21
29720	Repair of body cast		S	0058	1.0537	\$71.03		\$14.21
29730	Widening of cast		S	0058	1.0537	\$71.03		\$14.21
29740	Wedging of cast		S	0058	1.0537	\$71.03		\$14.21
29750	Wedging of clubfoot cast		S	0058	1.0537	\$71.03		\$14.21
29799	Casting/strapping procedure		T	0041	29.9198	\$2,016.77		\$403.36
29800	Jaw arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29804	Law arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29805	Shoulder arthroscopy, dx		T	0041	29.9198	\$2,016.77		\$403.36
29806	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29807	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29819	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29820	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29821	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29822	Shoulder arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29823	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29824	Shoulder arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29825	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29826	Shoulder arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29827	Arthroscopy rotator cuff repr		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29828	Arthroscopy biceps tenodesis		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29830	Elbow arthroscopy		T	0041	29.9198	\$2,016.77		\$403.36
29834	Elbow arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29835	Elbow arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29836	Elbow arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29837	Elbow arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29838	Elbow arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29840	Wrist arthroscopy		T	0041	29.9198	\$2,016.77		\$403.36
29843	Wrist arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29844	Wrist arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29846	Wrist arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29847	Wrist arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29848	Wrist arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29850	Knee arthroscopy/surgery		T	0041	29.9198	\$2,016.77		\$403.36
29851	Knee arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29855	Tibial arthroscopy/surgery		T	0042	48.8176	\$3,290.60	\$804.74	\$658.12

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
29010	Application of body cast		S	0426	2.3457	\$158.11		\$31.63
29015	Application of body cast		S	0426	2.3457	\$158.11		\$31.63
29020	Application of body cast		S	0058	1.0537	\$71.03		\$14.21
29025	Application of body cast		S	0058	1.0537	\$71.03		\$14.21
29035	Application of body cast		S	0426	2.3457	\$158.11		\$31.63
29040	Application of body cast		S	0058	1.0537	\$71.03		\$14.21
29044	Application of body cast		S	0426	2.3457	\$158.11		\$31.63
29046	Application of body cast		S	0426	2.3457	\$158.11		\$31.63
29049	Application of figure eight		S	0058	1.0537	\$71.03		\$14.21
29055	Application of shoulder cast		S	0426	2.3457	\$158.11		\$31.63
29058	Application of shoulder cast		S	0058	1.0537	\$71.03		\$14.21
29065	Application of long arm cast		S	0426	2.3457	\$158.11		\$31.63
29075	Application of forearm cast		S	0426	2.3457	\$158.11		\$31.63
29085	Apply hand/wrist cast		S	0058	1.0537	\$71.03		\$14.21
29086	Apply finger cast		S	0058	1.0537	\$71.03		\$14.21
29105	Apply long arm splint		S	0058	1.0537	\$71.03		\$14.21
29125	Apply forearm splint		S	0058	1.0537	\$71.03		\$14.21
29126	Apply forearm splint		S	0058	1.0537	\$71.03		\$14.21
29130	Application of finger splint		S	0058	1.0537	\$71.03		\$14.21
29131	Application of finger splint		S	0058	1.0537	\$71.03		\$14.21
29200	Strapping of chest		S	0058	1.0537	\$71.03		\$14.21
29220	Strapping of low back	CH	D					
29240	Strapping of shoulder		S	0058	1.0537	\$71.03		\$14.21
29260	Strapping of elbow or wrist		S	0058	1.0537	\$71.03		\$14.21
29280	Strapping of hand or finger		S	0058	1.0537	\$71.03		\$14.21
29305	Application of hip cast		S	0426	2.3457	\$158.11		\$31.63
29325	Application of hip casts		S	0426	2.3457	\$158.11		\$31.63
29345	Application of long leg cast		S	0426	2.3457	\$158.11		\$31.63
29355	Application of long leg cast		S	0426	2.3457	\$158.11		\$31.63
29358	Apply long leg cast brace		S	0426	2.3457	\$158.11		\$31.63
29365	Application of long leg cast		S	0426	2.3457	\$158.11		\$31.63
29405	Apply short leg cast		S	0426	2.3457	\$158.11		\$31.63
29425	Apply short leg cast		S	0426	2.3457	\$158.11		\$31.63
29435	Apply short leg cast		S	0058	1.0537	\$71.03		\$14.21
29440	Addition of walker to cast		S	0058	1.0537	\$71.03		\$14.21
29445	Apply rigid leg cast		S	0426	2.3457	\$158.11		\$31.63
29450	Application of leg cast		S	0058	1.0537	\$71.03		\$14.21
29505	Application, long leg splint		S	0058	1.0537	\$71.03		\$14.21
29515	Application lower leg splint		S	0058	1.0537	\$71.03		\$14.21
29520	Strapping of hip		S	0058	1.0537	\$71.03		\$14.21
29530	Strapping of knee		S	0058	1.0537	\$71.03		\$14.21
29540	Strapping of ankle and/or ft		S	0058	1.0537	\$71.03		\$14.21
29550	Strapping of toes		S	0058	1.0537	\$71.03		\$14.21

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
30020	Drainage of nose lesion	T	T	0251	3.425	\$230.87	\$109.16	\$46.18
30100	Intranasal biopsy	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
30110	Removal of nose polyp(s)	T	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30115	Removal of nose polyp(s)	T	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30117	Removal of intranasal lesion	T	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30118	Removal of intranasal lesion	T	T	0254	24.9637	\$1,682.70		\$336.54
30120	Revision of nose	CH	T	0254	24.9637	\$1,682.70		\$336.54
30124	Removal of nose lesion	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
30125	Removal of nose lesion	T	T	0256	42.9827	\$2,897.29		\$579.46
30130	Excise inferior turbinate	T	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30140	Resect inferior turbinate	T	T	0254	24.9637	\$1,682.70		\$336.54
30150	Partial removal of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30160	Removal of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30200	Injection treatment of nose	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
30210	Nasal sinus therapy	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
30220	Insert nasal septal button	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
30300	Remove nasal foreign body	X	X	0340	0.6693	\$45.11		\$9.03
30310	Remove nasal foreign body	T	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30320	Remove nasal foreign body	T	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30400	Reconstruction of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30410	Reconstruction of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30420	Reconstruction of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30430	Revision of nose	T	T	0254	24.9637	\$1,682.70		\$336.54
30435	Revision of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30450	Revision of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30460	Revision of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30462	Revision of nose	T	T	0256	42.9827	\$2,897.29		\$579.46
30520	Repair nasal stenosis	T	T	0254	24.9637	\$1,682.70		\$336.54
30540	Repair nasal defect	T	T	0256	42.9827	\$2,897.29		\$579.46
30545	Repair nasal defect	T	T	0256	42.9827	\$2,897.29		\$579.46
30560	Release of nasal adhesions	T	T	0251	3.425	\$230.87		\$46.18
30580	Repair upper jaw fistula	T	T	0256	42.9827	\$2,897.29		\$579.46
30600	Intranasal reconstruction	T	T	0256	42.9827	\$2,897.29		\$579.46
30630	Repair nasal septum defect	T	T	0254	24.9637	\$1,682.70		\$336.54
30801	Ablate inf turbinate, superl	T	T	0252	7.6196	\$513.61	\$109.16	\$102.73
30802	Ablate inf turbinate submuc	CH	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72
30901	Control of nosebleed	T	T	0250	1.1522	\$77.67	\$25.10	\$15.54
30905	Control of nosebleed	T	T	0250	1.1522	\$77.67	\$25.10	\$15.54
30906	Repeat control of nosebleed	T	T	0250	1.1522	\$77.67	\$25.10	\$15.54
30915	Ligation, nasal sinus artery	T	T	0092	26.5713	\$1,791.07		\$358.22

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
29856	Tibial arthroscopy, surgery	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29860	Hip arthroscopy, dx	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29861	Hip arthroscopy/surgery	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29862	Hip arthroscopy/surgery	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29863	Hip arthroscopy/surgery	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29866	Autlgrt implant, knee w/scope	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29867	Allgrt implant, knee w/scope	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29868	Miniscal frnspl, knee w/scope	T	T	0041	29.9198	\$2,016.77	\$804.74	\$658.12
29870	Knee arthroscopy, dx	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29871	Knee arthroscopy/drainage	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29873	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29874	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29875	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29876	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29877	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29878	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29880	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29881	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29882	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29883	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29884	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$804.74	\$658.12
29885	Knee arthroscopy/surgery	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29886	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29887	Knee arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29888	Knee arthroscopy/surgery	CH	T	0052	88.6521	\$5,975.68		\$1,195.14
29889	Knee arthroscopy/surgery	CH	T	0052	88.6521	\$5,975.68		\$1,195.14
29891	Ankle arthroscopy/surgery	CH	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29892	Ankle arthroscopy/surgery	CH	T	0052	88.6521	\$5,975.68		\$1,195.14
29893	Scope, plantar fasciotomy	T	T	0055	21.8439	\$1,472.41	\$355.34	\$294.49
29894	Ankle arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29895	Ankle arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29897	Ankle arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29898	Ankle arthroscopy/surgery	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29899	Ankle arthroscopy/surgery	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29900	Mcp joint arthroscopy, dx	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29901	Mcp joint arthroscopy, surg	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29902	Mcp joint arthroscopy, surg	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29904	Subtalar arthro w/ft rmtl	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29905	Subtalar arthro w/ft w/exc	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29906	Subtalar arthro w/ft w/exc	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
29907	Subtalar arthro w/fusion	T	T	0042	48.8176	\$3,290.60	\$804.74	\$658.12
29999	Arthroscopy of joint	T	T	0041	29.9198	\$2,016.77	\$403.36	\$403.36
30000	Drainage of nose lesion	T	T	0251	3.425	\$230.87		\$46.18

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
31294	Nasal/sinus endoscopy, surg	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31299	Sinus surgery procedure	T	0250	1,1522	\$77.67	\$25.10	\$15.54	
31300	Removal of larynx lesion	T	0254	24,9637	\$1,682.70		\$396.54	
31320	Diagnostic incision, larynx	T	0256	42,9827	\$2,897.29		\$579.46	
31360	Removal of larynx	C						
31367	Removal of larynx	C						
31368	Partial removal of larynx	C						
31369	Partial removal of larynx	C						
31370	Partial removal of larynx	C						
31375	Partial removal of larynx	C						
31380	Partial removal of larynx	C						
31382	Partial removal of larynx	C						
31390	Removal of larynx & pharynx	C						
31395	Reconstruct larynx & pharynx	C						
31400	Revision of larynx	T	0256	42,9827	\$2,897.29		\$579.46	
31420	Removal of epiglottitis	T	0256	42,9827	\$2,897.29		\$579.46	
31500	Insert emergency airway	S	0094	2,4553	\$165.50	\$46.29	\$33.10	
31502	Change of windpipe airway	S	0078	1,4142	\$95.33		\$19.07	
31505	Diagnostic laryngoscopy	T	0071	0,8007	\$53.97	\$11.14	\$10.80	
31510	Laryngoscopy with biopsy	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31511	Remove foreign body, larynx	T	0072	1,8425	\$124.20		\$24.84	
31512	Removal of larynx lesion	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31513	Injection into vocal cord	T	0072	1,8425	\$124.20		\$24.84	
31515	Laryngoscopy for aspiration	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31520	Dx laryngoscopy, new/bn	T	0072	1,8425	\$124.20		\$24.84	
31525	Dx laryngoscopy excl nb	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31526	Dx laryngoscopy w/oper scope	CH	T	0074	21,6572	\$1,459.83	\$292.25	
31527	Laryngoscopy for treatment	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31528	Laryngoscopy and dilation	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31529	Laryngoscopy with dilation	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31530	Laryngoscopy w/ft removal	CH	T	0074	21,6572	\$1,459.83	\$292.25	
31535	Laryngoscopy w/ft & op scope	CH	T	0074	21,6572	\$1,459.83	\$292.25	
31536	Laryngoscopy w/ft & op scope	CH	T	0074	21,6572	\$1,459.83	\$292.25	
31540	Laryngoscopy w/exc of tumor	CH	T	0074	21,6572	\$1,459.83	\$292.25	
31541	Laryngoscopy w/whr exc + scope	CH	T	0074	21,6572	\$1,459.83	\$292.25	
31545	Remove vc lesion w/scope	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31546	Remove vc lesion scope/graft	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31560	Laryngoscopy w/arytenoidectom	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31561	Laryngoscopy, remove cart + scop	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31570	Laryngoscopy w/vc int	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31571	Laryngoscopy w/vc int + scope	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31575	Diagnostic laryngoscopy	T	0072	1,8425	\$124.20		\$24.84	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
30920	Ligation, upper jaw artery	T	0092	26,5713	\$1,791.07	\$358.22		
30930	Ther fx, nasal int turbinate	T	0253	17,1879	\$1,156.57	\$282.29	\$231.72	
30999	Nasal surgery procedure	T	0250	1,1522	\$77.67	\$25.10	\$15.54	
31000	Irrigation, maxillary sinus	T	0251	3,425	\$230.87		\$46.18	
31002	Irrigation, sphenoid sinus	T	0252	7,6196	\$513.61	\$102.73	\$102.73	
31020	Exploration, maxillary sinus	T	0254	24,9637	\$1,682.70	\$336.54	\$336.54	
31030	Exploration, maxillary sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31032	Explore sinus, remove polyps	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31040	Exploration behind upper jaw	T	0254	24,9637	\$1,682.70	\$336.54	\$336.54	
31050	Exploration, sphenoid sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31051	Sphenoid sinus surgery	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31070	Exploration of frontal sinus	T	0254	24,9637	\$1,682.70	\$336.54	\$336.54	
31075	Exploration of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31080	Removal of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31081	Removal of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31084	Removal of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31085	Removal of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31086	Removal of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31087	Removal of frontal sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31090	Exploration of sinuses	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31200	Removal of ethmoid sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31201	Removal of ethmoid sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31205	Removal of ethmoid sinus	T	0256	42,9827	\$2,897.29	\$579.46	\$579.46	
31225	Removal of upper jaw	C						
31230	Removal of upper jaw	C						
31231	Nasal endoscopy, dx	T	0072	1,8425	\$124.20		\$24.84	
31233	Nasal/sinus endoscopy, dx	T	0072	1,8425	\$124.20		\$24.84	
31235	Nasal/sinus endoscopy, dx	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31237	Nasal/sinus endoscopy, surg	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31238	Nasal/sinus endoscopy, surg	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31239	Nasal/sinus endoscopy, surg	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31240	Nasal/sinus endoscopy, surg	T	0074	21,6572	\$1,459.83	\$292.25	\$291.97	
31254	Revision of ethmoid sinus	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31255	Removal of ethmoid sinus	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31256	Exploration maxillary sinus	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31267	Sinus endoscopy, maxillary sinus	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31276	Sinus endoscopy, surgical	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31287	Nasal/sinus endoscopy, surg	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31288	Nasal/sinus endoscopy, surg	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31290	Nasal/sinus endoscopy, surg	C						
31291	Nasal/sinus endoscopy, surg	C						
31292	Nasal/sinus endoscopy, surg	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	
31293	Nasal/sinus endoscopy, surg	T	0075	29,2718	\$1,973.09	\$445.92	\$394.62	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
31646	Bronchoscopy, reclear airway		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31656	Bronchoscopy, inj for x-ray		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31715	Injection for bronchus x-ray		N					
31717	Bronchial brush biopsy		T	0073	4.4119	\$297.39	\$69.15	\$59.48
31720	Clearance of airways		S	0077	0.4058	\$27.35	\$7.74	\$5.47
31725	Clearance of airways		C					
31730	Intro, windpipe wiretube		T	0073	4.4119	\$297.39	\$69.15	\$59.48
31750	Repair of windpipe		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31755	Repair of windpipe		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31760	Repair of windpipe		C					
31766	Reconstruction of windpipe		C					
31770	Repair/graft of bronchus		C					
31775	Reconstruct bronchus		C					
31780	Reconstruct windpipe		C					
31781	Reconstruct windpipe		C					
31785	Remove windpipe lesion		T	0254	24.9637	\$1,682.70		\$336.54
31786	Remove windpipe lesion		C					
31800	Repair of windpipe injury		C					
31805	Repair of windpipe injury		C					
31820	Closure of windpipe lesion	CH	T	0254	24.9637	\$1,682.70		\$336.54
31825	Repair of windpipe defect		T	0254	24.9637	\$1,682.70		\$336.54
31830	Revise windpipe scar		T	0254	24.9637	\$1,682.70		\$336.54
31899	Airways surgical procedure		T	0076	10.3707	\$699.05	\$189.82	\$139.81
32035	Exploration of chest		C					
32036	Exploration of chest		C					
32095	Biopsy through chest wall		C					
32100	Exploration/biopsy of chest		C					
32110	Explore/repair chest		C					
32120	Re-exploration of chest		C					
32124	Explore chest free adhesions		C					
32140	Removal of lung lesion(s)		C					
32141	Remove/treat lung lesions		C					
32150	Removal of lung lesion(s)		C					
32151	Remove lung foreign body		C					
32160	Open chest heart massage		C					
32200	Drain, open, lung lesion		C					
32201	Drain, percut, lung lesion		T	0070	5.5521	\$374.24		\$74.85
32215	Treat chest lining		C					
32220	Release of lung		C					
32225	Partial release of lung		C					
32310	Removal of chest lining		C					
32320	Free/remove chest lining		C					
32400	Needle biopsy chest lining		T	0685	9.6666	\$651.59		\$130.32

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
31576	Laryngoscopy with biopsy	CH	T	0074	21.6572	\$1,459.83	\$292.25	\$291.97
31577	Remove foreign body, larynx		T	0073	4.4119	\$297.39	\$69.15	\$59.48
31578	Removal of larynx lesion		T	0075	29.2718	\$1,973.09	\$445.92	\$394.82
31579	Diagnostic laryngoscopy		T	0073	4.4119	\$297.39	\$69.15	\$59.48
31580	Revision of larynx		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31582	Revision of larynx		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31584	Treat larynx fracture		C					
31587	Revision of larynx		C					
31588	Revision of larynx		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31590	Reinervate larynx		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31595	Larynx nerve surgery		T	0256	1.1522	\$77.67	\$25.10	\$15.54
31599	Larynx surgery procedure		T	0254	24.9637	\$1,682.70	\$336.54	\$336.54
31600	Incision of windpipe		T	0254	24.9637	\$1,682.70	\$336.54	\$336.54
31601	Incision of windpipe		T	0252	7.6196	\$513.61	\$109.16	\$102.73
31605	Incision of windpipe		T	0252	7.6196	\$513.61	\$109.16	\$102.73
31610	Incision of windpipe		T	0254	24.9637	\$1,682.70	\$336.54	\$336.54
31611	Surgery/speech prosthesis		T	0254	24.9637	\$1,682.70	\$336.54	\$336.54
31612	Puncture/clear windpipe		T	0254	24.9637	\$1,682.70	\$336.54	\$336.54
31613	Repair windpipe opening		T	0254	24.9637	\$1,682.70	\$336.54	\$336.54
31614	Repair windpipe opening		T	0256	42.9827	\$2,897.29	\$79.46	\$79.46
31615	Visualization of windpipe		T	0252	7.6196	\$513.61	\$109.16	\$102.73
31620	Endobronchial us add-on		N					
31622	Dx bronchoscope/wash		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31623	Dx bronchoscope/brush		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31624	Dx bronchoscope/lavage		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31625	Bronchoscopy w/biopsy(s)		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31626	Bronchoscopy w/markers	NI	T	0076	10.3707	\$699.05	\$189.82	\$139.81
31627	Navigational bronchoscopy	NI	N					
31628	Bronchoscopy/lung bx, each		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31629	Bronchoscopy/needle bx, each		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31630	Bronchoscopy dilate/fix repr		T	0415	25.9024	\$1,745.98	\$459.92	\$349.20
31631	Bronchoscopy, dilate w/silent		T	0415	25.9024	\$1,745.98	\$459.92	\$349.20
31632	Bronchoscopy/lung bx, addl		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31633	Bronchoscopy/needle bx addl		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31635	Bronchoscopy w/ib removal		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31636	Bronchoscopy, bronch stenosis		T	0415	25.9024	\$1,745.98	\$459.92	\$349.20
31637	Bronchoscopy, stent add-on		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31638	Bronchoscopy, revise stent		T	0415	25.9024	\$1,745.98	\$459.92	\$349.20
31640	Bronchoscopy w/tumor excise		T	0415	25.9024	\$1,745.98	\$459.92	\$349.20
31641	Bronchoscopy, treat blockage		T	0415	25.9024	\$1,745.98	\$459.92	\$349.20
31643	Diag bronchoscope/catheter		T	0076	10.3707	\$699.05	\$189.82	\$139.81
31645	Bronchoscopy, clear airways		T	0076	10.3707	\$699.05	\$189.82	\$139.81

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
32681	Thoracoscopy, surgical	C						
32682	Thoracoscopy, surgical	C						
32683	Thoracoscopy, surgical	C						
32684	Thoracoscopy, surgical	C						
32685	Thoracoscopy, surgical	C						
32800	Repair lung hernia	C						
32810	Close chest after drainage	C						
32815	Close bronchial fistula	C						
32820	Reconstruct injured chest	C						
32850	Donor pneumonectomy	C						
32851	Lung transplant, single	C						
32852	Lung transplant with bypass	C						
32853	Lung transplant, double	C						
32854	Lung transplant with bypass	C						
32855	Prepare donor lung, single	C						
32856	Prepare donor lung, double	C						
32900	Removal of rib(s)	C						
32905	Revise & repair chest wall	C						
32906	Revise & repair chest wall	C						
32940	Revision of lung	C						
32960	Therapeutic pneumothorax	T		0070	5.5521	\$374.24		\$74.85
32987	Total lung lavage	C						
32988	Perq tr ablate tx, pul tumor	T		0423	51.3618	\$9,462.09		\$692.42
32989	Chest surgery procedure	T		0070	5.5521	\$374.24		\$74.85
33010	Drainage of heart sac	T		0070	5.5521	\$374.24		\$74.85
33011	Repeat drainage of heart sac	T		0070	5.5521	\$374.24		\$74.85
33015	Incision of heart sac	C						
33020	Incision of heart sac	C						
33025	Incision of heart sac	C						
33030	Partial removal of heart sac	C						
33031	Partial removal of heart sac	C						
33050	Removal of heart sac lesion	C						
33120	Removal of heart lesion	C						
33130	Removal of heart lesion	C						
33140	Heart revascularize (tmr)	C						
33141	Heart tmr w/other procedure	C						
33202	Insert epicard eltrd, open	C						
33203	Insert epicard eltrd, endo	C						
33206	Insertion of heart pacemaker	T		0089	118.8826	\$8,013.40	\$1,682.28	\$1,602.68
33207	Insertion of heart pacemaker	T		0089	118.8826	\$8,013.40	\$1,682.28	\$1,602.68
33208	Insertion of heart pacemaker	T		0655	142.1623	\$9,582.59		\$1,916.52
33210	Insertion of heart electrode	T		0106	48.9361	\$3,298.59		\$659.72
33211	Insertion of heart electrode	T		0106	48.9361	\$3,298.59		\$659.72

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
32402	Open biopsy chest lining	C						
32405	Biopsy, lung or mediastinum	T		0685	9.6666	\$651.59		\$130.32
32420	Puncture/clear lung	T		0070	5.5521	\$374.24		\$74.85
32421	Thoracentesis for aspiration	T		0070	5.5521	\$374.24		\$74.85
32422	Thoracentesis w/tube insert	T		0070	5.5521	\$374.24		\$74.85
32440	Removal of lung	C						
32442	Sleeve pneumonectomy	C						
32445	Removal of lung	C						
32480	Partial removal of lung	C						
32482	Bilobectomy	C						
32484	Segmentectomy	C						
32486	Sleeve lobectomy	C						
32488	Completion pneumonectomy	C						
32491	Lung volume reduction	C						
32500	Partial removal of lung	C						
32501	Repair bronchus add-on	C						
32503	Resect apical lung tumor	C						
32504	Resect apical lung tum/chest	C						
32540	Removal of lung lesion	C						
32550	Insert pleural cath	T		0652	30.4602	\$2,053.20		\$410.64
32551	Insertion of chest tube	T		0070	5.5521	\$374.24		\$74.85
32552	Remove lung catheter	NI	S	0078	1.4142	\$95.33		\$19.07
32553	Ins mark thor for rt perq	NI	X	0310	13.7576	\$927.34	\$25.27	\$185.47
32560	Treat pleurodesis w/agent	T		0070	5.5521	\$374.24		\$74.85
32561	Lyse chest fibrin nit day	NI	T	0070	5.5521	\$374.24		\$74.85
32562	Lyse chest fibrin subq day	NI	T	0070	5.5521	\$374.24		\$74.85
32601	Thoracoscopy, diagnostic	T		0069	34.0084	\$2,292.37	\$591.64	\$458.48
32602	Thoracoscopy, diagnostic	T		0069	34.0084	\$2,292.37	\$591.64	\$458.48
32603	Thoracoscopy, diagnostic	T		0069	34.0084	\$2,292.37	\$591.64	\$458.48
32604	Thoracoscopy, diagnostic	T		0069	34.0084	\$2,292.37	\$591.64	\$458.48
32605	Thoracoscopy, diagnostic	T		0069	34.0084	\$2,292.37	\$591.64	\$458.48
32606	Thoracoscopy, diagnostic	T		0069	34.0084	\$2,292.37	\$591.64	\$458.48
32651	Thoracoscopy, surgical	C						
32652	Thoracoscopy, surgical	C						
32653	Thoracoscopy, surgical	C						
32654	Thoracoscopy, surgical	C						
32655	Thoracoscopy, surgical	C						
32656	Thoracoscopy, surgical	C						
32657	Thoracoscopy, surgical	C						
32658	Thoracoscopy, surgical	C						
32659	Thoracoscopy, surgical	C						
32660	Thoracoscopy, surgical	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010										
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment		
33322	Repair major blood vessel(s)		C							
33330	Insert major vessel graft		C							
33332	Insert major vessel graft		C							
33335	Insert major vessel graft		C							
33400	Repair of aortic valve		C							
33401	Valvuloplasty, open		C							
33403	Valvuloplasty, w/cp bypass		C							
33404	Prepare heart-aorta conduit		C							
33405	Replacement of aortic valve		C							
33406	Replacement of aortic valve		C							
33410	Replacement of aortic valve		C							
33411	Replacement of aortic valve		C							
33412	Replacement of aortic valve		C							
33413	Replacement of aortic valve		C							
33414	Repair of aortic valve		C							
33415	Revision, subvalvular tissue		C							
33416	Revise ventricle muscle		C							
33417	Repair of aortic valve		C							
33420	Revision of mitral valve		C							
33422	Revision of mitral valve		C							
33425	Repair of mitral valve		C							
33426	Repair of mitral valve		C							
33427	Repair of mitral valve		C							
33430	Replacement of mitral valve		C							
33460	Revision of tricuspid valve		C							
33463	Valvuloplasty, tricuspid		C							
33464	Valvuloplasty, tricuspid		C							
33465	Replace tricuspid valve		C							
33468	Revision of tricuspid valve		C							
33470	Revision of pulmonary valve		C							
33471	Valvotomy, pulmonary valve		C							
33472	Revision of pulmonary valve		C							
33474	Revision of pulmonary valve		C							
33475	Replacement, pulmonary valve		C							
33476	Revision of heart chamber		C							
33478	Revision of heart chamber		C							
33496	Repair, prosth valve clot		C							
33500	Repair heart vessel fistula		C							
33501	Repair heart vessel fistula		C							
33502	Coronary artery correction		C							
33503	Coronary artery graft		C							
33504	Coronary artery graft		C							
33505	Repair artery w/tunnel		C							

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010										
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment		
33212	Insertion of pulse generator		T	0090	98.1576	\$6,616.41	\$1,597.43	\$1,323.29		
33213	Insertion of pulse generator		T	0654	108.1692	\$7,291.25		\$1,458.25		
33214	Upgrade of pacemaker system		T	0655	142.1623	\$9,582.59		\$1,916.52		
33215	Reposition pacing-defib lead		T	0105	22.9412	\$1,546.37		\$309.28		
33216	Insert 1 electrode pm-defib		T	0106	48.9361	\$3,298.59		\$659.72		
33217	Insert 2 electrode pm-defib		T	0106	48.9361	\$3,298.59		\$659.72		
33218	Repair lead pace-defib, one		T	0105	22.9412	\$1,546.37		\$309.28		
33220	Repair lead pace-defib, dual		T	0105	22.9412	\$1,546.37		\$309.28		
33222	Revise pocket, pacemaker		T	0136	16.0542	\$1,082.15		\$216.43		
33223	Revise pocket for defib		T	0136	16.0542	\$1,082.15		\$216.43		
33224	Insert pacing lead & connect		T	0418	204.1256	\$13,759.29		\$2,751.86		
33225	L ventric pacing lead add-on		T	0418	204.1256	\$13,759.29		\$2,751.86		
33233	Removal of pacemaker system		T	0105	22.9412	\$1,546.37		\$309.28		
33234	Removal of pacemaker system		T	0105	22.9412	\$1,546.37		\$309.28		
33235	Removal pacemaker electrode		T	0105	22.9412	\$1,546.37		\$309.28		
33236	Remove electrode/thoracotomy		C							
33237	Remove electrode/thoracotomy		C							
33238	Remove electrode/thoracotomy		C							
33240	Insert pulse generator		T	0107	325.8288	\$21,962.82		\$4,392.57		
33241	Remove pulse generator		T	0105	22.9412	\$1,546.37		\$309.28		
33243	Remove eltrd/thoracotomy		C							
33244	Remove eltrd, transven		T	0105	22.9412	\$1,546.37		\$309.28		
33249	Eltrd/insert pace-defib		T	0108	412.3722	\$27,796.36		\$5,559.28		
33250	Ablate heart dysrhythm focus		C							
33251	Ablate heart dysrhythm focus		C							
33254	Ablate atria, lmid		C							
33255	Ablate atria w/o bypass, ext		C							
33256	Ablate atria w/bypass, exten		C							
33257	Ablate atria, lmid, add-on		C							
33258	Ablate atria, x10sv, add-on		C							
33259	Ablate atria w/bypass add-on		C							
33261	Ablate heart dysrhythm focus		C							
33265	Ablate atria, lmid, endo		C							
33266	Ablate atria, x10sv, endo		C							
33282	Implant pat-active ht record		S	0680	78.0661	\$5,262.12		\$1,052.43		
33284	Remove pat-active ht record		T	0020	8.2028	\$552.92		\$110.59		
33300	Repair of heart wound		C							
33305	Repair of heart wound		C							
33310	Exploratory heart surgery		C							
33315	Exploratory heart surgery		C							
33320	Repair major blood vessel(s)		C							
33321	Repair major blood vessel		C							

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
33681	Repair heart septum defect		C					
33684	Repair heart septum defect		C					
33688	Repair heart septum defect		C					
33690	Reinforce pulmonary artery		C					
33692	Repair of heart defects		C					
33694	Repair of heart defects		C					
33697	Repair of heart defects		C					
33702	Repair of heart defects		C					
33710	Repair of heart defects		C					
33720	Repair of heart defect		C					
33722	Repair of heart defect		C					
33724	Repair venous anomaly		C					
33726	Repair pul venous stenosis		C					
33730	Repair heart-vein defect(s)		C					
33732	Repair heart-vein defect		C					
33735	Revision of heart chamber		C					
33736	Revision of heart chamber		C					
33737	Revision of heart chamber		C					
33750	Major vessel shunt		C					
33755	Major vessel shunt		C					
33762	Major vessel shunt		C					
33764	Major vessel shunt & graft		C					
33766	Major vessel shunt		C					
33767	Major vessel shunt		C					
33768	Cavopulmonary shunting		C					
33770	Repair great vessels defect		C					
33771	Repair great vessels defect		C					
33774	Repair great vessels defect		C					
33775	Repair great vessels defect		C					
33776	Repair great vessels defect		C					
33777	Repair great vessels defect		C					
33778	Repair great vessels defect		C					
33779	Repair great vessels defect		C					
33780	Repair great vessels defect		C					
33781	Repair great vessels defect		C					
33782	Nikaidoh proc	NI	C					
33783	Nikaidoh proc w/ostia impit	NI	C					
33786	Repair arterial trunk		C					
33788	Revision of pulmonary artery		C					
33800	Aortic suspension		C					
33802	Repair vessel defect		C					
33803	Repair vessel defect		C					
33813	Repair septal defect		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
33506	Repair art, translocation		C					
33507	Repair art, intramural		C					
33508	Endoscopic vein harvest		N					
33510	CABG, vein, single		C					
33511	CABG, vein, two		C					
33512	CABG, vein, three		C					
33513	CABG, vein, four		C					
33514	CABG, vein, five		C					
33516	Cabg, vein, six or more		C					
33517	CABG, artery-vein, single		C					
33518	CABG, artery-vein, two		C					
33519	CABG, artery-vein, three		C					
33521	CABG, artery-vein, four		C					
33522	CABG, artery-vein, five		C					
33523	Cabg, art-vein, six or more		C					
33530	Coronary artery, bypass/reop.		C					
33533	CABG, arterial, single		C					
33534	CABG, arterial, two		C					
33535	CABG, arterial, three		C					
33536	Cabg, arterial, four or more		C					
33542	Removal of heart lesion		C					
33545	Repair of heart damage		C					
33548	Restore/remodel, ventricle		C					
33572	Open coronary endarterectomy		C					
33600	Closure of valve		C					
33602	Closure of valve		C					
33606	Anastomosis/artery-aorta		C					
33608	Repair anomaly w/conduit		C					
33610	Repair by enlargement		C					
33611	Repair double ventricle		C					
33612	Repair double ventricle		C					
33615	Repair, modified fontan		C					
33617	Repair single ventricle		C					
33619	Repair single ventricle		C					
33641	Repair heart septum defect		C					
33645	Revision of heart veins		C					
33647	Repair heart septum defects		C					
33660	Repair of heart defects		C					
33665	Repair of heart defects		C					
33670	Repair of heart chambers		C					
33675	Close mult vsd		C					
33676	Close mult vsd w/resection		C					
33677	Ci mult vsd w/rem pul band		C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
33971	Aortic circulation assist	C	C					
33973	Insert balloon device	C	C					
33974	Remove intra-aortic balloon	C	C					
33975	Implant ventricular device	C	C					
33976	Implant ventricular device	C	C					
33977	Remove ventricular device	C	C					
33979	Remove ventricular device	C	C					
33979	Insert intracorporeal device	C	C					
33980	Remove intracorporeal device	C	C					
33981	Replace vad pump ext	NI	C					
33982	Replace vad intra w/ bp	NI	C					
33983	Replace vad intra w/ bp	NI	C					
33989	Cardiac surgery procedure	NI	C					
34001	Removal of artery clot	T	C	0070	5.5521	\$574.24		\$74.85
34051	Removal of artery clot	C	C					
34101	Removal of artery clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34111	Removal of arm artery clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34151	Removal of artery clot	C	C					
34201	Removal of artery clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34203	Removal of leg artery clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34401	Removal of vein clot	C	C					
34421	Removal of vein clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34451	Removal of vein clot	C	C					
34471	Removal of vein clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34490	Removal of vein clot	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34501	Repair valve, femoral vein	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34502	Reconstruct vena cava	C	C					
34510	Transposition of vein valve	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34520	Cross-over vein graft	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34530	Leg vein fusion	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
34800	Endovasc aaa repr w/sm tube	C	C					
34802	Endovasc aaa repr w/2-p part	C	C					
34804	Endovasc aaa repr w/3-p part	C	C					
34805	Endovasc aaa repr w/long tube	C	C					
34806	Aneurysm press sensor add-on	C	C					
34808	Endovasc iliac a device add-on	C	C					
34812	Xpose for endoprosth, femor	C	C					
34813	Femoral endovasc graft add-on	C	C					
34820	Xpose for endoprosth, iliac	C	C					
34825	Endovasc extend prosth, init	C	C					
34826	Endovasc: exten prosth, addl	C	C					
34830	Open aortic tube prosth repr	C	C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
33814	Repair septal defect	C	C					
33820	Revised major vessel	C	C					
33822	Revised major vessel	C	C					
33824	Revised major vessel	C	C					
33840	Remove aorta constriction	C	C					
33845	Remove aorta constriction	C	C					
33851	Remove aorta constriction	C	C					
33852	Repair septal defect	C	C					
33853	Repair septal defect	C	C					
33860	Ascending aortic graft	C	C					
33861	Ascending aortic graft	C	C					
33863	Ascending aortic graft	C	C					
33864	Ascending aortic graft	C	C					
33870	Transverse aortic arch graft	C	C					
33875	Thoracic aortic graft	C	C					
33877	Thoracoabdominal graft	C	C					
33880	Endovasc taa repr incl subcl	C	C					
33881	Endovasc taa repr w/o subcl	C	C					
33883	Insert endovasc prosth, taa	C	C					
33884	Endovasc prosth, taa, add-on	C	C					
33886	Endovasc prosth, delayed	C	C					
33889	Artery transpose/endovasc taa	C	C					
33891	Car-car bp grt/endovasc taa	C	C					
33910	Remove lung artery emboli	C	C					
33915	Remove lung artery emboli	C	C					
33916	Surgery of great vessel	C	C					
33917	Repair pulmonary artery	C	C					
33920	Repair pulmonary atresia	C	C					
33922	Transsect pulmonary artery	C	C					
33924	Remove pulmonary shunt	C	C					
33925	Repr pul art unifocal w/cpb	C	C					
33926	Repr pul art, unifocal w/cpb	C	C					
33930	Removal of donor heart/lung	C	C					
33933	Prepare donor heart/lung	C	C					
33935	Transplantation, heart/lung	C	C					
33940	Removal of donor heart	C	C					
33944	Prepare donor heart	C	C					
33945	Transplantation of heart	C	C					
33960	External circulation assist	C	C					
33961	External circulation assist	C	C					
33967	Insert ia percut device	C	C					
33968	Remove aortic assist device	C	C					
33970	Aortic circulation assist	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
35236	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35241	Repair blood vessel lesion	C	C					
35246	Repair blood vessel lesion	C	C					
35251	Repair blood vessel lesion	C	C					
35256	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35261	Repair blood vessel lesion	T	C	0653	46.6883	\$3,147.07		\$629.42
35266	Repair blood vessel lesion	T	C	0653	46.6883	\$3,147.07		\$629.42
35271	Repair blood vessel lesion	C	C					
35276	Repair blood vessel lesion	C	C					
35281	Repair blood vessel lesion	C	C					
35286	Repair blood vessel lesion	T	C	0653	46.6883	\$3,147.07		\$629.42
35301	Rechanneling of artery	C	C					
35302	Rechanneling of artery	C	C					
35303	Rechanneling of artery	C	C					
35304	Rechanneling of artery	C	C					
35305	Rechanneling of artery	C	C					
35306	Rechanneling of artery	C	C					
35311	Rechanneling of artery	C	C					
35321	Rechanneling of artery	T	C	0093	35.5969	\$2,399.44		\$479.89
35331	Rechanneling of artery	C	C					
35341	Rechanneling of artery	C	C					
35351	Rechanneling of artery	C	C					
35355	Rechanneling of artery	C	C					
35361	Rechanneling of artery	C	C					
35363	Rechanneling of artery	C	C					
35371	Rechanneling of artery	C	C					
35372	Rechanneling of artery	C	C					
35390	Reoperation, carotid add-on	C	C					
35400	Angioscopy	C	C					
35450	Repair arterial blockage	C	C					
35452	Repair arterial blockage	C	C					
35456	Repair arterial blockage	C	C					
35458	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35459	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35460	Repair venous blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35470	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35471	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35472	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35473	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35474	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35475	Repair arterial blockage	T	C	0083	50.6809	\$3,416.20		\$683.24
35476	Repair venous blockage	T	C	0083	50.6809	\$3,416.20		\$683.24

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
34831	Open aortic/iliac prosth repr	C	C					
34832	Open aortofemor prosth repr	C	C					
34833	Xpose for endoprosth, iliac	C	C					
34834	Xpose, endoprosth, brachial	C	C					
34900	Endovasc iliac repr w/graft	C	C					
35001	Repair defect of artery	C	C					
35002	Repair artery rupture, neck	C	C					
35005	Repair defect of artery	C	C					
35011	Repair defect of artery	T	C	0653	46.6883	\$3,147.07		\$629.42
35013	Repair artery rupture, arm	C	C					
35021	Repair defect of artery	C	C					
35022	Repair artery rupture, chest	C	C					
35045	Repair defect of arm artery	C	C					
35081	Repair defect of artery	C	C					
35082	Repair artery rupture, aorta	C	C					
35091	Repair defect of artery	C	C					
35092	Repair artery rupture, aorta	C	C					
35102	Repair defect of artery	C	C					
35103	Repair artery rupture, groin	C	C					
35111	Repair defect of artery	C	C					
35112	Repair artery rupture, spleen	C	C					
35121	Repair defect of artery	C	C					
35122	Repair artery rupture, belly	C	C					
35131	Repair defect of artery	C	C					
35132	Repair artery rupture, groin	C	C					
35141	Repair defect of artery	C	C					
35142	Repair artery rupture, thigh	C	C					
35151	Repair defect of artery	C	C					
35152	Repair artery rupture, knee	C	C					
35180	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35182	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35188	Repair blood vessel lesion	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
35189	Repair blood vessel lesion	C	C					
35190	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35201	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35206	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89
35207	Repair blood vessel lesion	T	C	0088	40.9003	\$2,756.93	\$655.22	\$551.39
35211	Repair blood vessel lesion	C	C					
35216	Repair blood vessel lesion	C	C					
35221	Repair blood vessel lesion	C	C					
35226	Repair blood vessel lesion	CH	T	0020	8.2028	\$552.92		\$110.59
35231	Repair blood vessel lesion	T	C	0093	35.5969	\$2,399.44		\$479.89

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
35565	Artery bypass graft		C					
35566	Artery bypass graft		C					
35570	Artery bypass graft		C					
35571	Artery bypass graft		C					
35572	Harvest femoropopliteal vein		N					
35585	Vein bypass graft		C					
35587	Vein bypass graft		C					
35600	Harvest art for cabg add-on		C					
35601	Artery bypass graft		C					
35606	Artery bypass graft		C					
35612	Artery bypass graft		C					
35616	Artery bypass graft		C					
35621	Artery bypass graft		C					
35623	Bypass graft, not vein		C					
35626	Artery bypass graft		C					
35631	Artery bypass graft		C					
35632	Artery bypass graft		C					
35633	Artery bypass graft		C					
35634	Artery bypass graft		C					
35636	Artery bypass graft		C					
35637	Artery bypass graft		C					
35638	Artery bypass graft		C					
35642	Artery bypass graft		C					
35645	Artery bypass graft		C					
35646	Artery bypass graft		C					
35647	Artery bypass graft		C					
35650	Artery bypass graft		C					
35651	Artery bypass graft		C					
35654	Artery bypass graft		C					
35656	Artery bypass graft		C					
35661	Artery bypass graft		C					
35663	Artery bypass graft		C					
35665	Artery bypass graft		C					
35666	Artery bypass graft		C					
35671	Artery bypass graft		C					
35681	Composite bypass graft		C					
35682	Composite bypass graft		C					
35683	Composite bypass graft		C					
35685	Bypass graft patency/patch		T	0083	35.5969	\$2,399.44		\$479.89
35686	Bypass graft/av fist patency		T	0083	35.5969	\$2,399.44		\$479.89
35691	Arterial transposition		C					
35693	Arterial transposition		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
35480	Atherectomy, open		C					
35481	Atherectomy, open		C					
35482	Atherectomy, open		C					
35483	Atherectomy, open		C					
35484	Atherectomy, open		T	0082	93.3244	\$6,290.62		\$1,258.13
35485	Atherectomy, open		T	0082	93.3244	\$6,290.62		\$1,258.13
35490	Atherectomy, percutaneous		T	0082	93.3244	\$6,290.62		\$1,258.13
35491	Atherectomy, percutaneous		T	0082	93.3244	\$6,290.62		\$1,258.13
35492	Atherectomy, percutaneous		T	0082	93.3244	\$6,290.62		\$1,258.13
35493	Atherectomy, percutaneous		T	0082	93.3244	\$6,290.62		\$1,258.13
35494	Atherectomy, percutaneous		T	0082	93.3244	\$6,290.62		\$1,258.13
35495	Atherectomy, percutaneous		T	0082	93.3244	\$6,290.62		\$1,258.13
35500	Harvest vein for bypass		T	0103	17.8436	\$1,202.77		\$240.56
35501	Artery bypass graft		C					
35506	Artery bypass graft		C					
35508	Artery bypass graft		C					
35509	Artery bypass graft		C					
35510	Artery bypass graft		C					
35511	Artery bypass graft		C					
35512	Artery bypass graft		C					
35515	Artery bypass graft		C					
35516	Artery bypass graft		C					
35518	Artery bypass graft		C					
35521	Artery bypass graft		C					
35522	Artery bypass graft		C					
35523	Artery bypass graft		C					
35525	Artery bypass graft		C					
35526	Artery bypass graft		C					
35531	Artery bypass graft		C					
35533	Artery bypass graft		C					
35535	Artery bypass graft		C					
35536	Artery bypass graft		C					
35537	Artery bypass graft		C					
35538	Artery bypass graft		C					
35539	Artery bypass graft		C					
35540	Artery bypass graft		C					
35548	Artery bypass graft		C					
35549	Artery bypass graft		C					
35551	Artery bypass graft		C					
35556	Artery bypass graft		C					
35558	Artery bypass graft		C					
35560	Artery bypass graft		C					
35563	Artery bypass graft		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
36218	Place catheter in artery	N						
36245	Place catheter in artery	N						
36246	Place catheter in artery	N						
36247	Place catheter in artery	N						
36248	Place catheter in artery	N						
36260	Insertion of infusion pump	T	0623		30.4396	\$2,051.81		\$410.37
36261	Removal of infusion pump	T	0105		22.9412	\$1,546.37		\$309.28
36262	Removal of infusion pump	T	0105		22.9412	\$1,546.37		\$309.28
36299	Vessel injection procedure	N						
36400	Bl draw < 3 yrs fem/ligular	N						
36405	Bl draw < 3 yrs scalp vein	N						
36406	Bl draw < 3 yrs other vein	N						
36410	Non-routine bl draw > 3 yrs	N						
36415	Routine venipuncture	A						
36416	Capillary blood draw	N						
36420	Vein access cutdown < 1 yr	X	0035		0.232	\$15.64		\$3.13
36425	Vein access cutdown > 1 yr	X	0035		0.232	\$15.64		\$3.13
36430	Blood transfusion service	S	0110		3.3809	\$227.89		\$45.58
36440	Bl push transfuse, 2 yr or <	S	0110		3.3809	\$227.89		\$45.58
36450	Bl exchange/transfuse, nb	S	0110		3.3809	\$227.89		\$45.58
36460	Transfusion service, fetal	S	0110		3.3809	\$227.89		\$45.58
36468	Injection(s), spider veins	T	0013		0.8789	\$59.24		\$11.85
36470	Injection therapy of vein	T	0013		0.8789	\$59.24		\$11.85
36471	Injection therapy of veins	T	0013		0.8789	\$59.24		\$11.85
36475	Endovenous rf, 1st vein	T	0091		45.0292	\$3,035.24		\$607.05
36476	Endovenous laser, 1st vein	T	0092		26.5713	\$1,791.07		\$358.22
36478	Endovenous laser, 1st vein	T	0092		26.5713	\$1,791.07		\$358.22
36479	Endovenous laser vein add-on	T	0092		26.5713	\$1,791.07		\$358.22
36481	Insertion of catheter, vein	N						
36500	Insertion of catheter, vein	N						
36511	Apheresis wbc	S	0111		11.9424	\$804.99	\$198.40	\$161.00
36512	Apheresis rbc	S	0111		11.9424	\$804.99	\$198.40	\$161.00
36513	Apheresis platelets	S	0111		11.9424	\$804.99	\$198.40	\$161.00
36514	Apheresis plasma	S	0111		11.9424	\$804.99	\$198.40	\$161.00
36515	Apheresis, adscorp/reinfuse	S	0112		33.3206	\$2,246.01		\$449.21
36516	Apheresis, selective	S	0112		33.3206	\$2,246.01		\$449.21
36522	Photopheresis	S	0112		33.3206	\$2,246.01		\$449.21
36555	Insert non-tunnel cv cath	T	0621		11.166	\$752.66		\$150.54
36556	Insert non-tunnel cv cath	T	0621		11.166	\$752.66		\$150.54
36557	Insert tunneled cv cath	T	0622		25.3344	\$1,707.69		\$341.54

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
35694	Arterial transposition	C						
35695	Arterial transposition	C						
35697	Reimplant artery each	C						
35700	Reperation, bypass graft	C						
35701	Exploration, carotid artery	C						
35721	Exploration, femoral artery	C						
35741	Exploration popliteal artery	C						
35761	Exploration of artery/vein	C	0093		35.5969	\$2,399.44		\$479.89
35800	Explore neck vessels	C						
35820	Explore chest vessels	C						
35840	Explore abdominal vessels	C						
35860	Explore limb vessels	C	0093		35.5969	\$2,399.44		\$479.89
35870	Repair vessel graft defect	C						
35875	Removal of clot in graft	T	0088		40.9003	\$2,756.93	\$655.22	\$51.39
35876	Removal of clot in graft	T	0088		40.9003	\$2,756.93	\$655.22	\$51.39
35879	Revise graft w/vein	T	0088		40.9003	\$2,756.93	\$655.22	\$51.39
35881	Revise graft w/vein	T	0088		40.9003	\$2,756.93	\$655.22	\$51.39
35883	Revise graft w/nonauto graft	T	0088		40.9003	\$2,756.93	\$655.22	\$51.39
35884	Revise graft w/vein	T	0088		40.9003	\$2,756.93	\$655.22	\$51.39
35901	Excision, graft, neck	C						
35903	Excision, graft, extremity	C						
35905	Excision, graft, thorax	C						
35907	Excision, graft, abdomen	C						
36000	Place needle in vein	N						
36002	Pseudoaneurysm injection trt	S	0267		2.3005	\$155.07	\$60.50	\$31.02
36005	Injection ext venography	N						
36010	Place catheter in vein	N						
36011	Place catheter in vein	N						
36012	Place catheter in vein	N						
36013	Place catheter in artery	N						
36014	Place catheter in artery	N						
36015	Place catheter in artery	N						
36100	Establish access to artery	N						
36120	Establish access to artery	N						
36140	Establish access to artery	N						
36145	Artery to vein shunt	CH						
36147	Access av dial grft for eval	NI	0676		2.3954	\$161.46		\$32.30
36148	Access av dial grft for proc	NI						
36160	Establish access to aorta	N						
36200	Place catheter in aorta	N						
36215	Place catheter in artery	N						
36216	Place catheter in artery	N						
36217	Place catheter in artery	N						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
36825	Artery-vein autograft	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36830	Artery-vein nonautograft	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36831	Open thrombect av fistula	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36832	Av fistula revision, open	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36833	Av fistula revision	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36834	Repair A-V aneurysm	CH	D					
36835	Artery to vein shunt	T	0115	30.9784	\$2,088.13		\$417.63	
36838	Dist revas ligation, hemo	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36860	External cannula deoclotting	T	0676	2.3954	\$161.46		\$32.30	
36861	Cannula deoclotting	T	0115	30.9784	\$2,088.13		\$417.63	
36870	Percut thrombect av fistula	T	0653	46.6883	\$3,147.07		\$629.42	
37140	Revision of circulation	C						
37145	Revision of circulation	C						
37160	Revision of circulation	C						
37180	Revision of circulation	C						
37181	Splice spleen/kidney veins	C						
37182	Insert hepatic shunt (tips)	C						
37183	Remove hepatic shunt (tips)	T	0229	97.291	\$6,558.00		\$1,311.60	
37184	Prim art mech thrombectomy	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
37185	Prim art m-thrombect add-on	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
37186	Sec art m-thrombect add-on	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
37187	Venous mech thrombectomy	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
37188	Venous m-thrombectomy add-on	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
37195	Thrombolytic therapy, stroke	T	0676	2.3954	\$161.46		\$32.30	
37200	Transcatheter biopsy	T	0623	30.4396	\$2,051.81		\$410.37	
37201	Transcatheter therapy infuse	T	0103	17.8436	\$1,202.77		\$240.56	
37202	Transcatheter therapy infuse	T	0103	17.8436	\$1,202.77		\$240.56	
37203	Transcatheter retrieval	T	0623	30.4396	\$2,051.81		\$410.37	
37204	Transcatheter occlusion	T	0082	93.3244	\$6,290.62		\$1,258.13	
37205	Transcath iv stent, percut	T	0229	97.291	\$6,558.00		\$1,311.60	
37206	Transcath iv stent/tpicr addl	T	0229	97.291	\$6,558.00		\$1,311.60	
37207	Transcath iv stent, open	T	0229	97.291	\$6,558.00		\$1,311.60	
37208	Transcath iv stent/open addl	T	0229	97.291	\$6,558.00		\$1,311.60	
37209	Change iv cath at thromb tx	T	0623	30.4396	\$2,051.81		\$410.37	
37210	Embolization uterine fibroid	T	0229	97.291	\$6,558.00		\$1,311.60	
37215	Transcath stent, cca weeps	CH	T	0229	97.291	\$6,558.00		
37216	Transcath stent, cca w/o eps	E						
37250	Iv us first vessel add-on	N						
37251	Iv us each add vessel add-on	N						
37500	Endoscopy, ligate perf veins	T	0091	45.0292	\$3,035.24		\$607.05	
37501	Vascular endoscopy procedure	T	0092	26.5713	\$1,791.07		\$358.22	
37565	Ligation of neck vein	T	0093	35.5969	\$2,399.44		\$479.89	
37600	Ligation of neck artery	T	0093	35.5969	\$2,399.44		\$479.89	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
36558	Insert tunneled cv cath	T	0622	25.3344	\$1,707.69		\$341.54	
36560	Insert tunneled cv cath	T	0623	30.4396	\$2,051.81		\$410.37	
36561	Insert tunneled cv cath	T	0623	30.4396	\$2,051.81		\$410.37	
36565	Insert tunneled cv cath	T	0623	30.4396	\$2,051.81		\$410.37	
36566	Insert tunneled cv cath	T	0623	30.4396	\$2,051.81		\$410.37	
36568	Insert picc cath	T	0621	11.166	\$752.66		\$150.54	
36569	Insert picc cath	T	0621	11.166	\$752.66		\$150.54	
36570	Insert picvad cath	T	0622	25.3344	\$1,707.69		\$341.54	
36571	Insert picvad cath	T	0622	25.3344	\$1,707.69		\$341.54	
36575	Repair tunneled cv cath	T	0121	6.3742	\$429.66		\$85.94	
36576	Repair tunneled cv cath	T	0621	11.166	\$752.66		\$150.54	
36578	Replace tunneled cv cath	T	0622	25.3344	\$1,707.69		\$341.54	
36580	Replace cvad cath	T	0621	11.166	\$752.66		\$150.54	
36581	Replace tunneled cv cath	T	0622	25.3344	\$1,707.69		\$341.54	
36582	Replace tunneled cv cath	T	0623	30.4396	\$2,051.81		\$410.37	
36583	Replace picc cath	T	0623	30.4396	\$2,051.81		\$410.37	
36584	Replace picc cath	T	0621	11.166	\$752.66		\$150.54	
36585	Replace picvad cath	T	0622	25.3344	\$1,707.69		\$341.54	
36589	Removal tunneled cv cath	T	0121	6.3742	\$429.66		\$85.94	
36590	Removal tunneled cv cath	T	0621	11.166	\$752.66		\$150.54	
36591	Draw blood off venous device	Q1	0624	0.6132	\$41.33	\$12.65	\$8.27	
36592	Collect blood from picc	Q1	0624	0.6132	\$41.33	\$12.65	\$8.27	
36593	Declot vascular device	T	0676	2.3954	\$161.46		\$32.30	
36595	Mech remov tunneled cv cath	T	0622	25.3344	\$1,707.69		\$341.54	
36596	Mech remov tunneled cv cath	T	0621	11.166	\$752.66		\$150.54	
36597	Reposition venous catheter	T	0621	11.166	\$752.66		\$150.54	
36598	W/fluor, eval cv device	T	0676	2.3954	\$161.46		\$32.30	
36600	Withdrawal of arterial blood	Q1	0035	0.232	\$15.64		\$3.13	
36620	Insertion catheter, artery	N						
36625	Insertion catheter, artery	N						
36640	Insertion catheter, artery	C						
36660	Insertion catheter, artery	C						
36680	Insert needle, bone cavity	T	0002	1.5111	\$101.86		\$20.38	
36800	Insertion of cannula	T	0115	30.9784	\$2,088.13		\$417.63	
36810	Insertion of cannula	T	0115	30.9784	\$2,088.13		\$417.63	
36815	Insertion of cannula	T	0115	30.9784	\$2,088.13		\$417.63	
36818	Av fuse, upper arm, cephalic	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36819	Av fuse, upper arm, basilic	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36820	Av fusion/forearm vein	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36821	Av fusion direct any site	T	0088	40.9003	\$2,756.93	\$655.22	\$551.39	
36822	Insertion of cannula(s)	C						
36823	Insertion of cannula(s)	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
38220	Bone marrow aspiration	T	0003	0003	3.0998	\$208.95		\$41.79
38221	Bone marrow biopsy	T	0003	0003	3.0998	\$208.95		\$41.79
38230	Bone marrow collection	S	0112	0112	33.3206	\$2,246.01		\$449.21
38240	Bone marrow/stem transplant	S	0112	0112	33.3206	\$2,246.01		\$449.21
38241	Bone marrow/stem transplant	S	0112	0112	33.3206	\$2,246.01		\$449.21
38242	Lymphocyte infuse transplant	S	0111	0111	11.9424	\$804.99	\$198.40	\$161.00
38300	Drainage, lymph node lesion	T	0007	0007	12.6217	\$850.78		\$170.16
38305	Drainage, lymph node lesion	T	0008	0008	19.4063	\$1,308.10		\$261.62
38308	Incision of lymph channels	T	0113	0113	24.6146	\$1,659.17		\$331.84
38380	Thoracic duct procedure	C						
38381	Thoracic duct procedure	C						
38382	Thoracic duct procedure	C						
38500	Biopsy/removal, lymph nodes	T	0113	0113	24.6146	\$1,659.17		\$331.84
38505	Needle biopsy, lymph nodes	T	0005	0005	7.8145	\$526.74		\$105.35
38510	Biopsy/removal, lymph nodes	T	0113	0113	24.6146	\$1,659.17		\$331.84
38520	Biopsy/removal, lymph nodes	T	0113	0113	24.6146	\$1,659.17		\$331.84
38525	Biopsy/removal, lymph nodes	T	0113	0113	24.6146	\$1,659.17		\$331.84
38530	Biopsy/removal, lymph nodes	T	0113	0113	24.6146	\$1,659.17		\$331.84
38542	Explore deep node(s), neck	T	0114	0114	48.7561	\$3,286.45		\$657.29
38550	Removal, neck/arm/pit lesion	T	0113	0113	24.6146	\$1,659.17		\$331.84
38555	Removal, neck/arm/pit lesion	T	0113	0113	24.6146	\$1,659.17		\$331.84
38562	Removal, pelvic lymph nodes	C						
38564	Removal, abdomen lymph nodes	C						
38570	Laparoscopy, lymph node biop	T	0131	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
38571	Lymphadenectomy	T	0132	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
38572	Lymphadenectomy	T	0131	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
38589	Laparoscopy proc, lymphatic	T	0130	0130	38.054	\$2,565.07	\$659.53	\$513.02
38700	Removal of lymph nodes, neck	T	0113	0113	24.6146	\$1,659.17		\$331.84
38720	Removal of lymph nodes, neck	T	0113	0113	24.6146	\$1,659.17		\$331.84
38724	Removal of lymph nodes, neck	C						
38740	Remove armpit lymph nodes	T	0114	0114	48.7561	\$3,286.45		\$657.29
38745	Remove armpit lymph nodes	T	0114	0114	48.7561	\$3,286.45		\$657.29
38746	Remove thoracic lymph nodes	C						
38747	Remove abdominal lymph nodes	C						
38760	Remove groin lymph nodes	T	0113	0113	24.6146	\$1,659.17		\$331.84
38765	Remove groin lymph nodes	C						
38770	Remove pelvis lymph nodes	C						
38780	Remove abdomen lymph nodes	C						
38780	Inject for lymphatic x-ray	N						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
37605	Ligation of neck artery	T	0091	0091	45.0292	\$3,035.24		\$607.05
37606	Ligation of neck artery	T	0092	0092	26.5713	\$1,791.07		\$358.22
37607	Ligation of a-v fistula	T	0092	0092	26.5713	\$1,791.07		\$358.22
37609	Temporal artery procedure	T	0021	0021	17.4975	\$1,179.44		\$235.89
37615	Ligation of neck artery	T	0092	0092	26.5713	\$1,791.07		\$358.22
37616	Ligation of chest artery	C						
37617	Ligation of abdomen artery	C						
37618	Ligation of extremity artery	C						
37620	Revision of major vein	T	0091	0091	45.0292	\$3,035.24		\$607.05
37650	Revision of major vein	T	0092	0092	26.5713	\$1,791.07		\$358.22
37660	Revision of major vein	C						
37700	Revise leg vein	T	0092	0092	26.5713	\$1,791.07		\$358.22
37718	Ligate/strip short leg vein	T	0092	0092	26.5713	\$1,791.07		\$358.22
37722	Ligate/strip long leg vein	T	0091	0091	45.0292	\$3,035.24		\$607.05
37735	Removal of leg veins/lesion	T	0091	0091	45.0292	\$3,035.24		\$607.05
37760	Ligate leg veins/radical	NI						
37761	Ligate leg veins - extrem - to 20	T	0092	0092	26.5713	\$1,791.07		\$358.22
37766	Phleb veins - extrem 20+	T	0092	0092	26.5713	\$1,791.07		\$358.22
37780	Revision of leg vein	T	0092	0092	26.5713	\$1,791.07		\$358.22
37785	Ligate/divide/excise vein	T	0092	0092	26.5713	\$1,791.07		\$358.22
37788	Revascularization, penis	C						
37790	Penile venous occlusion	T	0181	0181	34.8837	\$2,351.37	\$621.82	\$470.28
37799	Vascular surgery procedure	CH	X	0624	0.6132	\$41.33	\$12.65	\$8.27
38100	Removal of spleen, total	C						
38101	Removal of spleen, partial	C						
38102	Removal of spleen, total	C						
38115	Repair of ruptured spleen	C						
38120	Laparoscopy, splenectomy	T	0131	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
38129	Laparoscopy proc, spleen	T	0130	0130	38.054	\$2,565.07	\$659.53	\$513.02
38200	Injection for spleen x-ray	N						
38204	Bi donor search management	N						
38205	Harvest allogeneic stem cells	CH	B					
38206	Harvest auto stem cells	S	0111	0111	11.9424	\$804.99	\$198.40	\$161.00
38207	Cryopreserve stem cells	S	0110	0110	3.3809	\$227.89		\$45.58
38208	Thaw preserved stem cells	S	0110	0110	3.3809	\$227.89		\$45.58
38209	Wash harvest stem cells	S	0110	0110	3.3809	\$227.89		\$45.58
38210	T-cell depletion of harvest	S	0393	0393	5.7873	\$390.10	\$79.97	\$78.02
38211	Tumor cell depletion of harvest	S	0393	0393	5.7873	\$390.10	\$79.97	\$78.02
38212	Rbc depletion of harvest	S	0393	0393	5.7873	\$390.10	\$79.97	\$78.02
38213	Platelet depletion of harvest	S	0393	0393	5.7873	\$390.10	\$79.97	\$78.02
38214	Volume depletion of harvest	S	0393	0393	5.7873	\$390.10	\$79.97	\$78.02
38215	Harvest stem cell concentrate	S	0393	0393	5.7873	\$390.10	\$79.97	\$78.02

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
40810	Excision of mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
40812	Excise/repair mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
40814	Excise/repair mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
40816	Excision of mouth lesion	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40818	Excise oral mucosa for graft	T	0251	3.425	\$230.87	\$46.18	\$46.18	
40819	Excise lip or cheek fold	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40820	Treatment of mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
40830	Repair mouth laceration	T	0251	3.425	\$230.87	\$46.18	\$46.18	
40831	Repair mouth laceration	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40840	Reconstruction of mouth	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40842	Reconstruction of mouth	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40843	Reconstruction of mouth	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40844	Reconstruction of mouth	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40845	Reconstruction of mouth	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40889	Mouth surgery procedure	T	0250	1.1522	\$77.67	\$25.10	\$15.54	
41000	Drainage of mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41005	Drainage of mouth lesion	T	0251	3.425	\$230.87	\$46.18	\$46.18	
41006	Drainage of mouth lesion	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
41007	Drainage of mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41008	Drainage of mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41010	Incision of tongue fold	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
41015	Drainage of mouth lesion	T	0251	3.425	\$230.87	\$46.18	\$46.18	
41016	Drainage of mouth lesion	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
41017	Drainage of mouth lesion	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
41018	Drainage of mouth lesion	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
41019	Place needles in for it	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
41100	Biopsy of tongue	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
41105	Biopsy of tongue	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41108	Biopsy of floor of mouth	CH	0019	4.3625	\$294.06	\$64.51	\$58.82	
41110	Excision of tongue lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41112	Excision of tongue lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41114	Excision of tongue lesion	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
41115	Excision of tongue fold	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
41116	Excision of mouth lesion	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
41120	Partial removal of tongue	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
41130	Partial removal of tongue	C						
41135	Tongue and neck surgery	C						
41140	Removal of tongue	C						
41145	Tongue removal, neck surgery	C						
41150	Tongue, mouth, jaw surgery	C						
41153	Tongue, mouth, neck surgery	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
38792	Identify sentinel node	O1	0392	2.6817	\$180.76	\$43.95	\$36.16	
38794	Access thoracic lymph duct	N						
38989	Blood/lymph system procedure	S	0110	3.3809	\$227.89		\$45.58	
39000	Exploration of chest	C						
39010	Exploration of chest	C						
39200	Removal chest lesion	C						
39220	Removal chest lesion	C						
39400	Visualization of chest	T	0069	34.0084	\$2,292.37	\$591.64	\$458.48	
39499	Chest procedure	C						
39501	Repair diaphragm laceration	C						
39502	Repair paroesophageal hernia	C						
39503	Repair of diaphragm hernia	C						
39520	Repair of diaphragm hernia	C						
39530	Repair of diaphragm hernia	C						
39531	Repair of diaphragm hernia	C						
39540	Repair of diaphragm hernia	C						
39541	Repair of diaphragm hernia	C						
39545	Revision of diaphragm	C						
39560	Resect diaphragm, simple	C						
39561	Resect diaphragm, complex	C						
39599	Diaphragm surgery procedure	C						
40490	Biopsy of lip	T	0251	3.425	\$230.87	\$46.18	\$46.18	
40500	Partial excision of lip	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
40510	Partial excision of lip	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40520	Partial excision of lip	T	0253	17.1879	\$1,158.57	\$282.29	\$231.72	
40525	Reconstruct lip with flap	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40527	Reconstruct lip with flap	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40530	Partial removal of lip	T	0254	24.9637	\$1,682.70	\$336.54	\$336.54	
40650	Repair lip	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40652	Repair lip	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40654	Repair lip	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40700	Repair cleft lip/nasal	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40701	Repair cleft lip/nasal	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40702	Repair cleft lip/nasal	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40720	Repair cleft lip/nasal	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40761	Repair cleft lip/nasal	T	0256	42.9827	\$2,897.29	\$579.46	\$579.46	
40799	Lip surgery procedure	T	0250	1.1522	\$77.67	\$25.10	\$15.54	
40800	Drainage of mouth lesion	T	0006	1.4557	\$98.12	\$19.53	\$19.53	
40801	Drainage of mouth lesion	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40804	Removal, foreign body, mouth	X	0340	0.6893	\$45.11	\$9.03	\$9.03	
40805	Removal, foreign body, mouth	T	0252	7.6196	\$513.61	\$109.16	\$109.16	
40806	Incision of lip fold	T	0251	3.425	\$230.87	\$46.18	\$46.18	
40808	Biopsy of mouth lesion	T	0251	3.425	\$230.87	\$46.18	\$46.18	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
42225	Reconstruct cleft palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42226	Lengthening of palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42227	Lengthening of palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42235	Repair palate	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42280	Preparation, palate mold	T	0251	3,425	\$230.87	\$46.18		\$46.18
42281	Insertion, palate prosthesis	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42289	Palate/uvula surgery	T	0250	1,152	\$77.67	\$15.54		\$15.54
42300	Drainage of salivary gland	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42305	Drainage of salivary gland	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42310	Drainage of salivary gland	T	0251	3,425	\$230.87	\$46.18		\$46.18
42320	Drainage of salivary gland	T	0251	3,425	\$230.87	\$46.18		\$46.18
42330	Removal of salivary stone	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42335	Removal of salivary stone	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42340	Removal of salivary stone	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42400	Biopsy of salivary gland	T	0005	7,814.5	\$526.74	\$105.35		\$105.35
42405	Biopsy of salivary gland	CH	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
42408	Excision of salivary cyst	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42409	Drainage of salivary cyst	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42410	Excise parotid gland/lesion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42415	Excise parotid gland/lesion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42420	Excise parotid gland/lesion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42425	Excise parotid gland/lesion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42426	Excise parotid gland/lesion	C						
42440	Excise sublingual gland	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42450	Repair salivary duct	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
42500	Parotid duct diversion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42507	Parotid duct diversion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42508	Parotid duct diversion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42509	Parotid duct diversion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42550	Injection for salivary x-ray	N						
42600	Closure of salivary fistula	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42650	Dilation of salivary duct	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
42660	Dilation of salivary duct	T	0251	3,425	\$230.87	\$46.18		\$46.18
42665	Ligation of salivary duct	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
42689	Salivary surgery procedure	T	0250	1,152	\$77.67	\$15.54		\$15.54
42700	Drainage of tonsil abscess	T	0251	3,425	\$230.87	\$46.18		\$46.18
42720	Drainage of throat abscess	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42725	Drainage of throat abscess	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42800	Biopsy of throat	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42802	Biopsy of throat	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
41155	Tongue, jaw, & neck surgery	C						
41250	Repair tongue laceration	T	0250	1,152	\$77.67	\$15.54		\$15.54
41251	Repair tongue laceration	T	0251	3,425	\$230.87	\$46.18		\$46.18
41252	Repair tongue laceration	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
41500	Fixation of tongue	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41510	Tongue to lip surgery	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41512	Tongue suspension	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
41520	Reconstruction, tongue fold	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
41530	Tongue base vol reduction	CH	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41599	Tongue and mouth surgery	T	0250	1,152	\$77.67	\$15.54		\$15.54
41800	Drainage of gum lesion	T	0006	1,455.7	\$98.12	\$19.63		\$19.63
41805	Removal foreign body, gum	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41806	Removal foreign body, jawbone	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41820	Excision, gum, each quadrant	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
41821	Excision of gum flap	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
41822	Excision of gum lesion	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41823	Excision of gum lesion	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41825	Excision of gum lesion	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41826	Excision of gum lesion	CH	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41827	Excision of gum lesion	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41828	Excision of gum lesion	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41830	Removal of gum tissue	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41850	Treatment of gum lesion	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41870	Gum graft	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41872	Repair gum	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
41874	Repair tooth socket	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
41899	Dental surgery procedure	T	0250	1,152	\$77.67	\$15.54		\$15.54
42000	Drainage mouth roof lesion	T	0251	3,425	\$230.87	\$46.18		\$46.18
42100	Biopsy roof of mouth	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
42104	Excision lesion, mouth roof	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42106	Excision lesion, mouth roof	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42107	Excision lesion, mouth roof	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
42120	Remove palate/lesion	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42140	Excision of uvula	T	0252	7,619.6	\$513.61	\$102.73		\$102.73
42145	Repair palate, pharynx/uvula	T	0254	24,963.7	\$1,682.70	\$336.54		\$336.54
42160	Treatment mouth roof lesion	T	0253	17,187.9	\$1,158.57	\$231.72		\$231.72
42180	Repair palate	T	0251	3,425	\$230.87	\$46.18		\$46.18
42182	Repair palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42200	Reconstruct cleft palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42205	Reconstruct cleft palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42210	Reconstruct cleft palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42215	Reconstruct cleft palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46
42220	Reconstruct cleft palate	T	0256	42,982.7	\$2,897.29	\$579.46		\$579.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
43117	Partial removal of esophagus	C						
43118	Partial removal of esophagus	C						
43121	Partial removal of esophagus	C						
43122	Partial removal of esophagus	C						
43123	Partial removal of esophagus	C						
43124	Removal of esophagus	C						
43130	Removal of esophagus pouch	T		0256	42.9827	\$2,897.29		\$579.46
43135	Removal of esophagus pouch	C						
43201	Esophagus endoscopy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43202	Esophagus endoscopy, biopsy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43205	Esophagus endoscopy, biopsy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43215	Esophagus endoscopy, biopsy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43216	Esophagus endoscopy, biopsy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43219	Esophagus endoscopy	T		0384	26.4913	\$1,785.67		\$357.14
43220	Esoph endoscopy, dilation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43226	Esoph endoscopy, dilation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43227	Esoph endoscopy, repair	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43228	Esoph endoscopy, repair	T		0422	24.2703	\$1,635.96	\$437.96	\$327.20
43231	Esoph endoscopy, ablation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43232	Esoph endoscopy, ablation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43234	Upper GI endoscopy, exam	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43235	Upper GI endoscopy, diagnosis	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43236	Upper GI endoscopy, diagnosis	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43237	Endoscopic us exam, esoph	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43238	Upper GI endoscopy w/us fn bx	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43239	Upper GI endoscopy, biopsy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43240	Upper GI endoscopy, biopsy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43241	Upper GI endoscopy with tube	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43242	Upper GI endoscopy w/us fn bx	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43243	Upper GI endoscopy & inject	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43244	Upper GI endoscopy/ligation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43245	Upper GI endoscopy, dilation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43246	Place gastrostomy tube	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43247	Operative upper GI endoscopy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43248	Upper GI endoscopy/guide wire	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43249	Esoph endoscopy, dilation	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43250	Upper GI endoscopy/tumor	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43251	Operative upper GI endoscopy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43255	Operative upper GI endoscopy	T		0141	8.7462	\$589.55	\$143.38	\$117.91
43256	Upper GI endoscopy w/stent	T		0384	26.4913	\$1,785.67		\$357.14

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
42804	Blopsy of upper nose/throat	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
42806	Blopsy of upper nose/throat	T		0254	24.9637	\$1,682.70		\$336.54
42808	Excise pharynx lesion	X		0254	24.9637	\$1,682.70		\$336.54
42809	Remove pharynx foreign body	T		0340	0.6893	\$45.11		\$9.03
42810	Excision of neck cyst	T		0254	24.9637	\$1,682.70		\$336.54
42815	Excision of neck cyst	T		0256	42.9827	\$2,897.29		\$579.46
42820	Remove tonsils and adenoids	T		0254	24.9637	\$1,682.70		\$336.54
42821	Remove tonsils and adenoids	T		0254	24.9637	\$1,682.70		\$336.54
42825	Removal of tonsils	T		0254	24.9637	\$1,682.70		\$336.54
42830	Removal of adenoids	T		0254	24.9637	\$1,682.70		\$336.54
42831	Removal of adenoids	T		0254	24.9637	\$1,682.70		\$336.54
42835	Removal of adenoids	T		0254	24.9637	\$1,682.70		\$336.54
42836	Removal of adenoids	T		0254	24.9637	\$1,682.70		\$336.54
42842	Extensive surgery of throat	T		0254	24.9637	\$1,682.70		\$336.54
42844	Extensive surgery of throat	T		0256	42.9827	\$2,897.29		\$579.46
42845	Extensive surgery of throat	C						
42846	Excision of tonsil tags	T		0254	24.9637	\$1,682.70		\$336.54
42870	Excision of lingual tonsil	T		0254	24.9637	\$1,682.70		\$336.54
42890	Partial removal of pharynx	T		0256	42.9827	\$2,897.29		\$579.46
42892	Revision of pharyngeal walls	T		0256	42.9827	\$2,897.29		\$579.46
42894	Revision of pharyngeal walls	C						
42900	Repair throat wound	T		0252	7.6196	\$513.61	\$109.16	\$102.73
42950	Reconstruction of throat	T		0254	24.9637	\$1,682.70		\$336.54
42953	Repair throat, esophagus	C						
42955	Surgical opening of throat	T		0254	24.9637	\$1,682.70		\$336.54
42960	Control throat bleeding	T		0250	1.1522	\$77.67	\$25.10	\$15.54
42961	Control throat bleeding	C						
42962	Control throat bleeding	T		0256	42.9827	\$2,897.29		\$579.46
42970	Control nose/throat bleeding	T		0250	1.1522	\$77.67	\$25.10	\$15.54
42971	Control nose/throat bleeding	C						
42972	Control nose/throat bleeding	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
42999	Throat surgery procedure	T		0250	1.1522	\$77.67	\$25.10	\$15.54
43020	Incision of esophagus	T		0252	7.6196	\$513.61	\$109.16	\$102.73
43030	Throat muscle surgery	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
43045	Incision of esophagus	C						
43100	Excision of esophagus lesion	C						
43101	Excision of esophagus lesion	C						
43107	Removal of esophagus	C						
43108	Removal of esophagus	C						
43112	Removal of esophagus	C						
43113	Removal of esophagus	C						
43116	Partial removal of esophagus	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
43415	Repair esophagus wound	T	C	0254	24.9637	\$1,682.70		\$336.54
43420	Repair esophagus opening	T	C	0140	5.9622	\$401.89	\$84.79	\$60.38
43425	Dilate esophagus	T	C	0140	5.9622	\$401.89	\$84.79	\$60.38
43450	Dilate esophagus	T	C	0140	5.9622	\$401.89	\$84.79	\$60.38
43456	Dilate esophagus	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43458	Pressure treatment esophagus	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43466	Free jejunum flap, microvasc	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43499	Esophagus surgery procedure	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43500	Surgical opening of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43501	Surgical repair of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43502	Surgical repair of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43510	Surgical opening of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43520	Incision of pyloric muscle	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43600	Biopsy of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43605	Biopsy of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43610	Excision of stomach lesion	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43611	Excision of stomach lesion	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43620	Removal of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43621	Removal of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43622	Removal of stomach	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43631	Removal of stomach, partial	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43632	Removal of stomach, partial	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43633	Removal of stomach, partial	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43634	Removal of stomach, partial	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43635	Removal of stomach, partial	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43640	Vagotomy & pylorus repair	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43641	Vagotomy & pylorus repair	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43644	Lap gastric bypass/roux-en-y	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43645	Lap gastric bypass incl simli	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43647	Lap impl electrode, antrum	S	C	0061	86.5171	\$5,831.77		\$1,166.36
43648	Lap revise/remv efltrd antrum	T	C	0130	38.054	\$2,565.07	\$659.53	\$513.02
43651	Laparoscopy, vagus nerve	T	C	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
43652	Laparoscopy, vagus nerve	T	C	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
43653	Laparoscopy, gastrostomy	T	C	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
43659	Laparoscopy proc, stom	T	C	0130	38.054	\$2,565.07	\$659.53	\$513.02
43732	Nasal/orogastric w/strnt	X	C	0272	1.2693	\$85.56	\$31.15	\$17.12
43760	Change gastrostomy tube	CH	T	0676	2.3954	\$161.48		\$32.30
43761	Reposition gastrostomy tube	CH	T	0676	2.3954	\$161.48		\$32.30
43770	Lap place gastr adj device	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43771	Lap revise gastr adj device	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43772	Lap rrmvl gastr adj device	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
43257	Upr gl scope w/rlml txmnt	T	C	0422	24.2703	\$1,635.96	\$437.96	\$327.20
43258	Operative upper GI endoscopy	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43259	Endoscopic ultrasound exam	T	C	0141	8.7462	\$589.55	\$143.38	\$117.91
43260	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43261	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43262	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43263	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43264	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43265	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43266	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43268	Endo cholangiopancreatograph	T	C	0384	26.4913	\$1,785.67	\$357.14	\$357.14
43269	Endo cholangiopancreatograph	T	C	0384	26.4913	\$1,785.67	\$357.14	\$357.14
43271	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43272	Endo cholangiopancreatograph	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43273	Endoscopic pancreatoscopy	T	C	0151	22.6111	\$1,524.12	\$304.83	\$304.83
43279	Lap myotomy, heller	C	C					
43280	Laparoscopy, fundoplasty	T	C	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
43281	Lap paraesophag hern repair	NI	C					
43289	Laparoscopy proc, esoph	NI	C					
43300	Repair of esophagus	T	C	0130	38.054	\$2,565.07	\$659.53	\$513.02
43305	Repair esophagus and fistula	T	C					
43310	Repair of esophagus	T	C					
43312	Repair esophagus and fistula	T	C					
43313	Esophagectomy congenital	T	C					
43314	Tracheo-esophagectomy cong	T	C					
43320	Fuse esophagus & stomach	T	C					
43324	Revise esophagus & stomach	T	C					
43325	Revise esophagus & stomach	T	C					
43326	Revise esophagus & stomach	T	C					
43330	Repair of esophagus	T	C					
43331	Repair of esophagus	T	C					
43340	Fuse esophagus & intestine	T	C					
43341	Fuse esophagus & intestine	T	C					
43350	Surgical opening, esophagus	T	C					
43351	Surgical opening, esophagus	T	C					
43352	Surgical opening, esophagus	T	C					
43360	Gastrointestinal repair	T	C					
43361	Gastrointestinal repair	T	C					
43400	Ligate esophagus veins	T	C					
43401	Esophagus surgery for veins	T	C					
43405	Ligate/staple esophagus	T	C					
43410	Repair esophagus wound	T	C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
44126	Enterectomy w/o taper, cong	C	C					
44127	Enterectomy w/taper, cong	C	C					
44128	Enterectomy cong, add-on	C	C					
44130	Bowel to bowel fusion	C	C					
44132	Enterectomy, cadaver donor	C	C					
44133	Enterectomy, live donor	C	C					
44135	Intestine transplant, cadaver	C	C					
44136	Intestine transplant, live	C	C					
44137	Remove intestinal allograft	C	C					
44139	Mobilization of colon	C	C					
44140	Partial removal of colon	C	C					
44141	Partial removal of colon	C	C					
44143	Partial removal of colon	C	C					
44144	Partial removal of colon	C	C					
44145	Partial removal of colon	C	C					
44146	Partial removal of colon	C	C					
44147	Partial removal of colon	C	C					
44150	Removal of colon	C	C					
44151	Removal of colon/ileostomy	C	C					
44155	Removal of colon/ileostomy	C	C					
44156	Removal of colon/ileostomy	C	C					
44157	Colectomy w/ileoanal anast	C	C					
44158	Colectomy w/ileo-rectum pouch	C	C					
44160	Removal of colon	C	C					
44180	Lap. enterolysis	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
44186	Lap. jejunostomy	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
44187	Lap. ileo/jejunostomy	C	C					
44188	Lap. colectomy	C	C					
44202	Lap. resect s/intestine	C	C					
44203	Lap. partial colectomy	C	C					
44204	Lap. colectomy part w/ileum	C	C					
44206	Lap part colectomy w/stoma	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
44207	L. colectomy/coloproctostomy	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
44208	L. colectomy/coloproctostomy	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
44210	Laparo total proctocolectomy	C	C					
44211	Lap colectomy w/proctectomy	C	C					
44212	Laparo total proctocolectomy	C	C					
44213	Lap. mobil splenic fl add-on	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
44227	Lap. close enterostomy	C	C					
44238	Laparoscopy proc. intestine	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
44300	Open bowel to skin	C	C					
44310	Ileostomy/jejunostomy	C	C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
43773	Lap replace gastr adj device	C	C					
43774	Lap rmyl gastr adj all parts	C	C					
43775	Lap sleeve gastrectomy	NI	C					
43800	Reconstruction of pylorus	C	C					
43810	Fusion of stomach and bowel	C	C					
43820	Fusion of stomach and bowel	C	C					
43825	Fusion of stomach and bowel	C	C					
43830	Piece gastrostomy tube	T	0422		24.2703	\$1,636.96	\$437.96	\$327.20
43831	Piece gastrostomy tube	T	0141		8.7462	\$589.55	\$143.38	\$117.91
43832	Piece gastrostomy tube	C	C					
43840	Repair of stomach lesion	C	C					
43842	V-band gastroplasty	E	C					
43843	Gastroplasty w/o v-band	C	C					
43845	Gastroplasty duodenal switch	C	C					
43846	Gastroplasty for obesity	C	C					
43847	Gastric bypass incl small I	C	C					
43848	Revision gastroplasty	C	C					
43850	Revise stomach-bowel fusion	C	C					
43855	Revise stomach-bowel fusion	C	C					
43860	Revise stomach-bowel fusion	C	C					
43870	Repair stomach opening	T	0141		8.7462	\$589.55	\$143.38	\$117.91
43880	Repair stomach-bowel fistula	C	C					
43881	Impl/retract electr, antrum	C	C					
43882	Revise/remove electr antrum	C	C					
43886	Revise gastric port, open	T	0137		23.9317	\$1,613.14	\$322.63	\$222.63
43887	Remove gastric port, open	T	0135		4.4386	\$299.19	\$59.84	\$59.84
43888	Change gastric port, open	T	0137		23.9317	\$1,613.14	\$322.63	\$222.63
43899	Stomach surgery procedure	T	0141		8.7462	\$589.55	\$143.38	\$117.91
44005	Freeing of bowel adhesion	C	C					
44010	Incision of small bowel	C	C					
44015	Insert needle cath bowel	C	C					
44020	Explore small intestine	C	C					
44021	Decompress small bowel	C	C					
44025	Incision of large bowel	C	C					
44050	Reduce bowel obstruction	C	C					
44055	Correct malrotation of bowel	C	C					
44100	Biopsy of bowel	T	0141		8.7462	\$589.55	\$143.38	\$117.91
44110	Excise intestine lesion(s)	C	C					
44111	Excision of bowel lesion(s)	C	C					
44120	Removal of small intestine	C	C					
44121	Removal of small intestine	C	C					
44125	Removal of small intestine	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
44626	Repair bowel opening	C	C					
44640	Repair bowel-skin fistula	C	C					
44650	Repair bowel fistula	C	C					
44660	Repair bowel-bladder fistula	C	C					
44661	Repair bowel-bladder fistula	C	C					
44680	Surgical revision, intestine	C	C					
44700	Suspend bowel w/prosthesis	C	C					
44701	Intraop colon lavage add-on	N	N					
44715	Prep donor intestine/venous	C	C					
44720	Prep donor intestine/venous	C	C					
44721	Prep donor intestine/artery	C	C					
44799	Unlisted procedure intestine	T	T	0153	27.1855	\$1,832.47	\$376.05	\$366.50
44800	Excision of bowel pouch	C	C					
44820	Excision of mesentery lesion	C	C					
44850	Repair of mesentery	C	C					
44899	Bowel surgery procedure	C	C					
44901	Drain app abscess, open	T	T	0037	15.5003	\$1,044.81	\$228.76	\$208.97
44950	Appendectomy	CH	T	0153	27.1855	\$1,832.47	\$376.05	\$366.50
44955	Appendectomy add-on	CH	T	0153	27.1855	\$1,832.47	\$376.05	\$366.50
44960	Appendectomy	C	C					
44970	Laparoscopy, appendectomy	T	T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
44979	Laparoscopy proc, app	T	T	0130	38.054	\$2,565.07	\$659.53	\$513.02
45000	Drainage of pelvic abscess	T	T	0155	14.114	\$951.37		\$190.28
45005	Drainage of rectal abscess	T	T	0155	14.114	\$951.37		\$190.28
45020	Drainage of rectal abscess	T	T	0155	14.114	\$951.37		\$190.28
45100	Biopsy of rectum	T	T	0149	23.9703	\$1,615.74		\$323.15
45108	Removal of anorectal lesion	T	T	0149	23.9703	\$1,615.74		\$323.15
45110	Removal of rectum	C	C					
45111	Partial removal of rectum	C	C					
45112	Removal of rectum	C	C					
45113	Partial proctectomy	C	C					
45114	Partial removal of rectum	C	C					
45116	Partial removal of rectum	C	C					
45119	Remove rectum w/reservoir	C	C					
45120	Removal of rectum	C	C					
45121	Removal of rectum and colon	C	C					
45123	Partial proctectomy	C	C					
45126	Pelvic exenteration	C	C					
45130	Excision of rectal prolapse	C	C					
45135	Excision of rectal prolapse	C	C					
45136	Excise ileoanal reservoir	C	C					
45150	Excision of rectal stricture	T	T	0149	23.9703	\$1,615.74		\$323.15

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
44312	Revision of ileostomy	C	C	0137	23.9317	\$1,613.14		\$322.63
44314	Revision of ileostomy	C	C					
44316	Devised bowel pouch	C	C					
44320	Colostomy	C	C					
44322	Colostomy with biopsies	C	C					
44340	Revision of colostomy	C	C	0137	23.9317	\$1,613.14		\$322.63
44346	Revision of colostomy	C	C					
44360	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44361	Small bowel endoscopy/biopsy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44363	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44364	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44365	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44366	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44369	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44370	Small bowel endoscopy/stent	T	T	0384	26.4913	\$1,785.67		\$357.14
44372	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44373	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44376	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44377	Small bowel endoscopy/biopsy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44378	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44379	S bowel endoscope w/stent	T	T	0384	26.4913	\$1,785.67		\$357.14
44380	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44382	Small bowel endoscopy	T	T	0142	9.8503	\$663.97	\$152.78	\$132.80
44383	Ileoscopy w/stent	T	T	0384	26.4913	\$1,785.67		\$357.14
44385	Endoscopy of bowel pouch	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44386	Endoscopy, bowel pouch/biops	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44388	Colonoscopy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44389	Colonoscopy with biopsy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44390	Colonoscopy for foreign body	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44391	Colonoscopy for bleeding	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44392	Colonoscopy & polypectomy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44393	Colonoscopy, lesion removal	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44394	Colonoscopy w/snares	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
44397	Colonoscopy w/stent	T	T	0384	26.4913	\$1,785.67		\$357.14
44500	Intra. gastrointestinal tube	T	T	0121	6.3742	\$429.66		\$85.94
44603	Suture, small intestine	C	C					
44604	Suture, large intestine	C	C					
44605	Repair of bowel lesion	C	C					
44615	Intestinal stricturoplasty	C	C					
44620	Repair bowel opening	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
45397	Lap. remove rectum w/pouch	C	C					
45400	Laparoscopic proc.	C	C					
45402	Lap proctopexy w/sig resect	C	C	0130	38.054	\$2,585.07	\$659.53	\$513.02
45499	Laparoscopic proc. rectum	T	T	0149	23.9703	\$1,615.74		\$323.15
45500	Repair of rectum	T	T	0150	32.1812	\$2,169.21	\$437.12	\$433.85
45505	Repair of rectum	T	T	0013	0.8789	\$69.24		\$11.85
45520	Treatment of rectal prolapse	C	C					
45540	Correct rectal prolapse	C	C	0150	32.1812	\$2,169.21	\$437.12	\$433.85
45550	Repair rectum/remove sigmoid	C	C	0150	32.1812	\$2,169.21	\$437.12	\$433.85
45560	Repair of rectocele	T	T	0150	32.1812	\$2,169.21	\$437.12	\$433.85
45562	Exploration/repair of rectum	C	C					
45563	Exploration/repair of rectum	C	C					
45800	Repair recti/bladder fistula	C	C					
45805	Repair fistula w/colostomy	C	C					
45820	Repair rectourethral fistula	C	C					
45825	Repair fistula w/colostomy	C	C					
45900	Reduction of rectal prolapse	C	C	0148	5.365	\$361.63		\$72.33
45905	Dilation of anal sphincter	T	T	0149	23.9703	\$1,615.74		\$323.15
45910	Dilation of rectal narrowing	T	T	0149	23.9703	\$1,615.74		\$323.15
45915	Remove rectal obstruction	T	T	0155	14.114	\$951.37		\$190.28
45990	Surg dx exam. anorectal	T	T	0149	23.9703	\$1,615.74		\$323.15
45999	Rectum surgery procedure	T	T	0148	5.365	\$361.63		\$72.33
46020	Placement of seton	T	T	0149	23.9703	\$1,615.74		\$323.15
46030	Removal of rectal marker	T	T	0148	5.365	\$361.63		\$72.33
46040	Incision of rectal abscess	T	T	0149	23.9703	\$1,615.74		\$323.15
46045	Incision of rectal abscess	T	T	0149	23.9703	\$1,615.74		\$323.15
46050	Incision of anal abscess	T	T	0155	14.114	\$951.37		\$190.28
46060	Incision of rectal abscess	T	T	0149	23.9703	\$1,615.74		\$323.15
46070	Incision of anal septum	T	T	0149	23.9703	\$1,615.74		\$323.15
46080	Incision of anal sphincter	T	T	0149	23.9703	\$1,615.74		\$323.15
46083	Incise external hemorrhoid	T	T	0164	2.0194	\$136.12		\$27.23
46200	Removal of anal fissure	T	T	0149	23.9703	\$1,615.74		\$323.15
46210	Removal of anal crypt	CH	D					
46211	Removal of anal crypt	CH	D					
46220	Excise anal ext tag/papilla	CH	T	0155	14.114	\$951.37		\$190.28
46221	Ligation of hemorrhoid(s)	CH	T	0148	5.365	\$361.63		\$72.33
46230	Removal of anal tags	T	T	0149	23.9703	\$1,615.74		\$323.15
46250	Remove ext hem. groups = 2	T	T	0149	23.9703	\$1,615.74		\$323.15
46255	Remove int/ext hem. 1 group	T	T	0149	23.9703	\$1,615.74		\$323.15
46257	Remove int/ext hem. grp. & fiss	T	T	0149	23.9703	\$1,615.74		\$323.15
46258	Remove int/ext hem. grp. w/fissu	T	T	0149	23.9703	\$1,615.74		\$323.15
46260	Remove int/ext hem. groups = 2	T	T	0149	23.9703	\$1,615.74		\$323.15

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
45160	Excision of rectal lesion	CH	D	0149	23.9703	\$1,615.74		\$323.15
45170	Excision of rectal lesion	CH	D	0149	23.9703	\$1,615.74		\$323.15
45171	Exc. rect. tum. transanal part	NI	T	0155	14.114	\$951.37		\$190.28
45172	Exc. rect. tum. transanal full	NI	T	0149	23.9703	\$1,615.74		\$323.15
45190	Destruction, rectal tumor	T	T	0149	23.9703	\$1,615.74		\$323.15
45300	Proctosigmoidoscopy dk	T	T	0146	5.7747	\$389.25		\$77.85
45303	Proctosigmoidoscopy dilate	T	T	0147	9.245	\$623.17		\$124.64
45305	Proctosigmoidoscopy w/bx	T	T	0147	9.245	\$623.17		\$124.64
45307	Proctosigmoidoscopy fb	T	T	0428	22.8208	\$1,538.26	\$307.66	\$307.66
45308	Proctosigmoidoscopy removal	T	T	0147	9.245	\$623.17		\$124.64
45309	Proctosigmoidoscopy removal	T	T	0147	9.245	\$623.17		\$124.64
45315	Proctosigmoidoscopy bleed	T	T	0147	9.245	\$623.17		\$124.64
45317	Proctosigmoidoscopy bleed	T	T	0147	9.245	\$623.17		\$124.64
45320	Proctosigmoidoscopy ablate	T	T	0428	22.8208	\$1,538.26	\$307.66	\$307.66
45321	Proctosigmoidoscopy w/vulv	T	T	0428	22.8208	\$1,538.26	\$307.66	\$307.66
45327	Proctosigmoidoscopy w/inst	T	T	0384	26.4913	\$1,785.67	\$357.14	\$357.14
45330	Diagnostic sigmoidoscopy	T	T	0146	5.7747	\$389.25		\$77.85
45331	Sigmoidoscopy and biopsy	T	T	0146	5.7747	\$389.25		\$77.85
45332	Sigmoidoscopy w/bf removal	T	T	0146	5.7747	\$389.25		\$77.85
45333	Sigmoidoscopy & polypectomy	T	T	0147	9.245	\$623.17		\$124.64
45334	Sigmoidoscopy for bleeding	T	T	0147	9.245	\$623.17		\$124.64
45335	Sigmoidoscopy w/submuc inj	T	T	0146	5.7747	\$389.25		\$77.85
45337	Sigmoidoscopy & decompress	T	T	0146	5.7747	\$389.25		\$77.85
45338	Sigmoidoscopy w/tumr remove	T	T	0147	9.245	\$623.17		\$124.64
45339	Sigmoidoscopy w/ablate tumr	T	T	0147	9.245	\$623.17		\$124.64
45340	Sig. w/ablation dilation	T	T	0147	9.245	\$623.17		\$124.64
45341	Sigmoidoscopy w/ultrasound	T	T	0147	9.245	\$623.17		\$124.64
45342	Sigmoidoscopy w/us guide bx	T	T	0384	26.4913	\$1,785.67	\$357.14	\$357.14
45345	Sigmoidoscopy w/inst	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45355	Surgical colonoscopy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45378	Diagnostic colonoscopy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45379	Colonoscopy w/bf removal	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45380	Colonoscopy and biopsy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45381	Colonoscopy, submucous inj	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45382	Colonoscopy/control bleeding	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45383	Lesion removal colonoscopy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45384	Lesion remove colonoscopy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45385	Lesion removal colonoscopy	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45386	Colonoscopy dilate stricture	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45387	Colonoscopy w/inst	T	T	0384	26.4913	\$1,785.67	\$357.14	\$357.14
45391	Colonoscopy w/endoscope us	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45392	Colonoscopy w/endoscopic fib	T	T	0143	9.1051	\$613.74	\$186.06	\$122.75
45395	Lap. removal of rectum	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
46916	Cryosurgery, anal lesion(s)	T		0015	1.5412	\$103.89		\$20.78
46917	Laser surgery, anal lesions	T		0017	21.2653	\$1,433.41		\$286.69
46922	Excision of anal lesion(s)	T		0017	21.2653	\$1,433.41		\$286.69
46924	Destruction, anal lesion(s)	T		0017	21.2653	\$1,433.41		\$286.69
46930	Destruct internal hemorrhoids	T		0148	5.365	\$361.63		\$72.33
46937	Cryotherapy of rectal lesion	CH	D					
46938	Treatment of anal fissure	CH	D					
46940	Treatment of anal fissure	T		0149	23.9703	\$1,615.74		\$323.15
46942	Remove by ligat int hem grp	T		0155	14.114	\$951.37		\$190.28
46945	Remove by ligat int hem grp	T		0155	14.114	\$951.37		\$190.28
46946	Hemorrhoidectomy by stapling	T		0150	32.1812	\$2,169.21		\$437.12
46947	Needle biopsy of liver	T		0685	9.6666	\$651.59		\$130.32
47001	Open drainage, liver add-on	N						
47010	Partial removal of liver	C		0037	15.5003	\$1,044.81		\$208.97
47011	Inject/aspirate liver cyst	C						
47015	Wedge biopsy of liver	C						
47100	Partial removal of liver	C						
47120	Extensive removal of liver	C						
47122	Partial removal of liver	C						
47130	Partial removal of liver	C						
47133	Removal of donor liver	C						
47135	Transplantation of liver	C						
47136	Partial removal, donor liver	C						
47140	Partial removal, donor liver	C						
47141	Partial removal, donor liver	C						
47142	Prep donor liver, whole	C						
47143	Prep donor liver, 3-segment	C						
47144	Prep donor liver, lobe split	C						
47145	Prep donor liver/venous	C						
47146	Prep donor liver/arterial	C						
47300	Surgery for liver lesion	C						
47360	Repair liver wound	C						
47361	Repair liver wound	C						
47362	Repair liver wound	C						
47370	Laparoscopic procedure, liver	T		0174	109.9237	\$7,409.52	\$2,069.30	\$1,481.91
47371	Laparoscopic procedure, liver	CH	T	0174	109.9237	\$7,409.52	\$2,069.30	\$1,481.91
47379	Open ablate liver tumor, r	T		0130	38.054	\$2,565.07	\$659.53	\$513.02
47380	Open ablate liver tumor, r	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
46261	Remove inflex hem grps & fiss	T		0149	23.9703	\$1,615.74		\$323.15
46262	Remove inflex hem grps w/fist	T		0149	23.9703	\$1,615.74		\$323.15
46270	Remove anal fist subq	T		0149	23.9703	\$1,615.74		\$323.15
46275	Remove anal fist inter	T		0149	23.9703	\$1,615.74		\$323.15
46280	Remove anal fist complex	T		0149	23.9703	\$1,615.74		\$323.15
46285	Remove anal fist 2 stage	T		0149	23.9703	\$1,615.74		\$323.15
46288	Repair anal fistula	T		0149	23.9703	\$1,615.74		\$323.15
46320	Removal of hemorrhoid clot	T		0149	23.9703	\$1,615.74		\$323.15
46500	Injection into hemorrhoid(s)	T		0155	14.114	\$951.37		\$190.28
46505	Chemodenervation anal musc	CH		0149	23.9703	\$1,615.74		\$323.15
46600	Diagnostic anoscopy	X		0340	0.6693	\$45.11		\$9.03
46604	Anoscopy and dilation	T		0147	9.245	\$623.17		\$124.64
46606	Anoscopy, remove lesion	T		0146	5.7747	\$389.25		\$77.85
46608	Anoscopy, remove for body	T		0147	9.245	\$623.17		\$124.64
46611	Anoscopy	T		0147	9.245	\$623.17		\$124.64
46612	Anoscopy, remove lesions	T		0428	22.8208	\$1,538.26		\$307.66
46614	Anoscopy, control bleeding	T		0146	5.7747	\$389.25		\$77.85
46615	Anoscopy	T		0428	22.8208	\$1,538.26		\$307.66
46700	Repair of anal stricture	T		0149	23.9703	\$1,615.74		\$323.15
46705	Repr of anal fistula w/glu	C						
46706	Repr of anal fistula w/glu	T		0150	32.1812	\$2,169.21	\$437.12	\$433.85
46707	Repair anorectal fist w/plug	NI		0150	32.1812	\$2,169.21	\$437.12	\$433.85
46710	Repr per/vag pouch sngl proc	C						
46712	Repr per/vag pouch dbl proc	C						
46715	Rep per/ anoper fistu	C						
46716	Rep per/ anoper/vesib fistu	C						
46730	Construction of absent anus	C						
46735	Construction of absent anus	C						
46740	Construction of absent anus	C						
46742	Repair of imperforated anus	C						
46744	Repair of cloacal anomaly	C						
46746	Repair of cloacal anomaly	C						
46748	Repair of cloacal anomaly	C						
46750	Repair of anal sphincter	C		0150	32.1812	\$2,169.21	\$437.12	\$433.85
46751	Repair of anal sphincter	C						
46753	Reconstruction of anus	T		0149	23.9703	\$1,615.74		\$323.15
46754	Removal of suture from anus	T		0149	23.9703	\$1,615.74		\$323.15
46760	Repair of anal sphincter	T		0150	32.1812	\$2,169.21	\$437.12	\$433.85
46761	Repair of anal sphincter	T		0150	32.1812	\$2,169.21	\$437.12	\$433.85
46762	Implant artificial sphincter	T		0150	32.1812	\$2,169.21	\$437.12	\$433.85
46800	Destruction, anal lesion(s)	T		0016	2.7982	\$188.62		\$37.73
46910	Destruction, anal lesion(s)	T		0017	21.2653	\$1,433.41		\$286.69

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
47760	Fuse bile ducts and bowel	C	C					
47765	Fuse liver ducts & bowel	C	C					
47780	Fuse bile ducts and bowel	C	C					
47785	Fuse bile ducts and bowel	C	C					
47800	Reconstruction of bile ducts	C	C					
47801	Placement, bile duct support	C	C					
47802	Fuse liver duct & intestine	C	C					
47900	Suture bile duct injury	C	C					
47999	Bile tract surgery procedure	T	0152		30.743	\$2,072.26		\$414.46
48000	Drainage of abdomen	C	C					
48001	Placement of drain, pancreas	C	C					
48020	Removal of pancreatic stone	C	C					
48100	Biopsy of pancreas, open	C	C					
48102	Needle biopsy, pancreas	T	0685		9.6666	\$661.59		\$130.32
48105	Resect/obtrude pancreas	C	C					
48120	Removal of pancreas lesion	C	C					
48140	Partial removal of pancreas	C	C					
48145	Partial removal of pancreas	C	C					
48146	Pancreatectomy	C	C					
48148	Removal of pancreatic duct	C	C					
48150	Partial removal of pancreas	C	C					
48152	Pancreatectomy	C	C					
48153	Pancreatectomy	C	C					
48154	Pancreatectomy	C	C					
48155	Removal of pancreas	C	C					
48160	Pancreas removal/transplant	E	E					
48400	Injection, intraop add-on	C	C					
48500	Surgery of pancreatic cyst	C	C					
48510	Drain pancreatic pseudocyst	C	C					
48511	Drain pancreatic pseudocyst	T	0037		15.5003	\$1,044.81	\$228.76	\$208.97
48520	Fuse pancreas cyst and bowel	C	C					
48540	Fuse pancreas cyst and bowel	C	C					
48545	Pancreatohaphy	C	C					
48547	Duodenal exclusion	C	C					
48548	Fuse pancreas and bowel	C	C					
48550	Donor pancreatectomy	E	E					
48551	Prep donor pancreas	C	C					
48552	Prep donor pancreas/venous	C	C					
48554	Transpl allograft pancreas	C	C					
48556	Removal, allograft pancreas	C	C					
48999	Pancreas surgery procedure	T	0004		4.5991	\$310.01		\$62.01
49000	Exploration of abdomen	C	C					
49002	Reopening of abdomen	C	C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
47381	Open ablate liver tumor cryo	C	C					
47382	Percut ablate liver rf	T	0423		51.3618	\$3,462.09		\$692.42
47399	Liver surgery procedure	T	0004		4.5991	\$310.01		\$62.01
47400	Incision of liver duct	C	C					
47420	Incision of bile duct	C	C					
47425	Incision of bile duct	C	C					
47460	Incise bile duct sphincter	C	C					
47480	Incision of gallbladder	C	C					
47490	Incision of gallbladder	T	0152		30.743	\$2,072.26		\$414.46
47500	Injection for liver x-rays	N	N					
47505	Injection for liver x-rays	N	N					
47510	Insert catheter, bile duct	T	0152		30.743	\$2,072.26		\$414.46
47511	Insert bile duct drain	T	0152		30.743	\$2,072.26		\$414.46
47525	Change bile duct catheter	T	0427		15.3103	\$1,032.01		\$206.41
47530	Reviser/reinsert bile tube	T	0427		15.3103	\$1,032.01		\$206.41
47550	Bile duct endoscopy add-on	C	C					
47552	Biliary endoscopy thru skin	T	0152		30.743	\$2,072.26		\$414.46
47553	Biliary endoscopy thru skin	T	0152		30.743	\$2,072.26		\$414.46
47554	Biliary endoscopy thru skin	T	0152		30.743	\$2,072.26		\$414.46
47555	Biliary endoscopy thru skin	T	0152		30.743	\$2,072.26		\$414.46
47560	Laparoscopy, w/cholangio	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
47561	Laparo w/cholangiobiopsy	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
47562	Laparoscopic cholecystectomy	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
47563	Laparo cholecystectomy/graph	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
47564	Laparo cholecystectomy/explr	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
47570	Laparo cholecystenterostomy	C	C					
47579	Laparoscopy proc, biliary	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
47600	Removal of gallbladder	C	C					
47605	Removal of gallbladder	C	C					
47610	Removal of gallbladder	C	C					
47620	Removal of gallbladder	C	C					
47630	Remove bile duct stone	T	0152		30.743	\$2,072.26		\$414.46
47700	Exploration of bile ducts	C	C					
47701	Bile duct revision	C	C					
47711	Excision of bile duct tumor	C	C					
47712	Excision of bile duct tumor	C	C					
47715	Excision of bile duct cyst	C	C					
47720	Fuse gallbladder & bowel	C	C					
47721	Fuse upper gi structures	C	C					
47740	Fuse gallbladder & bowel	C	C					
47741	Fuse gallbladder & bowel	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
49441	Place dirod/fej tube perc	T	0141	8.7462	\$589.55	\$143.38	\$17.91	
49442	Place cecostomy tube perc	T	0155	14.114	\$951.37	\$143.38	\$190.28	
49446	Change g-tube to g1 perc	T	0141	8.7462	\$589.55	\$143.38	\$117.91	
49450	Replace g1 tube perc	T	0121	6.3742	\$429.66	\$85.94	\$85.94	
49451	Replace g1 tube perc	T	0121	6.3742	\$429.66	\$85.94	\$85.94	
49452	Replace g1 tube perc	T	0121	6.3742	\$429.66	\$85.94	\$85.94	
49460	Fix g/colon tube widevics	T	0121	6.3742	\$429.66	\$85.94	\$85.94	
49465	Fluro exam of g/colon tube	Q1	0276	1.2985	\$87.53	\$34.74	\$17.51	
49491	Rpr hern premie reduc	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49492	Rpr hern premie, blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49495	Rpr ing hernia baby, reduc	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49496	Rpr ing hernia baby, blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49500	Rpr ing hernia, init, reduce	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49501	Rpr ing hernia, init blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49505	Pp /hern init reduc >5 yr	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49507	Pp /hern init reduc >5 yr	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49520	Rerepair ing hernia, reduce	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49521	Rerepair ing hernia, blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49525	Repair ing hernia, sliding	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49540	Repair lumbar hernia	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49550	Rpr rem hernia, init, reduce	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49553	Rpr fem hernia, init blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49555	Rerepair fem hernia, reduce	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49557	Rerepair fem hernia, blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49560	Rpr ventral hern init, reduc	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49561	Rpr ventral hern init, block	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49565	Rerepair ventri hern, reduce	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49566	Rerepair ventri hern, block	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49568	Hernia repair w/mesh	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49570	Rpr epigastric hern, reduce	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49572	Rpr epigastric hern, blocked	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49580	Rpr umbil hern, reduc < 5 yr	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49582	Rpr umbil hern, block < 5 yr	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49585	Rpr umbil hern, reduc > 5 yr	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49587	Rpr umbil hern, block > 5 yr	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49590	Repair spigelian hernia	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
49600	Repair umbilical lesion	C				\$464.85	\$430.97	
49605	Repair umbilical lesion	C				\$464.85	\$430.97	
49606	Repair umbilical lesion	C				\$464.85	\$430.97	
49610	Repair umbilical lesion	C				\$464.85	\$430.97	
49611	Repair umbilical lesion	C				\$464.85	\$430.97	
49650	Lap ing hernia repair init	T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46	
49651	Lap ing hernia repair recur	T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
49010	Exploration behind abdomen	C						
49020	Drain abdominal abscess	C						
49021	Drain abdominal abscess	T	0037	15.5003	\$1,044.81	\$228.76	\$208.97	
49040	Drain, open, abdom abscess	C						
49041	Drain, perc, abdom abscess	T	0037	15.5003	\$1,044.81	\$228.76	\$208.97	
49060	Drain, open, retroper abscess	C						
49061	Drain, perc, retroper abscess	T	0037	15.5003	\$1,044.81	\$228.76	\$208.97	
49062	Drain to peritoneal cavity	C						
49080	Puncture, peritoneal cavity	T	0070	5.5521	\$374.24	\$74.85	\$74.85	
49081	Removal of abdominal fluid	T	0070	5.5521	\$374.24	\$74.85	\$74.85	
49180	Biopsy, abdominal mass	T	0685	9.6666	\$651.59	\$130.32	\$130.32	
49203	Exc abd tum 5 cm or less	C						
49204	Exc abd tum over 5 cm	C						
49205	Exc abd tum over 10 cm	C						
49215	Excise sacral spine tumor	C						
49220	Multiple surgery, abdomen	C						
49250	Excision of umbilicus	T	0153	27.1855	\$1,832.47	\$376.05	\$366.50	
49255	Removal of omentum	C						
49320	Diag laparo separate proc	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49321	Laparoscopy, biopsy	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49322	Laparoscopy, aspiration	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49323	Laparo drain lymphocele	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49324	Lap insertion perm ip cath	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49325	Lap revision perm ip cath	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49326	Lap w/omentopexy add-on	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49329	Laparo proc, abdm/per/oment	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
49400	Air injection into abdomen	N						
	Remove foreign body,							
49402	abdomen	T	0153	27.1855	\$1,832.47	\$376.05	\$366.50	
49411	Ins mark abd/pe/ for rt perq	NI	X	0310	13.7576	\$927.34	\$185.47	
49419	Insrt abdom cath for chemox	T	0115	30.9784	\$2,088.13	\$325.27	\$417.63	
49420	Insert abdom drain, temp	T	0652	30.4602	\$2,053.20	\$410.64	\$410.64	
49421	Insert abdom drain, perm	T	0652	30.4602	\$2,053.20	\$410.64	\$410.64	
49422	Remove perm cannula/catheter	T	0105	22.9412	\$1,546.37	\$309.28	\$309.28	
49423	Exchange drainage catheter	T	0427	15.3103	\$1,032.01	\$206.41	\$206.41	
49424	Assess cyst, contrast inject	N						
49425	Insert abdomen-venous drain	C						
49426	Revise abdomen-venous shunt	T	0153	27.1855	\$1,832.47	\$376.05	\$366.50	
49427	Injection, abdominal shunt	N						
49428	Ligation of shunt	C						
49429	Removal of shunt	C						
49435	Insert subn exten to ip cath	T	0105	22.9412	\$1,546.37	\$309.28	\$309.28	
49436	Embedded ip cath exit-site	T	0427	15.3103	\$1,032.01	\$206.41	\$206.41	
49440	Place gastrostomy tube perc	T	0141	8.7462	\$589.55	\$143.38	\$117.91	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
49652	Lap vent/abd hernia repair	CH	T	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
49653	Lap vent/abd hernia repair comp	CH	T	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
49654	Lap inc hernia repair	CH	T	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
49655	Lap inc hernia repair comp	CH	T	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
49656	Lap inc hernia repair recur	CH	T	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
49657	Lap inc hern recur comp	CH	T	0132	72.9582	\$4,917.82	\$1,239.22	\$983.57
49659	Laparoscopic procedure, hernia repair		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
49800	Repair of abdominal wall		C					
49804	Omental flap, intra-abdom		C					
49805	Omental flap, extra-abdom		C					
49806	Free omental flap, microvasc		C					
49899	Abdomen surgery procedure		C					
50010	Exploration of kidney		C	0153	27.1855	\$1,832.47	\$376.05	\$366.50
50020	Renal abscess, open drain		T	0162	25.5223	\$1,720.36		\$344.08
50021	Renal abscess, percut drain		T	0037	15.5003	\$1,044.81	\$228.76	\$208.97
50040	Drainage of kidney		C					
50045	Exploration of kidney		C					
50060	Removal of kidney stone		C					
50065	Incision of kidney		C					
50070	Inclusion of kidney		C					
50075	Removal of kidney stone		C					
50080	Removal of kidney stone		T	0429	46.68	\$3,146.51		\$629.31
50081	Removal of kidney stone		T	0429	46.68	\$3,146.51		\$629.31
50100	Revised kidney blood vessels		C					
50120	Exploration of kidney		C					
50125	Explore and drain kidney		C					
50130	Removal of kidney stone		C					
50135	Exploration of kidney		C					
50200	Renal biopsy perq		C	0685	9.6666	\$651.59		\$130.32
50205	Renal biopsy open		C					
50220	Remove kidney, open		C					
50225	Remove kidney open, complex		C					
50230	Remove kidney open, radical		C					
50234	Removal of kidney & ureter		C					
50236	Removal of kidney & ureter		C					
50240	Partial removal of kidney		C					
50250	Cryosablate renal mass open		C					
50280	Removal of kidney lesion		C					
50290	Removal of kidney lesion		C					
50300	Remove cadaver donor kidney		C					
50320	Remove kidney, living donor		C					
50323	Prep cadaver renal allograft		C					
50325	Prep donor renal graft		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
50327	Prep renal graft/venous		C					
50328	Prep renal graft/arterial		C					
50329	Prep renal graft/ureteral		C					
50340	Removal of kidney		C					
50360	Transplantation of kidney		C					
50365	Transplantation of kidney		C					
50370	Remove transplanted kidney		C					
50380	Reimplantation of kidney		C					
50382	Change ureter stent, percut		T	0162	25.5223	\$1,720.36		\$344.08
50384	Remove ureter stent, percut		T	0161	17.0344	\$1,148.22		\$229.65
50385	Change stent via transureth	CH	T	0162	25.5223	\$1,720.36		\$344.08
50386	Remove stent via transureth		T	0160	7.1342	\$480.89		\$96.18
50387	Change ext/mt ureter stent		T	0427	15.3103	\$1,032.01		\$206.41
50389	Remove renal tube w/fluoro		T	0160	7.1342	\$480.89		\$96.18
50390	Drainage of kidney lesion		T	0685	9.6666	\$651.59		\$130.32
50391	Insitl rx agrit into renal tub		T	0126	1.0958	\$73.86	\$16.21	\$14.78
50392	Insert kidney drain		T	0161	17.0344	\$1,148.22		\$229.65
50393	Insert ureteral tube		T	0162	25.5223	\$1,720.36		\$344.08
50394	Injection for kidney x-ray		N					
50395	Create passage to kidney	CH	T	0162	25.5223	\$1,720.36		\$344.08
50396	Measure kidney pressure		T	0164	2.0194	\$136.12		\$27.23
50398	Change kidney tube		T	0427	15.3103	\$1,032.01		\$206.41
50400	Revision of kidney/ureter		C					
50405	Repair of kidney wound		C					
50500	Repair of kidney wound		C					
50520	Close kidney-skin fistula		C					
50525	Repair renal-abdomen fistula		C					
50526	Repair renal-abdomen fistula		C					
50540	Revision of horseshoe kidney		C					
50541	Laparo ablate renal cyst		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
50542	Laparo ablate renal mass		T	0174	109.9237	\$7,409.52	\$2,069.30	\$1,481.91
50543	Laparo partial nephrectomy		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
50544	Laparoscopy, pyeloplasty		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
50545	Laparo radical nephrectomy		C					
50546	Laparoscopic nephrectomy		C					
50547	Laparo removal donor kidney		C					
50548	Laparo remove w/ureter		C					
50549	Laparoscopy proc, renal		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
50551	Kidney endoscopy		T	0160	7.1342	\$480.89		\$96.18
50553	Kidney endoscopy		T	0162	25.5223	\$1,720.36		\$344.08
50555	Kidney endoscopy & biopsy		T	0160	7.1342	\$480.89		\$96.18
50557	Kidney endoscopy & treatment		T	0162	25.5223	\$1,720.36		\$344.08
50561	Kidney endoscopy & treatment		T	0162	25.5223	\$1,720.36		\$344.08

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
50860	Transplant ureter to skin		C					
50900	Repair of ureter		C					
50920	Closure ureter/skin fistula		C					
50930	Closure ureter/bowel fistula		C					
50940	Release of ureter		C					
50945	Laparoscopic ureterolithotomy		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
50947	Laparo new ureter/bladder		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
50948	Laparo new ureter/bladder		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
50949	Laparoscopic proc. ureter		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
50951	Endoscopy of ureter		T	0160	7.1342	\$480.89		\$96.18
50953	Endoscopy of ureter		T	0160	7.1342	\$480.89		\$96.18
50955	Ureter endoscopy & biopsy		T	0162	25.5223	\$1,720.36		\$344.08
50957	Ureter endoscopy & treatment		T	0162	25.5223	\$1,720.36		\$344.08
50961	Ureter endoscopy & treatment		T	0162	25.5223	\$1,720.36		\$344.08
50970	Ureter endoscopy & treatment		T	0160	7.1342	\$480.89		\$96.18
50972	Ureter endoscopy & catheter		T	0160	7.1342	\$480.89		\$96.18
50974	Ureter endoscopy & biopsy		T	0161	17.0344	\$1,148.22		\$229.65
50976	Ureter endoscopy & treatment		T	0161	17.0344	\$1,148.22		\$229.65
50980	Ureter endoscopy & treatment		T	0162	25.5223	\$1,720.36		\$344.08
51020	Incise & treat bladder		T	0162	25.5223	\$1,720.36		\$344.08
51030	Incise & treat bladder		T	0162	25.5223	\$1,720.36		\$344.08
51040	Incise & drain bladder		T	0162	25.5223	\$1,720.36		\$344.08
51045	Incise bladder/drain ureter		T	0160	7.1342	\$480.89		\$96.18
51050	Removal of bladder stone		T	0162	25.5223	\$1,720.36		\$344.08
51060	Removal of ureter stone	CH	T	0163	36.2009	\$2,440.16		\$488.04
51065	Remove ureter calculus		T	0162	25.5223	\$1,720.36		\$344.08
51080	Drainage of bladder abscess		T	0008	19.4063	\$1,308.10		\$261.62
51100	Drain bladder by needle		T	0164	2.0194	\$136.12		\$27.23
51101	Drain bladder by trocar/cath		T	0126	1.0958	\$73.86	\$16.21	\$14.78
51102	Drain bl w/cath insertion		T	0165	20.0243	\$1,349.76		\$269.96
51500	Removal of bladder cyst		T	0154	31.9682	\$2,154.65	\$464.85	\$430.97
51525	Removal of bladder lesion		T	0162	25.5223	\$1,720.36		\$344.08
51530	Removal of bladder lesion		C					
51535	Repair of ureter lesion		T	0162	25.5223	\$1,720.36		\$344.08
51550	Partial removal of bladder		C					
51555	Partial removal of bladder		C					
51565	Revise bladder & ureter(s)		C					
51570	Removal of bladder		C					
51575	Removal of bladder & nodes		C					
51580	Remove bladder/revise tract		C					
51585	Removal of bladder & nodes		C					
51590	Remove bladder/revise tract		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
50562	Renal scope w/tumor resect		T	0160	7.1342	\$480.89		\$96.18
50570	Kidney endoscopy		T	0160	7.1342	\$480.89		\$96.18
50572	Kidney endoscopy		T	0160	7.1342	\$480.89		\$96.18
50574	Kidney endoscopy & biopsy		T	0163	36.2009	\$2,440.16		\$488.04
50575	Kidney endoscopy		T	0161	17.0344	\$1,148.22		\$229.65
50576	Kidney endoscopy & treatment		T	0161	17.0344	\$1,148.22		\$229.65
50580	Kidney endoscopy & treatment		T	0169	41.3626	\$2,788.09	\$997.74	\$229.65
50590	Fragmenting of kidney stone		T	0423	51.3618	\$3,462.09	\$692.42	\$557.62
50592	Perc rfr ablate renal tumor		T	0423	51.3618	\$3,462.09	\$692.42	\$557.62
50593	Perc crvo ablate renal tum		T	0423	51.3618	\$3,462.09	\$692.42	\$557.62
50600	Exploration of ureter		C					
50605	Insert ureteral support		C					
50610	Removal of ureter stone		C					
50620	Removal of ureter stone		C					
50630	Removal of ureter stone		C					
50650	Removal of ureter		C					
50660	Removal of ureter		C					
50684	Injection for ureter x-ray		N					
50686	Measure ureter pressure		T	0126	1.0958	\$73.86	\$16.21	\$14.78
50688	Change of ureter tube/stent		T	0427	15.3103	\$1,032.01	\$206.41	\$206.41
50690	Injection for ureter x-ray		N					
50700	Revision of ureter		C					
50715	Release of ureter		C					
50722	Release of ureter		C					
50725	Release/revise ureter		C					
50727	Revise ureter		T	0165	20.0243	\$1,349.76		\$269.96
50728	Revise ureter		C					
50740	Fusion of ureter & kidney		C					
50750	Fusion of ureter & kidney		C					
50760	Fusion of ureter & kidney		C					
50770	Fusion of ureters		C					
50780	Splicing of ureters		C					
50782	Reimplant ureter in bladder		C					
50783	Reimplant ureter in bladder		C					
50785	Reimplant ureter in bladder		C					
50800	Implant ureter in bowel		C					
50810	Fusion of ureter & bowel		C					
50815	Urine shunt to intestine		C					
50820	Construct bowel bladder		C					
50825	Construct bowel bladder		C					
50830	Revise urine flow		C					
50840	Replace ureter by bowel		C					
50845	Appendico-vesicostomy		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
51992	Laparoscopic operation		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
51999	Laparoscopic proc. bla		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
52000	Cystoscopy		T	0160	7.1342	\$480.89		\$96.18
52005	Cystoscopy, removal of clots		T	0161	17.0344	\$1,148.22		\$229.65
52010	Cystoscopy, ureter catheter	CH	T	0162	25.5223	\$1,720.36		\$344.08
52015	Cystoscopy and biopsy		T	0162	25.5223	\$1,720.36		\$344.08
52020	Cystoscopy & duct catheter		T	0160	7.1342	\$480.89		\$96.18
52025	Cystoscopy w/biopsy(s)	CH	T	0162	25.5223	\$1,720.36		\$344.08
52214	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52219	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52224	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52229	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52234	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52239	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52244	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52249	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52254	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52259	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52264	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52269	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52274	Cystoscopy and treatment		T	0161	17.0344	\$1,148.22		\$229.65
52279	Cystoscopy and treatment		T	0160	7.1342	\$480.89		\$96.18
52284	Cystoscopy & revise urethra		T	0161	17.0344	\$1,148.22		\$229.65
52289	Cystoscopy & revise urethra		T	0162	25.5223	\$1,720.36		\$344.08
52294	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52299	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52304	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52309	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52314	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52319	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52324	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52329	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52334	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52339	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52344	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52349	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52354	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52359	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52364	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52369	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52374	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52379	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52384	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52389	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52394	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52399	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52404	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52409	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52414	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52419	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52424	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52429	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52434	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52439	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52444	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08
52449	Cystoscopy and treatment		T	0162	25.5223	\$1,720.36		\$344.08

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
51595	Remove bladder/revise tract		C					
51596	Remove bladder/create pouch		C					
51597	Removal of pelvic structures		C					
51600	Injection for bladder x-ray		N					
51605	Preparation for bladder x-ray		N					
51610	Injection for bladder x-ray		N					
51700	Irrigation of bladder		T	0164	2.0194	\$136.12		\$27.23
51701	Insert bladder catheter		X	0340	0.6693	\$45.11		\$9.03
51702	Insert temp bladder cath		X	0340	0.6693	\$45.11		\$9.03
51703	Insert bladder cath. complex		T	0126	1.0958	\$73.86	\$16.21	\$14.78
51705	Change of bladder tube		T	0164	2.0194	\$136.12		\$27.23
51710	Change of bladder tube	CH	T	0121	6.3742	\$429.66		\$85.94
51715	Endoscopic injection/implant		T	0168	31.377	\$2,115.00		\$423.00
51720	Treatment of bladder lesion		T	0156	3.0214	\$203.66		\$40.74
51725	Simple cystometrogram		T	0156	3.0214	\$203.66		\$40.74
51726	Complex cystometrogram		T	0156	3.0214	\$203.66		\$40.74
51727	Cystometrogram w/up	NI	T	0156	3.0214	\$203.66		\$40.74
51728	Cystometrogram w/p	NI	T	0156	3.0214	\$203.66		\$40.74
51729	Cystometrogram w/rp&up	NI	T	0156	3.0214	\$203.66		\$40.74
51736	Urine flow measurement		T	0126	1.0958	\$73.86	\$16.21	\$14.78
51741	Electro-uroflowmetry, first		T	0126	1.0958	\$73.86	\$16.21	\$14.78
51772	Urethra pressure profile	CH	D					
51784	Anal/urinary muscle study		T	0126	1.0958	\$73.86	\$16.21	\$14.78
51785	Anal/urinary muscle study		T	0164	2.0194	\$136.12		\$27.23
51792	Urinary reflex study		T	0126	1.0958	\$73.86	\$16.21	\$14.78
51795	Urine voiding pressure study	CH	D					
51797	Intraabdominal pressure test		T	0164	2.0194	\$136.12		\$27.23
51798	Us urine capacity measure		X	0340	0.6693	\$45.11		\$9.03
51800	Revision of bladder/urethra		C					
51820	Revision of urinary tract		C					
51840	Attach bladder/urethra		C					
51841	Attach bladder/urethra		C					
51845	Repair bladder neck		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
51860	Repair of bladder wound		T	0162	25.5223	\$1,720.36		\$344.08
51865	Repair of bladder wound		C					
51900	Repair bladder/vagina lesion		C					\$344.08
51920	Close bladder-urethra fistula		C					
51925	Hysterectomy/bladder repair		C					
51940	Correction of bladder defect		C					
51960	Revision of bladder & bowel		C					
51980	Construct bladder opening		C					
51990	Laparoscopic suspension		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
53430	Reconstruct urethra/bladder	T	0168	0168	31.377	\$2,115.00		\$423.00
53431	Reconstruct urethra/bladder	T	0168	0168	31.377	\$2,115.00		\$423.00
53440	Male sling procedure	S	0385	0385	98.0867	\$6,611.63		\$1,322.33
53442	Remove/revise male sling	T	0168	0168	31.377	\$2,115.00		\$423.00
53444	Insert tandem cuff	S	0385	0385	98.0867	\$6,611.63		\$1,322.33
53445	Insert uro/ves neck sphincter	S	0386	0386	164.32	\$11,076.15		\$2,212.23
53446	Remove uro sphincter	T	0168	0168	31.377	\$2,115.00		\$423.00
53447	Remove/replace ur sphincter	S	0386	0386	164.32	\$11,076.15		\$2,212.23
53448	Remove/replace ur sphincter comp	C						
53449	Repair ur sphincter	T	0168	0168	31.377	\$2,115.00		\$423.00
53450	Revision of urethra	T	0168	0168	31.377	\$2,115.00		\$423.00
53460	Revision of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53500	Urethry, transvag w/ scope	T	0168	0168	31.377	\$2,115.00		\$423.00
53502	Repair of urethra injury	T	0166	0166	20.3374	\$1,370.86		\$274.18
53505	Repair of urethra injury	T	0168	0168	31.377	\$2,115.00		\$423.00
53515	Repair of urethra injury	T	0166	0166	20.3374	\$1,370.86		\$274.18
53520	Repair of urethra defect	T	0168	0168	31.377	\$2,115.00		\$423.00
53600	Dilate urethra stricture	T	0156	0156	3.0214	\$203.66		\$40.74
53601	Dilate urethra stricture	T	0126	0126	1.0958	\$73.86	\$16.21	\$14.78
53605	Dilate urethra stricture	T	0161	0161	17.0344	\$1,148.22		\$229.65
53620	Dilate urethra stricture	T	0165	0165	20.0243	\$1,349.76		\$269.96
53621	Dilate urethra stricture	T	0164	0164	2.0194	\$136.12		\$27.23
53660	Dilation of urethra	T	0126	0126	1.0958	\$73.86	\$16.21	\$14.78
53665	Dilation of urethra	T	0126	0126	1.0958	\$73.86	\$16.21	\$14.78
53850	Prostatic microwave thermox	T	0429	0429	46.68	\$3,146.51		\$629.31
53852	Prostatic rf thermox	T	0429	0429	46.68	\$3,146.51		\$629.31
53855	Insert prost urethral stent	NI						
53899	Urology surgery, procedure	T	0126	0126	1.0958	\$73.86	\$16.21	\$14.78
54000	Sitting of prepuc	T	0166	0166	20.3374	\$1,370.86		\$274.18
54001	Sitting of prepuc	T	0166	0166	20.3374	\$1,370.86		\$274.18
54015	Drain penis lesion	T	0008	0008	19.4063	\$1,308.10		\$261.62
54050	Destruction, penis lesion(s)	T	0013	0013	0.8789	\$59.24		\$11.85
54055	Destruction, penis lesion(s)	T	0017	0017	21.2653	\$1,433.41		\$286.69
54056	Cryosurgery, penis lesion(s)	T	0013	0013	0.8789	\$59.24		\$11.85
54057	Laser surg, penis lesion(s)	T	0017	0017	21.2653	\$1,433.41		\$286.69
54060	Excision of penis lesion(s)	T	0017	0017	21.2653	\$1,433.41		\$286.69
54065	Destruction, penis lesion(s)	T	0017	0017	21.2653	\$1,433.41		\$286.69
54100	Biopsy of penis	T	0021	0021	17.4975	\$1,179.44		\$235.89
54105	Biopsy of penis	T	0022	0022	23.388	\$1,576.49		\$315.30
54110	Treatment of penis lesion	T	0181	0181	34.8837	\$2,351.37		\$470.28
54111	Treat penis lesion, graft	T	0181	0181	34.8837	\$2,351.37		\$470.28

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
52346	Cystouretero w/renal strict	T	0162	0162	25.5223	\$1,720.36		\$344.08
52351	Cystouretero & or pyeloscope	T	0162	0162	25.5223	\$1,720.36		\$344.08
52352	Cystouretero w/stone remove	T	0162	0162	25.5223	\$1,720.36		\$344.08
52353	Cystouretero w/ultrasony	T	0163	0163	36.2009	\$2,440.16		\$488.04
52354	Cystouretero w/biopsy	T	0162	0162	25.5223	\$1,720.36		\$344.08
52355	Cystouretero w/excise tumor	T	0162	0162	25.5223	\$1,720.36		\$344.08
52400	Cystouretero w/congen repr	T	0162	0162	25.5223	\$1,720.36		\$344.08
52402	Cystourethro cut ejac duct	T	0162	0162	25.5223	\$1,720.36		\$344.08
52450	Incision of prostate	T	0162	0162	25.5223	\$1,720.36		\$344.08
52500	Revision of bladder neck	T	0162	0162	25.5223	\$1,720.36		\$344.08
52601	Prostatectomy (TURP)	T	0163	0163	36.2009	\$2,440.16		\$488.04
52630	Remove prostate regrowth	T	0163	0163	36.2009	\$2,440.16		\$488.04
52640	Relieve bladder contracture	T	0162	0162	25.5223	\$1,720.36		\$344.08
52647	Laser surgery of prostate	T	0429	0429	46.68	\$3,146.51		\$629.31
52648	Laser surgery of prostate	T	0429	0429	46.68	\$3,146.51		\$629.31
52649	Prostate laser enucleation	T	0429	0429	46.68	\$3,146.51		\$629.31
52700	Drainage of prostate abscess	T	0162	0162	25.5223	\$1,720.36		\$344.08
53000	Incision of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53010	Incision of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53020	Incision of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53025	Incision of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53040	Drainage of urethra abscess	T	0166	0166	20.3374	\$1,370.86		\$274.18
53060	Drainage of urethra abscess	T	0166	0166	20.3374	\$1,370.86		\$274.18
53085	Drainage of urinary leakage	T	0166	0166	20.3374	\$1,370.86		\$274.18
53095	Drainage of urinary leakage	T	0166	0166	20.3374	\$1,370.86		\$274.18
53200	Biopsy of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53210	Removal of urethra	T	0168	0168	31.377	\$2,115.00		\$423.00
53215	Removal of urethra	T	0166	0166	20.3374	\$1,370.86		\$274.18
53220	Treatment of urethra lesion	T	0168	0168	31.377	\$2,115.00		\$423.00
53230	Removal of urethra lesion	T	0168	0168	31.377	\$2,115.00		\$423.00
53235	Removal of urethra lesion	T	0166	0166	20.3374	\$1,370.86		\$274.18
53240	Surgery for urethra pouch	T	0168	0168	31.377	\$2,115.00		\$423.00
53250	Removal of urethra gland	T	0166	0166	20.3374	\$1,370.86		\$274.18
53260	Treatment of urethra lesion	T	0166	0166	20.3374	\$1,370.86		\$274.18
53265	Treatment of urethra lesion	T	0166	0166	20.3374	\$1,370.86		\$274.18
53270	Repair of urethra gland	T	0166	0166	20.3374	\$1,370.86		\$274.18
53275	Repair of urethra defect	T	0166	0166	20.3374	\$1,370.86		\$274.18
53400	Revise urethra, stage 1	T	0168	0168	31.377	\$2,115.00		\$423.00
53405	Revise urethra, stage 2	T	0168	0168	31.377	\$2,115.00		\$423.00
53410	Reconstruction of urethra	C						
53415	Reconstruction of urethra	T	0168	0168	31.377	\$2,115.00		\$423.00
53420	Reconstruct urethra, stage 1	T	0168	0168	31.377	\$2,115.00		\$423.00
53425	Reconstruct urethra, stage 2	T	0168	0168	31.377	\$2,115.00		\$423.00

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
54406	Remove multi-comp penis pros	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54408	Repair multi-comp penis pros	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54410	Remove/replace penis prosth	S	0386	164.32	\$11,076.15		\$2,215.23	
54411	Remove/replace penis pros, comp	C						
54415	Remove self-cont'd penis pros	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54416	Remove self-cont'd penis pros	S	0386	164.32	\$11,076.15		\$2,215.23	
54417	Remove/replace penis pros, comp	C						
54420	Revision of penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54430	Revision of penis	C						
54435	Revision of penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54440	Repair of penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54450	Preputial stretching	T	0156	3,021.4	\$203.66		\$40.74	
54500	Biopsy of testis	T	0037	15,003	\$1,044.81	\$228.76	\$313.90	
54505	Biopsy of testis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54512	Excise lesion testis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54520	Removal of testis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54522	Orchiectomy, partial	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54530	Removal of testis	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
54535	Extensive testis surgery	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54550	Exploration for testis	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
54560	Exploration for testis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54600	Reduce testis torsion	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54620	Suspension of testis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54640	Suspension of testis	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
54650	Orchiopexy (Fowler-Stephens)	C						
54660	Revision of testis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54670	Repair testis injury	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54680	Relocation of testis(es)	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54690	Laparoscopy, orchiectomy	T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46	
54692	Laparoscopy, orchiopexy	T	0132	72.9882	\$4,917.82	\$1,239.22	\$983.57	
54699	Laparoscopy proc. testis	T	0130	38.054	\$2,565.07	\$659.53	\$513.02	
54700	Drainage of scrotum	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54800	Biopsy of epididymis	T	0004	4.5991	\$310.01		\$62.01	
54830	Remove epididymis lesion	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54840	Remove epididymis lesion	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54860	Removal of epididymis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54861	Removal of epididymis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54865	Explore epididymis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54900	Fusion of spermatic ducts	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
55000	Drainage of hydrocele	T	0004	4.5991	\$310.01		\$62.01	
55040	Insert self-cont'd prosthesis	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	
55041	Remove multi-comp penis pros	T	0154	31.9682	\$2,154.85	\$464.85	\$430.97	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
54112	Treat penis lesion, graft	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54115	Treatment of penis lesion	T	0008	19,4063	\$1,308.10	\$261.62	\$261.62	
54120	Partial removal of penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54125	Removal of penis	C						
54130	Remove penis & nodes	C						
54135	Remove penis & nodes	C						
54150	Circumcision w/regional block	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54160	Circumcision, neonate	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54161	Circum 28 days or older	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54162	Lysis penil circumic lesion	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54163	Repair of circumcision	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54164	Frenulotomy of penis	T	0183	23.2838	\$1,569.47	\$621.82	\$470.28	
54200	Treatment of penis lesion	T	0164	2,0194	\$136.12	\$27.23	\$27.23	
54205	Treatment of penis lesion	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54220	Treatment of penis lesion	T	0164	2,0194	\$136.12	\$27.23	\$27.23	
54230	Prepare penis study	N						
54231	Dynamic cavernosometry	T	0165	20,0243	\$1,349.76	\$269.96	\$269.96	
54235	Penile injection	T	0164	2,0194	\$136.12	\$27.23	\$27.23	
54240	Penis study	T	0126	1,0958	\$73.86	\$14.78	\$14.78	
54250	Penis study	T	0164	2,0194	\$136.12	\$27.23	\$27.23	
54300	Revision of penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54304	Revision of penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54308	Reconstruction of urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54312	Reconstruction of urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54316	Reconstruction of urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54322	Reconstruction of urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54326	Reconstruction of urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54328	Reconstruct urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54332	Reconstruct urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54336	Reconstruct urethra	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54340	Secondary urethral surgery	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54344	Secondary urethral surgery	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54348	Secondary urethral surgery	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54352	Reconstruct urethral/penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54360	Penis plastic surgery	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54380	Repair penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54385	Repair penis	T	0181	34.8837	\$2,351.37	\$621.82	\$470.28	
54390	Repair penis and bladder	C						
54400	Insert semi-rigid prosthesis	S	0385	98.0867	\$6,611.63		\$1,322.33	
54401	Insert self-cont'd prosthesis	S	0386	164.32	\$11,076.15		\$2,215.23	
54405	Insert multi-comp penis pros	S	0386	164.32	\$11,076.15		\$2,215.23	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
55875	Transpen needle place, pros		Q3	0163	36.2009	\$2,440.16		\$488.04
55876	Place rt device/marker, pros		X	0310	13.7576	\$927.34	\$325.27	\$185.47
55899	Genital surgery procedure		T	0126	1.0958	\$73.86	\$16.21	\$14.78
55920	Place needles pelvic for r		T	0153	27.1855	\$1,832.47	\$376.05	\$366.50
55970	Sex transformation, M to F		E					
55980	Sex transformation, F to M		E					
56405	I & D of vulva/perineum	CH	T	0188	1.5277	\$102.98		\$20.60
56420	Drainage of gland abscess		T	0188	1.5277	\$102.98		\$20.60
56440	Surgery for vulva lesion		T	0193	20.0452	\$1,351.17		\$270.24
56441	Lysis of labial lesion(s)		T	0193	20.0452	\$1,351.17		\$270.24
56442	Hymenotomy		T	0193	20.0452	\$1,351.17		\$270.24
56501	Destroy, vulva lesions, sim		T	0017	21.2653	\$1,433.41	\$286.69	\$286.69
56515	Destroy vulva lesion/s compl		T	0017	21.2653	\$1,433.41		\$286.69
56605	Biopsy of vulva/perineum		T	0189	3.4085	\$229.75		\$45.95
56606	Biopsy of vulva/perineum		T	0188	1.5277	\$102.98		\$20.60
56620	Partial removal of vulva		T	0193	20.0452	\$1,351.17		\$270.24
56625	Complete removal of vulva		T	0193	20.0452	\$1,351.17		\$270.24
56630	Extensive vulva surgery		C					
56631	Extensive vulva surgery		C					
56632	Extensive vulva surgery		C					
56633	Extensive vulva surgery		C					
56634	Extensive vulva surgery		C					
56637	Extensive vulva surgery		C					
56640	Extensive vulva surgery		C					
56700	Partial removal of hymen		T	0193	20.0452	\$1,351.17		\$270.24
56740	Remove vagina gland lesion		T	0193	20.0452	\$1,351.17		\$270.24
56800	Repair of vagina		T	0193	20.0452	\$1,351.17		\$270.24
56805	Repair clitoris		T	0193	20.0452	\$1,351.17		\$270.24
56810	Repair of perineum		T	0188	1.5277	\$102.98		\$20.60
56820	Exam of vulva w/scope		T	0188	1.5277	\$102.98		\$20.60
56821	Exam/biopsy of vulva w/scope		T	0193	20.0452	\$1,351.17		\$270.24
57000	Exploration of vagina		T	0193	20.0452	\$1,351.17		\$270.24
57010	Drainage of pelvic abscess		T	0193	20.0452	\$1,351.17		\$270.24
57020	Drainage of pelvic fluid		T	0192	6.7949	\$458.02		\$91.61
57022	I & d vaginal hematoma, pp		T	0007	12.6217	\$850.78		\$170.16
57023	I & d vag hematoma, non-ob		T	0008	19.4063	\$1,308.10		\$261.62
57061	Destroy vag lesions, simple		T	0193	20.0452	\$1,351.17		\$270.24
57065	Destroy vag lesions, complex		T	0193	20.0452	\$1,351.17		\$270.24
57100	Biopsy of vagina		T	0192	6.7949	\$458.02		\$91.61
57105	Biopsy of vagina		T	0193	20.0452	\$1,351.17		\$270.24
57106	Remove vagina wall, partial		T	0193	20.0452	\$1,351.17		\$270.24
57107	Remove vagina tissue, part		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57109	Vaginectomy partial w/nodes		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
55060	Repair of hydrocele		T	0183	23.2838	\$1,569.47		\$313.90
55100	Drainage of scrotum abscess		T	0007	12.6217	\$850.78		\$170.16
55110	Explore scrotum		T	0183	23.2838	\$1,569.47		\$313.90
55120	Removal of scrotum lesion		T	0183	23.2838	\$1,569.47		\$313.90
55150	Removal of scrotum		T	0183	23.2838	\$1,569.47		\$313.90
55175	Revision of scrotum		T	0183	23.2838	\$1,569.47		\$313.90
55180	Revision of scrotum		T	0183	23.2838	\$1,569.47		\$313.90
55200	Incision of sperm duct		T	0183	23.2838	\$1,569.47		\$313.90
55250	Removal of sperm duct(s)		T	0183	23.2838	\$1,569.47		\$313.90
55300	Prepare, sperm duct x-ray		N					
55400	Repair of sperm duct		T	0183	23.2838	\$1,569.47		\$313.90
55450	Ligation of sperm duct		T	0183	23.2838	\$1,569.47		\$313.90
55500	Removal of hydrocele		T	0183	23.2838	\$1,569.47		\$313.90
55520	Removal of sperm cord lesion		T	0183	23.2838	\$1,569.47		\$313.90
55530	Revise spermatic cord veins		T	0183	23.2838	\$1,569.47		\$313.90
55535	Revise spermatic cord veins		T	0154	31.9682	\$2,154.85	\$464.85	\$430.97
55540	Revise hernia & sperm veins		T	0154	31.9682	\$2,154.85	\$464.85	\$430.97
55550	Laparoscopic spermatic vein		T	0131	46.84	\$3,157.30	\$1,001.89	\$631.46
55559	Laparoscopic, spermatic cord		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
55605	Incise sperm duct pouch		T	0183	23.2838	\$1,569.47		\$313.90
55650	Remove sperm duct pouch		C					
55680	Remove sperm pouch lesion		C					
55700	Biopsy of prostate		T	0183	23.2838	\$1,569.47		\$313.90
55705	Biopsy of prostate		T	0184	12.4313	\$837.94		\$167.59
55706	Prostate saturation sampling		T	0184	12.4313	\$837.94		\$167.59
55720	Drainage of prostate abscess		T	0162	25.5223	\$1,720.36		\$344.08
55725	Drainage of prostate abscess		T	0162	25.5223	\$1,720.36		\$344.08
55801	Removal of prostate		C					
55810	Extensive prostate surgery		C					
55812	Extensive prostate surgery		C					
55815	Extensive prostate surgery		C					
55821	Removal of prostate		C					
55831	Removal of prostate		C					
55840	Extensive prostate surgery		C					
55842	Extensive prostate surgery		C					
55845	Extensive prostate surgery		C					
55860	Surgical exposure, prostate		T	0165	20.0243	\$1,349.76		\$269.96
55865	Extensive prostate surgery		C					
55866	Extensive prostate surgery		C					
55870	Laparoscopic prostatectomy		C					
55873	Cryoblation prostate		T	0674	113.8626	\$7,675.02		\$1,535.01

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
57400	Dilation of vagina		T	0193	20.0452	\$1,351.17		\$270.24
57410	Pelvic examination		T	0193	20.0452	\$1,351.17		\$270.24
57415	Remove vaginal foreign body		T	0193	20.0452	\$1,351.17		\$270.24
57420	Exam of vagina w/scope		T	0189	3.4085	\$229.75		\$45.95
57421	Exam/biopsy of vag w/scope		T	0189	3.4085	\$229.75		\$45.95
57423	Repair paravag defect, lap		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57425	Laparoscopy, surg, colpopoxy		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
57426	Revise prosth vag graft lap	NI	T	0193	20.0452	\$1,351.17		\$270.24
57452	Exam of cervix w/scope	CH	T	0188	1.5277	\$102.98		\$20.60
57454	Bx/curett of cervix w/scope		T	0189	3.4085	\$229.75		\$45.95
57455	Biopsy of cervix w/scope		T	0189	3.4085	\$229.75		\$45.95
57456	Endocerv curettage w/scope		T	0189	3.4085	\$229.75		\$45.95
57460	Bx of cervix w/scope, keep		T	0193	20.0452	\$1,351.17		\$270.24
57461	Conz of cervix w/scope, keep		T	0193	20.0452	\$1,351.17		\$270.24
57500	Biopsy of cervix		T	0192	6.7949	\$458.02		\$91.61
57505	Endocervical curettage		T	0192	6.7949	\$458.02		\$91.61
57510	Cauterization of cervix		T	0193	20.0452	\$1,351.17		\$270.24
57511	Cryocautery of cervix		T	0188	1.5277	\$102.98		\$20.60
57513	Laser surgery of cervix		T	0193	20.0452	\$1,351.17		\$270.24
57520	Conization of cervix		T	0193	20.0452	\$1,351.17		\$270.24
57522	Conization of cervix		T	0193	20.0452	\$1,351.17		\$270.24
57530	Removal of cervix		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57531	Removal of cervix, radical		C					
57540	Removal of residual cervix		C					
57545	Remove cervix/repair pelvis		C					
57550	Removal of residual cervix		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57555	Remove cervix/repair vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57556	Remove cervix, repair bowel		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57558	D&C of cervical stump		T	0193	20.0452	\$1,351.17		\$270.24
57700	Revision of cervix		T	0193	20.0452	\$1,351.17		\$270.24
57720	Revision of cervix		T	0193	20.0452	\$1,351.17		\$270.24
58100	Dilation of cervical canal		T	0193	20.0452	\$1,351.17		\$270.24
58110	Biopsy of uterus lining		T	0188	1.5277	\$102.98		\$20.60
58120	Bx done w/colposcopy add-on		N					
58140	Dilation and curettage		T	0193	20.0452	\$1,351.17		\$270.24
58145	Myomectomy abdom method		C					
58146	Myomectomy vag method		C					
58152	Myomectomy abdom complex		C					
58150	Total hysterectomy		C					
58152	Total hysterectomy		C					
58180	Partial hysterectomy		C					
58200	Extensive hysterectomy		C					
58210	Extensive hysterectomy		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
57110	Remove vagina wall, complete		C					
57111	Remove vagina tissue, compl		C					
57112	Vaginectomy w/nodes, compl		C					
57120	Closure of vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57130	Remove vagina lesion		T	0193	20.0452	\$1,351.17		\$270.24
57135	Remove vagina lesion		T	0193	20.0452	\$1,351.17		\$270.24
57150	Treat vagina infection		T	0188	1.5277	\$102.98		\$20.60
57155	Insert uter tandem/s/ovoids		T	0192	6.7949	\$458.02	\$91.61	\$20.60
57160	Insert pessary/other device		T	0188	1.5277	\$102.98	\$20.60	\$20.60
57170	Fitting of diaphragm/cap		T	0191	0.132	\$6.90	\$2.09	\$1.78
57180	Treat vaginal bleeding		T	0188	1.5277	\$102.98		\$20.60
57200	Repair of vagina		T	0193	20.0452	\$1,351.17		\$270.24
57210	Repair vagina/perineum		T	0193	20.0452	\$1,351.17		\$270.24
57220	Revision of urethra		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57230	Repair of urethral lesion		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57240	Repair bladder & vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57250	Repair rectum & vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57260	Repair of vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57265	Extensive repair of vagina		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57267	Insert mesh/pelvic fir addon		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57268	Repair of bowel bulge		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57270	Repair of bowel pouch		C					
57280	Suspension of vagina		C					
57282	Colpopoxy, extraperitoneal		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57283	Colpopoxy, intraperitoneal		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57284	Repair paravag defect, open		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57285	Repair paravag defect, vag	CH	T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57287	Revise/remove sling repair		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57288	Repair bladder defect		T	0202	44.9894	\$3,032.56	\$981.50	\$606.52
57289	Repair bladder & vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57291	Construction of vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57292	Construct vagina with graft		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57295	Revise vag graft via vagina		T	0193	20.0452	\$1,351.17		\$270.24
57296	Revise vag graft, open abd		C					
57300	Repair rectum-vagina fistula		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57305	Repair rectum-vagina fistula		C					
57307	Fistula repair & colostomy		C					
57308	Fistula repair, transperine		C					
57310	Repair urethrovaginal lesion		C					
57311	Repair urethrovaginal lesion		C					
57320	Repair bladder-vagina lesion		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57330	Repair bladder-vagina lesion		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44
57335	Repair vagina		T	0195	35.2666	\$2,377.18	\$483.80	\$475.44

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
58560	Hysteroscopy, resect septum	T	0387		37.2199	\$2,508.84	\$655.55	\$501.77
58561	Hysteroscopy, remove myoma	T	0387		37.2199	\$2,508.84	\$655.55	\$501.77
58562	Hysteroscopy, remove fb	T	0190		22.5926	\$1,522.88	\$424.28	\$304.58
58563	Hysteroscopy, ablation	T	0387		37.2199	\$2,508.84	\$655.55	\$501.77
58565	Hysteroscopy, sterilization	T	0202		44.9894	\$3,032.56	\$981.50	\$606.52
58570	Th, uterus 250 g or less	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58571	Th w/lo 250 g or less	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58572	Th, uterus over 250 g	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58573	Th w/lo uterus over 250 g	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58578	Laparo proc, uterus	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
58579	Hysteroscope procedure	T	0190		22.5926	\$1,522.88	\$424.28	\$304.58
58600	Division of fallopian tube	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58605	Ligate oviduct(s) add-on	C						
58611	Occlude fallopian tube(s)	T	0195		20.0452	\$1,351.17		\$270.24
58660	Laparoscopy, lysis	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58661	Laparoscopy, remove adnexa	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58662	Laparoscopy, excise lesions	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58670	Laparoscopy, tubal cautery	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58671	Laparoscopy, tubal block	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58672	Laparoscopy, fimbrioplasty	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58673	Laparoscopy, salpingostomy	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58700	Laparo proc, oviduct-ovary	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
58720	Removal of ovary/tube(s)	C						
58740	Adhesiolysis tube, ovary	C						
58750	Repair oviduct	C						
58752	Revise ovarian tube(s)	C						
58760	Fimbrioplasty	C						
58770	Creates new tubal opening	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58800	Drainage of ovarian cyst(s)	T	0193		20.0452	\$1,351.17		\$270.24
58805	Drainage of ovarian cyst(s)	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58820	Drain ovary abscess, open	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58822	Drain ovary abscess, percut	C						
58823	Drain pelvic abscess, percut	T	0193		20.0452	\$1,351.17		\$270.24
58825	Transposition, ovary(s)	C						
58900	Biopsy of ovary(s)	T	0193		20.0452	\$1,351.17		\$270.24
58920	Partial removal of ovary(s)	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58925	Removal of ovarian cyst(s)	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58940	Removal of ovary(s)	C						
58943	Removal of ovary(s)	C						
58950	Resect ovarian malignancy	C						
58951	Resect ovarian malignancy	C						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
58240	Removal of pelvic contents	C						
58260	Vaginal hysterectomy	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58262	Vag hyster including tv	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58263	Vag hyster w/uro & vag repair	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58267	Vag hyster w/urinary repair	C						
58270	Vag hyster w/enterocole repair	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58275	Hysterectomy/revise vagina	C						
58280	Hysterectomy/revise vagina	C						
58285	Extensive hysterectomy	C						
58290	Vag hyster complex	T	0202		44.9894	\$3,032.56	\$981.50	\$606.52
58291	Vag hyster incl tv, complex	T	0202		44.9894	\$3,032.56	\$981.50	\$606.52
58292	Vag hyster tv & repair, compl	T	0202		44.9894	\$3,032.56	\$981.50	\$606.52
58293	Vag hyster w/uro repair, compl	C						
58294	Vag hyster w/enterocole, compl	T	0202		44.9894	\$3,032.56	\$981.50	\$606.52
58300	Insert intrauterine device	E						
58301	Remove intrauterine device	T	0188		1.5277	\$102.98		\$20.60
58321	Artificial insemination	T	0189		3.4085	\$229.75	\$45.95	\$45.95
58322	Artificial insemination	T	0189		3.4085	\$229.75	\$45.95	\$45.95
58323	Sperm washing	T	0189		3.4085	\$229.75	\$45.95	\$45.95
58340	Catheter for hystero-graphy	N						
58345	Reopen fallopian tube	T	0193		20.0452	\$1,351.17		\$270.24
58346	Insert heyman uteri capsule	T	0193		20.0452	\$1,351.17		\$270.24
58350	Reopen fallopian tube	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58353	Endometr ablate, thermal	T	0195		35.2666	\$2,377.18	\$483.80	\$475.44
58356	Endometrial cryoablation	T	0202		44.9894	\$3,032.56	\$981.50	\$606.52
58400	Suspension of uterus	C						
58410	Suspension of uterus	C						
58520	Repair of ruptured uterus	C						
58540	Revision of uterus	C						
58541	Lsh, uterus 250 g or less	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
58542	Lsh w/lo ut 250 g or less	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
58543	Lsh uterus above 250 g	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
58544	Lsh w/lo uterus above 250 g	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
58545	Laparoscopic myomectomy	T	0130		38.054	\$2,565.07	\$659.53	\$513.02
58546	Laparo-myomectomy, complex	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58548	Lap radical hyster	C						
58550	Laparo-assst vag hysterectomy	T	0132		72.9582	\$4,917.82	\$1,239.22	\$983.57
58552	Laparo-vag hyster incl tv	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58553	Laparo-vag hyster, complex	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58554	Laparo-vag hyster w/lo, compl	T	0131		46.84	\$3,157.30	\$1,001.89	\$631.46
58555	Hysterectomy, dx, sep proc	T	0190		22.5926	\$1,522.88	\$424.28	\$304.58
58558	Hysterectomy, biopsy	T	0190		22.5926	\$1,522.88	\$424.28	\$304.58
58559	Hysterectomy, lysis	T	0190		22.5926	\$1,522.88	\$424.28	\$304.58

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
59414	Deliver placenta	T	0193	20.0452	\$1,351.17		\$270.24	
59425	Antepartum care only	B						
59426	Antepartum care only	B						
59430	Care after delivery	B						
59510	Cesarean delivery	B						
59514	Cesarean delivery only	C						
59515	Cesarean delivery	B						
59525	Remove uterus after cesarean	C						
59610	Vbac delivery	B						
59612	Vbac delivery only	T	0193	20.0452	\$1,351.17		\$270.24	
59614	Vbac care after delivery	B						
59618	Attempted vbac delivery	B						
59620	Attempted vbac delivery only	C						
59622	Attempted vbac after care	B						
59812	Treatment of miscarriage	T	0193	20.0452	\$1,351.17		\$270.24	
59820	Care of miscarriage	T	0193	20.0452	\$1,351.17		\$270.24	
59821	Treatment of miscarriage	T	0193	20.0452	\$1,351.17		\$270.24	
59830	Treat uterus infection	C						
59840	Abortion	T	0193	20.0452	\$1,351.17		\$270.24	
59841	Abortion	T	0193	20.0452	\$1,351.17		\$270.24	
59850	Abortion	C						
59851	Abortion	C						
59852	Abortion	C						
59855	Abortion	C						
59856	Abortion	C						
59857	Abortion	C						
59866	Abortion (mfp)	T	0189	3.4085	\$229.75		\$45.95	
59870	Evacuate mole of uterus	T	0193	20.0452	\$1,351.17		\$270.24	
59871	Remove cerclage suture	T	0193	20.0452	\$1,351.17		\$270.24	
59887	Fetal invas px w/us	T	0191	0.132	\$8.90		\$1.78	
59898	Laparoscopic care/deliver	T	0130	38.054	\$2,565.07		\$513.02	
60000	Maternity care procedure	T	0252	7.6196	\$513.61		\$102.73	
60100	Drain thyroid/tongue cyst	T	0004	4.5991	\$310.01		\$62.01	
60200	Biopsy of thyroid	T	0114	48.7561	\$3,286.45		\$657.29	
60210	Remove thyroid lesion	T	0114	48.7561	\$3,286.45		\$657.29	
60212	Partial thyroid excision	T	0114	48.7561	\$3,286.45		\$657.29	
60220	Partial thyroid excision	T	0114	48.7561	\$3,286.45		\$657.29	
60225	Partial removal of thyroid	T	0114	48.7561	\$3,286.45		\$657.29	
60240	Removal of thyroid	T	0114	48.7561	\$3,286.45		\$657.29	
60252	Removal of thyroid	T	0256	42.9827	\$2,897.29		\$579.46	
60254	Extensive thyroid surgery	C						
60260	Repeat thyroid surgery	T	0256	42.9827	\$2,897.29		\$579.46	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
58952	Resect ovarian malignancy	C						
58953	Tah, rad dissect for debulk	C						
58954	Tah rad debulk/lymph remove	C						
58956	Bso, omentectomy w/tah	C						
58957	Resect recurrent gyn mal	C						
58958	Resect recur gyn mal w/lym	C						
58960	Exploration of abdomen	C						
58970	Retrieval of oocyte	T	0189	3.4085	\$229.75		\$45.95	
58974	Transfer of embryo	T	0189	3.4085	\$229.75		\$45.95	
58976	Transfer of embryo	T	0189	3.4085	\$229.75		\$45.95	
58999	Genital surgery procedure	T	0191	0.132	\$8.90		\$1.78	
59000	Amniocentesis, diagnostic	T	0189	3.4085	\$229.75		\$45.95	
59001	Amniocentesis, therapeutic	T	0192	6.7949	\$458.02		\$91.61	
59012	Fetal cord puncture, prenatal	T	0189	3.4085	\$229.75		\$45.95	
59015	Chorion biopsy	T	0189	3.4085	\$229.75		\$45.95	
59020	Fetal contract stress test	T	0188	1.5277	\$102.98		\$20.60	
59025	Fetal non-stress test	T	0188	1.5277	\$102.98		\$20.60	
59030	Fetal scalp blood sample	T	0189	3.4085	\$229.75		\$45.95	
59050	Fetal monitor w/report	M						
59051	Fetal monitor/interpret only	B						
59070	Transabdom amniocent w/us	CH	0188	1.5277	\$102.98		\$20.60	
59072	Umbilical cord occlud w/us	T	0189	3.4085	\$229.75		\$45.95	
59074	Fetal fluid drainage w/us	T	0189	3.4085	\$229.75		\$45.95	
59076	Fetal shunt placement, w/us	T	0189	3.4085	\$229.75		\$45.95	
59100	Remove uterus lesion	T	0195	35.2666	\$2,377.18		\$483.80	
59120	Treat ectopic pregnancy	C						
59121	Treat ectopic pregnancy	C						
59130	Treat ectopic pregnancy	C						
59135	Treat ectopic pregnancy	C						
59136	Treat ectopic pregnancy	C						
59140	Treat ectopic pregnancy	C						
59150	Treat ectopic pregnancy	C						
59151	Treat ectopic pregnancy	T	0131	46.84	\$3,157.30		\$631.46	
59160	D & c after delivery	T	0193	20.0452	\$1,351.17		\$270.24	
59200	Insert cervical dilator	T	0188	1.5277	\$102.98		\$20.60	
59300	Episiotomy or vaginal repair	CH	0193	20.0452	\$1,351.17		\$270.24	
59320	Revision of cervix	T	0193	20.0452	\$1,351.17		\$270.24	
59325	Revision of cervix	C						
59350	Repair of uterus	C						
59400	Obstetrical care	B						
59409	Obstetrical care	T	0193	20.0452	\$1,351.17		\$270.24	
59410	Obstetrical care	B						
59412	Antepartum manipulation	T	0193	20.0452	\$1,351.17		\$270.24	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
61314	Open skull for drainage	C	C					
61315	Open skull for drainage	C	C					
61316	Implt cran bone flap to abdo	C	C					
61320	Open skull for drainage	C	C					
61321	Open skull for drainage	C	C					
61322	Decompressive craniotomy	C	C					
61323	Decompressive lobectomy	C	C					
61330	Decompress eye socket	T	T	0256	42.9827	\$2,897.29		\$579.46
61332	Explore/biopsy eye socket	C	C					
61333	Explore orbit/remove lesion	C	C					
61334	Explore orbit/remove object	T	T	0256	42.9827	\$2,897.29		\$579.46
61340	Subtemporal decompression	C	C					
61343	Incise skull (press relief)	C	C					
61345	Relieve cranial pressure	C	C					
61440	Incise skull for surgery	C	C					
61450	Incise skull for surgery	C	C					
61458	Incise skull for brain wound	C	C					
61470	Incise skull for surgery	C	C					
61480	Incise skull for surgery	C	C					
61490	Incise skull for surgery	C	C					
61500	Removal of skull lesion	C	C					
61501	Remove infected skull bone	C	C					
61510	Removal of brain lesion	C	C					
61512	Remove brain lining lesion	C	C					
61514	Removal of brain abscess	C	C					
61516	Removal of brain lesion	C	C					
61517	Implt brain chemotx add-on	C	C					
61518	Removal of brain lesion	C	C					
61519	Remove brain lining lesion	C	C					
61520	Removal of brain lesion	C	C					
61521	Removal of brain abscess	C	C					
61522	Removal of brain abscess	C	C					
61524	Removal of brain lesion	C	C					
61526	Removal of brain lesion	C	C					
61531	Implant brain electrodes	C	C					
61533	Implant brain electrodes	C	C					
61534	Removal of brain lesion	C	C					
61535	Remove brain electrodes	C	C					
61536	Removal of brain lesion	C	C					
61537	Removal of brain tissue	C	C					
61538	Removal of brain tissue	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
60270	Removal of thyroid	C	C					
60271	Removal of thyroid	T	T	0256	42.9827	\$2,897.29		\$579.46
60280	Remove thyroid duct lesion	T	T	0114	48.7561	\$3,286.45		\$657.29
60281	Remove thyroid duct lesion	T	T	0114	48.7561	\$3,286.45		\$657.29
60300	Aspirating thyroid cyst	T	T	0004	4.5991	\$310.01		\$62.01
60500	Explore parathyroid glands	T	T	0256	42.9827	\$2,897.29		\$579.46
60502	Re-explore parathyroids	T	T	0256	42.9827	\$2,897.29		\$579.46
60505	Explore parathyroid glands	C	C					
60512	Autotransplant parathyroid	T	T	0022	23.388	\$1,576.49		\$315.30
60520	Removal of thymus gland	T	T	0256	42.9827	\$2,897.29		\$579.46
60521	Removal of thymus gland	C	C					
60522	Removal of thymus gland	C	C					
60540	Explore adrenal gland	C	C					
60545	Explore adrenal gland	C	C					
60600	Remove carotid body lesion	C	C					
60605	Remove carotid body lesion	C	C					
60650	Laparoscopy adrenalectomy	C	C					
60659	Laparo proc, endocrine	T	T	0130	38.054	\$2,565.07	\$659.53	\$513.02
60699	Endocrine surgery procedure	T	T	0114	48.7561	\$3,286.45		\$657.29
61000	Remove cranial cavity fluid	T	T	0207	7.2002	\$485.34		\$97.07
61001	Remove cranial cavity fluid	T	T	0207	7.2002	\$485.34		\$97.07
61020	Remove brain cavity fluid	T	T	0207	7.2002	\$485.34		\$97.07
61026	Injection into brain canal	T	T	0207	7.2002	\$485.34		\$97.07
61050	Remove brain canal fluid	T	T	0207	7.2002	\$485.34		\$97.07
61055	Injection into brain canal	T	T	0207	7.2002	\$485.34		\$97.07
61070	Brain canal shunt procedure	T	T	0121	6.3742	\$429.66		\$85.94
61105	Twist drill hole	C	C					
61107	Drill skull for implantation	C	C					
61108	Drill skull for drainage	C	C					
61120	Burr hole for puncture	C	C					
61140	Pierce skull for biopsy	C	C					
61150	Pierce skull for drainage	C	C					
61151	Pierce skull for drainage	C	C					
61154	Pierce skull & remove clot	C	C					
61156	Pierce skull, implant device	C	C					
61210	Pierce skull, implant device	C	C					
61215	Insert brain-fluid device	T	T	0224	41.0288	\$2,765.59		\$553.12
61250	Pierce skull & explore	C	C					
61253	Pierce skull & explore	C	C					
61304	Open skull for exploration	C	C					
61305	Open skull for exploration	C	C					
61312	Open skull for drainage	C	C					
61313	Open skull for drainage	C	C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
61609	Transect artery, sinus	C	C					
61610	Transect artery, sinus	C	C					
61611	Transect artery, sinus	C	C					
61612	Transect artery, sinus	C	C					
61613	Remove aneurysm, sinus	C	C					
61615	Resect/excise lesion, skull	C	C					
61616	Resect/excise lesion, skull	C	C					
61618	Repair dura	C	C					
61619	Repair dura	C	C					
61623	Endovasc temporary vessel occl	T	0082		93.3244	\$6,290.62		\$1,258.13
61624	Transect occlusion, cns	C	C					
61626	Transect occlusion, non-cns	T	0082		93.3244	\$6,290.62		\$1,258.13
61630	Intracranial angioplasty	C	C					
61635	Intracran angioplasty w/steint	C	C					
61640	Dilate ic vasospasm, init	E	E					
61641	Dilate ic vasospasm add-on	E	E					
61642	Dilate ic vasospasm add-on	E	E					
61680	Intracranial vessel surgery	C	C					
61682	Intracranial vessel surgery	C	C					
61684	Intracranial vessel surgery	C	C					
61686	Intracranial vessel surgery	C	C					
61690	Intracranial vessel surgery	C	C					
61692	Intracranial vessel surgery	C	C					
61697	Brain aneurysm repr, complx	C	C					
61698	Brain aneurysm repr, complx	C	C					
61700	Brain aneurysm repr, simple	C	C					
61702	Inner skull vessel surgery	C	C					
61703	Clamp neck artery	C	C					
61705	Revise circulation to head	C	C					
61708	Revise circulation to head	C	C					
61710	Revise circulation to head	C	C					
61711	Fusion of skull arteries	C	C					
61720	Incise skull/brain surgery	T	0221		37.2806	\$2,512.94		\$502.59
61735	Incise skull/brain surgery	C	C					
61750	Incise skull/brain biopsy	C	C					
61751	Brain biopsy w/ct/mr guide	C	C					
61760	Implant brain electrodes	C	C					
61770	Incise skull for treatment	T	0221		37.2806	\$2,512.94		\$502.59
61790	Treat trigeminal nerve	T	0220		18.72	\$1,261.84		\$252.37
61791	Treat trigeminal tract	T	0203		13.2439	\$892.72	\$225.98	\$178.55
61795	Brain surgery using computer	N	N					
61796	Srs, cranial lesion simple	B	B					
61797	Srs, cran les simple, add	B	B					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
61539	Removal of brain tissue	C	C					
61540	Removal of brain tissue	C	C					
61541	Incision of brain tissue	C	C					
61542	Removal of brain tissue	C	C					
61543	Removal of brain tissue	C	C					
61544	Remove & treat brain lesion	C	C					
61545	Excision of brain tumor	C	C					
61546	Removal of pituitary gland	C	C					
61548	Removal of pituitary gland	C	C					
61550	Release of skull seams	C	C					
61552	Release of skull seams	C	C					
61556	Incise skull/sutures	C	C					
61557	Incise skull/sutures	C	C					
61558	Excision of skull/sutures	C	C					
61559	Excision of skull/sutures	C	C					
61563	Excision of skull tumor	C	C					
61564	Excision of skull tumor	C	C					
61566	Removal of brain tissue	C	C					
61567	Incision of brain tissue	C	C					
61570	Remove foreign body, brain	C	C					
61571	Incise skull for brain wound	C	C					
61575	Skull base/brainstem surgery	C	C					
61576	Skull base/brainstem surgery	C	C					
61580	Craniofacial approach, skull	C	C					
61581	Craniofacial approach, skull	C	C					
61582	Craniofacial approach, skull	C	C					
61583	Craniofacial approach, skull	C	C					
61584	Orbitocranial approach/skull	C	C					
61585	Orbitocranial approach/skull	C	C					
61586	Resect nasopharynx, skull	C	C					
61590	Infratemporal approach/skull	C	C					
61591	Infratemporal approach/skull	C	C					
61592	Orbitocranial approach/skull	C	C					
61595	Transmastoid approach/skull	C	C					
61596	Transmastoid approach/skull	C	C					
61597	Transcondylar approach/skull	C	C					
61598	Transpetrosal approach/skull	C	C					
61600	Resect/excise cranial lesion	C	C					
61601	Resect/excise cranial lesion	C	C					
61605	Resect/excise cranial lesion	C	C					
61606	Resect/excise cranial lesion	C	C					
61607	Resect/excise cranial lesion	C	C					
61608	Resect/excise cranial lesion	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
62201	Brain cavity shunt w/scope		C					
62202	Establish brain cavity shunt		C					
62223	Establish brain cavity shunt		C					
62225	Replace/irrigate catheter		T	0427	15.3103	\$1,032.01		\$206.41
62230	Replace/revise brain shunt		T	0224	41.0288	\$2,765.59		\$553.12
62252	Cst shunt reprogram		S	0691	2.5406	\$171.25	\$50.49	\$34.25
62256	Remove brain cavity shunt		C					
62258	Replace brain cavity shunt		C					
62263	Epidural lysis mult sessions		T	0207	7.2002	\$485.34		\$97.07
62264	Epidural lysis on single day		T	0203	13.2439	\$892.72	\$225.98	\$178.55
62267	Interfacial perq aspir. dk		T	0004	4.5991	\$310.01		\$62.01
62268	Drain spinal cord cyst		T	0207	7.2002	\$485.34		\$97.07
62269	Needle biopsy, spinal cord		T	0685	9.6866	\$651.59		\$130.32
62270	Spinal fluid tap, diagnostic		T	0206	3.7221	\$250.89	\$51.76	\$50.18
62272	Drain cerebro spinal fluid		T	0206	3.7221	\$250.89	\$51.76	\$50.18
62273	Inject epidural patch		T	0206	3.7221	\$250.89	\$51.76	\$50.18
62280	Treat spinal cord lesion		T	0207	7.2002	\$485.34		\$97.07
62281	Treat spinal cord lesion		T	0207	7.2002	\$485.34		\$97.07
62282	Treat spinal canal lesion		T	0207	7.2002	\$485.34		\$97.07
62284	Injection for myelogram		N					
62287	Percutaneous disectomy		T	0221	37.2806	\$2,512.94		\$502.59
62290	Inject for spine disk x-ray		N					
62291	Inject for spine disk x-ray		N					
62292	Injection into disk lesion		T	0207	7.2002	\$485.34		\$97.07
62294	Injection into spinal artery		T	0207	7.2002	\$485.34		\$97.07
62310	Inject spine c/t		T	0207	7.2002	\$485.34		\$97.07
62311	Inject spine l/s (cd)		T	0207	7.2002	\$485.34		\$97.07
62318	Inject spine w/cath, c/t		T	0207	7.2002	\$485.34		\$97.07
62319	Inject spine w/cath l/s (cd)		T	0224	41.0288	\$2,765.59		\$553.12
62350	Implant spinal canal cath		T	0208	49.2256	\$3,318.10		\$663.62
62351	Implant spinal canal cath		T	0203	13.2439	\$892.72	\$225.98	\$178.55
62360	Remove spinal canal catheter		T	0224	41.0288	\$2,765.59		\$553.12
62361	Insert spine infusion device		T	0227	198.6572	\$13,390.69		\$2,678.14
62362	Implant spine infusion pump		T	0227	198.6572	\$13,390.69		\$2,678.14
62365	Remove spine infusion pump		T	0221	37.2806	\$2,512.94		\$502.59
62367	Analyze spine infusion pump	CH	S	0691	2.5406	\$171.25	\$50.49	\$34.25
62368	Analyze spine infusion pump		S	0691	2.5406	\$171.25	\$50.49	\$34.25
63001	Removal of spinal lamina		T	0208	49.2256	\$3,318.10		\$663.62
63003	Removal of spinal lamina		T	0208	49.2256	\$3,318.10		\$663.62
63005	Removal of spinal lamina		T	0208	49.2256	\$3,318.10		\$663.62
63011	Removal of spinal lamina		T	0208	49.2256	\$3,318.10		\$663.62
63012	Removal of spinal lamina		T	0208	49.2256	\$3,318.10		\$663.62

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
61798	Srs. cranial lesion complex		B					
61799	Srs. cran les complex, addl		B					
61800	Apply srs headframe add-on		B					
61850	Implant neuroelectrodes		C					
61860	Implant neuroelectrodes		C					
61863	Implant neuroelectrode		C					
61864	Implant neuroelectrode, addl		C					
61867	Implant neuroelectrode		C					
61868	Implant neuroelectrode, addl		C					
61870	Implant neuroelectrodes		C					
61875	Implant neuroelectrodes		C					
61880	Revise/remove neuroelectrode		T	0687	19.6381	\$1,323.73	\$397.37	\$264.75
61885	Inst/redo neurostim 1 array		S	0039	206.1011	\$13,892.45		\$2,778.49
61886	Implant neurostim arrays		S	0315	274.7397	\$18,519.10		\$3,703.82
61888	Revise/remove neuroreceiver		T	0688	28.6636	\$1,932.10	\$770.83	\$386.42
62005	Treat skull fracture		T	0254	24.9637	\$1,682.70		\$336.54
62010	Treatment of head injury		C					
62100	Repair brain fluid leakage		C					
62115	Reduction of skull defect		C					
62116	Reduction of skull defect		C					
62117	Reduction of skull defect		C					
62120	Repair skull cavity lesion		C					
62121	Incise skull repair		C					
62140	Repair of skull defect		C					
62141	Repair of skull defect		C					
62142	Remove skull plate/flap		C					
62143	Replace skull plate/flap		C					
62145	Repair of skull & brain		C					
62146	Repair of skull with graft		C					
62147	Repair of skull with graft		C					
62148	Reir bone flap to fix skull		C					
62160	Neuroendoscopy add-on		N					
62161	Dissect brain w/scope		C					
62162	Neuroendoscopy w/scope		C					
62163	Neuroendoscopy w/ib removal		C					
62164	Remove brain tumor w/scope		C					
62165	Remove pituit tumor w/scope		C					
62180	Establish brain cavity shunt		C					
62190	Establish brain cavity shunt		C					
62192	Establish brain cavity shunt		C					
62194	Replace/irrigate catheter		T	0207	7.2002	\$485.34		\$97.07
62200	Establish brain cavity shunt		C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
63191	Incise spinal column/nerves	C	C					
63194	Incise spinal column & cord	C	C					
63195	Incise spinal column & cord	C	C					
63196	Incise spinal column & cord	C	C					
63197	Incise spinal column & cord	C	C					
63198	Incise spinal column & cord	C	C					
63199	Incise spinal column & cord	C	C					
63200	Release of spinal cord	C	C					
63201	Revise spinal cord vessels	C	C					
63251	Revise spinal cord vessels	C	C					
63252	Revise spinal cord vessels	C	C					
63265	Excise intraspinal lesion	C	C					
63266	Excise intraspinal lesion	C	C					
63267	Excise intraspinal lesion	C	C					
63268	Excise intraspinal lesion	C	C					
63270	Excise intraspinal lesion	C	C					
63271	Excise intraspinal lesion	C	C					
63272	Excise intraspinal lesion	C	C					
63273	Excise intraspinal lesion	C	C					
63275	Biopsy/excise spinal tumor	C	C					
63276	Biopsy/excise spinal tumor	C	C					
63277	Biopsy/excise spinal tumor	C	C					
63278	Biopsy/excise spinal tumor	C	C					
63280	Biopsy/excise spinal tumor	C	C					
63281	Biopsy/excise spinal tumor	C	C					
63282	Biopsy/excise spinal tumor	C	C					
63283	Biopsy/excise spinal tumor	C	C					
63285	Biopsy/excise spinal tumor	C	C					
63286	Biopsy/excise spinal tumor	C	C					
63287	Biopsy/excise spinal tumor	C	C					
63290	Biopsy/excise spinal tumor	C	C					
63295	Repair of laminectomy defect	C	C					
63300	Removal of vertebral body	C	C					
63301	Removal of vertebral body	C	C					
63302	Removal of vertebral body	C	C					
63303	Removal of vertebral body	C	C					
63304	Removal of vertebral body	C	C					
63305	Removal of vertebral body	C	C					
63306	Removal of vertebral body	C	C					
63307	Removal of vertebral body	C	C					
63308	Remove vertebral body add-on	C	C					
63600	Remove spinal cord lesion	T	T	0220	18.72	\$1,261.84		\$252.37
63610	Stimulation of spinal cord	T	T	0220	18.72	\$1,261.84		\$252.37

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
63015	Removal of spinal lamina	T	T	0208	49.2256	\$3,318.10		\$663.62
63016	Removal of spinal lamina	T	T	0208	49.2256	\$3,318.10		\$663.62
63017	Removal of spinal lamina	T	T	0208	49.2256	\$3,318.10		\$663.62
63020	Neck spine disk surgery	T	T	0208	49.2256	\$3,318.10		\$663.62
63030	Low back disk surgery	T	T	0208	49.2256	\$3,318.10		\$663.62
63035	Spinal disk surgery add-on	T	T	0208	49.2256	\$3,318.10		\$663.62
63040	Laminotomy, single cervical	T	T	0208	49.2256	\$3,318.10		\$663.62
63042	Laminotomy, single lumbar	T	T	0208	49.2256	\$3,318.10		\$663.62
63043	Laminotomy, addl cervical	C	C					
63044	Laminotomy, addl lumbar	C	C					
63045	Removal of spinal lamina	T	T	0208	49.2256	\$3,318.10		\$663.62
63046	Removal of spinal lamina	T	T	0208	49.2256	\$3,318.10		\$663.62
63047	Removal of spinal lamina	T	T	0208	49.2256	\$3,318.10		\$663.62
63048	Remove spinal lamina add-on	T	T	0208	49.2256	\$3,318.10		\$663.62
63050	Cervical laminoplasty	C	C					
63051	C-laminoplasty w/graft/plate	C	C					
63055	Decompress spinal cord	T	T	0208	49.2256	\$3,318.10		\$663.62
63056	Decompress spinal cord	T	T	0208	49.2256	\$3,318.10		\$663.62
63057	Decompress spine cord add-on	T	T	0208	49.2256	\$3,318.10		\$663.62
63064	Decompress spinal cord	T	T	0208	49.2256	\$3,318.10		\$663.62
63066	Decompress spine cord add-on	T	T	0208	49.2256	\$3,318.10		\$663.62
63075	Neck spine disk surgery	T	T	0208	49.2256	\$3,318.10		\$663.62
63076	Neck spine disk surgery	CH	T	0208	49.2256	\$3,318.10		\$663.62
63077	Spine disk surgery, thorax	C	C					
63078	Spine disk surgery, thorax	C	C					
63081	Removal of vertebral body	C	C					
63082	Remove vertebral body add-on	C	C					
63085	Removal of vertebral body	C	C					
63086	Remove vertebral body add-on	C	C					
63087	Removal of vertebral body	C	C					
63088	Remove vertebral body add-on	C	C					
63090	Removal of vertebral body	C	C					
63091	Remove vertebral body add-on	C	C					
63101	Removal of vertebral body	C	C					
63102	Removal of vertebral body	C	C					
63103	Remove vertebral body add-on	C	C					
63170	Incise spinal cord tract(s)	C	C					
63172	Drainage of spinal cyst	C	C					
63173	Drainage of spinal cyst	C	C					
63180	Revise spinal cord ligaments	C	C					
63182	Revise spinal cord ligaments	C	C					
63185	Incise spinal column/nerves	C	C					
63190	Incise spinal column/nerves	C	C					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
64449	N block inj, lumbar plexus	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64450	N block, other, peripheral	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64455	N block inj, plantar digit	T	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64470	Inj paravertebral c/t	CH	D					
64472	Inj paravertebral c/t add-on	CH	D					
64475	Inj paravertebral l/s	CH	D					
64476	Inj paravertebral l/s add-on	CH	D					
64479	Inj foramen epidural c/t	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64480	Inj foramen epidural add-on	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64483	Inj foramen epidural l/s	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64484	Inj foramen epidural add-on	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64490	Inj paravert f jnt c/t 1 lev	NI	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64491	Inj paravert f jnt c/t 2 lev	NI	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64492	Inj paravert f jnt c/t 3 lev	NI	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64493	Inj paravert f jnt l/s 1 lev	NI	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64494	Inj paravert f jnt l/s 2 lev	NI	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64495	Inj paravert f jnt l/s 3 lev	NI	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64505	N block, sphenolathine gangli	T	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64508	N block, carotid sinus s/p	T	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64510	N block, stellate ganglion	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64517	N block inj, hypogloss plus	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64520	N block, lumbar/thoracic	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64530	N block inj, celliac peltus	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64553	Apply neurostimulator	A						
64553	Implant neuroelectrodes	S	0040	0040	65.7095	\$4,429.21	\$885.85	\$885.85
64555	Implant neuroelectrodes	S	0040	0040	65.7095	\$4,429.21	\$885.85	\$885.85
64561	Implant neuroelectrodes	S	0040	0040	65.7095	\$4,429.21	\$885.85	\$885.85
64565	Implant neuroelectrodes	S	0040	0040	65.7095	\$4,429.21	\$885.85	\$885.85
64573	Implant neuroelectrodes	S	0225	0225	157.939	\$10,646.04	\$2,129.21	\$2,129.21
64575	Implant neuroelectrodes	S	0061	0061	86.5171	\$5,831.77	\$1,166.36	\$1,166.36
64580	Implant neuroelectrodes	S	0061	0061	86.5171	\$5,831.77	\$1,166.36	\$1,166.36
64581	Implant neuroelectrodes	S	0061	0061	86.5171	\$5,831.77	\$1,166.36	\$1,166.36
64585	Reviser/remove neuroelectrode	T	0687	0687	19.6381	\$1,323.73	\$397.37	\$397.37
64590	Instr/redo pn/gastr stimul	S	0039	0039	206.1011	\$13,892.45	\$2,778.49	\$2,778.49
64595	Reviser/mv pn/gastr stimul	T	0688	0688	28.6636	\$1,932.10	\$770.83	\$770.83
64600	Injection treatment of nerve	CH	0203	0203	13.2439	\$892.72	\$225.98	\$178.55
64605	Injection treatment of nerve	CH	0220	0220	18.72	\$1,261.84	\$252.37	\$252.37
64610	Injection treatment of nerve	CH	0220	0220	18.72	\$1,261.84	\$252.37	\$252.37
64612	Destroy nerve, face muscle	T	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64613	Destroy nerve, neck muscle	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64614	Destroy nerve, extrem musc	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
63615	Remove lesion of spinal cord	T	0220	0220	18.72	\$1,261.84	\$252.37	\$252.37
63620	Srs, spinal lesion	B						
63621	Srs, spinal lesion, addl	B						
63650	Implant neuroelectrodes	S	0040	0040	65.7095	\$4,429.21	\$885.85	\$885.85
63655	Implant neuroelectrodes	S	0061	0061	86.5171	\$5,831.77	\$1,166.36	\$1,166.36
63660	Reviser/remove neuroelectrode	CH	D					
63661	Remove spine eltrd perq aray	NI	0687	0687	19.6381	\$1,323.73	\$397.37	\$397.37
63662	Remove spine eltrd plate	NI	0687	0687	19.6381	\$1,323.73	\$397.37	\$397.37
63663	Reviser spine eltrd perq aray	NI	0687	0687	19.6381	\$1,323.73	\$397.37	\$397.37
63664	Reviser spine eltrd plate	NI	0687	0687	19.6381	\$1,323.73	\$397.37	\$397.37
63685	Instr/redo spine n generator	CH	S	0039	206.1011	\$13,892.45	\$2,778.49	\$2,778.49
63688	Reviser/remove neuroreover	T	0688	0688	28.6636	\$1,932.10	\$770.83	\$770.83
63700	Repair of spinal herniation	C						
63702	Repair of spinal herniation	C						
63704	Repair of spinal herniation	C						
63706	Repair of spinal herniation	C						
63707	Repair spinal fluid leakage	C						
63709	Repair spinal fluid leakage	C						
63710	Graft repair of spine defect	C						
63740	Install spinal shunt	C						
63741	Install spinal shunt	C	0224	0224	41.0288	\$2,765.59	\$553.12	\$553.12
63744	Revision of spinal shunt	T	0224	0224	41.0288	\$2,765.59	\$553.12	\$553.12
63746	Removal of spinal shunt	T	0203	0203	13.2439	\$892.72	\$225.98	\$178.55
64400	N block inj, trigeminal	T	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64402	N block inj, facial	T	0204	0204	2.5558	\$172.28	\$40.13	\$34.46
64405	N block inj, occipital	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64408	N block inj, vagus	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64410	N block inj, phrenic	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64412	N block inj, spinal accessor	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64413	N block inj, cervical plexus	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64415	N block inj, brachial plexus	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64416	N block cont infuse, b plex	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64417	N block inj, axillary	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64418	N block inj, suprascapular	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64420	N block inj, intercost, sng	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64421	N block inj, intercost, mlt	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64425	N block inj, ilio-ino/hyppooj	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64430	N block inj, pudendal	T	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64435	N block inj, paracervical	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64445	N block inj, sciatic, sng	CH	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64446	N blk inj, sciatic, cont inf	CH	0207	0207	7.2002	\$485.34	\$51.76	\$97.07
64447	N block inj fem, single	T	0206	0206	3.7221	\$250.89	\$51.76	\$50.18
64448	N block inj fem, cont inf	CH	0207	0207	7.2002	\$485.34	\$51.76	\$97.07

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
64778	Digit nerve surgery add-on	T		0220	18.72	\$1,261.84		\$252.37
64782	Remove limb nerve lesion	T		0220	18.72	\$1,261.84		\$252.37
64783	Limb nerve surgery add-on	T		0220	18.72	\$1,261.84		\$252.37
64784	Remove nerve lesion	T		0221	37.2806	\$2,512.94		\$502.59
64786	Remove sciatic nerve lesion	T		0220	18.72	\$1,261.84		\$252.37
64788	Remove skin nerve lesion	T		0220	18.72	\$1,261.84		\$252.37
64790	Removal of nerve lesion	T		0221	37.2806	\$2,512.94		\$502.59
64792	Removal of nerve lesion	T		0220	18.72	\$1,261.84		\$252.37
64795	Biopsy of nerve	T		0220	18.72	\$1,261.84		\$252.37
64802	Remove sympathetic nerves	T		0220	18.72	\$1,261.84		\$252.37
64804	Remove sympathetic nerves	T		0220	18.72	\$1,261.84		\$252.37
64809	Remove sympathetic nerves	C						
64818	Remove sympathetic nerves	C						
64820	Remove sympathetic nerves	T		0220	18.72	\$1,261.84		\$252.37
64821	Remove sympathetic nerves	T		0054	28.2376	\$1,903.38		\$380.68
64822	Remove sympathetic nerves	T		0054	28.2376	\$1,903.38		\$380.68
64823	Remove sympathetic nerves	T		0054	28.2376	\$1,903.38		\$380.68
64831	Repair of digit nerve	T		0221	37.2806	\$2,512.94		\$502.59
64832	Repair nerve add-on	T		0221	37.2806	\$2,512.94		\$502.59
64834	Repair of hand or foot nerve	T		0221	37.2806	\$2,512.94		\$502.59
64835	Repair of hand or foot nerve	T		0221	37.2806	\$2,512.94		\$502.59
64837	Repair of hand or foot nerve	T		0221	37.2806	\$2,512.94		\$502.59
64840	Repair of leg nerve	T		0221	37.2806	\$2,512.94		\$502.59
64856	Repair/transpose nerve	T		0221	37.2806	\$2,512.94		\$502.59
64857	Repair arm/leg nerve	T		0221	37.2806	\$2,512.94		\$502.59
64858	Repair sciatic nerve	T		0221	37.2806	\$2,512.94		\$502.59
64861	Repair of arm nerves	T		0221	37.2806	\$2,512.94		\$502.59
64862	Repair of low back nerves	T		0221	37.2806	\$2,512.94		\$502.59
64864	Repair of facial nerve	T		0221	37.2806	\$2,512.94		\$502.59
64865	Repair of facial nerve	T		0221	37.2806	\$2,512.94		\$502.59
64866	Fusion of facial/other nerve	C						
64868	Fusion of facial/other nerve	C						
64870	Fusion of facial/other nerve	T		0221	37.2806	\$2,512.94		\$502.59
64872	Subsequent repair of nerve	T		0221	37.2806	\$2,512.94		\$502.59
64874	Repair & revise nerve add-on	T		0221	37.2806	\$2,512.94		\$502.59
64875	Repair nerve/shorten bone	T		0221	37.2806	\$2,512.94		\$502.59
64885	Nerve graft, head or neck	T		0221	37.2806	\$2,512.94		\$502.59
64886	Nerve graft, head or neck	T		0221	37.2806	\$2,512.94		\$502.59
64890	Nerve graft, hand or foot	T		0221	37.2806	\$2,512.94		\$502.59
64891	Nerve graft, hand or foot	T		0221	37.2806	\$2,512.94		\$502.59

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
64620	Injection treatment of nerve	T		0207	7.2002	\$465.34	\$225.98	\$97.07
64622	Desr paravertebral nerve i/s	T		0203	13.2439	\$692.72		\$178.55
64623	Desr paravertebral n add-on	T		0207	7.2002	\$465.34		\$97.07
64626	Desr paravertebral nerve c/t	T		0204	2.5558	\$172.28	\$40.13	\$34.46
64630	Injection treatment of nerve	T		0207	7.2002	\$465.34		\$97.07
64632	N block inj, common digit	T		0204	2.5558	\$172.28	\$40.13	\$34.46
64640	Injection treatment of nerve	T		0207	7.2002	\$465.34		\$97.07
64650	Chemodenerv ecocrine glands	T		0204	2.5558	\$172.28	\$40.13	\$34.46
64653	Chemodenerv ecocrine glands	T		0204	2.5558	\$172.28	\$40.13	\$34.46
64680	Injection treatment of nerve	CH		0207	7.2002	\$465.34		\$97.07
64681	Injection treatment of nerve	T		0203	13.2439	\$692.72	\$225.98	\$178.55
64702	Revise finger/Toe nerve	T		0220	18.72	\$1,261.84		\$252.37
64704	Revise hand/foot nerve	T		0220	18.72	\$1,261.84		\$252.37
64708	Revise arm/leg nerve	T		0220	18.72	\$1,261.84		\$252.37
64712	Revision of sciatic nerve	T		0220	18.72	\$1,261.84		\$252.37
64713	Revision of arm nerve(s)	T		0220	18.72	\$1,261.84		\$252.37
64714	Revision low back nerve(s)	T		0220	18.72	\$1,261.84		\$252.37
64716	Revision of cranial nerve	T		0220	18.72	\$1,261.84		\$252.37
64718	Revise ulnar nerve at elbow	T		0220	18.72	\$1,261.84		\$252.37
64719	Revise ulnar nerve at wrist	T		0220	18.72	\$1,261.84		\$252.37
64721	Carpal tunnel surgery	T		0220	18.72	\$1,261.84		\$252.37
64722	Relieve pressure on nerve(s)	T		0220	18.72	\$1,261.84		\$252.37
64726	Release foot/toe nerve	T		0220	18.72	\$1,261.84		\$252.37
64727	Internal nerve revision	T		0220	18.72	\$1,261.84		\$252.37
64732	Incision of brow nerve	T		0220	18.72	\$1,261.84		\$252.37
64734	Incision of cheek nerve	T		0220	18.72	\$1,261.84		\$252.37
64736	Incision of chin nerve	T		0220	18.72	\$1,261.84		\$252.37
64738	Incision of jaw nerve	T		0220	18.72	\$1,261.84		\$252.37
64740	Incision of tongue nerve	T		0220	18.72	\$1,261.84		\$252.37
64742	Incision of facial nerve	T		0220	18.72	\$1,261.84		\$252.37
64744	Incise nerve, back of head	T		0220	18.72	\$1,261.84		\$252.37
64746	Incise diaphragm nerve	T		0220	18.72	\$1,261.84		\$252.37
64752	Incision of vagus nerve	C						
64755	Incision of stomach nerves	C						
64760	Incision of vagus nerve	C						
64761	Incision of pelvis nerve	T		0220	18.72	\$1,261.84		\$252.37
64763	Incise hip/ thigh nerve	T		0220	18.72	\$1,261.84		\$252.37
64766	Incise hip/ thigh nerve	T		0221	37.2806	\$2,512.94		\$502.59
64771	Sever cranial nerve	T		0220	18.72	\$1,261.84		\$252.37
64772	Incision of spinal nerve	T		0220	18.72	\$1,261.84		\$252.37
64774	Remove skin nerve lesion	T		0220	18.72	\$1,261.84		\$252.37
64776	Remove digit nerve lesion	T		0220	18.72	\$1,261.84		\$252.37

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
65400	Removal of eye lesion	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65410	Biopsy of cornea	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65420	Removal of eye lesion	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65426	Removal of eye lesion	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65430	Corneal smear	S		0698	0.9553	\$64.39		\$12.88
65435	Curette/treat cornea	T		0239	7.6421	\$515.12		\$103.03
65436	Curette/treat cornea	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65450	Treatment of corneal lesion	S		0231	1.9575	\$131.95		\$26.39
65600	Revision of cornea	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
65710	Corneal transplant	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65730	Corneal transplant	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65750	Corneal transplant	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65755	Corneal transplant	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65766	Corneal trnspl. endothelial	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65757	Prep corneal endo allograft	N						
65760	Revision of cornea	E						
65765	Revision of cornea	E						
65767	Corneal tissue transplant	E						
65770	Revise cornea with implant	T		0293	101.0161	\$6,809.09		\$1,361.82
65771	Radial keratotomy	E						
65772	Correction of astigmatism	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65775	Correction of astigmatism	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65780	Ocular reconst, transplant	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65781	Ocular reconst, transplant	T		0244	37.5388	\$2,530.34	\$803.26	\$506.07
65800	Drainage of eye	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65805	Drainage of eye	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65810	Drainage of eye	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65815	Drainage of eye	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65820	Relieve inner eye pressure	T		0232	4.5074	\$303.83	\$76.12	\$60.77
65850	Incision of eye	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65855	Laser surgery of eye	T		0247	5.3014	\$357.35	\$104.31	\$71.47
65860	Incise inner eye adhesions	T		0247	5.3014	\$357.35	\$104.31	\$71.47
65865	Incise inner eye adhesions	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65870	Incise inner eye adhesions	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65875	Incise inner eye adhesions	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65880	Incise inner eye adhesions	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65900	Remove eye lesion	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65920	Remove implant of eye	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65930	Remove blood clot from eye	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
66020	Injection treatment of eye	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
66030	Injection treatment of eye	T		0232	4.5074	\$303.83	\$76.12	\$60.77
66130	Remove eye lesion	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
64892	Nerve graft, arm or leg	T		0221	37.2806	\$2,512.94		\$502.59
64893	Nerve graft, arm or leg	T		0221	37.2806	\$2,512.94		\$502.59
64895	Nerve graft, hand or foot	T		0221	37.2806	\$2,512.94		\$502.59
64896	Nerve graft, hand or foot	T		0221	37.2806	\$2,512.94		\$502.59
64897	Nerve graft, arm or leg	T		0221	37.2806	\$2,512.94		\$502.59
64898	Nerve graft, arm or leg	T		0221	37.2806	\$2,512.94		\$502.59
64901	Nerve graft add-on	T		0221	37.2806	\$2,512.94		\$502.59
64902	Nerve graft add-on	T		0221	37.2806	\$2,512.94		\$502.59
64905	Nerve pedicle transfer	T		0221	37.2806	\$2,512.94		\$502.59
64907	Nerve pedicle transfer	T		0221	37.2806	\$2,512.94		\$502.59
64910	Nerve repair w/allograft	T		0221	37.2806	\$2,512.94		\$502.59
64911	Neurograft w/vein autograft	T		0221	37.2806	\$2,512.94		\$502.59
64999	Neurovascular system surgery	T		0204	2.5558	\$172.28	\$40.13	\$34.46
65091	Revise eye	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65093	Revise eye with implant	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65101	Removal of eye	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65103	Remove eye/insert implant	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65105	Remove eye/attach implant	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65110	Removal of eye	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65112	Remove eye/revise socket	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65114	Remove eye/revise socket	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65125	Revise ocular implant	T		0241	26.8396	\$1,809.15	\$361.83	\$361.83
65130	Insert ocular implant	T		0241	26.8396	\$1,809.15	\$361.83	\$361.83
65135	Insert ocular implant	T		0241	26.8396	\$1,809.15	\$361.83	\$361.83
65140	Attach ocular implant	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65150	Revise ocular implant	T		0241	26.8396	\$1,809.15	\$361.83	\$361.83
65155	Reinsert ocular implant	T		0242	39.0115	\$2,629.61	\$597.36	\$255.93
65175	Removal of ocular implant	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
65205	Remove foreign body from eye	S		0698	0.9553	\$64.39		\$12.88
65210	Remove foreign body from eye	S		0698	0.9553	\$64.39		\$12.88
65220	Remove foreign body from eye	S		0698	0.9553	\$64.39		\$12.88
65222	Remove foreign body from eye	S		0698	0.9553	\$64.39		\$12.88
65235	Remove foreign body from eye	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
65260	Remove foreign body from eye	T		0235	5.8498	\$394.31		\$78.87
65265	Remove foreign body from eye	T		0237	20.7145	\$1,396.28		\$279.26
65270	Repair of eye wound	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
65272	Repair of eye wound	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65273	Repair of eye wound	C						
65275	Repair of eye wound	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
65280	Repair of eye wound	T		0237	20.7145	\$1,396.28		\$279.26
65285	Repair of eye wound	T		0672	39.9643	\$2,693.83		\$538.77
65286	Repair of eye wound	T		0232	4.5074	\$303.83	\$76.12	\$60.77
65290	Repair of eye socket wound	T		0243	23.4694	\$1,581.98	\$418.00	\$316.40

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
66990	Ophthalmic endoscope add-on		N	0232	4.5074	\$303.83	\$76.12	\$60.77
66999	Eye surgery procedure		T	0232	20.7145	\$1,396.28		\$279.26
67005	Partial removal of eye fluid		T	0672	39.9643	\$2,693.83		\$538.77
67010	Partial removal of eye fluid	CH	T	0672	39.9643	\$2,693.83		\$538.77
67015	Release of eye fluid		T	0672	20.7145	\$1,396.28		\$279.26
67025	Replace eye fluid		T	0672	39.9643	\$2,693.83		\$538.77
67027	Implant eye drug system		T	0672	3.2267	\$217.50		\$43.50
67028	Injection eye drug		T	0238	5.3014	\$357.35	\$104.31	\$71.47
67030	Incise inner eye strands		T	0237	39.9643	\$2,693.83		\$538.77
67031	Laser surgery, eye strands		T	0672	39.9643	\$2,693.83		\$538.77
67036	Removal of inner eye fluid		T	0672	39.9643	\$2,693.83		\$538.77
67039	Laser treatment of retina		T	0672	39.9643	\$2,693.83		\$538.77
67040	Laser treatment of retina		T	0672	39.9643	\$2,693.83		\$538.77
67041	Vit for macular pucker		T	0672	39.9643	\$2,693.83		\$538.77
67042	Vit for macular hole		T	0672	39.9643	\$2,693.83		\$538.77
67043	Vit for membrane dissect		T	0672	39.9643	\$2,693.83		\$538.77
67101	Repair detached retina	CH	T	0237	20.7145	\$1,396.28		\$279.26
67105	Repair detached retina		T	0247	5.3014	\$357.35	\$104.31	\$71.47
67107	Repair detached retina		T	0672	39.9643	\$2,693.83		\$538.77
67108	Repair detached retina		T	0672	39.9643	\$2,693.83		\$538.77
67110	Repair detached retina		T	0237	20.7145	\$1,396.28		\$279.26
67112	Repair detached retina		T	0672	39.9643	\$2,693.83		\$538.77
67113	Repair detached retina		T	0672	39.9643	\$2,693.83		\$538.77
67115	Release encircling material		T	0237	20.7145	\$1,396.28		\$279.26
67120	Remove eye implant material		T	0237	20.7145	\$1,396.28		\$279.26
67121	Remove eye implant material		T	0237	20.7145	\$1,396.28		\$279.26
67141	Treatment of retina		T	0235	5.8498	\$394.31		\$78.87
67145	Treatment of retina		T	0247	5.3014	\$357.35	\$104.31	\$71.47
67208	Treatment of retinal lesion		T	0235	5.8498	\$394.31		\$78.87
67210	Treatment of retinal lesion		T	0247	5.3014	\$357.35	\$104.31	\$71.47
67218	Treatment of retinal lesion		T	0237	20.7145	\$1,396.28		\$279.26
67220	Treatment of retinal lesion		T	0235	5.8498	\$394.31		\$78.87
67221	Ocular photodynamic ther		T	0235	5.8498	\$394.31		\$78.87
67225	Eye photodynamic ther add-on		T	0235	5.8498	\$394.31		\$78.87
67228	Treatment of retinal lesion		T	0237	20.7145	\$1,396.28		\$279.26
67229	Treatment of retinal lesion		T	0247	5.3014	\$357.35	\$104.31	\$71.47
67230	Tr retinal les preterm inf		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67255	Reinforce eye wall		T	0237	20.7145	\$1,396.28		\$279.26
67299	Reinforce/graft eye wall		T	0235	5.8498	\$394.31		\$78.87
67311	Eye surgery procedure		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67312	Revise eye muscle		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67314	Revise two eye muscles		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
66155	Glaucoma surgery		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66160	Glaucoma surgery		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66165	Glaucoma surgery		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66170	Glaucoma surgery		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66172	Incision of eye		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66180	Implant eye shunt		T	0673	41.884	\$2,823.23	\$649.56	\$564.85
66185	Revise eye shunt	CH	T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66220	Repair eye lesion		T	0672	39.9643	\$2,693.83		\$538.77
66225	Repair/graft eye lesion		T	0673	41.884	\$2,823.23	\$649.56	\$564.85
66250	Follow-up surgery of eye		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
66500	Incision of iris		T	0232	4.5074	\$303.83	\$76.12	\$60.77
66505	Incision of iris		T	0232	4.5074	\$303.83	\$76.12	\$60.77
66600	Remove iris and lesion		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66605	Removal of iris		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66625	Removal of iris		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
66630	Removal of iris		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66635	Removal of iris		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66680	Repair iris & ciliary body		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66682	Repair iris & ciliary body		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66700	Destruction, ciliary body		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
66710	Ciliary transscleral therapy		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
66711	Ciliary endoscopic ablation		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
66720	Destruction, ciliary body		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
66740	Destruction, ciliary body		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66761	Revision of iris		T	0247	5.3014	\$357.35	\$104.31	\$71.47
66762	Revision of iris		T	0247	5.3014	\$357.35	\$104.31	\$71.47
66770	Removal of inner eye lesion		T	0247	5.3014	\$357.35	\$104.31	\$71.47
66820	Incision, secondary cataract		T	0232	4.5074	\$303.83	\$76.12	\$60.77
66821	After cataract laser surgery		T	0247	5.3014	\$357.35	\$104.31	\$71.47
66825	Reposition intraocular lens		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
66830	Removal of lens lesion		T	0232	4.5074	\$303.83	\$76.12	\$60.77
66840	Removal of lens material		T	0245	15.8065	\$1,065.59	\$214.11	\$213.12
66850	Removal of lens material		T	0249	30.3293	\$2,044.38	\$518.26	\$408.88
66852	Removal of lens material		T	0249	30.3293	\$2,044.38	\$518.26	\$408.88
66890	Extraction of lens		T	0249	30.3293	\$2,044.38	\$518.26	\$408.88
66940	Extraction of lens		T	0245	15.8065	\$1,065.59	\$214.11	\$213.12
66982	Cataract surgery, complex		T	0246	24.2879	\$1,637.15	\$495.96	\$327.43
66983	Cataract surg w/inf, 1 stage		T	0246	24.2879	\$1,637.15	\$495.96	\$327.43
66984	Cataract surg w/inf, 1 stage		T	0246	24.2879	\$1,637.15	\$495.96	\$327.43
66985	Insert lens prosthesis		T	0246	24.2879	\$1,637.15	\$495.96	\$327.43
66986	Exchange lens prosthesis		T	0246	24.2879	\$1,637.15	\$495.96	\$327.43

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
67850	Treat eyelid lesion		T	0239	7.6421	\$515.12		\$103.03
67850	Closure of eyelid by suture		T	0239	7.6421	\$515.12		\$103.03
67880	Revision of eyelid		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
67882	Revision of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67900	Repair brow defect		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
67901	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67902	Repair eyelid defect		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
67903	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67904	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67906	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67908	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67909	Revise eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67911	Revise eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67912	Correction eyelid w/inplant		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67914	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67915	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67916	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67917	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67921	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67922	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67923	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67924	Repair eyelid defect		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67930	Repair eyelid wound		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67935	Repair eyelid wound		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67938	Remove eyelid foreign body		S	0231	1.9575	\$131.95		\$28.39
67950	Revision of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67961	Revision of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67966	Revision of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67971	Reconstruction of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67973	Reconstruction of eyelid		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
67974	Reconstruction of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67975	Reconstruction of eyelid		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67999	Revision of eyelid		T	0238	3.2267	\$217.50		\$43.50
68020	Incise/drain eyelid lining		T	0238	3.2267	\$217.50		\$43.50
68040	Treatment of eyelid lesions		S	0698	0.9553	\$64.39		\$12.88
68100	Biopsy of eyelid lining		T	0232	4.5074	\$303.83	\$76.12	\$60.77
68110	Remove eyelid lining lesion		T	0699	15.5407	\$1,047.54	\$296.93	\$209.51
68115	Remove eyelid lining lesion		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
68135	Remove eyelid lining lesion		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
68135	Remove eyelid lining lesion		T	0239	7.6421	\$515.12		\$103.03
68200	Treat eyelid by injection		S	0698	0.9553	\$64.39		\$12.88
68320	Revise/graft eyelid lining		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
68325	Revise/graft eyelid lining		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
67316	Revise two eye muscles		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67316	Revise eye muscle(s)		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67320	Revise eye muscle(s) add-on		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67331	Eye surgery follow-up add-on		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67332	Revise eye muscles add-on		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67334	Revise eye muscle w/suture		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67335	Eye suture during surgery		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67940	Revise eye muscle add-on		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67943	Release eye tissue		T	0243	23.4694	\$1,581.98	\$418.00	\$316.40
67945	Destroy nerve of eye muscle		T	0238	3.2267	\$217.50	\$43.50	\$209.51
67946	Biopsy, eye muscle		T	0699	15.5407	\$1,047.54	\$296.93	\$244.80
67999	Eye muscle surgery procedure		T	0240	23.4694	\$1,581.98	\$418.00	\$316.40
67400	Explore/biopsy eye socket		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67405	Explore/drain eye socket		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
67412	Explore/treat eye socket		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67413	Explore/treat eye socket		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
67414	Expir/decompress eye socket		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67415	Aspiration, orbital contents		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67420	Explore/treat eye socket		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67430	Explore/treat eye socket		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67440	Explore/drain eye socket		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67445	Expir/decompress eye socket		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67450	Explore/biopsy eye socket		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67500	Inject/treat eye socket		S	0231	1.9575	\$131.95	\$26.39	\$26.39
67505	Inject/treat eye socket		T	0238	3.2267	\$217.50	\$43.50	\$43.50
67550	Insert eye socket implant		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67560	Revise eye socket implant		T	0241	26.8396	\$1,809.15	\$383.45	\$361.83
67570	Decompress optic nerve		T	0242	39.0115	\$2,629.61	\$597.36	\$525.93
67599	Orbit surgery procedure		T	0238	3.2267	\$217.50	\$43.50	\$43.50
67700	Drainage of eyelid abscess		T	0239	3.2267	\$217.50	\$43.50	\$43.50
67710	Incision of eyelid		T	0239	7.6421	\$515.12	\$103.03	\$103.03
67715	Incision of eyelid fold		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67800	Remove eyelid lesion		T	0238	3.2267	\$217.50	\$43.50	\$43.50
67801	Remove eyelid lesions		T	0239	7.6421	\$515.12	\$103.03	\$103.03
67805	Remove eyelid lesions		T	0238	3.2267	\$217.50	\$43.50	\$43.50
67808	Remove eyelid lesion(s)		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67810	Biopsy of eyelid		T	0238	3.2267	\$217.50	\$43.50	\$43.50
67820	Revise eyelashes		S	0698	0.9553	\$64.39	\$12.88	\$12.88
67825	Revise eyelashes		T	0238	3.2267	\$217.50	\$43.50	\$43.50
67830	Revise eyelashes		T	0239	7.6421	\$515.12	\$103.03	\$103.03
67835	Revise eyelashes		T	0240	18.158	\$1,223.96	\$296.93	\$244.80
67840	Remove eyelid lesion		T	0239	7.6421	\$515.12	\$103.03	\$103.03

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
69120	Remove of external ear	T		0254	24.9637	\$1,682.70		\$336.54
69140	Remove ear canal lesion(s)	T		0254	24.9637	\$1,682.70		\$336.54
69145	Remove ear canal lesion(s)	T		0021	17.4975	\$1,179.44		\$235.89
69150	Extensive ear canal surgery	T		0252	7.6196	\$513.61	\$109.16	\$102.73
69200	Clear outer ear canal	C						
69205	Clear outer ear canal	X		0340	0.6693	\$45.11		\$9.03
69210	Remove impacted ear wax	T		0222	23.388	\$1,576.49	\$354.45	\$315.30
69220	Remove impacted ear wax	X		0340	0.6693	\$45.11		\$9.03
69222	Clean out mastoid cavity	T		0013	0.8789	\$59.24		\$11.85
69300	Clean out mastoid cavity	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
69310	Revise external ear	T		0254	24.9637	\$1,682.70		\$336.54
69320	Rebuild outer ear canal	T		0256	42.9827	\$2,897.29		\$579.46
69330	Rebuild outer ear canal	T		0256	42.9827	\$2,897.29		\$579.46
69399	Outer ear surgery procedure	T		0250	1.1522	\$77.67	\$25.10	\$15.54
69400	Inflate middle ear canal	T		0251	3.425	\$230.87		\$46.18
69405	Inflate middle ear canal	T		0251	3.425	\$230.87		\$46.18
69420	Catheterize middle ear canal	T		0252	7.6196	\$513.61	\$109.16	\$102.73
69421	Incision of eardrum	T		0251	3.425	\$230.87		\$46.18
69424	Remove ventilating tube	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
69433	Create eardrum opening	T		0252	7.6196	\$513.61	\$109.16	\$102.73
69436	Create eardrum opening	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
69440	Exploration of middle ear	T		0254	24.9637	\$1,682.70		\$336.54
69501	Mastoidectomy	T		0256	42.9827	\$2,897.29		\$579.46
69502	Mastoidectomy	T		0254	24.9637	\$1,682.70		\$336.54
69505	Remove mastoid structures	T		0256	42.9827	\$2,897.29		\$579.46
69511	Extensive mastoid surgery	T		0256	42.9827	\$2,897.29		\$579.46
69530	Extensive mastoid surgery	T		0256	42.9827	\$2,897.29		\$579.46
69535	Remove part of temporal bone	C						
69540	Remove ear lesion	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
69550	Remove ear lesion	T		0256	42.9827	\$2,897.29		\$579.46
69552	Remove ear lesion	T		0256	42.9827	\$2,897.29		\$579.46
69554	Remove ear lesion	C						
69601	Mastoid surgery revision	T		0256	42.9827	\$2,897.29		\$579.46
69602	Mastoid surgery revision	T		0256	42.9827	\$2,897.29		\$579.46
69603	Mastoid surgery revision	T		0256	42.9827	\$2,897.29		\$579.46
69604	Mastoid surgery revision	T		0256	42.9827	\$2,897.29		\$579.46
69605	Mastoid surgery revision	T		0256	42.9827	\$2,897.29		\$579.46
69610	Repair of eardrum	T		0254	24.9637	\$1,682.70		\$336.54
69620	Repair of eardrum	T		0254	24.9637	\$1,682.70		\$336.54
69631	Repair eardrum structures	T		0256	42.9827	\$2,897.29		\$579.46
69632	Rebuild eardrum structures	T		0256	42.9827	\$2,897.29		\$579.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
68326	Revise/graft eyelid lining	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68328	Revise/graft eyelid lining	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68330	Revise eyelid lining	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
68335	Revise/graft eyelid lining	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68340	Separate eyelid adhesions	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68360	Revise eyelid lining	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
68362	Revise eyelid lining	T		0234	24.4021	\$1,644.85	\$511.31	\$328.97
68371	Harvest eye tissue, alograft	T		0233	16.2485	\$1,095.25	\$263.77	\$219.05
68399	Eyelid lining surgery	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68400	Incise/drain tear gland	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68420	Incise/drain tear sac	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68440	Incise tear duct opening	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68500	Removal of tear gland	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68505	Partial removal, tear gland	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68510	Biopsy of tear gland	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68520	Removal of tear sac	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68525	Biopsy of tear sac	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68530	Clearance of tear duct	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68540	Remove tear gland lesion	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68550	Remove tear gland lesion	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68700	Repair tear ducts	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68705	Revise tear duct opening	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68720	Create tear duct opening	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68745	Create tear duct drain	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68750	Create tear duct drain	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68760	Close tear duct opening	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68761	Close tear duct opening	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68770	Close tear system fistula	T		0241	26.8396	\$1,809.15	\$383.45	\$361.83
68801	Dilate tear duct opening	S		0698	0.9553	\$64.39	\$12.88	\$12.88
68810	Probe nasolacrimal duct	CH						
68811	Probe nasolacrimal duct	T		0238	3.2267	\$217.50	\$43.50	\$43.50
68815	Probe nasolacrimal duct	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68816	Probe n duct w/balloon	T		0240	18.158	\$1,223.96	\$296.93	\$244.80
68840	Explore/irrigate tear ducts	S		0231	1.9575	\$131.95	\$26.39	\$26.39
68850	Injection for tear sac x-ray	N						
68899	Tear duct system surgery	T		0238	3.2267	\$217.50	\$43.50	\$43.50
69000	Drain external ear lesion	T		0006	1.4557	\$88.12	\$19.83	\$19.83
69005	Drain external ear lesion	T		0008	19.4063	\$1,308.10	\$261.62	\$261.62
69020	Drain outer ear canal lesion	T		0006	1.4557	\$88.12	\$19.83	\$19.83
69090	Pierce earlobes	E						
69100	Biopsy of external ear	T		0251	3.425	\$230.87	\$46.18	\$46.18
69105	Biopsy of external ear canal	T		0253	17.1879	\$1,158.57	\$282.29	\$231.72
69110	Remove external ear, partial	T		0021	17.4975	\$1,179.44		\$235.89

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
69960	Release inner ear canal	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69970	Remove inner ear lesion	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69979	Temporal bone surgery	T	0250	1.1522	\$77.67	\$15.54	\$25.10	\$15.54
69990	Microsurgery add-on	N						
70010	Contrast x-ray of brain	Q2	0274	7.226	\$487.08	\$97.42		\$97.42
70015	Contrast x-ray of brain	Q2	0274	7.226	\$487.08	\$97.42		\$97.42
70030	X-ray eye for foreign body	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70100	X-ray exam of jaw	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70110	X-ray exam of jaw	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70120	X-ray exam of mastoids	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70130	X-ray exam of mastoids	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70134	X-ray exam of middle ear	X	0261	1.1161	\$75.23	\$15.05		\$15.05
70140	X-ray exam of facial bones	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70150	X-ray exam of facial bones	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70160	X-ray exam of nasal bones	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70170	X-ray exam of ear duct	Q2	0263	3.1192	\$210.25	\$42.05		\$42.05
70180	X-ray exam of eye sockets	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70200	X-ray exam of eye sockets	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70210	X-ray exam of sinuses	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70220	X-ray exam of sinuses	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70240	X-ray exam, pituitary saddle	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70250	X-ray exam of skull	X	0261	1.1161	\$75.23	\$15.05		\$15.05
70300	X-ray exam of teeth	X	0262	0.4469	\$30.12	\$6.03		\$6.03
70310	X-ray exam of teeth	X	0262	0.4469	\$30.12	\$6.03		\$6.03
70320	Full mouth x-ray of teeth	X	0262	0.4469	\$30.12	\$6.03		\$6.03
70328	X-ray exam of jaw joint	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70330	X-ray exam of jaw joints	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70332	X-ray exam of jaw joint	Q2	0275	3.9361	\$265.32	\$53.07	\$69.07	\$53.07
70336	Magnetic image, jaw joint	Q3	0336	5.1855	\$349.53	\$69.91	\$137.40	\$69.91
70350	X-ray head for otitidis media	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70355	Panoramic x-ray of jaws	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70360	X-ray exam of neck	X	0260	0.6661	\$44.90	\$6.98		\$6.98
70370	Throat x-ray & fluoroscopy	X	0272	1.2693	\$85.56	\$17.12		\$17.12
70371	Speech evaluation, complex	X	0272	1.2693	\$85.56	\$17.12		\$17.12
70373	Contrast x-ray of larynx	Q2	0263	3.1192	\$210.25	\$42.05		\$42.05
70380	X-ray exam of salivary gland	Q2	0263	3.1192	\$210.25	\$42.05		\$42.05
70390	X-ray exam of salivary duct	Q3	0332	2.894	\$195.07	\$39.02	\$75.24	\$39.02
70450	Ct head/brain w/o dye	Q3	0283	4.4066	\$297.03	\$59.41	\$96.86	\$59.41
70470	Ct head/brain w/o dye	Q3	0333	4.9479	\$333.52	\$66.71	\$116.41	\$66.71
70480	Ct orbit/ear/fossa w/o dye	Q3	0332	2.894	\$195.07	\$39.02	\$75.24	\$39.02
70481	Ct orbit/ear/fossa w/o dye	Q3	0283	4.4066	\$297.03	\$59.41	\$96.86	\$59.41

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
69633	Rebuild eardrum structures	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69635	Rebuild eardrum structures	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69636	Rebuild eardrum structures	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69637	Rebuild eardrum structures	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69641	Revise middle ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69642	Revise middle ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69643	Revise middle ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69644	Revise middle ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69645	Revise middle ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69646	Revise middle ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69650	Release middle ear bone	T	0254	24.9637	\$1,682.70	\$336.54		\$336.54
69660	Revise middle ear bone	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69661	Revise middle ear bone	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69662	Revise middle ear bone	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69666	Repair middle ear structures	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69667	Repair middle ear structures	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69670	Remove mastoid air cells	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69676	Remove middle ear nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69700	Close mastoid fistula	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69710	Implant/replace hearing aid	E						
69711	Remove/repair hearing aid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69714	Implant temple bone w/stimul	T	0425	118.767	\$8,005.61	\$1,601.13		\$1,601.13
69715	Temple bone implant w/stimulat	T	0425	118.767	\$8,005.61	\$1,601.13		\$1,601.13
69717	Temple bone implant revision	T	0425	118.767	\$8,005.61	\$1,601.13		\$1,601.13
69718	Revise temple bone implant	T	0425	118.767	\$8,005.61	\$1,601.13		\$1,601.13
69720	Release facial nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69725	Release facial nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69740	Repair facial nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69745	Repair facial nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69799	Middle ear surgery procedure	T	0250	1.1522	\$77.67	\$15.54	\$25.10	\$15.54
69801	Incise inner ear	CH	0254	24.9637	\$1,682.70	\$336.54		\$336.54
69802	Incise inner ear	CH	0254	24.9637	\$1,682.70	\$336.54		\$336.54
69805	Explore inner ear	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69806	Explore inner ear	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69840	Establish inner ear window	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69805	Remove inner ear	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69910	Remove inner ear & mastoid	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69915	Incise inner ear nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46
69930	Implant cochlear device	T	0259	428.8363	\$28,906.14	\$8,543.66		\$8,543.66
69949	Inner ear surgery procedure	T	0250	1.1522	\$77.67	\$15.54	\$25.10	\$15.54
69950	Incise inner ear nerve	C						
69955	Release facial nerve	T	0256	42.9827	\$2,897.29	\$579.46		\$579.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
71130	X-ray exam of breastbone	X	0260	0260	0.6661	\$44.90	\$75.24	\$9.98
71150	Ct thorax w/o dye	Q3	0283	0283	2.894	\$195.07	\$195.07	\$39.02
71260	Ct thorax w/dye	Q3	0283	0283	4.4066	\$297.03	\$297.03	\$59.41
71270	Ct thorax w/o & w/dye	Q3	0333	0333	4.9479	\$333.52	\$116.41	\$66.71
71275	Ct angiography, chest	Q3	0662	0662	5.0511	\$340.47	\$115.03	\$68.10
71550	Mri chest w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
71551	Mri chest w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
71552	Mri chest w/o & w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
71555	Mri angio chest w/ w/o dye	B	0261	0261	1.1161	\$75.23		\$15.05
72020	X-ray exam of spine	X	0260	0260	0.6661	\$44.90		\$9.98
72040	X-ray exam of neck spine	X	0260	0260	0.6661	\$44.90		\$9.98
72050	X-ray exam of neck spine	X	0261	0261	1.1161	\$75.23		\$15.05
72052	X-ray exam of neck spine	X	0261	0261	1.1161	\$75.23		\$15.05
72069	X-ray exam of trunk spine	X	0260	0260	0.6661	\$44.90		\$9.98
72070	X-ray exam of thoracic spine	X	0260	0260	0.6661	\$44.90		\$9.98
72074	X-ray exam of thoracic spine	X	0260	0260	0.6661	\$44.90		\$9.98
72080	X-ray exam of trunk spine	X	0260	0260	0.6661	\$44.90		\$9.98
72100	X-ray exam of lower spine	X	0260	0260	0.6661	\$44.90		\$9.98
72110	X-ray exam of lower spine	X	0261	0261	1.1161	\$75.23		\$15.05
72114	X-ray exam of lower spine	X	0261	0261	1.1161	\$75.23		\$15.05
72120	X-ray exam of lower spine	X	0261	0261	1.1161	\$75.23		\$15.05
72125	Ct neck spine w/o dye	Q3	0332	0332	2.894	\$195.07	\$75.24	\$39.02
72126	Ct neck spine w/dye	Q3	0283	0283	4.4066	\$297.03	\$96.86	\$59.41
72127	Ct neck spine w/o & w/dye	Q3	0333	0333	4.9479	\$333.52	\$116.41	\$66.71
72128	Ct chest spine w/o dye	Q3	0332	0332	2.894	\$195.07	\$75.24	\$39.02
72129	Ct chest spine w/dye	Q3	0283	0283	4.4066	\$297.03	\$96.86	\$59.41
72130	Ct chest spine w/o & w/dye	Q3	0333	0333	4.9479	\$333.52	\$116.41	\$66.71
72131	Ct lumbar spine w/o dye	Q3	0332	0332	2.894	\$195.07	\$75.24	\$39.02
72132	Ct lumbar spine w/dye	Q3	0283	0283	4.4066	\$297.03	\$96.86	\$59.41
72141	Mri neck spine w/o & w/dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
72142	Mri neck spine w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
72146	Mri chest spine w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
72149	Mri chest spine w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
72148	Mri lumbar spine w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
72156	Mri lumbar spine w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
72157	Mri chest spine w/o & w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
72158	Mri lumbar spine w/o & w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
72159	Mri angio spine w/o&w/dye	E						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
70482	Ct orbit/ear/fossa w/o&w/dye	Q3	0333	0333	4.9479	\$333.52	\$116.41	\$66.71
70486	Ct maxillofacial w/o dye	Q3	0332	0332	2.894	\$195.07	\$75.24	\$39.02
70487	Ct maxillofacial w/dye	Q3	0283	0283	4.4066	\$297.03	\$96.86	\$59.41
70488	Ct maxillofacial w/o & w/dye	Q3	0333	0333	4.9479	\$333.52	\$116.41	\$66.71
70490	Ct soft tissue neck w/o dye	Q3	0332	0332	2.894	\$195.07	\$75.24	\$39.02
70491	Ct soft tissue neck w/dye	Q3	0283	0283	4.4066	\$297.03	\$96.86	\$59.41
70492	Ct soft tissue neck w/o & w/dye	Q3	0333	0333	4.9479	\$333.52	\$116.41	\$66.71
70498	Ct angiography, head	Q3	0662	0662	5.0511	\$340.47	\$115.03	\$68.10
70540	Mri orbit/face/neck w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70542	Mri orbit/face/neck w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
70543	Mri orbit/face/neck w/o & w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
70544	Mri angiography, head w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70545	Mri angiography, head w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
70546	Mri angiography, head w/o&w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
70547	Mri angiography, neck w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70548	Mri angiography, neck w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
70549	Mri angiography, neck w/o&w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
70551	Mri brain w/o dye	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70552	Mri brain w/dye	Q3	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
70553	Mri brain w/o & w/dye	Q3	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
70554	Fmri brain by tech	Q3	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70555	Fmri brain by phys/psych	S	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70557	Mri brain w/o dye	S	0336	0336	5.1855	\$349.53	\$137.40	\$69.91
70558	Mri brain w/dye	S	0284	0284	6.2901	\$423.99	\$147.21	\$84.80
70559	Mri brain w/o & w/dye	S	0337	0337	7.9432	\$535.42	\$198.32	\$107.09
71010	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71015	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71020	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71021	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71022	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71023	Chest x-ray and fluoroscopy	X	0272	0272	1.2693	\$85.56	\$31.15	\$17.12
71030	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71034	Chest x-ray and fluoroscopy	X	0272	0272	1.2693	\$85.56	\$31.15	\$17.12
71035	Chest x-ray	X	0260	0260	0.6661	\$44.90		\$9.98
71040	Contrast x-ray of bronchi	Q2	0263	0263	3.1192	\$210.25	\$42.05	\$21.03
71060	Contrast x-ray of bronchi	O2	0263	0263	3.1192	\$210.25	\$42.05	\$21.03
71090	X-ray & pacemaker insertion	N						
71100	X-ray exam of ribs	X	0260	0260	0.6661	\$44.90		\$9.98
71101	X-ray exam of ribs/chest	X	0260	0260	0.6661	\$44.90		\$9.98
71110	X-ray exam of ribs	X	0260	0260	0.6661	\$44.90		\$9.98
71111	X-ray exam of ribs/chest	X	0261	0261	1.1161	\$75.23		\$15.05
71120	X-ray exam of breastbone	X	0260	0260	0.6661	\$44.90		\$9.98

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
73206	Ct angio upr extrm w/o&w/dye		Q3	0662	5.0511	\$340.47	\$115.03	\$68.10
73218	Mri upper extrem w/o&w/dye		Q3	0336	5.1855	\$349.53	\$137.40	\$69.91
73219	Mri upper extrem w/dye		Q3	0284	6.2901	\$423.99	\$147.21	\$84.80
73220	Mri uppr extrem w/o&w/dye		Q3	0337	7.9432	\$535.42	\$198.32	\$107.09
73221	Mri joint upr extrm w/o dye		Q3	0336	5.1855	\$349.53	\$137.40	\$69.91
73222	Mri joint upr extrm w/dye		Q3	0284	6.2901	\$423.99	\$147.21	\$84.80
73223	Mri joint upr extr w/o&w/dye		Q3	0337	7.9432	\$535.42	\$198.32	\$107.09
73225	Mri angio upr extr w/o&w/dye		E					
73500	X-ray exam of hip		X	0260	0.6661	\$44.90		\$8.98
73510	X-ray exam of hip		X	0260	0.6661	\$44.90		\$8.98
73520	X-ray exam of hips		X	0261	1.1161	\$75.23		\$15.05
73525	Contrast x-ray of hip		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73530	X-ray exam of hip		N					
73540	X-ray exam of pelvis & hips		X	0260	0.6661	\$44.90		\$8.98
73542	X-ray exam, sacroiliac joint		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73550	X-ray exam of thigh		X	0260	0.6661	\$44.90		\$8.98
73560	X-ray exam of knee, 1 or 2		X	0260	0.6661	\$44.90		\$8.98
73562	X-ray exam of knee, 3		X	0260	0.6661	\$44.90		\$8.98
73564	X-ray exam, knee, 4 or more		X	0260	0.6661	\$44.90		\$8.98
73565	X-ray exam of knees		X	0260	0.6661	\$44.90		\$8.98
73580	Contrast x-ray of knee joint		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73590	X-ray exam of lower leg		X	0260	0.6661	\$44.90		\$8.98
73592	X-ray exam of leg, infant		X	0260	0.6661	\$44.90		\$8.98
73610	X-ray exam of ankle		X	0260	0.6661	\$44.90		\$8.98
73615	Contrast x-ray of ankle		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73620	X-ray exam of foot		X	0260	0.6661	\$44.90		\$8.98
73630	X-ray exam of heel		X	0260	0.6661	\$44.90		\$8.98
73650	X-ray exam of heel		X	0260	0.6661	\$44.90		\$8.98
73660	X-ray exam of toes(s)		X	0260	0.6661	\$44.90		\$8.98
73700	Ct lower extremity w/o dye		Q3	0332	2.894	\$195.07	\$75.24	\$39.02
73701	Ct lower extremity w/dye		Q3	0283	4.4066	\$297.03	\$96.86	\$59.41
73702	Ct lwr extremity w/o&w/dye		Q3	0333	4.9479	\$333.52	\$116.41	\$66.71
73706	Ct angio lwr extr w/o&w/dye		Q3	0662	5.0511	\$340.47	\$115.03	\$68.10
73718	Mri lower extremity w/o dye		Q3	0336	5.1855	\$349.53	\$137.40	\$69.91
73719	Mri lower extremity w/dye		Q3	0284	6.2901	\$423.99	\$147.21	\$84.80
73720	Mri lwr extremity w/o&w/dye		Q3	0337	7.9432	\$535.42	\$198.32	\$107.09
73721	Mri jnt of lwr extre w/o dye		Q3	0336	5.1855	\$349.53	\$137.40	\$69.91
73722	Mri joint of lwr extre w/dye		Q3	0284	6.2901	\$423.99	\$147.21	\$84.80
73723	Mri joint lwr extr w/o&w/dye		Q3	0337	7.9432	\$535.42	\$198.32	\$107.09
73725	Mri ang lwr ext w/ w/o dye		B					
74000	X-ray exam of abdomen		X	0260	0.6661	\$44.90		\$8.98
74010	X-ray exam of abdomen		X	0260	0.6661	\$44.90		\$8.98

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
72170	X-ray exam of pelvis		X	0260	0.6661	\$44.90		\$8.98
72190	X-ray exam of pelvis		X	0260	0.6661	\$44.90		\$8.98
72191	Ct angiograph pelv w/o&w/dye		Q3	0662	5.0511	\$340.47	\$115.03	\$68.10
72192	Ct pelvis w/o dye		Q3	0332	2.894	\$195.07	\$75.24	\$39.02
72193	Ct pelvis w/dye		Q3	0283	4.4066	\$297.03	\$96.86	\$59.41
72194	Ct pelvis w/o & w/dye		Q3	0333	4.9479	\$333.52	\$116.41	\$66.71
72195	Mri pelvis w/o dye		Q3	0336	5.1855	\$349.53	\$137.40	\$69.91
72196	Mri pelvis w/dye		Q3	0284	6.2901	\$423.99	\$147.21	\$84.80
72197	Mri pelvis w/o & w/dye		Q3	0337	7.9432	\$535.42	\$198.32	\$107.09
72198	Mri angio pelvis w/o & w/dye		B					
72200	X-ray exam sacroiliac joints		X	0260	0.6661	\$44.90		\$8.98
72202	X-ray exam sacroiliac joints		X	0260	0.6661	\$44.90		\$8.98
72220	X-ray exam of tailbone		X	0260	0.6661	\$44.90		\$8.98
72240	Contrast x-ray of neck spine		Q2	0274	7.226	\$487.08	\$97.42	\$53.07
72255	Contrast x-ray, thorax spine		Q2	0274	7.226	\$487.08	\$97.42	\$53.07
72265	Contrast x-ray, lower spine		Q2	0274	7.226	\$487.08	\$97.42	\$53.07
72270	Contrast x-ray, spine		Q2	0274	7.226	\$487.08	\$97.42	\$53.07
72275	Epidurography		N					
72285	X-ray c/t spine disk		Q2	0388	25.8604	\$1,743.15	\$348.63	\$216.41
72291	Perq verte/sacroplasty, ilior		N					
72292	Perq verte/sacroplasty, ct		N					
72295	X-ray of lower spine disk		Q2	0388	25.8604	\$1,743.15	\$348.63	\$216.41
73000	X-ray exam of collar bone		X	0260	0.6661	\$44.90		\$8.98
73010	X-ray exam of shoulder blade		X	0260	0.6661	\$44.90		\$8.98
73020	X-ray exam of shoulder		X	0260	0.6661	\$44.90		\$8.98
73030	X-ray exam of shoulder		X	0260	0.6661	\$44.90		\$8.98
73040	Contrast x-ray of shoulder		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73050	X-ray exam of shoulders		X	0260	0.6661	\$44.90		\$8.98
73060	X-ray exam of humerus		X	0260	0.6661	\$44.90		\$8.98
73070	X-ray exam of elbow		X	0260	0.6661	\$44.90		\$8.98
73080	X-ray exam of elbow		X	0260	0.6661	\$44.90		\$8.98
73085	Contrast x-ray of elbow		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73090	X-ray exam of forearm		X	0260	0.6661	\$44.90		\$8.98
73092	X-ray exam of arm, infant		X	0260	0.6661	\$44.90		\$8.98
73100	X-ray exam of wrist		X	0260	0.6661	\$44.90		\$8.98
73110	X-ray exam of wrist		X	0260	0.6661	\$44.90		\$8.98
73115	Contrast x-ray of wrist		Q2	0275	3.9361	\$265.32	\$69.07	\$53.07
73120	X-ray exam of hand		X	0260	0.6661	\$44.90		\$8.98
73130	X-ray exam of hand		X	0260	0.6661	\$44.90		\$8.98
73140	X-ray exam of finger(s)		X	0260	0.6661	\$44.90		\$8.98
73200	Ct upper extremity w/o dye		Q3	0332	2.894	\$195.07	\$75.24	\$39.02
73201	Ct upper extremity w/dye		Q3	0283	4.4066	\$297.03	\$96.86	\$59.41
73202	Ct uppr extremity w/o&w/dye		Q3	0333	4.9479	\$333.52	\$116.41	\$66.71

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
74363	X-ray, bile duct dilation		N	0278	2.546	\$171.62	\$58.59	\$34.33
74400	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74410	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74415	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74420	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74425	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74430	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74435	Contrast x-ray, urinary tract		S	0278	2.546	\$171.62	\$58.59	\$34.33
74440	X-ray, male genital tract		O2	0278	2.546	\$171.62	\$58.59	\$34.33
74445	X-ray exam of penis		O2	0278	2.546	\$171.62	\$58.59	\$34.33
74450	X-ray, urethra/bladder		O2	0278	2.546	\$171.62	\$58.59	\$34.33
74455	X-ray, urethra/bladder		O2	0278	2.546	\$171.62	\$58.59	\$34.33
74470	X-ray exam of kidney lesion		O2	0263	3.1192	\$210.25	\$42.05	\$42.05
74475	X-ray control, cath insert	CH	O2	0161	17.0344	\$1,148.22	\$229.65	\$229.65
74480	X-ray control, cath insert	CH	O2	0161	17.0344	\$1,148.22	\$229.65	\$229.65
74485	X-ray guide, GU dilation	CH	O2	0161	17.0344	\$1,148.22	\$229.65	\$229.65
74710	X-ray measurement of pelvis		X	0261	1.1161	\$75.23	\$15.05	\$15.05
74740	X-ray, female genital tract		O2	0263	3.1192	\$210.25	\$42.05	\$42.05
74742	X-ray, fallopian tube		N					
74775	X-ray exam of perineum		S	0278	2.546	\$171.62	\$58.59	\$34.33
75557	Cardiac mri for morph		O3	0336	5.1855	\$349.53	\$137.40	\$69.91
75558	Cardiac mri flow/velocity	CH	D					
75559	Cardiac mri w/stress img		O3	0336	5.1855	\$349.53	\$137.40	\$69.91
75560	Cardiac mri flow/vel/stress	CH	D					
75561	Cardiac mri for morph w/dye		O3	0337	7.9432	\$535.42	\$198.32	\$107.09
75562	Card mri flow/vel w/dye	CH	D					
75563	Card mri w/stress img & dye		O3	0337	7.9432	\$535.42	\$198.32	\$107.09
75564	Ht mri w/flow/vel/stress & dye	CH	D					
75565	Card mri vel flow map add-on	NI	N					
75571	Ct hrt w/dye w/ca test	NI	X	0340	0.6693	\$45.11	\$9.03	\$9.03
75572	Ct hrt w/d3d image	NI	S	0383	3.9929	\$269.15	\$107.66	\$53.83
75573	Ct hrt w/d3d image, congen	NI	S	0383	3.9929	\$269.15	\$107.66	\$53.83
75574	Ct angio hrt w/d3d image	NI	S	0383	3.9929	\$269.15	\$107.66	\$53.83
75600	Contrast x-ray exam of aorta		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48
75605	Contrast x-ray exam of aorta		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48
75625	Contrast x-ray exam of aorta		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48
75630	X-ray aorta, leg arteries		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48
75635	Ct angio abdominal arteries		O2	0682	5.0511	\$340.47	\$115.03	\$68.10
75650	Artery x-rays, head & neck		O2	0280	45.8793	\$3,092.54	\$618.51	\$618.51
75658	Artery x-rays, arm		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48
75660	Artery x-rays, head & neck		O2	0280	45.8793	\$3,092.54	\$618.51	\$618.51
75662	Artery x-rays, head & neck		O2	0280	45.8793	\$3,092.54	\$618.51	\$618.51
75665	Artery x-rays, head & neck		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48
75671	Artery x-rays, head & neck		O2	0280	45.8793	\$3,092.54	\$618.51	\$618.51

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
74020	X-ray exam of abdomen		X	0260	0.6661	\$44.90	\$8.98	\$8.98
74022	X-ray exam series, abdomen		X	0261	1.1161	\$75.23	\$15.05	\$15.05
74150	Ct abdomen w/o dye		O3	0332	2.894	\$195.07	\$75.24	\$39.02
74160	Ct abdomen w/dye		O3	0283	4.4086	\$297.03	\$96.86	\$59.41
74170	Ct abdomen w/o & w/dye		O3	0333	4.9479	\$333.52	\$116.41	\$66.71
74175	Ct angio abdomen w/o & w/dye		O3	0662	5.0511	\$340.47	\$115.03	\$68.10
74181	Mri abdomen w/o dye		O3	0336	5.1855	\$349.53	\$137.40	\$69.91
74182	Mri abdomen w/dye		O3	0284	6.2901	\$423.99	\$147.21	\$84.90
74183	Mri abdomen w/o & w/dye		O3	0337	7.9432	\$535.42	\$198.32	\$107.09
74185	Mri angio, abdom, w onw/o dye		B					
74190	X-ray exam of peritoneum		O2	0263	3.1192	\$210.25	\$42.05	\$42.05
74210	Contrast x-ray exam of throat		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74220	Contrast x-ray, esophagus		O2	0276	1.2985	\$87.53	\$34.74	\$17.51
74230	Cine/wid x-ray, throat/esoph		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74235	Remove esophagus obstruction		N					
74240	X-ray exam, upper gi tract		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74241	X-ray exam, upper gi tract		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74245	X-ray exam, upper gi tract		S	0277	2.1001	\$141.56	\$54.04	\$28.32
74246	Contrast x-ray upper gi tract		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74247	Contrast x-ray upper gi tract		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74249	Contrast x-ray upper gi tract		S	0277	2.1001	\$141.56	\$54.04	\$28.32
74250	X-ray exam of small bowel		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74251	X-ray exam of small bowel		S	0277	2.1001	\$141.56	\$54.04	\$28.32
74260	X-ray exam of small bowel		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74261	Ct colonography, w/o dye	NI	O3	0332	2.894	\$195.07	\$75.24	\$39.02
74262	Ct colonography, w/dye	NI	O3	0283	4.4066	\$297.03	\$96.86	\$59.41
74263	Ct colonography, screen	NI	E					
74270	Contrast x-ray exam of colon		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74280	Contrast x-ray exam of colon		S	0277	2.1001	\$141.56	\$54.04	\$28.32
74283	Contrast x-ray exam of colon		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74290	Contrast x-ray, gallbladder		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74291	Contrast x-rays, gallbladder		S	0276	1.2985	\$87.53	\$34.74	\$17.51
74300	X-ray bile ducts/pancreas		N					
74301	X-rays at surgery add-on		N					
74305	X-ray bile ducts/pancreas		O2	0263	3.1192	\$210.25	\$42.05	\$42.05
74320	Contrast x-ray of bile ducts		O2	0317	5.6012	\$377.55	\$75.51	\$75.51
74327	X-ray bile stone removal		N					
74328	X-ray bile duct endoscopy		N					
74329	X-ray for pancreas endoscopy		N					
74330	X-ray bile/panc endoscopy		N					
74340	X-ray guide for GI tube		N					
74355	X-ray guide, intestinal tube		N					
74360	X-ray guide, GI dilation		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010												
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment				
75896	X-rays, transcath therapy	CH	Q1	0261	1.1161	\$75.23		\$15.05				
75898	Follow-up angiography											
75900	Intravascular cath exchange		C									
75901	Remove cva device obstruct		N									
75902	Remove cva lumen obstruct		N									
75940	X-ray placement, vein filter		N									
75945	Intravascular us		O2	0267	2.3005	\$155.07	\$60.50	\$31.02				
75946	Intravascular us add-on		N									
75952	Endovasc repair abdom aorta		C									
75953	Abdom aneurysm endovas rpr		C									
75954	Iliac aneurysm endovas rpr		C									
75956	X-ray, endovasc thor ac repr		C									
75957	X-ray, endovasc thor ac repr		C									
75958	X-ray, place prox ext thor ao		C									
75959	X-ray, place dist ext thor ao		C									
75961	Retrieval, broken catheter		N									
75962	Repair, arterial blockage		O2	0083	50.6809	\$3,416.20		\$683.24				
75964	Repair artery blockage, each		N									
75966	Repair arterial blockage		O2	0083	50.6809	\$3,416.20		\$683.24				
75968	Repair artery blockage, each		N									
75970	Vascular biopsy		N									
75978	Repair venous blockage	CH	O2	0093	35.5969	\$2,399.44		\$479.89				
75980	Contrast x-ray exam bile duct		N									
75982	Contrast x-ray exam bile duct		N									
75989	X-ray control catheter change		N									
75992	Abscess drainage under x-ray		N									
75993	Atherectomy, x-ray exam		N									
75994	Atherectomy, x-ray exam		N									
75995	Atherectomy, x-ray exam		N									
75996	Atherectomy, x-ray exam		N									
76000	Fluoroscope examination		Q1	0272	1.2693	\$85.56	\$31.15	\$17.12				
76001	Fluoroscope exam, extensive		N									
76010	X-ray, nose to rectum		X	0260	0.6661	\$44.90		\$9.98				
76080	X-ray exam, breast specimen	CH	O2	0263	3.1192	\$210.25		\$42.05				
76098	X-ray exam, breast specimen		O2	0317	5.6012	\$377.55		\$75.51				
76100	X-ray exam of body section		X	0261	1.1161	\$75.23		\$15.05				
76101	Complex body section x-ray		X	0263	3.1192	\$210.25		\$42.05				
76102	Complex body section x-rays		X	0263	3.1192	\$210.25		\$42.05				
76120	Cine/video x-rays		X	0272	1.2693	\$85.56	\$31.15	\$17.12				
76125	Cine/video x-rays add-on		N									
76140	X-ray consultation		E									

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010												
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment				
75676	Artery x-rays, neck		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75680	Artery x-rays, neck		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75685	Artery x-rays, spine		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75700	Artery x-rays, spine		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75710	Artery x-rays, arm/leg		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75716	Artery x-rays, arms/legs		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75722	Artery x-rays, kidney		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75724	Artery x-rays, kidneys		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75731	Artery x-rays, adrenal gland		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75733	Artery x-rays, adrenals		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75736	Artery x-rays, pelvis		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75741	Artery x-rays, lung		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75743	Artery x-rays, lungs		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75746	Artery x-rays, lung		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75756	Artery x-rays, chest		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75774	Artery x-ray, each vessel		N									
75790	Visualize A-V shunt	CH	D									
75791	Av dialysis shunt imaging	NI	O2	0676	2.3954	\$161.46	\$32.30	\$32.30				
75801	Lymph vessel x-ray, arm/leg		O2	0317	5.6012	\$377.55	\$75.51	\$75.51				
75803	Lymph vessel x-ray, arms/legs		O2	0317	5.6012	\$377.55	\$75.51	\$75.51				
75805	Lymph vessel x-ray, trunk		O2	0317	5.6012	\$377.55	\$75.51	\$75.51				
75807	Lymph vessel x-ray, trunk		O2	0317	5.6012	\$377.55	\$75.51	\$75.51				
75809	Nonvascular shunt, x-ray		O2	0261	1.1161	\$75.23	\$15.05	\$15.05				
75810	Vein x-ray, spleen/liver		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75820	Vein x-ray, arm/leg		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75822	Vein x-ray, arms/legs		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75825	Vein x-ray, trunk		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75827	Vein x-ray, chest		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75831	Vein x-ray, kidney		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75833	Vein x-ray, kidneys		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75840	Vein x-ray, adrenal gland		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75842	Vein x-ray, adrenal glands		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75860	Vein x-ray, neck		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75870	Vein x-ray, skull		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75872	Vein x-ray, skull		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75880	Vein x-ray, eye socket		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75885	Vein x-ray, liver		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75887	Vein x-ray, liver		O2	0668	10.2978	\$694.13	\$138.83	\$138.83				
75889	Vein x-ray, liver		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75891	Vein x-ray, liver		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75893	Venous sampling by catheter		O2	0279	29.1126	\$1,962.36	\$392.48	\$392.48				
75894	X-rays, transcath therapy		N									

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
76825	Echo exam of fetal heart	CH	S	0270	8.8425	\$596.04	\$141.32	\$119.21
76826	Echo exam of fetal heart	CH	S	0269	6.6903	\$450.97		\$90.20
76827	Echo exam of fetal heart		S	0265	0.9267	\$62.47		\$12.50
76828	Echo exam of fetal heart		S	0265	0.9267	\$62.47		\$12.50
76830	Transvaginal us, non-ob		S	0266	1.4434	\$97.29		\$19.46
76831	Echo exam, uterus	O3	Q3	0267	2.3005	\$155.07		\$31.02
76856	Us exam, pelvic, complete	O3	Q3	0266	1.4434	\$97.29		\$19.46
76857	Us exam, pelvic, limited	O3	Q3	0265	0.9267	\$62.47		\$12.50
76870	Us exam, scrotum	O3	Q3	0266	1.4434	\$97.29		\$19.46
76872	Us, transrectal		S	0266	1.4434	\$97.29		\$19.46
76873	Echograp trans r, pros study		N	0266	1.4434	\$97.29		\$19.46
76880	Us exam, extremity		S	0265	0.9267	\$62.47		\$12.50
76885	Us exam infant hips, dynamic		S	0265	0.9267	\$62.47		\$12.50
76886	Us exam infant hips, static		S	0265	0.9267	\$62.47		\$12.50
76930	Echo guide, cardiocentesis		N					
76932	Echo guide for heart biopsy		N					
76936	Echo guide for artery repair		N					
76937	Us guide, vascular access		N					
76940	Us guide, tissue ablation		N					
76941	Echo guide for transfusion		N					
76942	Echo guide for biopsy		N					
76945	Echo guide, vilus sampling		N					
76946	Echo guide for amniocentesis		N					
76948	Echo guide, ova aspiration		N					
76950	Echo guidance radiotherapy		N					
76965	Echo guidance radiotherapy		N					
76970	Ultrasound exam follow-up		S	0265	0.9267	\$62.47		\$12.50
76975	GI endoscopic ultrasound		Q2	0267	2.3005	\$155.07		\$31.02
76977	Us bone density measure		X	0340	0.6693	\$45.11		\$9.03
76998	Us guide, intracp		N					
76999	Echo examination procedure		S	0265	0.9267	\$62.47		\$12.50
77001	Fluoroguide for vein device		N					
77002	Needle localization by xray		N					
77003	Fluoroguide for spine inject		N					
77011	Ct scan for localization		N					
77012	Ct scan for needle biopsy		N					
77013	Ct guide for tissue ablation		N					
77014	Ct scan for therapy guide		N					
77021	Mr guidance for needle place		N					
77022	Mr for tissue ablation		N					
77031	Stereotact guide for brst bx		N					
77032	Guidance for needle, breast		N					
77051	Computer dx mammogram add-on		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
76150	X-ray exam, dry process	X		0260	0.6661	\$44.90		\$8.98
76350	Special x-ray contrast study		N					
76376	3d rendering w/postprocess		N					
76377	3d rendering w/postprocess		N					
76380	CAT scan follow-up study		N	0282	1.6291	\$109.81		\$21.97
76390	Mr spectroscopy		E					
76496	Fluoroscopic procedure	X	Q272	1.2693	\$85.56		\$31.15	\$17.12
76497	Ct procedure	X	Q282	1.6291	\$109.81		\$37.81	\$21.97
76498	Mr procedure	X	Q336	5.1855	\$349.53		\$137.40	\$69.91
76499	Radiographic procedure	X	Q260	0.6661	\$44.90		\$8.98	
76506	Echo exam of head		S	0265	0.9267	\$62.47		\$12.50
76510	Ophth us, b & quant a	T	Q232	4.5074	\$303.63		\$76.12	\$60.77
76511	Ophth us, b w/non-quant a	S	Q266	1.4434	\$97.29		\$37.61	\$19.46
76512	Echo exam of eye, water bath		S	0266	1.4434	\$97.29		\$19.46
76513	Echo exam of eye, thickness	X	Q035	0.232	\$15.64		\$3.13	\$19.46
76516	Echo exam of eye		S	0265	0.9267	\$62.47		\$12.50
76519	Echo exam of eye		S	0266	1.4434	\$97.29		\$19.46
76529	Echo exam of eye		S	0265	0.9267	\$62.47		\$12.50
76536	Us exam of head and neck		S	0266	1.4434	\$97.29		\$19.46
76604	Us exam, chest	O3	Q3	0265	0.9267	\$62.47		\$12.50
76645	Us exam, breast(s)		S	0265	0.9267	\$62.47		\$12.50
76700	Us exam, abdom, complete		O3	0266	1.4434	\$97.29		\$19.46
76705	Echo exam of abdomen		O3	0266	1.4434	\$97.29		\$19.46
76770	Us exam abdo back wall, comp		O3	0266	1.4434	\$97.29		\$19.46
76775	Us exam abdo back wall, lim		O3	0266	1.4434	\$97.29		\$19.46
76776	Us exam k, transp w/doppler		O3	0266	1.4434	\$97.29		\$19.46
76800	Ob us < 14 wks, single fetus		S	0266	1.4434	\$97.29		\$19.46
76801	Ob us >= 14 wks, addl fetus		S	0265	0.9267	\$62.47		\$12.50
76802	Ob us < 14 wks, addl fetus		S	0266	1.4434	\$97.29		\$19.46
76805	Ob us >= 14 wks, singl fetus		S	0266	1.4434	\$97.29		\$19.46
76810	Ob us >= 14 wks, addl fetus		S	0265	0.9267	\$62.47		\$12.50
76811	Ob us, detailed, singl fetus		S	0267	2.3005	\$155.07		\$31.02
76812	Ob us, detailed, addl fetus		S	0265	0.9267	\$62.47		\$12.50
76813	Ob us nuchal meas, 1 gest		S	0265	0.9267	\$62.47		\$12.50
76814	Ob us nuchal meas, add-on		S	0265	0.9267	\$62.47		\$12.50
76815	Ob us, limited, fetus(s)		S	0265	0.9267	\$62.47		\$12.50
76816	Ob us, follow-up, per fetus		S	0265	0.9267	\$62.47		\$12.50
76817	Transvaginal us, obstetric		S	0265	0.9267	\$62.47		\$12.50
76818	Fetal biophys profile w/ntc		S	0266	1.4434	\$97.29		\$19.46
76819	Fetal biophys profil w/o nst		S	0266	1.4434	\$97.29		\$19.46
76820	Umbilical artery echo	CH	S	0265	0.9267	\$62.47		\$12.50
76821	Middle cerebral artery echo	CH	S	0265	0.9267	\$62.47		\$12.50

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
77336	Radiation physics consult	X	X	0304	1.5271	\$102.94	\$34.71	\$20.59
77338	Design nlc device for imrt	NI	X	0303	2.8279	\$190.62	\$66.95	\$38.13
77370	Radiation physics consult	X	X	0304	1.5271	\$102.94	\$34.71	\$20.59
77371	Srs, multisource	S	0127		108.9558	\$7,344.27		\$1,468.86
77372	Srs, linear based	B						
77373	Sbrt delivery	B						
77389	External radiation dosimetry	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77401	Radiation treatment delivery	S	0300		1.3764	\$92.78		\$18.56
77402	Radiation treatment delivery	S	0300		1.3764	\$92.78		\$18.56
77403	Radiation treatment delivery	S	0300		1.3764	\$92.78		\$18.56
77404	Radiation treatment delivery	S	0300		1.3764	\$92.78		\$18.56
77406	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77407	Radiation treatment delivery	S	0300		1.3764	\$92.78		\$18.56
77408	Radiation treatment delivery	S	0300		1.3764	\$92.78		\$18.56
77409	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77411	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77412	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77413	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77414	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77416	Radiation treatment delivery	S	0301		2.303	\$155.24		\$31.05
77417	Radiology port film(s)	N						
77418	Radiation tx delivery, imrt	S	0412		6.249	\$421.22		\$84.25
77421	Stereoscopic x-ray guidance	N						
77422	Neutron beam tx, simple	S	0301		2.303	\$155.24		\$31.05
77423	Neutron beam tx, complex	S	0301		2.303	\$155.24		\$31.05
77427	Radiation tx management, x5	B						
77431	Radiation therapy management	B						
77432	Stereotactic radiation trmt	N						
77435	Sbrt management	N						
77470	Special radiation treatment	S	0299		5.6461	\$380.58		\$76.12
77489	Radiation therapy management	B						
77520	Proton trmt, simple w/comp	S	0664		13.9796	\$942.31		\$188.47
77523	Proton trmt, simple w/comp	S	0664		13.9796	\$942.31		\$188.47
77529	Proton trmt, intermediate	S	0667		18.2873	\$1,232.67		\$246.54
77525	Proton treatment, complex	S	0667		18.2873	\$1,232.67		\$246.54
77605	Hyperthermia treatment	S	0299		5.6461	\$380.58		\$76.12
77610	Hyperthermia treatment	S	0299		5.6461	\$380.58		\$76.12
77615	Hyperthermia treatment	S	0299		5.6461	\$380.58		\$76.12
77620	Hyperthermia treatment	S	0299		5.6461	\$380.58		\$76.12
77750	Intracav radioactive materials	S	0301		2.303	\$155.24		\$31.05
77761	Apply intracav radiat simple	S	0312		4.4846	\$302.29		\$60.46
77762	Apply intracav radiat intern	S	0312		4.4846	\$302.29		\$60.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
77052	Comp screen mammogram add-on	A						
77053	X-ray of mammary duct	Q2	0263		3.1192	\$210.25		\$42.05
77054	X-ray of mammary ducts	Q2	0263		3.1192	\$210.25		\$42.05
77055	Mammogram, one breast	A						
77056	Mammogram, both breasts	A						
77057	Mammogram, screening	A						
77058	Mfi, one breast	B						
77059	Mfi, both breasts	B						
77071	X-ray stress view	X	0260		0.6661	\$44.90		\$8.98
77072	X-rays for bone age	X	0260		0.6661	\$44.90		\$8.98
77073	X-rays, bone length studies	X	0260		0.6661	\$44.90		\$8.98
77074	X-rays, bone survey, limited	X	0261		1.1161	\$75.23		\$15.05
77075	X-rays, bone survey complete	X	0261		1.1161	\$75.23		\$15.05
77076	X-rays, bone survey, infant	X	0261		1.1161	\$75.23		\$15.05
77077	Joint survey, single view	X	0260		0.6661	\$44.90		\$8.98
77078	Cl bone density, axial	S	0288		1.0755	\$72.50	\$28.66	\$14.50
77079	Cl bone density, peripheral	S	0282		1.6291	\$109.81	\$37.81	\$21.97
77080	Dxa bone density, axial	S	0288		1.0755	\$72.50	\$28.66	\$14.50
77081	Dxa bone density/peripheral	S	0665		0.4318	\$29.11	\$11.63	\$5.83
77082	Dxa bone density, vert fx	X	0260		0.6661	\$44.90		\$8.98
77083	Radiographic absorptiometry	X	0261		1.1161	\$75.23		\$15.05
77084	Magnetic image, bone marrow	S	0336		5.1855	\$349.53	\$137.40	\$69.91
77261	Radiation therapy planning	B						
77262	Radiation therapy planning	B						
77263	Radiation therapy planning	B						
77280	Set radiation therapy field	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77285	Set radiation therapy field	X	0305		3.951	\$266.32	\$91.38	\$53.27
77290	Set radiation therapy field	X	0305		3.951	\$266.32	\$91.38	\$53.27
77295	Set radiation therapy field	X	0310		13.7576	\$927.34	\$325.27	\$185.47
77299	Radiation therapy planning	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77300	Radiation therapy, dose plan	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77301	Radiotherapy dose plan, imrt	X	0310		13.7576	\$927.34	\$325.27	\$185.47
77305	Telex isodose plan simple	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77310	Telex isodose plan intermed	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77315	Telex isodose plan complex	X	0305		3.951	\$266.32	\$91.38	\$53.27
77321	Special telex port plan	X	0305		3.951	\$266.32	\$91.38	\$53.27
77326	Brachytx isodose calc simp	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77327	Brachytx isodose calc intern	X	0305		3.951	\$266.32	\$91.38	\$53.27
77328	Brachytx isodose plan compl	X	0305		3.951	\$266.32	\$91.38	\$53.27
77331	Special radiation dosimetry	X	0304		1.5271	\$102.94	\$34.71	\$20.59
77332	Radiation treatment aid(s)	X	0303		2.8279	\$190.62	\$66.95	\$38.13
77333	Radiation treatment aid(s)	X	0303		2.8279	\$190.62	\$66.95	\$38.13
77334	Radiation treatment aid(s)	X	0303		2.8279	\$190.62	\$66.95	\$38.13

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
78206	Liver image (3D) with flow	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78215	Liver and spleen imaging	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78216	Liver & spleen image/flow	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78220	Liver function study	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78223	Hepatobiliary imaging	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78230	Salivary gland imaging	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78231	Serial salivary imaging	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78232	Salivary gland function exam	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78258	Esophageal motility study	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78261	Gastric mucosa imaging	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78262	Gastroesophageal reflux exam	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78264	Gastric emptying study	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78267	Breath test attain/anal c-14	A	A					
78268	Breath test analysis, c-14	A	A					
78270	Vit B-12 absorption exam	S	S	0392	2.6817	\$180.76	\$43.95	\$36.16
78271	Vit B-12 absorp exam, int fac	S	S	0392	2.6817	\$180.76	\$43.95	\$36.16
78272	Vit B-12 absorp, combined	S	S	0392	2.6817	\$180.76	\$43.95	\$36.16
78278	Acute GI blood loss imaging	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78282	GI protein loss exam	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78290	Meckels divert exam	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78291	Lewent/shunt patency exam	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78299	GI nuclear procedure	S	S	0395	3.6702	\$247.39	\$89.73	\$49.48
78300	Bone imaging, limited area	S	S	0396	3.6707	\$247.43	\$95.02	\$49.49
78305	Bone imaging, multiple areas	S	S	0396	3.6707	\$247.43	\$95.02	\$49.49
78306	Bone imaging, whole body	S	S	0396	3.6707	\$247.43	\$95.02	\$49.49
78315	Bone imaging, 3 phase	S	S	0396	3.6707	\$247.43	\$95.02	\$49.49
78320	Bone imaging (3D)	S	S	0396	3.6707	\$247.43	\$95.02	\$49.49
78350	Bone mineral, dual photon	E	E					
78351	Bone mineral, single photon	E	E					
78399	Musculoskeletal nuclear exam	S	S	0396	3.6707	\$247.43	\$95.02	\$49.49
78414	Non-imaging heart function	S	S	0398	4.5366	\$305.79	\$97.94	\$61.16
78428	Cardiac shunt imaging	S	S	0398	4.5366	\$305.79	\$97.94	\$61.16
78445	Vascular flow imaging	S	S	0397	2.9679	\$200.05	\$46.29	\$40.01
78451	HI muscle image spect, sing	NI	NI	0377	11.4989	\$775.09	\$158.84	\$155.02
78452	HI muscle image spect, mult	NI	NI	0377	11.4989	\$775.09	\$158.84	\$155.02
78453	HI muscle image planar, sing	NI	NI	0377	11.4989	\$775.09	\$158.84	\$155.02
78454	HI muscle image, planar, mult	NI	NI	0377	11.4989	\$775.09	\$158.84	\$155.02
78456	Acute venous thrombus image	S	S	0397	2.9679	\$200.05	\$46.29	\$40.01
78457	Venous thrombosis imaging	S	S	0397	2.9679	\$200.05	\$46.29	\$40.01
78458	Ven thrombosis images, bilat	S	S	0397	2.9679	\$200.05	\$46.29	\$40.01
78459	Heart muscle imaging (PET)	CH	D	0307	21.2573	\$1,432.87		\$286.58
78460	Heart muscle blood, single	CH	D					
78461	Heart muscle blood, multiple	CH	D					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
77763	Apply intracv radiat compl	S	S	0312	4.4846	\$302.29	\$60.46	\$60.46
77776	Apply intersit radiat compl	S	S	0312	4.4846	\$302.29	\$60.46	\$60.46
77777	Apply intersit radiat inter	S	S	0312	4.4846	\$302.29	\$60.46	\$60.46
77778	Apply intersit radiat compl	O3	S	0651	13.2555	\$993.50	\$178.70	\$178.70
77785	Hdr brachytx, 1 channel	S	S	0313	11.5353	\$777.55	\$293.30	\$155.51
77786	Hdr brachytx, 2-12 channel	S	S	0313	11.5353	\$777.55	\$293.30	\$155.51
77787	Hdr brachytx over 12 chan	S	S	0313	11.5353	\$777.55	\$293.30	\$155.51
77789	Apply surface radiation	S	S	0300	1.3764	\$92.78	\$18.56	\$18.56
77790	Radiation handling	N	N					
77799	Radium/radioisotope therapy	S	S	0312	4.4846	\$302.29	\$60.46	\$60.46
78001	Thyroid, single uptake	S	S	0389	1.6695	\$112.53	\$30.02	\$22.51
78003	Thyroid, multiple uptakes	S	S	0389	1.6695	\$112.53	\$30.02	\$22.51
78004	Thyroid suppress/stimul	CH	S	0389	1.6695	\$112.53	\$30.02	\$22.51
78006	Thyroid imaging with uptake	S	S	0391	3.2894	\$221.73	\$66.13	\$44.35
78007	Thyroid image, mult uptakes	S	S	0391	3.2894	\$221.73	\$66.13	\$44.35
78010	Thyroid imaging	S	S	0390	2.1595	\$145.56	\$52.15	\$29.12
78011	Thyroid imaging with flow	S	S	0390	2.1595	\$145.56	\$52.15	\$29.12
78015	Thyroid met imaging	S	S	0406	4.3034	\$290.07	\$88.22	\$58.02
78018	Thyroid met imaging/studies	S	S	0406	4.3034	\$290.07	\$88.22	\$58.02
78019	Thyroid met imaging, body	S	S	0406	4.3034	\$290.07	\$88.22	\$58.02
78020	Thyroid met uptake	N	N					
78070	Parathyroid nuclear imaging	S	S	0391	3.2894	\$221.73	\$66.13	\$44.35
78075	Adrenal nuclear imaging	S	S	0408	14.2874	\$963.06	\$192.62	\$192.62
78099	Endocrine nuclear procedure	S	S	0390	2.1595	\$145.56	\$52.15	\$29.12
78102	Bone marrow imaging, fld	S	S	0400	3.8342	\$258.45	\$91.75	\$51.69
78103	Bone marrow imaging, mult	S	S	0400	3.8342	\$258.45	\$91.75	\$51.69
78104	Bone marrow imaging, body	S	S	0400	3.8342	\$258.45	\$91.75	\$51.69
78110	Plasma volume, single	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78111	Plasma volume, multiple	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78120	Red cell mass, single	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78121	Red cell mass, multiple	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78122	Blood volume	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78130	Red cell survival study	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78135	Red cell survival kinetics	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78140	Red cell sequestration	S	S	0393	5.7873	\$390.10	\$79.97	\$78.02
78185	Spleen imaging	S	S	0400	3.8342	\$258.45	\$91.75	\$51.69
78190	Platelet survival, kinetics	S	S	0392	2.6817	\$180.76	\$43.95	\$36.16
78191	Platelet survival	S	S	0392	2.6817	\$180.76	\$43.95	\$36.16
78195	Lymph system imaging	S	S	0400	3.8342	\$258.45	\$91.75	\$51.69
78201	Blood/lymph nuclear exam	S	S	0400	3.8342	\$258.45	\$91.75	\$51.69
78202	Liver imaging	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78205	Liver imaging with flow	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16
78205	Liver imaging (3D)	S	S	0394	4.3139	\$290.78	\$99.32	\$58.16

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
78701	Kidney imaging with flow	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78707	K flow/func image w/o drug	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78708	K flow/func image w/drug	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78709	K flow/func image, multiple	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78710	Kidney imaging (3D)	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78725	Kidney function study	S	0392	2.6817	\$180.76	\$43.95	\$36.16	
78730	Urinary bladder retention	S	0389	1.6695	\$112.53	\$30.02	\$22.51	
78740	Ureteral reflux study	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78761	Testicular imaging w/flow	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78799	Genitourinary nuclear exam	S	0404	4.8221	\$325.04	\$83.10	\$65.01	
78800	Tumor imaging, limited area	S	0406	4.3034	\$290.07	\$88.22	\$58.02	
78801	Tumor imaging, mult areas	S	0414	7.5722	\$510.41		\$102.09	
78802	Tumor imaging, whole body	S	0414	7.5722	\$510.41		\$102.09	
78803	Tumor imaging (3D)	CH	S	0414	7.5722	\$510.41	\$102.09	
78804	Tumor imaging, whole body	S	0408	14.2874	\$963.06		\$192.62	
78805	Abscess imaging, lid area	S	0308	15.3895	\$1,037.34		\$207.47	
78806	Abscess imaging, whole body	S	0414	7.5722	\$510.41		\$102.09	
78807	Nuclear localization/abscess	CH	S	0406	4.3034	\$290.07	\$58.02	
78808	iv/in ra drug bx study	QT	0392	2.6817	\$180.76	\$43.95	\$36.16	
78811	Pet image, lid area	S	0308	15.3895	\$1,037.34		\$207.47	
78812	Pet image, skull-thigh	S	0308	15.3895	\$1,037.34		\$207.47	
78813	Pet image, full body	S	0308	15.3895	\$1,037.34		\$207.47	
78814	Pet image w/ct, limd	S	0308	15.3895	\$1,037.34		\$207.47	
78815	Pet image w/ct, skull-thigh	S	0308	15.3895	\$1,037.34		\$207.47	
78816	Pet image w/ct, full body	S	0308	15.3895	\$1,037.34		\$207.47	
78899	Nuclear diagnostic exam	S	0389	1.6695	\$112.53	\$30.02	\$22.51	
79005	Nuclear rx, oral admin	S	0407	3.2356	\$218.10	\$78.13	\$43.62	
79101	Nuclear rx, iv admin	S	0407	3.2356	\$218.10	\$78.13	\$43.62	
79200	Nuclear rx, intracav admin	S	0413	5.2007	\$350.56	\$78.13	\$43.62	
79300	Nucir rx, intersit colloid	S	0407	3.2356	\$218.10	\$78.13	\$43.62	
79403	Hematopoietic nuclear tx	S	0413	5.2007	\$350.56	\$78.13	\$43.62	
79445	Nuclear rx, intra-arterial	S	0407	3.2356	\$218.10	\$78.13	\$43.62	
79999	Nuclear medicine therapy	S	0407	3.2356	\$218.10	\$78.13	\$43.62	
80047	Metabolic panel ionized ca	A						
80048	Metabolic panel total ca	A						
80050	General health panel	E						
80051	Electrolyte panel	A						
80053	Comprehens metaboic panel	A						
80055	Obstetric panel	E						
80061	Lipid panel	A						
80069	Renal function panel	A						
80074	Acute hepatitis panel	A						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
78464	Heart image (3d), single	CH	D					
78465	Heart image (3d), multiple	CH	D					
78466	Heart infarct image	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78468	Heart infarct image (ef)	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78469	Heart infarct image (3D)	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78472	Gated heart, planar, single	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78473	Gated heart, multiple	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78478	Heart wall motion add-on	CH	D					
78480	Heart function add-on	CH	D					
78481	Heart first pass, single	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78483	Heart first pass, multiple	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78491	Heart image (pet), single	S	0307	21.2573	\$1,432.87	\$286.58	\$286.58	
78492	Heart image (pet), multiple	S	0307	21.2573	\$1,432.87	\$286.58	\$286.58	
78494	Heart image, spect	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78496	Heart first pass add-on	N						
78499	Cardiovascular nuclear exam	S	0398	4.5366	\$305.79	\$97.94	\$61.16	
78580	Lung perfusion imaging	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78584	Lung V/Q image single breath	S	0378	4.8032	\$323.76	\$124.94	\$64.76	
78585	Lung V/Q imaging	S	0378	4.8032	\$323.76	\$124.94	\$64.76	
78586	Aerosol lung image, single	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78587	Aerosol lung image, multiple	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78588	Perfusion lung image	S	0378	4.8032	\$323.76	\$124.94	\$64.76	
78591	Vent image, 1 breath, 1 proj	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78593	Vent image, 1 proj, gas	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78594	Vent image, mult proj, gas	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78596	Lung differential function	S	0378	4.8032	\$323.76	\$124.94	\$64.76	
78599	Respiratory nuclear exam	S	0401	3.0625	\$206.43	\$74.63	\$41.29	
78600	Brain image < 4 views	S	0403	2.9181	\$196.70	\$72.42	\$39.34	
78601	Brain image w/flow < 4 views	CH	S	0402	8.5845	\$578.65	\$115.73	
78605	Brain image 4+ views	S	0403	2.9181	\$196.70	\$72.42	\$39.34	
78606	Brain image w/flow 4+ views	S	0402	8.5845	\$578.65	\$115.73	\$115.73	
78607	Brain imaging (3D)	S	0402	8.5845	\$578.65	\$115.73	\$115.73	
78608	Brain imaging (PET)	S	0308	15.3895	\$1,037.34	\$207.47	\$207.47	
78609	Brain imaging (PET)	E						
78610	Brain flow imaging only	CH	S	0403	2.9181	\$196.70	\$72.42	\$39.34
78630	Cerebrospinal fluid scan	S	0402	8.5845	\$578.65	\$115.73	\$115.73	
78635	CSF ventriculography	S	0402	8.5845	\$578.65	\$115.73	\$115.73	
78645	CSF shunt evaluation	S	0403	2.9181	\$196.70	\$72.42	\$39.34	
78647	Cerebrospinal fluid scan	S	0402	8.5845	\$578.65	\$115.73	\$115.73	
78650	CSF leakage imaging	S	0402	8.5845	\$578.65	\$115.73	\$115.73	
78660	Nuclear exam of tear flow	S	0403	2.9181	\$196.70	\$72.42	\$39.34	
78699	Nervous system nuclear exam	S	0403	2.9181	\$196.70	\$72.42	\$39.34	
78700	Kidney imaging, morphol	S	0404	4.8221	\$325.04	\$83.10	\$65.01	

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
80412	CRH stimulation panel	A	A					
80414	Testosterone response	A	A					
80415	Estradiol response panel	A	A					
80416	Renin stimulation panel	A	A					
80417	Renin stimulation panel	A	A					
80418	Pituitary evaluation panel	A	A					
80420	Dexamethasone panel	A	A					
80422	Glucagon tolerance panel	A	A					
80424	Glucagon tolerance panel	A	A					
80426	Gonadotropin hormone panel	A	A					
80428	Growth hormone panel	A	A					
80430	Growth hormone panel	A	A					
80432	Insulin suppression panel	A	A					
80434	Insulin tolerance panel	A	A					
80435	Insulin tolerance panel	A	A					
80436	Metyrapone panel	A	A					
80438	TRH stimulation panel	A	A					
80439	TRH stimulation panel	A	A					
80440	TRH stimulation panel	A	A					
80500	Lab pathology consultation	X	0433		0.2482	\$16.73	\$5.17	\$3.35
80502	Lab pathology consultation	X	0342		0.1546	\$10.42		\$2.09
81000	Urinalysis, nonauto w/scope	A	A					
81001	Urinalysis, auto w/scope	A	A					
81002	Urinalysis nonauto w/o scope	A	A					
81003	Urinalysis, auto, w/o scope	A	A					
81005	Urinalysis	A	A					
81007	Urine screen for bacteria	A	A					
81015	Microscopic exam of urine	A	A					
81020	Urinalysis, glass test	A	A					
81025	Urine pregnancy test	A	A					
81099	Urinalysis, volume measure	A	A					
82000	Urinalysis test procedure	A	A					
82003	Assay of blood acetaldehyde	A	A					
82009	Assay of acetaminophen	A	A					
82010	Test for acetone/ketones	A	A					
82013	Acetone assay	A	A					
82016	Acetylcholinesterase assay	A	A					
82017	Acylcarbitrines, qual	A	A					
82024	Acylcarbitrines, quant	A	A					
82030	Assay of acth	A	A					
82040	Assay of adp & amp	A	A					
82042	Assay of serum albumin	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
80076	Hepatic function panel	A	A					
80100	Drug screen, qualitative/multi	CH	E					
80101	Drug screen, single	CH	E					
80102	Drug confirmation							
80103	Drug analysis, tissue prep	N	N					
80150	Assay of amikacin	A	A					
80152	Assay of amitritypyline	A	A					
80154	Assay of benzodiazepines	A	A					
80156	Assay, carbamazepine, total	A	A					
80157	Assay, carbamazepine, free	A	A					
80158	Assay of cyclosporine	A	A					
80160	Assay of desipramine	A	A					
80162	Assay of digoxin	A	A					
80164	Assay, dipropylacetic acid	A	A					
80166	Assay of doxepin	A	A					
80168	Assay of ethosuximide	A	A					
80170	Assay of gentamicin	A	A					
80172	Assay of gold	A	A					
80173	Assay of haloperidol	A	A					
80174	Assay of imipramine	A	A					
80176	Assay of lidocaine	A	A					
80178	Assay of lithium	A	A					
80182	Assay of nortriptyline	A	A					
80184	Assay of phenobarbital	A	A					
80185	Assay of phenytoin, total	A	A					
80186	Assay of phenytoin, free	A	A					
80188	Assay of primateone	A	A					
80190	Assay of procainamide	A	A					
80192	Assay of procainamide	A	A					
80194	Assay of quinidine	A	A					
80195	Assay of salicylate	A	A					
80196	Assay of sirolimus	A	A					
80197	Assay of tacrolimus	A	A					
80198	Assay of theophylline	A	A					
80200	Assay of tobramycin	A	A					
80201	Assay of topramate	A	A					
80202	Assay of vancomycin	A	A					
80299	Quantitative assay, drug	A	A					
80400	Acth stimulation panel	A	A					
80402	Acth stimulation panel	A	A					
80406	Acth stimulation panel	A	A					
80408	Aldosterone suppression eval	A	A					
80410	Calcitonin stimul panel	A	A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
82271	Occult blood, other sources	A	A					
82272	Occult bid feces, 1-3 tests	A	A					
82274	Assay bid feces, fecal	A	A					
82286	Assay of bradykinin	A	A					
82300	Assay of cadmium	A	A					
82306	Vitamin d, 25 hydroxy	CH	D					
82307	Assay of vitamin D							
82308	Assay of calcitonin	A	A					
82310	Assay of calcium	A	A					
82330	Assay of calcium	A	A					
82331	Calcium infusion test	A	A					
82340	Assay of calcium in urine	A	A					
82355	Calculus analysis, qual	A	A					
82360	Calculus assay, quant	A	A					
82365	Calculus spectroscopy	A	A					
82370	X-ray assay, calculus	A	A					
82373	Assay, c-d transfer measure	A	A					
82374	Assay, blood carbon dioxide	A	A					
82375	Assay, carboxyhb, quant	A	A					
82376	Assay, carboxyhb, qual	A	A					
82378	Carcinoembryonic antigen	A	A					
82379	Assay of carnitine	A	A					
82380	Assay of cardene	A	A					
82382	Assay, urine catecholamines	A	A					
82383	Assay, blood catecholamines	A	A					
82384	Assay, three catecholamines	A	A					
82387	Assay of cathepsin-d	A	A					
82390	Assay of ceruloplasmin	A	A					
82397	Chemiluminescent assay	A	A					
82415	Assay of chloramphenicol	A	A					
82435	Assay of blood chloride	A	A					
82436	Assay, other fluid chloride	A	A					
82438	Assay, urine chloride	A	A					
82441	Test for chlorohydrocarbons	A	A					
82465	Assay, bid/serum cholesterol	A	A					
82480	Assay, serum cholinesterase	A	A					
82482	Assay, rbc cholinesterase	A	A					
82485	Assay, chondroitin sulfate	A	A					
82486	Gas/liquid chromatography	A	A					
82487	Paper chromatography	A	A					
82488	Paper chromatography	A	A					
82489	Thin layer chromatography	A	A					
82491	Chromatography, quant, sing	A	A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
82043	Microalbumin, quantitative	A	A					
82044	Microalbumin, semiquant	A	A					
82045	Albumin, ischemia modified	A	A					
82055	Assay of ethanol	A	A					
82075	Assay of breath ethanol	A	A					
82085	Assay of aldolase	A	A					
82088	Assay of aldosterone	A	A					
82101	Assay of urine alkaloids	A	A					
82103	Alpha-1-antitrypsin, total	A	A					
82104	Alpha-1-antitrypsin, pheno	A	A					
82105	Alpha-fetoprotein, serum	A	A					
82106	Alpha-fetoprotein, amniotic	A	A					
82107	Alpha-fetoprotein I3	A	A					
82108	Assay of aluminum	A	A					
82120	Amines, vaginal fluid qual	A	A					
82127	Amino acid, single qual	A	A					
82128	Amino acids, mult qual	A	A					
82131	Amino acids, single quant	A	A					
82135	Assay, aminolevulinic acid	A	A					
82136	Amino acids, quant, 2-5	A	A					
82139	Amino acids, quant, 6 or more	A	A					
82140	Assay of ammonia	A	A					
82143	Amniotic fluid scan	A	A					
82145	Assay of amphetamines	A	A					
82150	Assay of amylase	A	A					
82154	Androstenediol glucuronide	A	A					
82157	Assay of androstenedione	A	A					
82160	Assay of androstereone	A	A					
82163	Assay of angiotensin II	A	A					
82164	Angiotensin I enzyme test	A	A					
82172	Assay of apolipoprotein	A	A					
82175	Assay of arsenic	A	A					
82180	Assay of ascorbic acid	A	A					
82190	Atomic absorption	A	A					
82205	Assay of barbiturates	A	A					
82232	Assay of beta-2 protein	A	A					
82239	Bile acids, total	A	A					
82240	Bile acids, cholyglycine	A	A					
82247	Bilirubin, total	A	A					
82248	Bilirubin, direct	A	A					
82252	Fecal bilirubin test	A	A					
82261	Assay of biotinidase	A	A					
82270	Occult blood, feces	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010									
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment	
82668	Assay of erythropoietin	A	A						
82670	Assay of estradiol	A	A						
82671	Assay of estrogens	A	A						
82672	Assay of estrin	A	A						
82677	Assay of estrin	A	A						
82679	Assay of estrone	A	A						
82690	Assay of ethionynol	A	A						
82693	Assay of ethylene glycol	A	A						
82696	Assay of etiocholanolone	A	A						
82705	Fats/lipids, feces, qual	A	A						
82710	Fats/lipids, feces, quant	A	A						
82715	Assay of fecal fat	A	A						
82725	Assay of blood fatty acids	A	A						
82726	Long chain fatty acids	A	A						
82728	Assay of ferritin	A	A						
82731	Assay of fetal fibronectin	A	A						
82735	Assay of fluoride	A	A						
82742	Assay of flurazepam	A	A						
82746	Blood folic acid serum	A	A						
82747	Assay of folic acid, rbc	A	A						
82757	Assay of semen fructose	A	A						
82759	Assay of rbc galactokinase	A	A						
82760	Assay of galactose	A	A						
82775	Assay galactose transferase	A	A						
82776	Galactose transferase test	A	A						
82784	Assay, igel/ig/ig/igm each	A	A						
82785	Assay of lge	A	A						
82787	Igg 1, 2, 3 or 4, each	A	A						
82800	Blood pH	A	A						
82803	Blood gases: pH, pO2 & pCO2	A	A						
82805	Blood gases w/O2 saturation	A	A						
82810	Blood gases, O2 sat only	A	A						
82820	Hemoglobin-oxygen affinity	A	A						
82926	Assay of gastric acid	A	A						
82928	Assay of gastric acid	A	A						
82938	Gastrin test	A	A						
82941	Assay of gastrin	A	A						
82943	Assay of glucagon	A	A						
82945	Glucose other fluid	A	A						
82946	Glucagon tolerance test	A	A						
82947	Assay, glucose, blood quant	A	A						
82948	Reagent strip/blood glucose	A	A						
82950	Glucose test	A	A						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010									
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment	
82492	Chromatography, quant, mult	A	A						
82495	Assay of chromium	A	A						
82507	Assay of citrate	A	A						
82520	Assay of cocaine	A	A						
82523	Collagen crosslinks	A	A						
82525	Assay of copper	A	A						
82528	Assay of corticosterone	A	A						
82530	Cortisol, free	A	A						
82533	Total cortisol	A	A						
82540	Assay of creatine	A	A						
82541	Column chromatography, qual	A	A						
	Column chromatography, quant	A	A						
82542	Column chromatography/isotope	A	A						
82543	Column chromatography/isotope	A	A						
82544	Column chromatography/isotope	A	A						
82550	Assay of ck (cpk)	A	A						
82552	Assay of cpk in blood	A	A						
82553	Creatine, MB fraction	A	A						
82554	Creatine, isoforms	A	A						
82565	Assay of creatinine	A	A						
82570	Assay of urine creatinine	A	A						
82575	Creatinine clearance test	A	A						
82585	Assay of cryofibrinogen	A	A						
82595	Assay of cryoglobulin	A	A						
82600	Assay of cyanide	A	A						
82607	Vitamin B-12	A	A						
82608	B-12 binding capacity	A	A						
82610	Cystatin c	A	A						
82615	Test for urine cystines	A	A						
82626	Dehydroepiandrosterone	A	A						
82627	Dehydroepiandrosterone	A	A						
82633	Deoxycorticosterone	A	A						
82634	Deoxycortisol	A	A						
82638	Assay of dibucaine number	A	A						
82646	Assay of dihydrocodeinone	A	A						
82649	Assay of dihydromorphine	A	A						
82651	Assay of dihydrotestosterone	A	A						
82652	Vit d 1, 25-dihydroxy	A	A						
82654	Assay of dimethadione	A	A						
82656	Pancreatic elastase, fecal	A	A						
82657	Enzyme cell activity	A	A						
82658	Enzyme cell activity, ra	A	A						
82664	Electrophoretic test	A	A						
82666	Assay of epiandrosterone	A	A						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
83088	Assay of histamine		A					
83090	Assay of homocysteine		A					
83150	Assay of for hva		A					
83491	Assay of corticosteroids		A					
83497	Assay of 5-hiaa		A					
83498	Assay of progesterone		A					
83499	Assay of progesterone		A					
83500	Assay, free hydroxyproline		A					
83505	Assay, total hydroxyproline		A					
83516	Immunoassay, nonantibody		A					
83518	Immunoassay, dipstick		A					
83520	Immunoassay, quant, nos nonab		A					
83525	Assay of insulin		A					
83527	Assay of insulin		A					
83528	Assay of intrinsic factor		A					
83540	Assay of iron		A					
83550	Iron binding test		A					
83570	Assay of idh enzyme		A					
83582	Assay of ketogenic steroids		A					
83586	Assay 17- ketosteroids		A					
83593	Fractionation, ketosteroids		A					
83605	Assay of lactic acid		A					
83615	Lactate (LD) (LDH) enzyme		A					
83625	Assay of ldh enzymes		A					
83630	Lactoferrin, fecal (quat)		A					
83631	Lactoferrin, fecal (quant)		A					
83632	Placental lactogen		A					
83633	Test urine for lactose		A					
83634	Assay of urine for lactose		A					
83655	Assay of lead		A					
83661	L/s ratio, fetal lung		A					
83662	Foam stability, fetal lung		A					
83663	Fluoro polarize, fetal lung		A					
83664	Lamellar bdy, fetal lung		A					
83670	Assay of lap enzyme		A					
83690	Assay of lipase		A					
83695	Assay of lipoprotein(a)		A					
83698	Assay lipoprotein pla2		A					
83700	Lipopro bid, electrophoretic		A					
83701	Lipoprotein bid, hr fraction		A					
83704	Lipoprotein, bid, by nmr		A					
83718	Assay of lipoprotein		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
82951	Glucose tolerance test (GTT)		A					
82952	GTT-added samples		A					
82953	Glucose-tolbutamide test		A					
82955	Assay of g6pd enzyme		A					
82960	Test for G6PD enzyme		A					
82962	Glucose blood test		A					
82963	Assay of glucosidase		A					
82965	Assay of gdh enzyme		A					
82975	Assay of glutamine		A					
82977	Assay of GGT		A					
82978	Assay of glutathione		A					
82979	Assay, roc, glutathione		A					
82980	Assay of glutethimide		A					
82985	Glycated protein		A					
83001	Gonadotropin (FSH)		A					
83002	Gonadotropin (LH)		A					
83003	Assay, growth hormone (hgh)		A					
83008	Assay of guanosine		A					
83009	H pylori (c-13), blood		A					
83010	Assay of haptoglobin, quant		A					
83012	Assay of haptoglobins		A					
83013	H pylori (c-13), breath		A					
83014	H pylori drug admin		A					
83015	Heavy metal screen		A					
83018	Quantitative screen, metals		A					
83020	Hemoglobin electrophoresis		A					
83021	Hemoglobin chromatography		A					
83026	Hemoglobin, copper sulfate		A					
83030	Fetal hemoglobin, chemical		A					
83033	Fetal hemoglobin assay, qual		A					
83036	Glycosylated hemoglobin test		A					
83037	Glycosylated hb, home device		A					
83045	Blood methemoglobin test		A					
83050	Blood methemoglobin assay		A					
83051	Assay of plasma hemoglobin		A					
83055	Blood sulfhemoglobin test		A					
83060	Blood sulfhemoglobin assay		A					
83065	Assay of hemoglobin heat		A					
83068	Hemoglobin stability screen		A					
83069	Assay of urine hemoglobin		A					
83070	Assay of hemosiderin, qual		A					
83071	Assay of hemosiderin, quant		A					
83080	Assay of b hexosaminidase		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
83913	Molecular, rna stabilization	A	A					
83914	Mutation ident oia/sboe/aspe	A	A					
83915	Assay of nucleic acid	A	A					
83916	Oligoclonal bands	A	A					
83918	Organic acids, total, quant	A	A					
83919	Organic acids, qual, each	A	A					
83921	Organic acid, single, quant	A	A					
83925	Assay of opiates	A	A					
83930	Assay of blood osmolality	A	A					
83935	Assay of urine osmolality	A	A					
83937	Assay of osteocalcin	A	A					
83945	Assay of oxalate	A	A					
83950	Oncoprotein, her-2/neu	A	A					
83951	Oncoprotein, dcp	A	A					
83970	Assay of parathormone	A	A					
83986	Assay ph body fluid nos	A	A					
83987	Exhaled breath condensate	NI	A					
83992	Assay for phenacyclidine	A	A					
83993	Assay for calprotectin fecal	A	A					
84022	Assay of phenothiazine	A	A					
84030	Assay of blood pku	A	A					
84035	Assay of phenylketones	A	A					
84060	Assay acid phosphatase	A	A					
84061	Phosphatase, forensic exam	A	A					
84066	Assay prostate phosphatase	A	A					
84075	Assay alkaline phosphatase	A	A					
84078	Assay alkaline phosphatase	A	A					
84080	Assay alkaline phosphatases	A	A					
84081	Amniotic fluid enzyme test	A	A					
84085	Assay of rbc po6d enzyme	A	A					
84087	Assay phosphohexose enzymes	A	A					
84100	Assay of phosphorus	A	A					
84105	Assay of urine phosphorus	A	A					
84106	Test for porphobilinogen	A	A					
84110	Assay of porphobilinogen	A	A					
84119	Test urine for porphyrins	A	A					
84120	Assay of urine porphyrins	A	A					
84126	Assay of feces porphyrins	A	A					
84127	Assay of feces porphyrins	A	A					
84132	Assay of serum potassium	A	A					
84133	Assay of urine potassium	A	A					
84134	Assay of prealbumin	A	A					
84135	Assay of pregnanediol	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
83719	Assay of blood lipoprotein	A	A					
83721	Assay of blood lipoprotein	A	A					
83727	Assay of lth hormone	A	A					
83735	Assay of magnesium	A	A					
83775	Assay of md enzyme	A	A					
83785	Assay of manganese	A	A					
83788	Mass spectrometry qual	A	A					
83789	Mass spectrometry quart	A	A					
83805	Assay of meprobamate	A	A					
83825	Assay of mercury	A	A					
83835	Assay of metanephrites	A	A					
83840	Assay of methadone	A	A					
83857	Assay of methalbumin	A	A					
83858	Assay of methaximide	A	A					
83864	Micopolysaccharides	A	A					
83866	Micopolysaccharides screen	A	A					
83872	Assay synovial fluid mucin	A	A					
83873	Assay of csf protein	A	A					
83874	Assay of myoglobin	A	A					
83876	Assay, myeloperoxidase	A	A					
83880	Natriuretic peptide	A	A					
83883	Assay, nephelometry not spec	A	A					
83885	Assay of nickel	A	A					
83887	Assay of nicotine	A	A					
83890	Molecule isolate	A	A					
83891	Molecule isolate nucleic	A	A					
83893	Molecular diagnostics	A	A					
83894	Molecule qof/sic/biot	A	A					
83894	Molecule gel electrophor	A	A					
83896	Molecular diagnostics	A	A					
83897	Molecule nucleic transfer	A	A					
83898	Molecule nucleic ampli, each	A	A					
83900	Molecule nucleic ampli 2 seq	A	A					
83901	Molecule nucleic ampli addon	A	A					
83902	Molecular diagnostics	A	A					
83903	Molecule mutation scan	A	A					
83904	Molecule mutation identify	A	A					
83905	Molecule mutation identify	A	A					
83906	Molecule mutation identify	A	A					
83907	Lyse cells for nucleic ext	A	A					
83908	Nucleic acid, signal ampli	A	A					
83909	Nucleic acid, high resolute	A	A					
83912	Genetic examination	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
84315	Body fluid specific gravity		A					
84375	Chromatogram assay, sugars		A					
84376	Sugars, single, qual		A					
84377	Sugars, multiple, qual		A					
84378	Sugars, single, quant		A					
84379	Sugars multiple quant		A					
84392	Assay of urine sulfate		A					
84402	Assay of testosterone		A					
84403	Assay of total testosterone		A					
84425	Assay of vitamin b-1		A					
84430	Assay of thiocyanate		A					
84431	Thromboxane, urine	NI	A					
84432	Assay of thyroglobulin		A					
84436	Assay of total thyroxine		A					
84437	Assay of neonatal thyroxine		A					
84439	Assay of free thyroxine		A					
84442	Assay of thyroid activity		A					
84443	Assay thyroid stim hormone		A					
84445	Assay of tsi		A					
84446	Assay of vitamin e		A					
84449	Assay of transcorin		A					
84450	Transferase (AST) (SGOT)		A					
84460	Alanine amino (ALT) (SGPT)		A					
84466	Assay of transferrin		A					
84478	Assay of triglycerides		A					
84479	Assay of thyroid (t3 or t4)		A					
84480	Assay, triiodothyronine (t3)		A					
84481	Free assay (FT-3)		A					
84482	T3 reverse		A					
84484	Assay of troponin, quant		A					
84485	Assay duodenal fluid trypsin		A					
84488	Test feces for trypsin		A					
84490	Assay of feces for trypsin		A					
84510	Assay of tyrosine		A					
84512	Assay of troponin, qual		A					
84520	Assay of urea nitrogen		A					
84525	Urea nitrogen semi-quant		A					
84540	Assay of urine/urea-n		A					
84545	Urea-N clearance test		A					
84550	Assay of blood/lactic acid		A					
84560	Assay of urine/lactic acid		A					
84577	Assay of feces/urobilinogen		A					
84578	Test urine urobilinogen		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
84138	Assay of pregnanetriol		A					
84140	Assay of pregnenolone		A					
84143	Assay of 17-hydroxypregnenone		A					
84144	Assay of progesterone		A					
84145	Procalcitonin (pct)	NI	A					
84146	Assay of prolactin		A					
84150	Assay of prostaglandin		A					
84152	Assay of psa, complexed		A					
84153	Assay of psa, total		A					
84154	Assay of psa, free		A					
84155	Assay of protein, serum		A					
84156	Assay of protein, urine		A					
84157	Assay of protein, other		A					
84160	Assay of protein, any source		A					
84163	Pappa, serum		A					
84165	Protein e-phoresis, serum		A					
84166	Protein e-phoresis/urine/csf		A					
84181	Western blot test		A					
84182	Protein, western blot test		A					
84202	Assay HBC protoporphyrin		A					
84203	Test RBC protoporphyrin		A					
84206	Assay of proinsulin		A					
84207	Assay of vitamin b-6		A					
84210	Assay of pyruvate		A					
84220	Assay of pyruvate kinase		A					
84228	Assay of quinine		A					
84233	Assay of estrogen		A					
84234	Assay of progesterone		A					
84235	Assay of endocrine hormone		A					
84238	Assay, nonendocrine receptor		A					
84244	Assay of renin		A					
84252	Assay of vitamin b-2		A					
84255	Assay of selenium		A					
84260	Assay of serotonin		A					
84270	Assay of sex hormone globul		A					
84275	Assay of sialic acid		A					
84285	Assay of silica		A					
84295	Assay of serum sodium		A					
84300	Assay of urine sodium		A					
84302	Assay of sweat sodium		A					
84305	Assay of somatomedin		A					
84307	Assay of somatostatin		A					
84311	Spectrophotometry		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
85240	Blood clot factor VIII test		A					
85244	Blood clot factor VIII test		A					
85245	Blood clot factor VIII test		A					
85246	Blood clot factor VIII test		A					
85247	Blood clot factor VIII test		A					
85250	Blood clot factor IX test		A					
85260	Blood clot factor X test		A					
85270	Blood clot factor XI test		A					
85280	Blood clot factor XII test		A					
85290	Blood clot factor XIII test		A					
85291	Blood clot factor XIII test		A					
85292	Blood clot factor assay		A					
85293	Blood clot factor assay		A					
85300	Antithrombin III test		A					
85301	Antithrombin III test		A					
85302	Blood clot inhibitor antigen		A					
85303	Blood clot inhibitor test		A					
85305	Blood clot inhibitor assay		A					
85306	Blood clot inhibitor test		A					
85307	Assay activated protein c		A					
85335	Factor inhibitor test		A					
85337	Thrombomodulin		A					
85345	Coagulation time		A					
85347	Coagulation time		A					
85348	Coagulation time		A					
85360	Euglobulin lysis		A					
85362	Fibrin degradation products		A					
85366	Fibrinogen test		A					
85370	Fibrinogen test		A					
85378	Fibrin degrade semiquant		A					
85379	Fibrin degradation, quant		A					
85380	Fibrin degradation, vte		A					
85384	Fibrinogen		A					
85385	Fibrinogen		A					
85390	Fibrinolysis screen		A					
85396	Clotting assay, whole blood		N					
85397	Clotting funct activity		A					
85400	Fibrinolytic plasmin		A					
85410	Fibrinolytic aminplasmin		A					
85415	Fibrinolytic plasminogen		A					
85420	Fibrinolytic plasminogen		A					
85421	Fibrinolytic plasminogen		A					
85441	Heinz bodies, direct		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
84580	Assay of urine urobilinogen		A					
84583	Assay of urine urobilinogen		A					
84585	Assay of urine vma		A					
84586	Assay of vip		A					
84588	Assay of vasopressin		A					
84590	Assay of vitamin a		A					
84591	Assay of nos vitamin		A					
84597	Assay of vitamin k		A					
84600	Assay of volatiles		A					
84620	Xylose tolerance test		A					
84630	Assay of zinc		A					
84681	Assay of c-peptide		A					
84702	Chorionic gonadotropin test		A					
84703	Chorionic gonadotropin assay		A					
84704	Hcg, free beta chain test		A					
84830	Ovulation tests		A					
84999	Clinical chemistry test		A					
85002	Bleeding time test		A					
85004	Automated diff wbc count		A					
85007	Bi smear w/diff wbc count		A					
85008	Bi smear w/o diff wbc count		A					
85009	Manual diff wbc count b-coat		A					
85013	Spun microhematocrit		A					
85014	Hematocrit		A					
85018	Hemoglobin		A					
85025	Complete cbc w/auto diff wbc		A					
85027	Complete cbc, automated		A					
85032	Manual cell count, each		A					
85041	Automated fbc count		A					
85044	Manual reticulocyte count		A					
85045	Automated reticulocyte count		A					
85046	Retiocyte/hgb concentrate		A					
85048	Automated leukocyte count		A					
85049	Automated platelet count		A					
85055	Reticulated platelet assay		A					
85060	Blood smear interpretation		B					
85097	Bone marrow interpretation		X	0343	0.5301	\$35.73	\$10.84	\$7.15
85130	Chromogenic substrate assay		A					
85170	Blood clot retraction		A					
85175	Blood clot lysis time		A					
85210	Blood clot factor II test		A					
85220	Blood clot factor V test		A					
85230	Blood clot factor VII test		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
86079	Physician blood bank service	X	X	0433	0.2482	\$16.73	\$5.17	\$3.35
86140	C-reactive protein		A					
86141	C-reactive protein, hs		A					
86146	Glycoprotein antibody		A					
86147	Cardiolipin antibody		A					
86148	Phospholipid antibody		A					
86155	Chemotaxis assay		A					
86156	Iron stain peripheral blood		A					
86157	Wbc alkaline phosphatase		A					
86160	RBC mechanical fragility		A					
86161	Muramidase		A					
86162	RBC osmotic fragility		A					
86162	RBC osmotic fragility		A					
86171	Blood platelet aggregation		A					
86185	Platelet neutralization		A					
86200	Prothrombin time		A					
86215	Prothrombin test		A					
86225	Viper venom prothrombin time		A					
86226	Russell viper venom, diluted		A					
86235	Reptilase test		A					
86243	Rbc sed rate, nonautomated		A					
86255	Rbc sed rate, automated		A					
86256	RBC sickle cell test		A					
86277	Thrombin time, plasma		A					
86294	Thrombin time, titer		A					
86300	Thromboplasmin inhibition		A					
86300	Thromboplasmin time, partial		A					
86304	Thromboplasmin time, partial		A					
86305	Blood viscosity examination		A					
86308	Hematology procedure		A					
86309	Agglutinins, febrile		A					
86310	Allergen specific IgG		A					
86316	Allergen specific IgE		A					
86317	Allergen specific IgE		A					
86318	Platelet antibodies		A					
86320	Platelet antibodies		A					
86325	Immunoglobulin assay		A					
86327	Antinuclear antibodies		A					
86329	Antinuclear antibodies (ANA)		A					
86331	Antistreptolysin o, titer		A					
86332	Antistreptolysin o, screen		A					
86334	Physician blood bank service	X	X	0433	0.2482	\$16.73	\$5.17	\$3.35
86335	Physician blood bank service	X	X	0943	0.5301	\$35.73	\$10.84	\$7.15

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
85445	Heinz bodies, induced		A					
85460	Hemoglobin, fetal		A					
85461	Hemoglobin, fetal		A					
85475	Hemolysin		A					
85520	Heparin assay		A					
85525	Heparin neutralization		A					
85530	Heparin-protamine tolerance		A					
85536	Iron stain peripheral blood		A					
85540	Wbc alkaline phosphatase		A					
85547	RBC mechanical fragility		A					
85549	Muramidase		A					
85555	RBC osmotic fragility		A					
85557	RBC osmotic fragility		A					
85576	Blood platelet aggregation		A					
85597	Platelet neutralization		A					
85610	Prothrombin time		A					
85611	Prothrombin test		A					
85612	Viper venom prothrombin time		A					
85613	Russell viper venom, diluted		A					
85635	Reptilase test		A					
85651	Rbc sed rate, nonautomated		A					
85652	Rbc sed rate, automated		A					
85660	RBC sickle cell test		A					
85670	Thrombin time, plasma		A					
85675	Thrombin time, titer		A					
85705	Thromboplasmin inhibition		A					
85730	Thromboplasmin time, partial		A					
85732	Thromboplasmin time, partial		A					
85810	Blood viscosity examination		A					
85999	Hematology procedure		A					
86000	Agglutinins, febrile		A					
86001	Allergen specific IgG		A					
86003	Allergen specific IgE		A					
86005	Allergen specific IgE		A					
86021	WBC antibody identification		A					
86022	Platelet antibodies		A					
86023	Immunoglobulin assay		A					
86038	Antinuclear antibodies		A					
86039	Antinuclear antibodies (ANA)		A					
86060	Antistreptolysin o, titer		A					
86063	Antistreptolysin o, screen		A					
86077	Physician blood bank service	X	X	0433	0.2482	\$16.73	\$5.17	\$3.35
86078	Physician blood bank service	X	X	0943	0.5301	\$35.73	\$10.84	\$7.15

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
86625	Campylobacter antibody		A					
86628	Candida antibody		A					
86631	Chlamydia antibody		A					
86632	Chlamydia igm antibody		A					
86635	Coccidioides antibody		A					
86638	Q fever antibody		A					
86641	Cryptococcus antibody		A					
86644	CMV antibody		A					
86645	CMV antibody, Igm		A					
86648	Diphtheria antibody		A					
86651	Encephalitis antibody		A					
86652	Encephalitis antibody		A					
86653	Encephalitis antibody		A					
86654	Encephalitis antibody		A					
86656	Enterovirus antibody		A					
86663	Epstein-barr antibody		A					
86664	Epstein-barr antibody		A					
86665	Epstein-barr antibody		A					
86666	Ehrlichia antibody		A					
86668	Francisella tularensis		A					
86671	Fungus antibody		A					
86674	Giardia lamblia antibody		A					
86677	Helicobacter pylori		A					
86682	Helminth antibody		A					
86684	Hemophilus influenza		A					
86687	Hiv-i antibody		A					
86689	Hiv-ii antibody		A					
86689	HTLV/HIV confirmatory test		A					
86692	Hepatitis, delta agent		A					
86694	Herpes simplex test		A					
86695	Herpes simplex test		A					
86696	Herpes simplex type 2		A					
86698	Histoplasma		A					
86701	HIV-1		A					
86702	HIV-2		A					
86703	HIV-1/HIV-2, single assay		A					
86704	Hep b core antibody, total		A					
86705	Hep b core antibody, igm		A					
86706	Hep b surface antibody		A					
86707	Hep be antibody		A					
86708	Hep a antibody, total		A					
86709	Hep a antibody, igm		A					
86710	Influenza virus antibody		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
86336	Inhibin A		A					
86337	Insulin antibodies		A					
86340	Intrinsic factor antibody		A					
86341	Islet cell antibody		A					
86343	Leukocyte histamine release		A					
86344	Leukocyte phagocytosis		A					
86352	Cell function assay w/strim	NI						
86353	Lymphocyte transformation		A					
86355	B cells, total count		A					
86356	Mononuclear cell antigen		A					
86357	Nk cells, total count		A					
86359	T cells, total count		A					
86360	T cell, absolute count/ratio		A					
86361	T cell, absolute count		A					
86367	Stem cells, total count		A					
86376	Microsomal antibody		A					
86378	Micralton inhibitory factor		A					
86382	Neutralization test, viral		A					
86384	Nitroblue tetrazolium dye		A					
86403	Particle agglutination test		A					
86406	Particle agglutination test		A					
86430	Rheumatoid factor test		A					
86431	Rheumatoid factor, quant		A					
86480	Tb test, cell immun measure		A					
86485	Skin test, candida		X	0341	0.0799	\$5.39	\$2.09	\$1.08
86486	Skin test, nos antigen		X	0341	0.0799	\$5.39	\$2.09	\$1.08
86490	Coccidioidomycosis skin test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
86510	Histoplasmosis skin test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
86580	TB intradermal test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
86590	Streptokinase, antibody		A					
86592	Syphilis test non-trep qual		A					
86593	Syphilis test non-trep quant		A					
86602	Antinomyces antibody		A					
86603	Adenovirus antibody		A					
86606	Aspergillus antibody		A					
86609	Bacterium antibody		A					
86611	Bartonella antibody		A					
86612	Blastomyces antibody		A					
86615	Bordetella antibody		A					
86617	Lyme disease antibody		A					
86618	Lyme disease antibody		A					
86619	Borrelia antibody		A					
86622	Brucella antibody		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
86821	Lymphocyte culture, mixed	A	A					
86822	Lymphocyte culture, primed	NI	A					
86825	Hla x-match, non-cytotoxic	NI	A					
86826	Hla x-match, non-cyt add-on	NI	A					
86849	Immunology procedure		A					
86850	RBC antibody screen	X	X	0345	0.2195	\$14.80		\$2.96
86860	RBC antibody elution	X	X	0346	0.3734	\$25.17		\$5.04
86870	RBC antibody identification	X	X	0345	0.2195	\$25.17		\$5.04
86880	Coombs test, direct	X	X	0409	0.1161	\$7.83	\$2.20	\$1.57
86885	Coombs test, indirect, qual	X	X	0409	0.1161	\$7.83	\$2.20	\$1.57
86886	Coombs test, indirect, titer	X	X	0409	0.1161	\$7.83	\$2.20	\$1.57
86890	Autologous blood process	X	X	0347	0.7384	\$49.77		\$9.96
86891	Autologous blood, op salvage	X	X	0345	0.2195	\$14.80		\$2.96
86900	Blood typing, ABO	X	X	0409	0.1161	\$7.83	\$2.20	\$1.57
86901	Blood typing, Rh (D)	X	X	0409	0.1161	\$7.83	\$2.20	\$1.57
86903	Blood typing, antigen screen	X	X	0345	0.2195	\$14.80		\$2.96
86904	Blood typing, patient serum	X	X	0345	0.2195	\$14.80		\$2.96
86905	Blood typing, RBC antigens	X	X	0345	0.2195	\$14.80		\$2.96
86906	Blood typing, Rh phenotype	X	X	0345	0.2195	\$14.80		\$2.96
86910	Blood typing, paternity test	E	E					
86920	Blood typing, antigen system	X	X	0345	0.2195	\$14.80		\$2.96
86921	Compatibility test, skin	X	X	0345	0.2195	\$14.80		\$2.96
86922	Compatibility test, incubate	X	X	0346	0.3734	\$25.17		\$5.04
86923	Compatibility test, antiglob	X	X	0345	0.2195	\$14.80		\$2.96
86927	Compatibility test, electric	X	X	0345	0.2195	\$14.80		\$2.96
86928	Plasma, fresh frozen	X	X	0345	0.2195	\$14.80		\$2.96
86930	Frozen blood prep	X	X	0347	0.7384	\$49.77		\$9.96
86931	Frozen blood thaw	X	X	0347	0.7384	\$49.77		\$9.96
86932	Frozen blood freeze/thaw	X	X	0347	0.7384	\$49.77		\$9.96
86940	Hemolysis/agglutinins, auto	A	A					
86941	Hemolysis/agglutinins	A	A					
86945	Blood product/irradiation	X	X	0345	0.2195	\$14.80		\$2.96
86950	Leukocyte transfusion	X	X	0345	0.2195	\$14.80		\$2.96
86960	Vol reduction of blood/prod	X	X	0346	0.3734	\$25.17		\$5.04
86970	RBC pretreatment	X	X	0345	0.2195	\$14.80		\$2.96
86971	RBC pretreatment	X	X	0345	0.2195	\$14.80		\$2.96
86972	RBC pretreatment	X	X	0345	0.2195	\$14.80		\$2.96
86975	RBC pretreatment, serum	X	X	0346	0.3734	\$25.17		\$5.04
86976	RBC pretreatment, serum	X	X	0345	0.2195	\$14.80		\$2.96
86977	RBC pretreatment, serum	X	X	0347	0.7384	\$49.77		\$9.96
86978	RBC pretreatment, serum	X	X	0346	0.3734	\$25.17		\$5.04
86985	Split blood or products	X	X	0345	0.2195	\$14.80		\$2.96

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
86713	Legionella antibody	A	A					
86717	Leishmania antibody	A	A					
86720	Leptospira antibody	A	A					
86723	Listeria monocytogenes ab	A	A					
86727	Lymph choriomeningitis ab	A	A					
86729	Lymph venereum antibody	A	A					
86732	Mucormycosis antibody	A	A					
86735	Mumps antibody	A	A					
86738	Mycoplasma antibody	A	A					
86741	Neisseria meningitidis	A	A					
86744	Nocardia antibody	A	A					
86747	Parvovirus antibody	A	A					
86750	Malaria antibody	A	A					
86753	Protozoa antibody nos	A	A					
86756	Respiratory virus antibody	A	A					
86757	Rickettsia antibody	A	A					
86759	Rotavirus antibody	A	A					
86762	Rubella antibody	A	A					
86765	Rubella antibody	A	A					
86768	Salmonella antibody	A	A					
86771	Shigella antibody	A	A					
86774	Tetanus antibody	A	A					
86777	Toxoplasma antibody	A	A					
86778	Toxoplasma antibody, igm	A	A					
86780	Treponema pallidum	NI	A					
86781	Treponema pallidum, confirm	CH	D					
86784	Trichinella antibody	A	A					
86787	Vaccella-zoster antibody	A	A					
86788	West Nile virus ab, igm	A	A					
86789	West Nile virus antibody	A	A					
86790	Virus antibody nos	A	A					
86793	Yersinia antibody	A	A					
86800	Thyroglobulin antibody	A	A					
86803	Hepatitis c ab test	A	A					
86804	Hep c ab test, confirm	A	A					
86805	Lymphocytotoxicity assay	A	A					
86806	Lymphocytotoxicity assay	A	A					
86807	Cytotoxic antibody screening	A	A					
86808	Cytotoxic antibody screening	A	A					
86812	HLA typing, A, B, or C	A	A					
86813	HLA typing, A, B, or C	A	A					
86816	HLA typing, DR/DQ	A	A					
86817	HLA typing, DR/DQ	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
87185	Microbe susceptible, enzyme		A					
87186	Microbe susceptible, mic		A					
87187	Microbe susceptible, mic		A					
87188	Microbe suscept, macrobroth		A					
87190	Microbe suscept, mycobacteri		A					
87197	Bactericidal level, serum		A					
87205	Smear, gram stain		A					
87206	Smear, fluorescein/acid stai		A					
87207	Smear, special stain		A					
87209	Smear, complex stain		A					
87210	Smear, wet mount, saline/ink		A					
87220	Tissue exam for fungi		A					
87230	Assay, toxin or antitoxin		A					
87250	Virus inoculate, eggs/animal		A					
87252	Virus inoculation, tissue		A					
87253	Virus inoculate tissue, addl		A					
87254	Virus inoculation, shell via		A					
87265	Genet virus isolate, hsv		A					
87260	Adenovirus ag, if		A					
87265	Pertussis ag, if		A					
87267	Enterovirus antibody, dfa		A					
87269	Giardia ag, if		A					
87270	Chlamydia trachomatis ag, if		A					
87271	Cytomegalovirus dfa		A					
87272	Cryptosporidium ag, if		A					
87273	Herpes simplex 2, ag, if		A					
87274	Herpes simplex 1, ag, if		A					
87275	Influenza b, ag, if		A					
87276	Influenza a, ag, if		A					
87277	Legionella micdadei, ag, if		A					
87278	Legion pneumophila ag, if		A					
87279	Parainfluenza, ag, if		A					
87280	Respiratory syncytial ag, if		A					
87281	Pneumocystis carinii, ag, if		A					
87283	Rubella, ag, if		A					
87285	Treponema pallidum, ag, if		A					
87290	Varicella zoster, ag, if		A					
87299	Antibody detection, nos, if		A					
87300	Ag detection, polyval, if		A					
87301	Adenovirus ag, eia		A					
87305	Aspergillus ag, eia		A					
87320	Chyind trach ag, eia		A					
87324	Clostridium ag, eia		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
86999	Transfusion procedure	X	X	0345	0.2195	\$14.80		\$2.96
87001	Small animal inoculation		A					
87003	Small animal inoculation		A					
87015	Specimen concentration		A					
87040	Blood culture for bacteria		A					
87045	Feces culture, bacteria		A					
87046	Stool cult, bacteria, each		A					
87070	Culture, bacteria, other		A					
87071	Culture bacteri aerobic othr		A					
87073	Culture bacteria anaerobic		A					
87075	Cult bacteria, except blood		A					
87076	Culture anaerobe ident, each		A					
87077	Culture aerobic identify		A					
87081	Culture screen only		A					
87084	Culture of specimen by kit		A					
87086	Urine culture/colony count		A					
87088	Urine bacteria culture		A					
87101	Skin fungi culture		A					
87102	Fungus isolation culture		A					
87103	Blood fungus culture		A					
87106	Fungi identification, yeast		A					
87107	Fungi identification, mold		A					
87109	Mycoplasma		A					
87110	Chlamydia culture		A					
87116	Mycobacteria culture		A					
87118	Mycobacteric identification		A					
87140	Culture type immunofluoresc		A					
87143	Culture typing, gtc/hpic		A					
87147	Culture type, immunologic		A					
87149	Dna/rna direct probe		A					
87150	Dna/rna, amplified probe		A					
87152	Culture type pulse field gel		NI					
87153	Dna/rna sequencing		NI					
87158	Culture typing, added method		A					
87164	Dark field examination		A					
87166	Dark field examination		A					
87168	Macroscopic exam arthropod		A					
87169	Macroscopic exam parasite		A					
87172	Pinworm exam		A					
87176	Tissue homogenization, cult		A					
87177	Ova and parasites smears		A					
87181	Microbe susceptible, diffuse		A					
87184	Microbe susceptible, disk		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
87498	Enterovirus, dna, amp probe	A	A					
87500	Vanomycin, dna, amp probe	A	A					
87510	Gardner vag, dna, dir probe	A	A					
87511	Gardner vag, dna, amp probe	A	A					
87512	Gardner vag, dna, quant	A	A					
87515	Hepatitis b, dna, dir probe	A	A					
87516	Hepatitis b, dna, amp probe	A	A					
87517	Hepatitis b, dna, quant	A	A					
87520	Hepatitis c, rna, dir probe	A	A					
87521	Hepatitis c, rna, amp probe	A	A					
87522	Hepatitis c, rna, quant	A	A					
87525	Hepatitis g, dna, dir probe	A	A					
87526	Hepatitis g, dna, amp probe	A	A					
87527	Hepatitis g, dna, quant	A	A					
87528	Hsv, dna, dir probe	A	A					
87529	Hsv, dna, amp probe	A	A					
87530	Hsv, dna, quant	A	A					
87531	Hiv-6, dna, dir probe	A	A					
87532	Hiv-6, dna, amp probe	A	A					
87533	Hiv-6, dna, quant	A	A					
87534	Hiv-1, dna, dir probe	A	A					
87535	Hiv-1, dna, amp probe	A	A					
87536	Hiv-1, dna, quant	A	A					
87537	Hiv-2, dna, dir probe	A	A					
87538	Hiv-2, dna, amp probe	A	A					
87539	Hiv-2, dna, quant	A	A					
87540	Legion pneumo, dna, dir prob	A	A					
87541	Legion pneumo, dna, amp prob	A	A					
87542	Legion pneumo, dna, quant	A	A					
87550	Mycobacteria, dna, dir probe	A	A					
87551	Mycobacteria, dna, amp probe	A	A					
87552	Mycobacteria, dna, quant	A	A					
87555	M.tuberculo, dna, dir probe	A	A					
87556	M.tuberculo, dna, amp probe	A	A					
87557	M.tuberculo, dna, quant	A	A					
87560	M.avium-intra, dna, dir prob	A	A					
87561	M.avium-intra, dna, amp prob	A	A					
87562	M.avium-intra, dna, quant	A	A					
87580	M.pneumon, dna, dir probe	A	A					
87581	M.pneumon, dna, amp probe	A	A					
87582	M.pneumon, dna, quant	A	A					
87590	N.gonorrhoeae, dna, dir prob	A	A					
87591	N.gonorrhoeae, dna, amp prob	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
87327	Cryptococcus neoform ag, eia	A	A					
87328	Cryptosporidium ag, eia	A	A					
87329	Giardia ag, eia	A	A					
87332	Cytomegalovirus ag, eia	A	A					
87335	E.coli 0157 ag, eia	A	A					
87336	Entamoeb hist dispr, ag, eia	A	A					
87337	Entamoeb hist group, ag, eia	A	A					
87338	Hpylori, stool, eia	A	A					
87339	H pylori ag, eia	A	A					
87340	Hepatitis b surface ag, eia	A	A					
87341	Hepatitis b surface, ag, eia	A	A					
87350	Hepatitis be ag, eia	A	A					
87380	Hepatitis delta ag, eia	A	A					
87385	Histoplasma capsul ag, eia	A	A					
87390	Hiv-1 ag, eia	A	A					
87391	Hiv-2 ag, eia	A	A					
87400	Influenza a/b, ag, eia	A	A					
87420	Resp syncytial ag, eia	A	A					
87425	Rotavirus ag, eia	A	A					
87427	Shiga-like toxin ag, eia	A	A					
87430	Strep a ag, eia	A	A					
87449	Ag detect nos, eia, mult	A	A					
87450	Ag detect nos, eia, single	A	A					
87451	Ag detect polyval, eia, mult	A	A					
87470	Bartonella, dna, dir probe	A	A					
87471	Bartonella, dna, amp probe	A	A					
87472	Bartonella, dna, quant	A	A					
87475	Lyme dis, dna, dir probe	A	A					
87476	Lyme dis, dna, amp probe	A	A					
87477	Lyme dis, dna, quant	A	A					
87480	Candida, dna, dir probe	A	A					
87481	Candida, dna, amp probe	A	A					
87482	Candida, dna, quant	A	A					
87485	Chyimd pneum, dna, dir probe	A	A					
87486	Chyimd pneum, dna, amp probe	A	A					
87487	Chyimd pneum, dna, quant	A	A					
87490	Chyimd trach, dna, dir probe	A	A					
87491	Chyimd trach, dna, amp probe	A	A					
87492	Chyimd trach, dna, quant	A	A					
87493	C diff amplified probe	NI	A					
87495	Cytomeg, dna, dir probe	A	A					
87496	Cytomeg, dna, amp probe	A	A					
87497	Cytomeg, dna, quant	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
88029	Autopsy (necropsy), complete		E					
88036	Limited autopsy		E					
88037	Limited autopsy		E					
88040	Forensic autopsy (necropsy)		E					
88045	Coroners autopsy (necropsy)		E					
88099	Necropsy (autopsy) procedure		E					
88104	Cytopath fl nongyn, smears		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88106	Cytopath fl nongyn, filter		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88107	Cytopath fl nongyn, sm/fltr		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88108	Cytopath, concentrate tech		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88112	Cytopath, cell enhance tech		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88125	Forensic cytopathology		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88130	Sex chromatin identification		A					
88140	Sex chromatin identification		A					
88141	Cytopath, c/v, interpret		N					
88142	Cytopath, c/v, thin layer		A					
88143	Cytopath c/v thin layer redo		A					
88147	Cytopath, c/v, automated		A					
88148	Cytopath, c/v, auto rescreen		A					
88150	Cytopath, c/v, manual		A					
88152	Cytopath, c/v, auto redo		A					
88153	Cytopath, c/v, redo		A					
88154	Cytopath, c/v, select		A					
88155	Cytopath, c/v, index add-on		A					
88160	Cytopath smear, other source		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88161	Cytopath smear, other source		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88162	Cytopath smear, other source	CH	X	0343	0.5301	\$35.73	\$10.84	\$7.15
88164	Cytopath tbs, c/v, manual		A					
88165	Cytopath tbs, c/v, redo		A					
88166	Cytopath tbs, c/v, auto redo		A					
88167	Cytopath tbs, c/v, select		A					
88172	Cytopathology eval of fna		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88173	Cytopath eval, fna, report		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88174	Cytopath, c/v auto, in fluid		A					
88175	Cytopath c/v auto fluid redo		A					
88182	Cell marker study		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88184	Flowcytometry/ lc, 1 marker		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88185	Flowcytometry/lc, add-on		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88187	Flowcytometry/read, 2-8		X	0342	0.1546	\$10.42	\$2.09	\$2.09
88188	Flowcytometry/read, 9-15		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88189	Flowcytometry/read, 16 & >		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88199	Cytopathology procedure		X	0342	0.1546	\$10.42	\$2.09	\$2.09
88230	Tissue culture, lymphocyte		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
87592	N.gonorrhoeae, dna, quant		A					
87620	Hpv, dna, dir probe		A					
87621	Hpv, dna, amp probe		A					
87622	Hpv, dna, quant		A					
87640	Staph a, dna, amp probe		A					
87641	Mr-staph, dna, amp probe		A					
87650	Strep a, dna, dir probe		A					
87651	Strep a, dna, amp probe		A					
87652	Strep a, dna, quant		A					
87653	Strep b, dna, amp probe		A					
87660	Trichomonas vagin, dir probe		A					
87797	Detect agent nos, dna, dir		A					
87798	Detect agent nos, dna, amp		A					
87799	Detect agent nos, dna, quant		A					
87800	Detect agent mult, dna, direc		A					
87801	Detect agent mult, dna, amplic		A					
87802	Strep b assay w/optic		A					
87803	Clostridium toxin a w/optic		A					
87804	Influenza assay w/optic		A					
87807	Rsv assay w/optic		A					
87808	Trichomonas assay w/optic		A					
87809	Adenovirus assay w/optic		A					
87810	Chyimd trach assay w/optic		A					
87850	N. gonorrhoeae assay w/optic		A					
87880	Strep a assay w/optic		A					
87899	Agent nos assay w/optic		A					
87900	Phenotype, infect agent drug		A					
87901	Genotype, dna, hiv reverse t		A					
87902	Genotype, dna, hepatitis C		A					
87903	Phenotype, dna hiv w/culture		A					
87904	Phenotype, dna hiv w/ict add		A					
87905	Sialidase enzyme assay		A					
87999	Microbiology procedure		A					
88000	Autopsy (necropsy), gross		E					
88005	Autopsy (necropsy), gross		E					
88007	Autopsy (necropsy), gross		E					
88012	Autopsy (necropsy), gross		E					
88014	Autopsy (necropsy), gross		E					
88016	Autopsy (necropsy), gross		E					
88020	Autopsy (necropsy), complete		E					
88025	Autopsy (necropsy), complete		E					
88027	Autopsy (necropsy), complete		E					
88028	Autopsy (necropsy), complete		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
88332	Path consult intraop, addl		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88333	Intraop cyto path consult, 1		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88334	Intraop cyto path consult, 2		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88342	Immunohistochemistry		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88347	Immunofluorescent study		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88348	Immunofluorescent study		X	0661	2.445	\$164.81	\$67.69	\$32.97
88349	Electron microscopy		X	0661	2.445	\$164.81	\$67.69	\$32.97
88355	Scanning electron microscopy		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88356	Analysis, skeletal muscle		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88356	Analysis, nerve		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88360	Analysis, tumor		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88360	Immunohistochem/manual		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88361	Tumor		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88362	Immunohistochem/comput	CH	X	0344	0.8028	\$54.11	\$15.60	\$10.83
88365	Nerve teasing preparations		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88365	In situ hybridization (fish)		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88367	In situ hybridization, auto		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88368	In situ hybridization, manual	CH	X	0344	0.8028	\$54.11	\$15.60	\$10.83
88371	Protein, western blot tissue		A					
88372	Protein analysis w/probe		A					
88380	Microdissection, laser		N					
88381	Microdissection, manual		N					
88384	Eval molec probab, 11-50		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88385	Eval molec probab, 51-250		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88386	Eval molec probab, 251-500		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88387	Tiss exam molecular study	NI	N					
88388	Tiss ex molecular study add-on	NI	N					
88399	Surgical pathology procedure		X	0342	0.1546	\$10.42		\$2.09
88720	Bilirubin total transitcut		A					
88738	Hgb quant transcutaneous	NI	A					
88740	Transcutaneous carboxyhb		A					
88741	Transcutaneous methb		A					
89049	Chct for mal hyperthermia		X	0342	0.1546	\$10.42		\$2.09
89050	Body fluid cell count		A					
89051	Body fluid cell count		A					
89055	Leukocyte assessment, fecal		A					
89060	Exam synovial fluid crystals		A					
89100	Sample intestinal contents		X	0360	1.5203	\$102.48	\$33.88	\$20.50
89105	Sample intestinal contents		X	0360	1.5203	\$102.48	\$33.88	\$20.50
89125	Specimen lat stain		A					
89130	Sample stomach contents		X	0360	1.5203	\$102.48	\$33.88	\$20.50
89132	Sample stomach contents		X	0360	1.5203	\$102.48	\$33.88	\$20.50
89135	Sample stomach contents		X	0360	1.5203	\$102.48	\$33.88	\$20.50

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
88233	Tissue culture, skin/biopsy		A					
88235	Tissue culture, placenta		A					
88237	Tissue culture, bone marrow		A					
88239	Tissue culture, tumor		A					
88240	Cell cryopreserve/storage		A					
88241	Frozen cell preparation		A					
88245	Chromosome analysis, 20-25		A					
88248	Chromosome analysis, 50-100		A					
88249	Chromosome analysis, 100		A					
88261	Chromosome analysis, 5		A					
88262	Chromosome analysis, 15-20		A					
88263	Chromosome analysis, 45		A					
88264	Chromosome analysis, 20-25		A					
88267	Chromosome analysis, placenta		A					
88269	Chromosome analysis, amniotic		A					
88271	Cytogenetics, dna probe		A					
88272	Cytogenetics, 3-5		A					
88273	Cytogenetics, 10-30		A					
88274	Cytogenetics, 25-99		A					
88275	Cytogenetics, 100-300		A					
88280	Chromosome karyotype study		A					
88283	Chromosome banding study		A					
88285	Chromosome count, additional		A					
88289	Chromosome study, additional		A					
88291	Cyto/molecular report		M					
88299	Cytogenetic study		X	0342	0.1546	\$10.42		\$2.09
88300	Surgical path, gross		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88302	Tissue exam by pathologist		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88304	Tissue exam by pathologist		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88305	Tissue exam by pathologist		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88307	Tissue exam by pathologist		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88309	Tissue exam by pathologist		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88311	Decalcify tissue		X	0342	0.1546	\$10.42		\$2.09
88312	Special stains group 1		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88313	Special stains group 2		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88314	Histochemical stain acid-on		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88318	Chemical histochemistry		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88319	Enzyme histochemistry		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88321	Microslide consultation		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88323	Microslide consultation		X	0343	0.5301	\$35.73	\$10.84	\$7.15
88325	Comprehensive review of data		X	0344	0.8028	\$54.11	\$15.60	\$10.83
88329	Path consult introp		X	0433	0.2482	\$16.73	\$5.17	\$3.35
88331	Path consult intraop, 1 bloc		X	0343	0.5301	\$35.73	\$10.84	\$7.15

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
89354	Thaw cryopresv'd; reprod tiss	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89356	Thawing cryopresv'd; oocyte	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89398	Unlisted reprod med lab proc	NI	X	0342	0.1546	\$10.42		\$2.09
90281	Human ig, im	E						
90283	Human ig, iv	E						
90284	Human ig, sc	E						
90287	Botulinum antitoxin	E						
90288	Botulinem ig, iv	E						
90291	Crmy ig, iv	E						
90296	Diphtheria antitoxin	CH	E					
90371	Hep b ig, im	K	K	1630	\$111.20			\$22.24
90375	Rabies ig, im/sc	K	K	9133	\$142.79			\$28.56
90376	Rabies ig, heat treated	K	K	9134	\$130.16			\$26.04
90378	Rsv, mab, im, 50mg	K	K	9003	\$997.29			\$187.46
90379	Rsv, ig, iv	CH	D					
90384	Rh ig, full-dose, im	E	E					
90385	Rh ig, minidose, im	N	N					
90386	Rh ig, iv	E	E					
90389	Tetanus ig, im	E	E					
90393	Vaccina ig, im	CH	E					
90396	Vartcella-zoster ig, im	K	K	9135	\$130.49			\$26.10
90399	Immune globulin	E	E					
90465	Immune admin 1 inj, < 8 yrs	B	B					
90466	Immune admin addl inj, < 8 y	B	B					
90467	Immune admin o r n, < 8 yrs	B	B					
90468	Immune admin o/r, addl < 8 y	B	B					
90470	Immune admin HTNI im/nasal	NI	E					
90471	Immunization admin	S	S	0436	\$25.67			\$5.14
90472	Immunization admin, each add	S	S	0436	\$25.67			\$5.14
90473	Immune admin oral/nasal	S	S	0436	\$25.67			\$5.14
90474	Immune admin oral/nasal addl	S	S	0436	\$25.67			\$5.14
90476	Adenovirus vaccine, type 4	CH	K	1254	\$72.17			\$14.44
90477	Adenovirus vaccine, type 7	CH	E					
90581	Anthrax vaccine, sc	CH	E					
90585	Bcg vaccine, percut	K	K	9137	\$111.66			\$22.34
90586	Bcg vaccine, intravesical	B	N					
90632	Hep a vaccine, adult, im	N	N					
90633	Hep a vacc, ped/adol, 2 dose	N	N					
90634	Hep a vacc, ped/adol, 3 dose	N	N					
90636	Hep a/hep b vacc, adult im	N	N					
90644	HIB/men/tf vaccine, im	NI	E					
90645	Hib vaccine, hboc, im	N	N					
90646	Hib vaccine, prp-d, im	N	N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
89136	Sample stomach contents	X	X	0360	1.5203	\$102.48	\$33.88	\$20.50
89140	Sample stomach contents	X	X	0360	1.5203	\$102.48	\$33.88	\$20.50
89141	Sample stomach contents	X	X	0360	1.5203	\$102.48	\$33.88	\$20.50
89160	Exam feces for meat fibers	A						
89190	Nasal smear for eosinophils	A						
89220	Sputum specimen collection	X	A	0433	0.2482	\$16.73	\$5.17	\$3.35
89225	Starch granules, feces	X	A					
89230	Collect sweat for test	X	X	0343	0.5301	\$35.73	\$10.84	\$7.15
89235	Water load test	A						
89240	Pathology lab procedure	X	A	0342	0.1546	\$10.42		\$2.09
89250	Cultr oocyte/embryo <4 days	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89251	Cultr oocyte/embryo <4 days	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89253	Embryo hatching	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89254	Oocyte identification	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89255	Prepare embryo for transfer	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89257	Sperm identification	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89258	Cryopreservation; embryo(s)	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89259	Cryopreservation; sperm	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89260	Sperm isolation, simple	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89261	Sperm isolation, complex	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89264	Identify sperm tissue	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89268	Insemination of oocytes	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89272	Extended culture of oocytes	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89280	Assist oocyte fertilization	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89281	Assist oocyte fertilization	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89290	Biopsy, oocyte polar body	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89291	Biopsy, oocyte polar body	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89300	Semen analysis whitnmer	A						
89310	Semen analysis w/count	A						
89320	Semen anal. vol/count/mot	A						
89321	Semen anal, sperm detection	A						
89322	Semen anal, strict criteria	A						
89325	Sperm antibody test	A						
89329	Sperm evaluation test	A						
89330	Evaluation, cervical mucus	A						
89331	Retrograde ejaculation anal	A						
89335	Cryopreserve testicular tiss	A	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89342	Storage/year, embryo(s)	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89343	Storage/year, sperm/semn	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89344	Storage/year, reprod tissue	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89346	Storage/year, oocyte(s)	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89352	Thawing cryopresv'd; embryo	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83
89353	Thawing cryopresv'd; sperm	X	X	0344	0.8028	\$54.11	\$15.60	\$10.83

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
90720	Diph/hib vaccine, im	N	N					
90721	Diph/hib vaccine, im	N	N					
90723	Dtap-hep b-ipv vaccine, im	E	E	1271		\$0.16		\$0.04
90725	Cholera vaccine, injectable	CH	K					
90727	Plague vaccine, im	CH	E					
90732	Pneumococcal vaccine	L	L					
90733	Meningococcal vaccine, sc	K	K	9143		\$96.66		\$19.34
90734	Meningococcal vaccine, im	K	K	9145		\$102.46		\$20.50
90735	Encephalitis vaccine, sc	CH	K	9144		\$100.15		\$20.03
90736	Zoster vacc, sc	M	M					
90738	Inactivated je vacc, im	M	M					
90740	Hepb vacc, ill pat 3 dose im	F	F					
90743	Hep b vacc, adol, 2 dose, im	F	F					
90744	Hepb vacc ped/adol 3 dose im	F	F					
90746	Hep b vaccine, adult, im	F	F					
90747	Hepb vacc, ill pat 4 dose im	F	F					
90748	Hep b/hib vaccine, im	E	E					
90749	Vaccine toxoid	N	N					
90801	Psy dx interview	O3	O3	0323	1.669	\$112.50		\$22.50
90802	Intac psy dx interview	O3	O3	0323	1.669	\$112.50		\$22.50
90804	Psytx, office, 20-30 min	O3	O3	0322	1.1958	\$80.60		\$16.12
90805	Psytx, off, 20-30 min w/e&m	O3	O3	0322	1.1958	\$80.60		\$16.12
90806	Psytx, off, 45-50 min	O3	O3	0323	1.669	\$112.50		\$22.50
90807	Psytx, off, 45-50 min w/e&m	O3	O3	0323	1.669	\$112.50		\$22.50
90808	Psytx, office, 75-80 min	O3	O3	0323	1.669	\$112.50		\$22.50
90809	Psytx, off, 75-80 w/e&m	O3	O3	0322	1.1958	\$80.60		\$16.12
90810	Intac psytx, off, 20-30 min	O3	O3	0322	1.1958	\$80.60		\$16.12
90811	Intac psytx, 20-30, w/e&m	O3	O3	0322	1.1958	\$80.60		\$16.12
90812	Intac psytx, off, 45-50 min	O3	O3	0323	1.669	\$112.50		\$22.50
90813	Intac psytx, 45-50 min w/e&m	O3	O3	0323	1.669	\$112.50		\$22.50
90814	Intac psytx, off, 75-80 min	O3	O3	0323	1.669	\$112.50		\$22.50
90815	Intac psytx, 75-80 w/e&m	O3	O3	0323	1.669	\$112.50		\$22.50
90816	Psytx, hosp, 20-30 min w/e&m	P	P					
90817	Psytx, hosp, 20-30 min	P	P					
90818	Psytx, hosp, 45-50 min	P	P					
90819	Psytx, hosp, 45-50 min w/e&m	P	P					
90821	Psytx, hosp, 75-80 min	P	P					
90822	Psytx, hosp, 75-80 min w/e&m	P	P					
90823	Intac psytx, hosp, 20-30 min	P	P					
90824	Intac psytx, hsp 20-30 w/e&m	P	P					
90826	Intac psytx, hosp, 45-50 min	P	P					
90827	Intac psytx, hsp 45-50 w/e&m	P	P					
90828	Intac psytx, hosp, 75-80 min	P	P					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
90647	Hib vaccine, prp-omp, im	N	N					
90648	Hib vaccine, prp-1, im	N	N					
90649	Hpv vaccine 4 valent, im	M	M					
90650	Hpv vaccine 2 valent, im	CH	M					
90655	Flu vaccine no preserv 6-35m	L	L					
90656	Flu vaccine no preserv 3 & >	L	L					
90657	Flu vaccine, 3 yrs, im	L	L					
90658	Flu vaccine, 3 yrs & >, im	L	L					
90660	Flu vaccine, nasal	L	L					
90661	Flu vacc cell cult, prsv free	E	E					
90662	Flu vacc prsv free inc antig	E	E					
90663	Flu vacc pandemic H1N1	E	E			\$0.93		\$0.19
90665	Lyme disease vaccine, im	K	K	1216				
90669	Pneumococcal vacc, 7 val, im	L	L					
90670	Pneumococcal vacc, 13 val, im	E	E					
90675	Rabies vaccine, im	K	K	9139		\$151.97		\$30.40
90676	Rabies vaccine, id	K	K	9140		\$96.27		\$19.26
90680	Rotavirus vacc 3 dose, oral	CH	K	1255		\$72.37		\$14.48
90681	Rotavirus vacc 2 dose oral	K	K	1239		\$106.60		\$21.32
90690	Typhoid vaccine, oral	N	N					
90691	Typhoid vaccine, im	N	N					
90692	Typhoid vaccine, h-p, sc/id	N	N					
90693	Typhoid vaccine, akd, sc	B	B					
90696	Dtap-ipv vacc 4-6 yr, im	CH	N					
90698	Dtap-hib-ipv vaccine, im	N	N					
90700	Dtap vaccine, < 7 yrs, im	N	N					
90701	Dtp vaccine, im	N	N					
90702	Dt vaccine < 7, im	N	N					
90703	Tetanus vaccine, im	N	N					
90704	Mumps vaccine, sc	N	N					
90705	Measles vaccine, sc	N	N					
90706	Rubella vaccine, sc	N	N					
90707	Mmr vaccine, sc	N	N					
90708	Measles-rubella vaccine, sc	N	N					
90710	Mmr vaccine, sc	N	N					
90712	Oral poliovirus vaccine	N	N					
90713	Poliovirus, ipv, sc/im	N	N					
90714	Td vaccine no prsv >= 7 im	N	N					
90715	Tdap vaccine >7 im	N	N					
90716	Chicken pox vaccine, sc	M	M					
90717	Yellow fever vaccine, sc	N	N					
90718	Td vaccine > 7, im	N	N					
90719	Diphtheria vaccine, im	N	N					

APPENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
90869	Esrd home pt serv p day 12-19		M					
90970	Esrd home pt serv p day, 20+		M					
90989	Dialysis training, complete		B					
90993	Dialysis training, incompl		B					
90997	Hemoperfusion		B					
90999	Dialysis procedure		X					
91000	Esophageal intubation		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91010	Esophagus motility study		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91011	Esophagus motility study		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91012	Esophagus motility study		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91020	Gastric motility studies		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91022	Duodenal motility study		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91030	Acid perfusion of esophagus		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91034	Gastroesophageal reflux test		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91035	G-esoph reflux tst w/electrod		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91037	Esoph impeded function test		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91038	Esoph impeded funct test > 1h		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91040	Esoph balloon distension tst		X	0360	1.5203	\$102.48	\$33.88	\$20.50
91052	Gastric analysis test		X	0361	4.0839	\$275.28	\$83.23	\$55.06
91065	Breath hydrogen test		X	0360	1.5203	\$102.48	\$33.88	\$20.50
91105	Gastric intubation treatment		X	0360	1.5203	\$102.48	\$33.88	\$20.50
91110	Gt tract capsule endoscopy		T	0142	9.8503	\$663.97	\$192.78	\$132.80
91111	Esophageal capsule endoscopy		T	0141	8.7462	\$589.55	\$143.38	\$117.91
91120	Rectal sensation test		T	0126	1.0958	\$73.86	\$16.21	\$14.78
91122	Anal pressure record		T	0156	3.0214	\$203.66		\$40.74
91123	Irrigate fecal impaction		N					
91133	Electrogastrography		X	0360	1.5203	\$102.48	\$33.88	\$20.50
91289	Electrogastrography w/leat		X	0360	1.5203	\$102.48	\$33.88	\$20.50
92002	Gastroenterology procedure		CH					
92004	Eye exam, new patient		V	0606	1.3222	\$89.12		\$17.83
92012	Eye exam, established pat		V	0604	0.8593	\$57.92		\$11.59
92014	Eye exam & treatment		V	0605	1.0337	\$69.68		\$13.94
92015	Refraction		E					
92018	New eye exam & treatment		T	0699	15.5407	\$1,047.54		\$209.51
92019	Eye exam & treatment		T	0698	15.5407	\$1,047.54		\$209.51
92020	Special eye evaluation		S	0230	0.5945	\$40.07		\$8.02
92060	Corneal topography		S	0698	0.9553	\$64.39		\$12.88
92065	Special eye evaluation		S	0698	0.9553	\$64.39		\$12.88
92070	Orthoptic/pleoptic training		S	0698	0.9553	\$64.39		\$12.88
92071	Fitting of contact lens		N					
92081	Visual field examination(s)		S	0230	0.5945	\$40.07		\$8.02

APPENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
90829	Intac psytX, hsp 75-80 w/e&rn		P					
90845	Psychoanalysis		O3	0323	1.669	\$112.50		\$22.50
90846	Family psytX w/o patient		O3	0324	2.2103	\$148.99		\$29.80
90847	Family psytX w/patient		O3	0324	2.2103	\$148.99		\$29.80
90849	Multiple family group psytX		O3	0325	0.8905	\$60.03	\$12.81	\$12.01
90853	Group psychotherapy		O3	0325	0.8905	\$60.03	\$12.81	\$12.01
90857	Intac group psytX		O3	0325	0.8905	\$60.03	\$12.81	\$12.01
90862	Medication management		O3	0606	1.3222	\$89.12		\$17.83
90865	Electroconvulsive therapy		O3	0323	1.669	\$112.50		\$22.50
90870	Narcosynthesis		S	0320	5.862	\$395.13	\$80.06	\$79.03
90875	Psychophysiological therapy		E					
90876	Psychophysiological therapy		E					
90880	Hypnotherapy		O3	0323	1.669	\$112.50		\$22.50
90882	Environmental manipulation		E					
90885	Psy evaluation of records		N					
90887	Consultation with family		N					
90889	Preparation of report		N					
90899	Psychiatric service/therapy		O3	0322	1.1958	\$80.60		\$16.12
90901	Biobfeedback train, any meth		A					
90911	Biobfeedback per/uro/rectal		T	0126	1.0958	\$73.86	\$16.21	\$14.78
90935	Hemodialysis, one evaluation		S	0170	6.8212	\$459.79		\$91.96
90937	Hemodialysis, repeated eval		B					
90940	Hemodialysis access study		N					
90945	Dialysis, one evaluation		CH					
90947	Dialysis, repeated eval		B	0608	2.4853	\$167.52		\$33.51
90951	Esrd serv, 4 visits p mo, <2		M					
90952	Esrd serv, 2-3 vists p mo, <2		M					
90953	Esrd serv, 1 visit p mo, <2		M					
90954	Esrd serv, 4 vists p mo, 2-11		M					
90955	Esrd srv 2-3 vists p mo, 2-11		M					
90956	Esrd srv, 1 visit p mo, 2-11		M					
90957	Esrd srn, 4 vists p mo, 12-19		M					
90958	Esrd srn 2-3 vists p mo 12-19		M					
90959	Esrd serv, 1 vstl p mo, 12-19		M					
90961	Esrd srn, 2-3 vists p mo, 20+		M					
90962	Esrd serv, 1 visit p mo, 20+		M					
90963	Esrd home pt, serv p mo, <2		M					
90964	Esrd home pt serv p mo, 2-11		M					
90965	Esrd home pt serv p mo 12-19		M					
90966	Esrd home pt, serv p mo, 20+		M					
90967	Esrd home pt serv p day, <2		M					
90968	Esrd home pt serv p day, 2-11		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
92499	Eye service or procedure		S	0230	0.5945	\$40.07		\$8.02
92502	Ear and throat examination		T	0251	3.425	\$230.87		\$46.18
92504	Ear microscopy examination		N					
92506	Speech/hearing evaluation		A					
92507	Speech/hearing therapy		A					
92508	Speech/hearing therapy		A					
92511	Nasopharyngoscopy		T	0071	0.8007	\$53.97		\$10.80
92512	Nasal function studies		X	0363	0.9061	\$61.08		\$12.22
92516	Facial nerve function test		X	0660	1.5019	\$101.24		\$20.25
92520	Laryngeal function studies		X	0660	1.5019	\$101.24		\$20.25
92526	Oral function therapy		A					
92531	Spontaneous nystagmus study		N					
92532	Positional nystagmus test		N					
92533	Caloric vestibular test		N					
92534	Optokinetic nystagmus test		N					
92540	Basic vestibular evaluation	NI	X	0660	1.5019	\$101.24		\$20.25
92541	Spontaneous nystagmus test		X	0363	0.9061	\$61.08		\$12.22
92542	Positional nystagmus test		X	0363	0.9061	\$61.08		\$12.22
92543	Caloric vestibular test		X	0660	1.5019	\$101.24		\$20.25
92544	Optokinetic nystagmus test		X	0363	0.9061	\$61.08		\$12.22
92545	Oscillating tracking test		X	0363	0.9061	\$61.08		\$12.22
92546	Sinusoidal rotational test		X	0660	1.5019	\$101.24		\$20.25
92547	Supplemental electrical test		N					
92548	Posturography		X	0660	1.5019	\$101.24		\$20.25
92550	Tympanometry & reflex thresh	NI	X	0364	0.47	\$31.68		\$6.34
92551	Pure tone hearing test, air		E					
92552	Pure tone audiometry, air		X	0364	0.47	\$31.68		\$6.34
92553	Audiometry, air & bone		X	0365	1.2675	\$85.44		\$17.09
92555	Speech threshold audiometry		X	0364	0.47	\$31.68		\$6.34
92556	Speech audiometry, complete		X	0364	0.47	\$31.68		\$6.34
92557	Comprehensive hearing test		X	0365	1.2675	\$85.44		\$17.09
92559	Group audiometric testing		E					
92560	Bekesy audiometry, screen		E					
92561	Bekesy audiometry, diagnosis		X	0364	0.47	\$31.68		\$6.34
92562	Loudness balance test		X	0364	0.47	\$31.68		\$6.34
92563	Tone decay hearing test		X	0364	0.47	\$31.68		\$6.34
92564	Sisi hearing test		X	0364	0.47	\$31.68		\$6.34
92565	Stenger test, pure tone		X	0364	0.47	\$31.68		\$6.34
92567	Tympanometry		X	0364	0.47	\$31.68		\$6.34
92568	Acoustic refl threshold tst		X	0364	0.47	\$31.68		\$6.34
92569	Acoustic reflex decay test	CH	D					
92570	Acoustic immittance testing	NI	X	0364	0.47	\$31.68		\$6.34
92571	Filtered speech hearing test		X	0364	0.47	\$31.68		\$6.34

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
92082	Visual field examination(s)		S	0698	0.9553	\$64.39		\$12.88
92083	Visual field examination(s)		S	0698	0.9553	\$64.39		\$12.88
92100	Serial tonometry exam(s)		N					
92120	Tonography & eye evaluation		S	0698	0.9553	\$64.39		\$12.88
92130	Water provocation tonography		S	0230	0.5945	\$40.07		\$8.02
92135	Optith dx imaging post seg		S	0230	0.5945	\$40.07		\$8.02
92136	Ophthalmic biometry		S	0698	0.9553	\$64.39		\$12.88
92140	Glaucoma provocative tests		S	0230	0.5945	\$40.07		\$8.02
92225	Special eye exam, initial		S	0230	0.5945	\$40.07		\$8.02
92226	Special eye exam, subsequent		S	0698	0.9553	\$64.39		\$12.88
92230	Eye exam with photos		S	0231	1.9575	\$131.95		\$26.39
92235	Eye exam with photos		S	0231	1.9575	\$131.95		\$26.39
92240	Icg angiography		S	0231	1.9575	\$131.95		\$26.39
92250	Eye exam with photos		S	0698	0.9553	\$64.39		\$12.88
92260	Ophthalmoscopy/dynamometry		S	0230	0.5945	\$40.07		\$8.02
92265	Eye muscle evaluation		S	0698	0.9553	\$64.39		\$12.88
92270	Electro-oculography		S	0230	0.5945	\$40.07		\$8.02
92275	Electroretinography		S	0231	1.9575	\$131.95		\$26.39
92283	Color vision examination		S	0230	0.5945	\$40.07		\$8.02
92284	Dark adaptation eye exam		S	0698	0.9553	\$64.39		\$12.88
92285	Eye photography		S	0698	0.9553	\$64.39		\$12.88
92286	Internal eye photography		S	0231	1.9575	\$131.95		\$26.39
92287	Internal eye photography		S	0231	1.9575	\$131.95		\$26.39
92310	Contact lens fitting		E					
92311	Contact lens fitting		S	0698	0.9553	\$64.39		\$12.88
92312	Contact lens fitting		S	0698	0.9553	\$64.39		\$12.88
92313	Contact lens fitting		S	0230	0.5945	\$40.07		\$8.02
92314	Prescription of contact lens		E					
92315	Prescription of contact lens		S	0230	0.5945	\$40.07		\$8.02
92316	Prescription of contact lens		S	0698	0.9553	\$64.39		\$12.88
92317	Prescription of contact lens		S	0230	0.5945	\$40.07		\$8.02
92325	Modification of contact lens		S	0230	0.5945	\$40.07		\$8.02
92326	Replacement of contact lens		S	0698	0.9553	\$64.39		\$12.88
92340	Fitting of spectacles		E					
92341	Fitting of spectacles		E					
92342	Fitting of spectacles		E					
92362	Special spectacles fitting		S	0698	0.9553	\$64.39		\$12.88
92363	Special spectacles fitting		S	0230	0.5945	\$40.07		\$8.02
92364	Special spectacles fitting		S	0230	0.5945	\$40.07		\$8.02
92365	Special spectacles fitting		S	0230	0.5945	\$40.07		\$8.02
92368	Eye prosthesis service		S	0230	0.5945	\$40.07		\$8.02
92370	Repair & adjust spectacles		E					
92371	Repair & adjust spectacles		S	0230	0.5945	\$40.07		\$8.02

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
92633	Aud rehab postling hear loss		E	X	0365	1.2675	\$85.44	\$17.09
92640	Aud brainstem impit program		X	0364	0.47	\$31.68	\$7.06	\$6.34
92700	Ert procedure/service		S	0094	2.4553	\$165.50	\$46.29	\$33.10
92950	Heart/lung resuscitation opr		S	0094	2.4553	\$165.50	\$46.29	\$33.10
92953	Temporary external pacing		S	0679	5.5047	\$371.05	\$95.30	\$74.21
92960	Cardioversion electric, ext		S	0679	5.5047	\$371.05	\$95.30	\$74.21
92961	Cardioversion, electric, int		C					
92970	Cardioassist, internal		C					
92971	Cardioassist, external		C					
92973	Percut coronary thrombectomy		T	0088	40.9003	\$2,756.93	\$655.22	\$551.39
92974	Cath place, cardio brachyox		T	0103	17.8436	\$1,202.77		\$240.56
92977	Dissolve clot, heart vessel		C					
92978	Intravasc us, heart add-on		N	0676	2.3954	\$161.46		
92979	Intravasc us, heart add-on		N					
92980	Insert intracoronary stent		T	0104	84.7773	\$5,714.50		\$1,142.90
92981	Insert intracoronary stent		T	0104	84.7773	\$5,714.50		\$1,142.90
92982	Coronary artery dilation		T	0083	50.6809	\$3,416.20		\$683.24
92984	Coronary artery dilation		T	0083	50.6809	\$3,416.20		\$683.24
92986	Revision of aortic valve		T	0083	50.6809	\$3,416.20		\$683.24
92987	Revision of mitral valve		T	0083	50.6809	\$3,416.20		\$683.24
92990	Revision of pulmonary valve		T	0083	50.6809	\$3,416.20		\$683.24
92992	Revision of heart chamber		C					
92993	Revision of heart chamber		C					
92995	Coronary atherectomy		T	0082	93.3244	\$6,290.62		\$1,258.13
92996	Coronary atherectomy add-on		T	0082	93.3244	\$6,290.62		\$1,258.13
92997	Pul art balloon repr, percut		T	0083	50.6809	\$3,416.20		\$683.24
92998	Pul art balloon repr, percut		T	0083	50.6809	\$3,416.20		\$683.24
93000	Electrocardiogram, complete		M					
93005	Electrocardiogram, tracing		S	0099	0.394	\$26.56		\$5.92
93010	Electrocardiogram report		B					
93012	Transmission of ecg		N					
93014	Report on transmitted ecg		B					
93015	Cardiovascular stress test		B					
93016	Cardiovascular stress test		B					
93017	Cardiovascular stress test		B					
93018	Cardiovascular stress test		B	0100	2.6136	\$176.17	\$41.44	\$35.24
93024	Cardiac drug stress test		X	0100	2.6136	\$176.17	\$41.44	\$35.24
93025	Microvolt t-wave assess		X	0100	2.6136	\$176.17	\$41.44	\$35.24
93040	Rhythm ECG with report		B					
93041	Rhythm ECG, tracing		X	0035	0.232	\$15.64		\$3.13
93042	Rhythm ECG, report		B					
93224	ECG monitor/report, 24 hrs		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
92572	Slaggered spondaic word test		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92575	Sensorineural acuity test		X	0364	0.47	\$31.68	\$7.06	\$6.34
92576	Synthetic sentence test		X	0364	0.47	\$31.68	\$7.06	\$6.34
92577	Stenger test, speech		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92579	Visual audiometry (vra)		X	0365	1.2675	\$85.44	\$18.52	\$17.09
92582	Conditioning play audiometry		X	0365	1.2675	\$85.44	\$18.52	\$17.09
92583	Select picture audiometry		X	0364	0.47	\$31.68	\$7.06	\$6.34
92584	Electrocochleography		S	0216	2.6831	\$180.86	\$36.18	\$36.18
92585	Auditor evoke potent, compre		S	0216	2.6831	\$180.86		\$36.18
92586	Auditor evoke potent, limit		S	0218	1.1965	\$80.65	\$16.13	\$16.13
92587	Evoled auditory test		X	0363	0.9061	\$61.08	\$17.10	\$12.22
92588	Evoled auditory test	CH	X	0363	0.9061	\$61.08	\$17.10	\$12.22
92590	Hearing aid exam, one ear		E					
92591	Hearing aid exam, both ears		E					
92592	Hearing aid check, one ear		E					
92593	Hearing aid check, both ears		E					
92594	Electro hearing aid test, one		E					
92595	Electro hearing aid test, both		E					
92596	Ear protector evaluation		X	0364	0.47	\$31.68	\$7.06	\$6.34
92597	Oral speech device eval		A					
92601	Cochlear impit /rup exam < 7		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92602	Reprogram cochlear impit < 7		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92603	Cochlear impit /rup exam 7 >		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92604	Reprogram cochlear impit 7 >		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92605	Eval for nonspeech device rx		A					
92606	Non-speech device service		A					
92607	Ex for speech device rx, 1hr		A					
92608	Ex for speech device rx addl		A					
92609	Use of speech device service		A					
92610	Evaluate swallowing function		A					
92611	Motion fluoroscopy/swallow		A					
92612	Endoscopy swallow tst (fees)		A					
92613	Endoscopy swallow tst (fees)		B					
92614	Laryngoscopic sensory test		A					
92615	Eval laryngoscopic sense tst		E					
92616	Fees w/laryngeal sense test		A					
92617	Interprt fees/laryngeal test		E					
92620	Auditory function, 60 min		X	0365	1.2675	\$85.44	\$18.52	\$17.09
92621	Auditory function, + 15 min		N					
92625	Tinnitus assessment		X	0365	1.2675	\$85.44	\$18.52	\$17.09
92626	Eval aud rehab status		X	0366	1.6243	\$109.49	\$25.00	\$21.90
92627	Eval aud status rehab add-on		N					
92630	Aud rehab pre-ling hear loss		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
93312	Echo transesophageal	S	S	0270	8.8425	\$596.04	\$141.32	\$119.21
93313	Echo transesophageal	CH	S	0269	6.6903	\$450.97		\$90.20
93314	Echo transesophageal	N	N					
93315	Echo transesophageal	S	S	0270	8.8425	\$596.04	\$141.32	\$119.21
93316	Echo transesophageal	S	S	0270	8.8425	\$596.04	\$141.32	\$119.21
93317	Echo transesophageal	N	N					
93318	Echo transesophageal intraop	S	S	0270	8.8425	\$596.04	\$141.32	\$119.21
93320	Doppler echo exam, heart	N	N					
93321	Doppler echo exam, heart	N	N					
93325	Doppler color flow add-on	N	N					
93350	Stress tie only	S	S	0269	6.6903	\$450.97		\$90.20
93351	Stress tie complete	CH	S	0270	8.8425	\$596.04	\$141.32	\$119.21
93352	Admin ecg contrast agent	M	M					
93501	Right heart catheterization	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93503	Insert/replace heart catheter	T	T	0103	17.8436	\$1,202.77		\$240.56
93505	Biopsy of heart lining	T	T	0103	17.8436	\$1,202.77		\$240.56
93508	Cath placement, angiography	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93510	Left heart catheterization	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93511	Left heart catheterization	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93514	Left heart catheterization	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93524	Left heart catheterization	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93526	Rt & Lt heart catheters	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93527	Rt & Lt heart catheters	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93528	Rt & Lt heart catheters	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93529	Rt heart cath, congenital	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93530	Rt heart cath, congenital	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93532	R & l heart cath, congenital	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93533	R & l heart cath, congenital	T	T	0080	39.8099	\$2,683.43	\$838.92	\$536.69
93539	Injection, cardiac cath	N	N					
93540	Injection, cardiac cath	N	N					
93541	Injection for heart x-rays	N	N					
93542	Injection for heart x-rays	N	N					
93543	Injection for heart x-rays	N	N					
93544	Inject for coronary x-rays	N	N					
93545	Imaging, cardiac cath	N	N					
93556	Imaging, cardiac cath	N	N					
93561	Cardiac output measurement	N	N					
93562	Cardiac output measurement	N	N					
93571	Heart flow reserve measure	N	N					
93572	Heart flow reserve measure	N	N					
93580	Transcath closure of asd	T	T	0434	147.3728	\$9,933.81		\$1,986.77

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
93225	ECG monitor/record, 24 hrs	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93226	ECG monitor/record, 24 hrs	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93227	ECG monitor/review, 24 hrs	M	M					
93228	Remote 30 day ecg rev/report	M	M					
93229	Remote 30 day ecg tech supp	M	M					
93230	ECG monitor/record, 24 hrs	S	S	0209	11.4315	\$770.55	\$268.73	\$154.11
93231	ECG monitor/record, 24 hrs	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93232	ECG monitor/record, 24 hrs	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93233	ECG monitor/review, 24 hrs	M	M					
93235	ECG monitor/record, 24 hrs	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93236	ECG monitor/record, 24 hrs	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93237	ECG monitor/review, 24 hrs	M	M					
93268	ECG record/review	M	M					
93270	ECG recording	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93271	Ecg/monitoring and analysis	S	S	0692	1.6	\$107.85	\$21.57	\$21.57
93272	Ecg/review, interpret only	M	M					
93278	ECG/signal-averaged	CH	X	0035	0.232	\$15.64		\$3.13
93279	Pm device progr eval, singl	S	S	0690	0.3506	\$23.63	\$8.67	\$4.73
93280	Pm device progr eval, dual	S	S	0690	0.3506	\$23.63	\$8.67	\$4.73
93281	Pm device progr eval, multi	S	S	0690	0.3506	\$23.63	\$8.67	\$4.73
93282	Icd device progr eval, 1 singl	S	S	0689	0.568	\$38.29	\$8.67	\$7.66
93283	Icd device progr eval, dual	S	S	0689	0.568	\$38.29	\$8.67	\$7.66
93284	Icd device progr eval, mult	S	S	0689	0.568	\$38.29	\$8.67	\$7.66
93285	Iir device eval progr	N	N	0690	0.3506	\$23.63	\$8.67	\$4.73
93286	Pre-op lcd device eval	N	N					
93288	Pm device eval in person	S	S	0690	0.3506	\$23.63	\$8.67	\$4.73
93289	Icd device interrogate	S	S	0689	0.568	\$38.29	\$8.67	\$7.66
93290	Icm device eval	S	S	0690	0.3506	\$23.63	\$8.67	\$4.73
93291	Iir device interrogate	S	S	0690	0.3506	\$23.63	\$8.67	\$4.73
93292	Wcd device interrogate	S	S	0689	0.568	\$38.29	\$8.67	\$7.66
93293	Pm phone r-strip device eval	M	M					
93294	Pm device interrogat remote	M	M					
93295	Icd device interrogat remote	M	M					
93296	Pm/icc remote tech serv	S	S	0689	0.568	\$38.29	\$8.67	\$7.66
93297	Icm device interrogat remote	M	M					
93298	Iir device interrogat remote	M	M					
93299	Icm/ir remote tech serv	CH	S	0689	0.568	\$38.29	\$8.67	\$7.66
93303	Echo transthoracic	CH	S	0270	8.8425	\$596.04	\$141.32	\$119.21
93304	Echo transthoracic	CH	S	0269	6.6903	\$450.97		\$90.20
93306	Tte w/doppler, complete	S	S	0269	6.6903	\$450.97		\$90.20
93307	Tte w/o doppler, complete	S	S	0697	3.9223	\$264.39		\$52.88
93308	Tte, 1-up or lmtd	S	S	0697	3.9223	\$264.39		\$52.88

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
93875	Extracranial study		S	0096	1.6075	\$108.36	\$37.42	\$31.02
93880	Extracranial study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93882	Extracranial study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93886	Intracranial study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93888	Intracranial study		S	0265	0.9267	\$62.47	\$22.34	\$12.50
93890	Tcd, vasoreactivity study		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93892	Tcd, emboli detect w/o inj		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93893	Tcd, emboli detect w/inj		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93922	Extremity study	CH	S	0097	0.9848	\$66.38	\$23.79	\$13.28
93923	Extremity study		S	0096	1.6075	\$108.36	\$37.42	\$21.68
93924	Extremity study		S	0096	1.6075	\$108.36	\$37.42	\$21.68
93925	Lower extremity study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93926	Lower extremity study		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93930	Upper extremity study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93931	Upper extremity study		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93965	Extremity study		S	0096	1.6075	\$108.36	\$37.42	\$21.68
93970	Extremity study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93971	Extremity study		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93975	Vascular study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93976	Vascular study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93977	Vascular study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93979	Vascular study		S	0266	1.4434	\$97.29	\$37.61	\$19.46
93980	Pentle vascular study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93981	Pentle vascular study		S	0267	2.3005	\$155.07	\$60.50	\$31.02
93982	Aneurysm pressure sens study		S	0097	0.9848	\$66.38	\$23.79	\$13.28
93990	Doppler flow testing		S	0266	1.4434	\$97.29	\$37.61	\$19.46
94002	Vent mgmt input, init day		S	0079	3.0865	\$208.05	\$68.05	\$11.46
94003	Vent mgmt input, subq day		S	0079	3.0865	\$208.05	\$68.05	\$11.46
94004	Vent mgmt, nt per day		B					
94005	Home vent mgmt supervision		M					
94010	Breathing capacity test		X	0368	0.8495	\$57.26	\$20.93	\$11.46
94011	Up to 2 yrs old, spirometry	NI	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94012	= 2 yrs, spirometry, w/tiltator	NI	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94013	= 2 yrs, lung volumes	NI	X	0369	2.6228	\$176.79	\$42.29	\$5.36
94014	Patient recorded spirometry		X	0367	0.5968	\$40.23	\$13.76	\$8.05
94015	Patient recorded spirometry		X	0367	0.5968	\$40.23	\$13.76	\$8.05
94016	Review patient spirometry		A					
94080	Evaluation of wheezing		S	0078	1.4142	\$95.33	\$31.07	\$19.07
94070	Evaluation of wheezing		X	0369	2.6228	\$176.79	\$42.29	\$5.36
94150	Vital capacity test		X	0367	0.5968	\$40.23	\$13.76	\$8.05
94200	Lung function test (MBC/MNV)		X	0367	0.5968	\$40.23	\$13.76	\$8.05
94240	Residual lung capacity		X	0368	0.8495	\$57.26	\$20.93	\$11.46
94250	Expired gas collection		X	0368	0.8495	\$57.26	\$20.93	\$11.46

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
93581	Transcath closure of vsd	T	0434	147.3728	\$9,933.81	\$1,986.77		
93600	Bundle of His recording	S	0084	10.811	\$728.73	\$145.75		
93602	Intra-atrial recording	S	0084	10.811	\$728.73	\$145.75		
93603	Right ventricular recording	S	0084	10.811	\$728.73	\$145.75		
93609	Map tachycardia, add-on	N						
93610	Intra-atrial pacing	S	0084	10.811	\$728.73	\$145.75		
93612	Intraventricular pacing	S	0084	10.811	\$728.73	\$145.75		
93613	Electrophys map 3d, add-on	N						
93615	Esophageal recording	S	0084	10.811	\$728.73	\$145.75		
93616	Esophageal recording	S	0084	10.811	\$728.73	\$145.75		
93618	Heart rhythm pacing	S	0084	10.811	\$728.73	\$145.75		
93619	Electrophysiology evaluation	O3	0085	52.3882	\$3,531.28	\$706.26		
93620	Electrophysiology evaluation	O3	0085	52.3882	\$3,531.28	\$706.26		
93621	Electrophysiology evaluation	N						
93622	Electrophysiology evaluation	N						
93623	Stimulation, pacing heart	N						
93624	Electrophysiology study	N	0085	52.3882	\$3,531.28	\$706.26		
93631	Heart pacing, mapping	N						
93640	Evaluation heart device	N						
93641	Electrophysiology evaluation	N						
93642	Electrophysiology evaluation	N	0084	10.811	\$728.73	\$145.75		
93650	Ablate heart dysrhythm focus	O3	0085	52.3882	\$3,531.28	\$706.26		
93651	Ablate heart dysrhythm focus	O3	0086	106.9978	\$7,212.29	\$1,442.46		
93652	Ablate heart dysrhythm focus	O3	0086	106.9978	\$7,212.29	\$1,442.46		
93660	Tilt table evaluation	S	0101	4.3705	\$294.60	\$58.92	\$100.24	
93662	Intracardiac eeg (ice)	N						
93668	Peripheral vascular rehab	E						
93701	Bioimpedance, cv analysis	S	0099	0.394	\$26.56	\$5.32		
93720	Total body plethysmography	B						
93721	Plethysmography tracing	X	0368	0.8495	\$57.26	\$20.93		
93722	Plethysmography report	B						
93724	Analyze pacemaker system	S	0690	0.3506	\$23.63	\$8.67		
93740	Temperature gradient studies	X	0368	0.8495	\$57.26	\$20.93		
93745	Set-up cardiovert-defibrill	S	0689	0.568	\$38.29	\$7.66		
93750	Interrogation vad, in person	NI	S	0692	1.6	\$107.85	\$21.57	
93770	Measure venous pressure	N						
93784	Ambulatory BP monitoring	E						
93786	Ambulatory BP recording	S	0097	0.9848	\$66.38	\$23.79		
93788	Ambulatory BP analysis	S	0097	0.9848	\$66.38	\$23.79		
93790	Review/report BP recording	M						
93797	Cardiac rehab	S	0095	0.5691	\$38.36	\$7.68		
93798	Cardiac rehab/monitor	S	0095	0.5691	\$38.36	\$7.68		
93799	Cardiovascular procedure	S	0097	0.9848	\$66.38	\$23.79		

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
95028	Id allergy test-delayed type	X	X	0381	0.4312	\$29.07		\$5.82
95044	Allergy patch tests	X	X	0381	0.4312	\$29.07		\$5.82
95052	Photo patch test	X	X	0381	0.4312	\$29.07		\$5.82
95056	Photosensitivity tests	X	X	0370	1.4598	\$98.40		\$19.68
95060	Eye allergy tests	X	X	0370	1.4598	\$98.40		\$19.68
95065	Nose allergy test	X	X	0381	0.4312	\$29.07		\$5.82
95070	Bronchial allergy tests	X	X	0369	2.6228	\$176.79	\$42.29	\$35.36
95071	Immunotherapy tests	X	X	0369	2.6228	\$176.79	\$42.29	\$35.36
95075	Ingestion challenge test	X	X	0361	4.0639	\$275.28	\$83.23	\$55.06
95115	Immunotherapy, one injection	S	S	0436	0.3809	\$25.67		\$5.14
95117	Immunotherapy injections	S	S	0436	0.3809	\$25.67		\$5.14
95120	Immunotherapy, one injection	E	E					
95120	Immunotherapy, many	E	E					
95125	antigens	E	E					
95130	Immunotherapy, insect venom	E	E					
95131	Immunotherapy, insect venoms	E	E					
95132	Immunotherapy, insect venoms	E	E					
95133	Immunotherapy, insect venoms	E	E					
95134	Immunotherapy, insect venoms	E	E					
95144	Antigen therapy services	S	S	0437	0.5555	\$37.44		\$7.49
95145	Antigen therapy services	CH	S	0437	0.5555	\$37.44		\$7.49
95146	Antigen therapy services	S	S	0438	1.1229	\$75.69		\$15.14
95147	Antigen therapy services	S	S	0438	1.1229	\$75.69		\$15.14
95148	Antigen therapy services	S	S	0437	0.5555	\$37.44		\$7.49
95149	Antigen therapy services	CH	S	0437	0.5555	\$37.44		\$7.49
95165	Antigen therapy services	S	S	0436	0.3809	\$25.67		\$5.14
95170	Antigen therapy services	CH	S	0437	0.5555	\$37.44		\$7.49
95180	Rapid desensitization	X	X	0370	1.4598	\$98.40		\$19.68
95199	Allergy immunology services	X	X	0381	0.4312	\$29.07		\$5.82
95250	Glucose monitoring, cont	V	V	0607	1.663	\$113.44		\$22.69
95251	Gluc monitor, cont, phys i&r	B	B					
95803	Actigraphy testing	S	S	0218	1.1965	\$80.65		\$16.13
95805	Multiple sleep latency test	S	S	0209	11.4315	\$770.55	\$268.73	\$154.11
95806	Sleep study unatt&resp efft	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95807	Sleep study, attended	S	S	0209	11.4315	\$770.55	\$268.73	\$154.11
95808	Polysomnography, 1-3	S	S	0209	11.4315	\$770.55	\$268.73	\$154.11
95810	Polysomnography, 4 or more	S	S	0209	11.4315	\$770.55	\$268.73	\$154.11
95811	Polysomnography w/cpap	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95812	Eeg, 41-60 minutes	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95813	Eeg, over 1 hour	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95816	Eeg, awake and drowsy	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95819	Eeg, awake and asleep	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95822	Eeg, coma or sleep only	S	S	0213	2.4043	\$162.06	\$53.58	\$32.42
95824	Eeg, cerebral death only	S	S	0216	2.6831	\$180.86		\$36.18

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
94260	Thoracic gas volume	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94350	Lung nitrogen washout curve	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94360	Measure airflow resistance	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
94370	Breath airflow closing volume	CH	X	0035	0.232	\$15.64		\$3.13
94375	Respiratory flow volume loop	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94400	CO2 breathing response curve	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
94450	Hypoxia response curve	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94452	Hast w/oreport	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94453	Surfactant admin thru tube	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94610	Pulmonary stress test/simple	S	S	0077	0.4058	\$27.35	\$7.74	\$5.47
94620	Pulmonary stress test/complex	X	X	0369	2.6228	\$176.79	\$42.29	\$35.36
94640	Airway inhalation treatment	S	S	0077	0.4058	\$27.35	\$7.74	\$5.47
94642	Aerosol inhalation treatment	S	S	0078	1.4142	\$95.33		\$19.07
94644	Cbr, each addl hour	X	X	0340	0.6693	\$45.11		\$9.03
94645	Pos airway pressure, CPAP	S	S	0078	1.4142	\$95.33		\$19.07
94660	Neg press ventilation, cnp	S	S	0079	3.0865	\$208.05		\$41.61
94664	Evaluate pt use of inhaler	S	S	0077	0.4058	\$27.35	\$7.74	\$5.47
94667	Chest wall manipulation	S	S	0077	0.4058	\$27.35	\$7.74	\$5.47
94668	Chest wall manipulation	S	S	0077	0.4058	\$27.35	\$7.74	\$5.47
94680	Exhaled air analysis, o2	CH	X	0369	2.6228	\$176.79	\$42.29	\$35.36
94681	Exhaled air analysis, o2/co2	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94690	Exhaled air analysis	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
94720	Monoxide diffusing capacity	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94725	Membrane diffusion capacity	X	X	0368	0.8495	\$57.26	\$20.93	\$11.46
94750	Pulmonary compliance study	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
94760	Measure blood oxygen level	N	N					
94761	Measure blood oxygen level	N	N					
94762	Measure blood oxygen level	Q1	Q1	0097	0.9848	\$66.38	\$23.79	\$13.28
94770	Exhaled carbon dioxide test	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
94772	Breath recording, infant	X	X	0369	2.6228	\$176.79	\$42.29	\$35.36
94774	Ped home apnea rec, compl	B	B					
94775	Ped home apnea rec, h-up	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
94776	Ped home apnea rec, downid	S	S	0097	0.9848	\$66.38	\$23.79	\$13.28
94777	Ped home apnea rec, report	B	B					
94799	Pulmonary service/procedure	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
95004	Percut allergy skin tests	X	X	0381	0.4312	\$29.07		\$5.82
95010	Percut allergy titrate test	X	X	0381	0.4312	\$29.07		\$5.82
95012	Exhaled nitric oxide meas	X	X	0367	0.5968	\$40.23	\$13.76	\$8.05
95015	Id allergy titrate-drug/bug	X	X	0381	0.4312	\$29.07		\$5.82
95024	Id allergy test, drug/bug	X	X	0381	0.4312	\$29.07		\$5.82
95027	Id allergy titrate-airborne	X	X	0381	0.4312	\$29.07		\$5.82

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010										
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment		
95951	EEG monitoring/monitoring	S	0209	11.4315	\$770.55	\$268.73	\$154.11	\$154.11		
95952	EEG monitoring/monitoring	S	0209	11.4315	\$770.55	\$268.73	\$154.11	\$154.11		
95953	EEG monitoring/computer	S	0209	11.4315	\$770.55	\$268.73	\$154.11	\$154.11		
95954	EEG monitoring/giving drugs	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95955	EEG during surgery	N								
95956	Eeg monitoring, cable/radio	S	0209	11.4315	\$770.55	\$268.73	\$154.11	\$154.11		
95957	EEG digital analysis	N								
95958	EEG monitoring/function test	S	0213	2.4043	\$162.06	\$53.58	\$32.42	\$32.42		
95961	Electrode stimulation, brain	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95962	Electrode stim. brain add-on	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95965	Meg, spontaneous	S	0067	52.9891	\$3,571.78		\$714.36	\$714.36		
95966	Meg, evoked, single	S	0065	14.2808	\$962.61		\$192.53	\$192.53		
95967	Meg, evoked, each addl	S	0065	14.2808	\$962.61		\$192.53	\$192.53		
95970	Analyze neurostim, no prog	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95971	Analyze neurostim, simple	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95972	Analyze neurostim, complex	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95973	Analyze neurostim, complex	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95974	Cranial neurostim, complex	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95975	Cranial neurostim, complex	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95976	Analyze neurostim brain/ih	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95977	Analyze neurostim brain add-on	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95979	Analyz neurostim brain add-on	N								
95980	lo anal gast n-stim init	N								
95981	lo anal gast n-stim subseq	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95982	lo ga n-stim subseq w/reprog	S	0692	1.6	\$107.85		\$21.57	\$21.57		
95990	Spin/brain pump refl & main	CH	S	0439	1.8808	\$126.78	\$25.36	\$25.36		
95991	Spin/brain pump refl & main	CH	S	0439	1.8808	\$126.78	\$25.36	\$25.36		
95992	Caralith repositioning proc	CH	E							
95999	Neurological procedure	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
96000	Motion analysis, video/qd	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
96001	Motion test w/ft press meas	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
96002	Dynamic surface emg	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
96003	Dynamic fine wire emg	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
96004	Phys review of motion tests	B								
96020	Functional brain mapping	N								
96040	Genetic counseling, 30 min	B								
96101	Psycho testing by psych/phys	Q3	0382	2.6683	\$179.86		\$35.98	\$35.98		
96102	Psycho testing by technician	Q3	0382	2.6683	\$179.86		\$35.98	\$35.98		
96103	Psycho testing admin by comp	Q3	0373	0.9758	\$85.77		\$13.16	\$13.16		
96105	Assessment of aphasia	A								
96110	Developmental test, lim	Q3	0373	0.9758	\$85.77		\$13.16	\$13.16		
96111	Developmental test, extend	CH	Q3	0373	\$85.77		\$13.16	\$13.16		
96116	Neurobehavioral status exam	Q3	0382	2.6683	\$179.86		\$35.98	\$35.98		
96118	Neuropsych tst by psych/phys	Q3	0382	2.6683	\$179.86		\$35.98	\$35.98		

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010										
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95827	Eeg, all night recording	S	0213	2.4043	\$162.06	\$53.58	\$32.42	\$32.42		
95829	Surgery electrocorticogram	N								
95830	Insert electrodes for EEG	B								
95831	Limb muscle testing, manual	A								
95832	Hand muscle testing, manual	A								
95833	Body muscle testing, manual	A								
95834	Body muscle testing, manual	A								
95851	Range of motion measurements	A								
95852	Range of motion measurements	A								
95857	Tension test	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95860	Muscle test, one limb	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95861	Muscle test, 2 limbs	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95863	Muscle test, 3 limbs	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95864	Muscle test, 4 limbs	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95865	Muscle test, larynx	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95866	Muscle test, hemidiaphragm	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95867	Muscle test cran nerve unilat	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95868	Muscle test cran nerve bilat	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95869	Muscle test, thor paraspinal	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95870	Muscle test, nonparaspinal	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95872	Muscle test, one fiber	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95873	Guide nerv destr, elec stim	N								
95874	Guide nerv destr, needle emg	N								
95875	Limb exercise test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95900	Motor nerve conduction test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95903	Motor nerve conduction test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95904	Sense nerve conduction test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95905	Motor/sense nve conduct test	N								
95920	Intrap nerve test add-on	N								
95921	Autonomic nerv function test	CH	S	0218	1.1965	\$80.65	\$16.13	\$16.13		
95922	Autonomic nerv function test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95923	Autonomic nerv function test	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95925	Somatosensory testing	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95926	Somatosensory testing	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95927	Somatosensory testing	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95928	C motor evoked, uppr limbs	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95929	C motor evoked, lwr limbs	S	0218	1.1965	\$80.65		\$16.13	\$16.13		
95930	Visual evoked potential test	S	0216	2.6831	\$180.86		\$36.18	\$36.18		
95933	Blink reflex test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95934	H-reflex test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95936	H-reflex test	S	0215	0.6135	\$41.35		\$8.27	\$8.27		
95937	Neuromuscular junction test	S	0218	1.1965	\$80.65		\$16.13	\$16.13		

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
96523	Irig drug delivery device	CH	S	0624	0.6132	\$41.33	\$12.65	\$8.27
96542	Chemotherapy injection	CH	S	0436	1.1229	\$75.69		\$15.14
96549	Chemotherapy, unspecified	CH	T	0016	2.7982	\$188.62		\$37.73
96570	Photodynamic tx, skin	CH	T	0015	1.5412	\$103.89		\$20.78
96571	Photodynamic tx, 30 min add-on	CH	T	0015	1.5412	\$103.89		\$20.78
96571	Photodynamic tx, addl 15 min	CH	T	0015	1.5412	\$103.89		\$20.78
96900	Ultrasound therapy	S	S	0001	0.5302	\$35.74		\$7.15
96902	Trichogram	N	N					
96904	Whole body photography	N	N					
96910	Photochemotherapy with UV-B	S	S	0001	0.5302	\$35.74		\$7.15
96912	Photochemotherapy with UV-A	S	S	0001	0.5302	\$35.74		\$7.15
96913	Photochemotherapy, UV-A or B	S	S	0683	2.679	\$180.58		\$36.12
96920	Laser tx, skin < 250 sq cm	T	T	0015	1.5412	\$103.89		\$20.78
96921	Laser tx, skin 250-500 sq cm	T	T	0015	1.5412	\$103.89		\$20.78
96922	Laser tx, skin > 500 sq cm	T	T	0015	1.5412	\$103.89		\$20.78
96999	Dermatological procedure	T	T	0012	0.4436	\$29.90		\$5.98
97001	Pt evaluation	A	A					
97002	Pt re-evaluation	A	A					
97003	Ot evaluation	A	A					
97004	Ot re-evaluation	A	A					
97005	Athletic train eval	E	E					
97006	Athletic train reeval	E	E					
97010	Hot or cold packs therapy	A	A					
97012	Mechanical traction therapy	A	A					
97014	Electric stimulation therapy	E	E					
97016	Vasopneumatic device therapy	A	A					
97018	Paraffin bath therapy	A	A					
97022	Whirlpool therapy	A	A					
97024	Diathermy eq, microwave	A	A					
97026	Infrared therapy	A	A					
97028	Ultraviolet therapy	A	A					
97032	Electrical stimulation	A	A					
97033	Electric current therapy	A	A					
97034	Contrast bath therapy	A	A					
97035	Ultrasound therapy	A	A					
97036	Hydrotherapy	A	A					
97039	Physical therapy treatment	A	A					
97110	Therapeutic exercises	A	A					
97112	Neuromuscular reeducation	A	A					
97113	Aquatic therapy/exercises	A	A					
97116	Gait training therapy	A	A					
97124	Massage therapy	A	A					
97139	Physical medicine procedure	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
96119	Neuropsych testing by tec	CH	O3	0382	2.6683	\$179.86	\$35.98	\$35.98
96120	Neuropsych tst admin w/comp	CH	O3	0382	2.6683	\$179.86	\$35.98	\$35.98
96125	Cognitive test by hc pro	A	A					
96150	Assess hlt/behav, init	O3	O3	0432	0.6013	\$40.53		\$8.11
96151	Assess hlt/behav, subseq	O3	O3	0432	0.6013	\$40.53		\$8.11
96152	Intervene hlt/behav, indiv	O3	O3	0432	0.6013	\$40.53		\$8.11
96153	Intervene hlt/behav, group	O3	O3	0432	0.6013	\$40.53		\$8.11
96154	Interv hlt/behav, fam w/pt	O3	O3	0432	0.6013	\$40.53		\$8.11
96155	Interv hlt/behav, fam no pt	E	E					
96360	Hydration iv infusion, init	S	S	0438	1.1229	\$75.69		\$15.14
96361	Hydrate iv infusion, add-on	S	S	0436	0.3809	\$25.67		\$5.14
96365	Ther/proph/diag iv inf, init	S	S	0439	1.8808	\$126.78		\$25.36
96366	Ther/proph/diag iv inf addon	S	S	0436	0.3809	\$25.67		\$5.14
96367	Tx/proph/dg addl seq iv inf	S	S	0437	0.5555	\$37.44		\$7.49
96368	Ther/diag concurrent inf	N	N					
96369	Sc ther infusion, up to 1 hr	CH	S	0439	1.8808	\$126.78		\$25.36
96370	Sc ther infusion, addl hr	S	S	0437	0.5555	\$37.44		\$7.49
96371	Sc ther infusion, reset pump	S	S	0436	0.3809	\$25.67		\$5.14
96372	Ther/proph/diag inj, sc/fm	S	S	0436	0.3809	\$25.67		\$5.14
96373	Ther/proph/diag inj, ia	S	S	0437	0.5555	\$37.44		\$7.49
96374	Ther/proph/diag inj, iv push	S	S	0437	0.5555	\$37.44		\$7.49
96375	Tx/pro/ox inj new drug addon	S	S	0437	0.5555	\$37.44		\$7.49
96376	Tx/pro/ox inj new drug addn	N	N					
96379	Ther/proph/diag inj/inf proc	S	S	0436	0.3809	\$25.67		\$5.14
96401	Chemo, anti-neopl, sq/fm	S	S	0437	0.5555	\$37.44		\$7.49
96402	Chemo, anti-neopl, sq/fm	S	S	0437	0.5555	\$37.44		\$7.49
96405	Chemo intravesical, up to 7	S	S	0437	0.5555	\$37.44		\$7.49
96406	Chemo intravesical over 7	CH	S	0439	1.8808	\$126.78		\$25.36
96409	Chemo, iv push, singl drug	S	S	0439	1.8808	\$126.78		\$25.36
96411	Chemo, iv push, addl drug	S	S	0438	1.1229	\$75.69		\$15.14
96413	Chemo, iv infusion, 1 hr	S	S	0440	3.2632	\$219.96		\$44.00
96415	Chemo, iv infusion, addl hr	S	S	0437	0.5555	\$37.44		\$7.49
96416	Chemo prolong infuse w/pump	S	S	0440	3.2632	\$219.96		\$44.00
96420	Chemo iv infus each addl seq	S	S	0438	1.1229	\$75.69		\$15.14
96422	Chemo ia, push technique	CH	S	0438	1.1229	\$75.69		\$15.14
96423	Chemo ia infusion up to 1 hr	S	S	0440	3.2632	\$219.96		\$44.00
96425	Chemo ia infuse each addl hr	S	S	0438	1.1229	\$75.69		\$15.14
96425	Chemotherapy/infusion method	S	S	0440	3.2632	\$219.96		\$44.00
96440	Chemotherapy, intracavitary	CH	S	0439	1.8808	\$126.78		\$25.36
96445	Chemotherapy, intracavitary	S	S	0440	3.2632	\$219.96		\$44.00
96450	Chemotherapy, into CNS	S	S	0440	3.2632	\$219.96		\$44.00
96521	Refill/maint, portable pump	CH	S	0439	1.8808	\$126.78		\$25.36
96522	Refill/maint pump/resvr syst	S	S	0439	1.8808	\$126.78		\$25.36

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
98969	Online service by hc pro		E					
99000	Specimen handling		E					
99001	Specimen handling		E					
99002	Device handling		B					
99024	Postop follow-up visit		B					
99026	In-hospital on call service		E					
99027	Out-of-hosp on call service		E					
99050	Medical services after hrs		B					
99051	Med serv, eve/wknd/holiday		B					
99053	Med serv 10pm-8am, 24 hr fac		B					
99056	Med service out of office		B					
99058	Office emergency care		B					
99060	Out of office emerg med serv		B					
99070	Special supplies		B					
99071	Patient education materials		B					
99075	Medical testimony		E					
99078	Group health education		N					
99080	Special reports or forms		B					
99082	Unusual physician travel		B					
99090	Computer data analysis		B					
99091	Collect/review data from pt		N					
99100	Special anesthesia service		B					
99116	Anesthesia with hypothermia		B					
99135	Special anesthesia procedure		B					
99140	Emergency anesthesia		B					
99143	Mod cs by same phys, < 5 yrs		N					
99144	Mod cs by same phys, 5 yrs +		N					
99145	Mod cs by same phys add-on		N					
99148	Mod cs diff phys < 5 yrs		N					
99149	Mod cs diff phys 5 yrs +		N					
99170	Anogenital exam, child		T	0191	0.132	\$8.90	\$2.09	\$1.78
99172	Ocular function screen		E					
99173	Visual acuity screen		E					
99174	Ocular photoscreening		E					
99175	Induction of vomiting		N					
99183	Hyperbaric oxygen therapy		B					
99185	Regional hypothermia	CH	D					
99186	Total body hypothermia	CH	D					
99190	Special pump services		C					
99191	Special pump services		C					
99192	Special pump services		C					
99195	Phlebotomy		X	0624	0.6132	\$41.33	\$12.65	\$8.27

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
97140	Manual therapy		A					
97150	Group therapeutic procedures		A					
97530	Therapeutic activities		A					
97532	Cognitive skills development		A					
97533	Sensory integration		A					
97535	Self care mgmt training		A					
97537	Community/work reintegration		A					
97542	Wheelchair mgmt training		A					
97545	Work hardening		A					
97546	Work hardening add-on		A					
97597	Active wound care/20 cm or <		T	0015	1.5412	\$103.89		\$20.78
97598	Active wound care > 20 cm		T	0015	1.5412	\$103.89		\$20.78
97602	Wound(s) care non-selective		T	0013	0.8789	\$59.24		\$11.85
97605	Neg press wound bx, < 50 cm		T	0013	0.8789	\$59.24		\$11.85
97606	Neg press wound bx, > 50 cm	CH	T	0015	1.5412	\$103.89		\$20.78
97750	Physical performance test		A					
97755	Assistive technology assess		A					
97760	Orthotic mgmt and training		A					
97761	Prosthetic training		A					
97762	C/o for orthotic/prosth use		A					
97799	Physical medicine procedure		A					
97802	Medical nutrition, indiv, in		A					
97803	Med nutrition, indiv, subseq		A					
97804	Medical nutrition, group		A					
97810	Acupunct w/o stimu 15 min		E					
97811	Acupunct w/o stimu addl 15m		E					
97813	Acupunct w/stimu 15 min		E					
97814	Acupunct w/stimu addl 15m		E					
98925	Osteopathic manipulation		S	0060	0.3808	\$25.67		\$5.14
98926	Osteopathic manipulation		S	0060	0.3808	\$25.67		\$5.14
98927	Osteopathic manipulation		S	0060	0.3808	\$25.67		\$5.14
98928	Osteopathic manipulation		S	0060	0.3808	\$25.67		\$5.14
98929	Osteopathic manipulation		S	0060	0.3808	\$25.67		\$5.14
98940	Chiropractic manipulation		S	0060	0.3808	\$25.67		\$5.14
98941	Chiropractic manipulation		S	0060	0.3808	\$25.67		\$5.14
98942	Chiropractic manipulation		S	0060	0.3808	\$25.67		\$5.14
98943	Chiropractic manipulation		E					
98960	Self-mgmt educ & train, 1 pt		E					
98961	Self-mgmt educ/train, 2-4 pt		E					
98962	Self-mgmt educ/train, 5-8 pt		E					
98966	Hc pro phone call 5-10 min		E					
98967	Hc pro phone call 11-20 min		E					
98968	Hc pro phone call 21-30 min		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
99392	Critical care, addl 30 min		N					
99304	Nursing facility care, init		B					
99305	Nursing facility care, init		B					
99306	Nursing facility care, init		B					
99307	Nursing facility care, subseq		B					
99308	Nursing facility care, subseq		B					
99309	Nursing facility care, subseq		B					
99310	Nursing facility care, subseq		B					
99311	Nursing facility care, subseq		B					
99312	Nursing facility care, subseq		B					
99313	Nursing facility care, subseq		B					
99314	Nursing facility care, subseq		B					
99315	Nursing facility care, subseq		B					
99316	Nursing facility care, subseq		B					
99317	Nursing facility care, subseq		B					
99318	Nursing facility care, subseq		B					
99319	Nursing facility care, subseq		B					
99320	Nursing facility care, subseq		B					
99321	Nursing facility care, subseq		B					
99322	Nursing facility care, subseq		B					
99323	Nursing facility care, subseq		B					
99324	Nursing facility care, subseq		B					
99325	Nursing facility care, subseq		B					
99326	Nursing facility care, subseq		B					
99327	Nursing facility care, subseq		B					
99328	Nursing facility care, subseq		B					
99329	Nursing facility care, subseq		B					
99330	Nursing facility care, subseq		B					
99331	Nursing facility care, subseq		B					
99332	Nursing facility care, subseq		B					
99333	Nursing facility care, subseq		B					
99334	Nursing facility care, subseq		B					
99335	Nursing facility care, subseq		B					
99336	Nursing facility care, subseq		B					
99337	Nursing facility care, subseq		B					
99338	Nursing facility care, subseq		B					
99339	Nursing facility care, subseq		B					
99340	Nursing facility care, subseq		B					
99341	Nursing facility care, subseq		B					
99342	Nursing facility care, subseq		B					
99343	Nursing facility care, subseq		B					
99344	Nursing facility care, subseq		B					
99345	Nursing facility care, subseq		B					
99346	Nursing facility care, subseq		B					
99347	Nursing facility care, subseq		B					
99348	Nursing facility care, subseq		B					
99349	Nursing facility care, subseq		B					
99350	Nursing facility care, subseq		B					
99351	Nursing facility care, subseq		B					
99352	Nursing facility care, subseq		B					
99353	Nursing facility care, subseq		B					
99354	Nursing facility care, subseq		B					
99355	Nursing facility care, subseq		B					
99356	Nursing facility care, subseq		B					
99357	Nursing facility care, subseq		B					
99358	Nursing facility care, subseq		B					
99359	Nursing facility care, subseq		B					
99360	Nursing facility care, subseq		B					
99361	Nursing facility care, subseq		B					
99362	Nursing facility care, subseq		B					
99363	Nursing facility care, subseq		B					
99364	Nursing facility care, subseq		B					
99365	Nursing facility care, subseq		B					
99366	Nursing facility care, subseq		B					
99367	Nursing facility care, subseq		B					
99368	Nursing facility care, subseq		B					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
99199	Special service/proc/report		B	0604	0.8593	\$57.92		\$11.59
99201	Office/outpatient visit, new		V	0605	1.0337	\$69.68		\$13.94
99202	Office/outpatient visit, new		V	0605	1.0337	\$69.68		\$13.94
99203	Office/outpatient visit, new		V	0606	1.3222	\$89.12		\$17.83
99204	Office/outpatient visit, new		V	0607	1.663	\$113.44		\$22.69
99205	Office/outpatient visit, new		Q3	0608	2.4853	\$167.52		\$33.51
99211	Office/outpatient visit, est		V	0604	0.8593	\$57.92		\$11.59
99212	Office/outpatient visit, est		V	0605	1.0337	\$69.68		\$13.94
99213	Office/outpatient visit, est		V	0606	1.3222	\$89.12		\$17.83
99214	Office/outpatient visit, est		V	0606	1.3222	\$89.12		\$17.83
99215	Office/outpatient visit, est		Q3	0607	1.663	\$113.44		\$22.69
99217	Observation care discharge		B					
99218	Observation care		B					
99219	Observation care		B					
99220	Observation care		B					
99221	Initial hospital care		B					
99222	Initial hospital care		B					
99223	Initial hospital care		B					
99224	Initial hospital care		B					
99231	Subsequent hospital care		B					
99232	Subsequent hospital care		B					
99233	Subsequent hospital care		B					
99234	Subsequent hospital care		B					
99235	Subsequent hospital care		B					
99236	Subsequent hospital care		B					
99237	Subsequent hospital care		B					
99238	Subsequent hospital care		B					
99239	Subsequent hospital care		B					
99241	Office consultation		CH					
99242	Office consultation		CH					
99243	Office consultation		CH					
99244	Office consultation		CH					
99245	Office consultation		CH					
99251	Inpatient consultation		CH					
99252	Inpatient consultation		CH					
99253	Inpatient consultation		CH					
99254	Inpatient consultation		CH					
99255	Inpatient consultation		CH					
99281	Emergency dept visit		V	0609	0.7886	\$53.16	\$12.70	\$10.64
99282	Emergency dept visit		V	0613	1.3033	\$87.85	\$21.06	\$17.57
99283	Emergency dept visit		V	0614	2.0796	\$140.18	\$34.50	\$28.04
99284	Emergency dept visit		Q3	0615	3.3109	\$223.17	\$48.49	\$44.64
99285	Emergency dept visit		Q3	0616	4.8917	\$329.73	\$72.86	\$65.95
99288	Direct advanced life support		B					
99291	Critical care, first hour		Q3	0617	7.3492	\$495.38	\$111.59	\$99.08

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
99464	Attendance at delivery	N	S	0094	2.4553	\$165.50	\$46.29	\$33.10
99465	Nb resuscitation	N	N					
99466	Ped crit care transport	N	N					
99467	Ped crit care transport addl	N	N					
99468	Neonate crit care, initial	C	C					
99469	Neonate crit care, subseq	C	C					
99471	Ped critical care, initial	C	C					
99472	Ped critical care, subseq	C	C					
99475	Ped crit care age 2-5, init	C	C					
99476	Ped crit care age 2-5, subseq	C	C					
99477	Init day hosp neonate care	C	C					
99478	lc, lbw inf < 1500 gm subseq	C	C					
99479	lc, lbw inf 1500-2500 g subseq	C	C					
99480	lc, inf pbw 2501-5000 g subseq	C	C					
99489	Unlisted e&m service	B						
99500	Home visit, prenatal	E	E					
99501	Home visit, postnatal	E	E					
99502	Home visit, nb care	E	E					
99503	Home visit, resp therapy	E	E					
99504	Home visit mech ventilator	E	E					
99505	Home visit, stoma care	E	E					
99506	Home visit, im injection	E	E					
99507	Home visit, cath maintain	E	E					
99509	Home visit day life activity	E	E					
99510	Home visit, sing/m/fam couns	E	E					
99511	Home visit, fecal/enema mgmt	E	E					
99512	Home visit for hemodialysis	E	E					
99600	Home visit nos	E	E					
99601	Home infusion/visit, 2 hrs	E	E					
99602	Home infusion, each addtl hr	E	E					
99605	Mims by pharm, nb, 15 min	E	E					
99606	Mims by pharm, est, 15 min	E	E					
99607	Mims by pharm, addl 15 min	E	E					
0001F	Heart failure composite	M						
0005F	Osteoarthritis composite	M						
0012F	Cap bacterial assess	M						
0014F	Comp preop assess cat surg	M						
0015F	Melan follow-up complete	M						
0016T	Thermtx otoroid vasc lesion	T	T	0235	5.8498	\$394.31		\$78.87
0017T	Photocoagulat macular drusen	T	T	0235	5.8498	\$394.31		\$78.87
0019T	Extracorp shock ww tx, nrs, nos	A	A					
0030T	Antiprothrombin antibody	A	A					
0042T	Ct perfusion w/contrast, cbf	N	N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
99374	Home health care supervision	B	B					
99375	Home health care supervision	E	E					
99377	Hospice care supervision	B	B					
99378	Hospice care supervision	E	E					
99379	Nursing fac care supervision	B	B					
99380	Nursing fac care supervision	E	E					
99381	Init pm e/m, new pat, inf	E	E					
99382	Init pm e/m, new pat 1-4 yrs	E	E					
99383	Prev visit, new, age 5-11	E	E					
99384	Prev visit, new, age 12-17	E	E					
99385	Prev visit, new, age 18-29	E	E					
99386	Prev visit, new, age 40-64	E	E					
99387	Init pm e/m, new pat 65+ yrs	E	E					
99391	Per pm reeval, est pat, inf	E	E					
99392	Prev visit, est, age 1-4	E	E					
99393	Prev visit, est, age 5-11	E	E					
99394	Prev visit, est, age 12-17	E	E					
99395	Prev visit, est, age 18-39	E	E					
99396	Prev visit, est, age 40-64	E	E					
99397	Per pm reeval est pat 65+ yr	E	E					
99401	Preventive counseling, indiv	E	E					
99402	Preventive counseling, indiv	E	E					
99403	Preventive counseling, indiv	E	E					
99404	Preventive counseling, group	E	E					
99406	Behav chng smoking 3-10 min	X	X	0031	0.3448	\$23.24		\$4.65
99407	Behav chng smoking > 10 min	X	X	0031	0.3448	\$23.24		\$4.65
99408	Audit/dest, 15-30 min	E	E					
99409	Audit/dest, over 30 min	E	E					
99411	Preventive counseling, group	E	E					
99412	Preventive counseling, group	E	E					
99420	Health risk assessment test	E	E					
99429	Unlisted preventive service	E	E					
99441	Phone e/m by phys 5-10 min	E	E					
99442	Phone e/m by phys 11-20 min	E	E					
99443	Phone e/m by phys 21-30 min	E	E					
99444	Online e/m by phys	E	E					
99450	Basic life disability exam	E	E					
99455	Work related disability exam	B	B					
99456	Disability examination	B	B					
99460	Init nb em per day, hosp	V	V	0605	1.0337	\$69.68		\$13.94
99461	Init nb em per day, non-fac	M	M					
99462	Sbseq nb em per day, hosp	C	C					
99463	Same day nb discharge	V	V	0605	1.0337	\$69.68		\$13.94

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0110T	Nes quant sensory test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
0111T	Rbc membranes fatty acids		A					
0123T	Scleral fistulization		T	0234	24.4021	\$1,644.85	\$511.31	\$228.97
0124T	Conjunctival drug placement		T	0232	4.5074	\$303.83	\$76.12	\$60.77
0126T	Chd risk limit study		Q1	0340	0.6693	\$45.11		\$9.03
0130T	Chron care drug investigatn		B					
0140T	Exhaled breath condensate ph		A					
0141T	Perq islet transplant		E					
0142T	Open islet transplant		E					
0143T	Laparoscopic islet transplant		E					
0144T	CT heart w/o dye; qual calc	CH	D					
0145T	CT heart w/w dye funct	CH	D					
0146T	CCTA w/w dye	CH	D					
0147T	CCTA w/w, quan calcium	CH	D					
0148T	CCTA w/w, strx	CH	D					
0149T	CCTA w/w, strx quan calc	CH	D					
0150T	CCTA w/w, disease strx	CH	D					
0151T	CT heart funct add-on	CH	D					
0155T	Lap impl gast curve electrd		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
0156T	Lap rmv gast curve electrd		T	0130	38.054	\$2,565.07	\$659.53	\$513.02
0157T	Open impl gast curve electrd		C					
0158T	Open rmv gast curve electrd		C					
0159T	Cad breast mri		N					
0160T	Tcranial magn strim tx plan		S	0216	2.6831	\$180.86		\$36.18
0161T	Tcranial magn strim tx deliv		S	0216	2.6831	\$180.86		\$36.18
0163T	Lumb arifr diskectomy addl		C					
0164T	Remove lumb arifr disc addl		C					
0165T	Revise lumb arifr disc addl		C					
0166T	Tcath vsd close w/o bypass		C					
0167T	Tcath vsd close w bypass		C					
0168T	Rhinophotox light app bilat		T	0251	3.425	\$230.87		\$46.18
0169T	Place stereo cath brain		C					
0170T	Anorectal fistula plug rpr	CH	D					
0171T	Aqu canal dilat w/o retent		T	0052	88.6521	\$5,975.68		\$1,195.14
0172T	Lumbar spine process addl		T	0052	88.6521	\$5,975.68		\$1,195.14
0173T	lop mont to pressure		N					
0174T	Cad cxr with interp		N					
0175T	Cad cxr remote		N					
0176T	Aqu canal dilat w/o retent		T	0673	41.884	\$2,823.23	\$649.56	\$564.65
0177T	Aqu canal dilat w retent		T	0673	41.884	\$2,823.23	\$649.56	\$564.65
0178T	64 lead ecg w i&r		B					
0179T	64 lead ecg w tracing		X	0100	2.6136	\$176.17	\$41.44	\$35.24
0180T	64 lead ecg w i&r only		B					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0048T	Implant ventricular device		C					
0050T	Removal circulation assist		C					
0051T	Implant total heart system		C					
0052T	Replace component heart syst		C					
0053T	Replace component heart syst		C					
0054T	Bone surgery using computer		N					
0055T	Bone surgery using computer		N					
0062T	Rep intradisc annulus;1 lev	CH	D					
0063T	Rep intradisc annulus;>1lev	CH	D					
0064T	Spectrosc eval expired gas	CH	D					
0066T	Ct colonography;screen	CH	D					
0067T	Ct colonography;dx	CH	D					
0068T	Interp/rept heart sound	CH	D					
0069T	Analysis only heart sound	CH	D					
0070T	Interp only heart sound	CH	D					
0071T	U/s leiomyomata ablate <200		S	0067	52.9891	\$3,571.78		\$714.36
0072T	U/s leiomyomata ablate >200		S	0067	52.9891	\$3,571.78		\$714.36
0073T	Delivery comp imrt		S	0412	6.249	\$421.22		\$84.25
0075T	Perq stent/chest vert art		C					
0076T	S&i stent/chest vert art		C					
0077T	Cereb therm perfusion probe	CH	D					
0078T	Endovasc aort repr widevise		C					
0079T	Endovasc visc extnsh repr		C					
0080T	Endovasc aort repr rad s&i		C					
0081T	Endovasc visc extnsh s&i		C					
0084T	Temp prostate urethral stent	CH	D					
0085T	Breath test heart reject	CH	D					
0086T	L ventricle fill pressure	CH	D					
0087T	Sperm eval hyaluronan	CH	D					
0092T	Artific disc addl		C					
0095T	Artific disc addl		C					
0098T	Flav artific disc addl		C					
0099T	Implant corneal ring		T	0233	16.2485	\$1,095.25	\$263.77	\$219.05
0100T	Prosth retina receive&gen		T	0672	39.9643	\$2,693.83		\$538.77
0101T	Extracorp shockwv tx,r/i enrg		T	0050	31.7717	\$2,141.60		\$428.32
0102T	Extracorp shockwv tx, anesth		T	0050	31.7717	\$2,141.60		\$428.32
0103T	Holotranscobalamin		A					
0104T	At rest cardio gas rebreathe		A					
0105T	Exerc cardio gas rebreathe		A					
0106T	Touch quant sensory test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
0107T	Vibrate quant sensory test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
0108T	Cool quant sensory test		X	0341	0.0799	\$5.39	\$2.09	\$1.08
0109T	Heat quant sensory test		X	0341	0.0799	\$5.39	\$2.09	\$1.08

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0501F	Prenatal flow sheet		M					
0502F	Subsequent prenatal care		M					
0503F	Postpartum care visit		M					
0505F	Hemodialysis plan docd		M					
0507F	Periton dialysis plan docd		M					
0509F	Urine incom plan docd		M					
0513F	Elev bp plan of care docd		M					
0514F	Care plan hgb docd esa pt		M					
0516F	Anemia plan of care docd		M					
0517F	Glaucoma plan of care docd		M					
0518F	Fall plan of care docd		M					
0519F	Plan chemo docd b/4 txmnt		M					
0520F	Rad dos limits b/4 3d rad		M					
0521F	Plan of care 4 pain docd		M					
0525F	Initial visit for episode		M					
0526F	Subs visit for episode		M					
0528F	Rcmdnd flw-up 10 yrs docd		E					
0529F	Intrl 3-yrs pls chncsp docd		M					
0535F	Dyspnea mgmnt plan docd		E					
0540F	Gluc mgmnt plan docd		E					
0545F	Follow up care plan mdd docd	NI	E					
0575F	HIV rna plan care docd		E					
1000F	Tobacco use assessed		M					
1002F	Assess anginal symptom/level		M					
1003F	Level of activity assess		M					
1004F	Clin symp vol ovrd assess		M					
1005F	Asthma symptoms evaluate		M					
1006F	Osteoarthritis assess		M					
1007F	Anti-inflm/antisc otc assess		M					
1008F	GI/renal risk assess		M					
1015F	Copd symptoms assess		M					
1018F	Assess dyspnea not present		M					
1019F	Assess dyspnea present		M					
1022F	Pneumo imm status assess		M					
1026F	Co-morbid condition assess		M					
1030F	Influenza imm status assess		M					
1034F	Current tobacco smoker		M					
1035F	Smokeless tobacco user		M					
1036F	Tobacco non-user		M					
1038F	Persistent asthma		M					
1039F	Intermittent asthma		M					
1040F	Dsm-ivm info mdd docd		M					
1050F	History of mole changes		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
0181T	Corneal hysterisis		S	0230	0.5945	\$40.07		\$8.02
0182T	Hdr elect brachytherapy	CH	S	0313	11.5353	\$777.55	\$293.30	\$155.51
0183T	Wound ultrasound	CH	T	0013	0.8789	\$59.24		\$11.85
0184T	Exc rectal tumor endoscopic		C					
0185T	Comstr probability analysis		N					
0186T	Suprachoroidal drug delivery		T	0237	20.7145	\$1,396.28		\$279.26
0187T	Ophthalmic dx image anterior		S	0230	0.5945	\$40.07		\$8.02
0188T	Videocent crit care 74 min		M					
0189T	Videocent crit care addl 30		M					
0190T	Place intrac. radiation src		T	0237	20.7145	\$1,396.28		\$279.26
0191T	Insert ant segment drain int		T	0234	24.4021	\$1,644.85	\$511.31	\$328.97
0192T	Insert ant segment drain ext		T	0673	41.884	\$2,823.23	\$649.56	\$564.65
0193T	Rf bladder neck microremodel		T	0165	20.0243	\$1,349.76		\$269.96
0194T	Procalcitonin (ppt)	CH	D					
0195T	Arthro presac interbody		C					
0196T	Arthro presac interbody eac		C					
0197T	Intrafraction track motion		C					
0198T	Ocular blood flow measure		N					
0199T	Physiologic tremor record		S	0230	0.5945	\$40.07		\$8.02
0200T	Perq sacral augmt unilat inj		S	0215	0.6135	\$41.35		\$8.27
0201T	Perq sacral augmt bilat inj		T	0049	22.0149	\$1,483.94		\$296.79
0202T	Post vert arthropl 1 lumbar		T	0050	31.7717	\$2,141.60		\$428.32
0203T	Unattend sleep study w/ftme		C					
0204T	Unattended sleep study	NI	S	0213	2.4043	\$162.06	\$53.58	\$32.42
0205T	Inlris each vessel add-on	NI	S	0213	2.4043	\$162.06	\$53.58	\$32.42
0206T	Remote algorithm analys eeg	NI	O1					
0207T	Clear eyelid gland wheat	NI	S	0340	0.6693	\$45.11		\$9.03
0208T	Automated audiometry air	NI	S	0230	0.5945	\$40.07		\$8.02
0209T	Auto audiometry airborne	NI	X	0035	0.232	\$15.64		\$3.13
0210T	Auto audiometry sp thresh	NI	X	0035	0.232	\$15.64		\$3.13
0211T	Auto audiometry sp thresh	NI	X	0035	0.232	\$15.64		\$3.13
0212T	Comprehen auto audiometry	NI	X	0364	0.47	\$31.68	\$7.06	\$6.34
0213T	Us facet jt inj cervlt 1 lev	NI	T	0207	7.2002	\$485.34		\$97.07
0214T	Us facet jt inj cervlt 2 lev	NI	T	0204	2.5558	\$172.28	\$40.13	\$34.46
0215T	Us facet jt inj cervlt 3 lev	NI	T	0204	2.5558	\$172.28	\$40.13	\$34.46
0216T	Us facet jt inj ls 1 level	NI	T	0207	7.2002	\$485.34		\$97.07
0217T	Us facet jt inj ls 2 level	NI	T	0204	2.5558	\$172.28	\$40.13	\$34.46
0218T	Us facet jt inj ls 3 level	NI	T	0204	2.5558	\$172.28	\$40.13	\$34.46
0219T	Fuse spine facet jt cerv	NI	C					
0220T	Fuse spine facet jt thor	NI	C					
0221T	Fuse spine facet jt lumbar	NI	T	0050	31.7717	\$2,141.60		\$428.32
0222T	Fuse spine facet jt add seg	NI	T	0050	31.7717	\$2,141.60		\$428.32
0500F	Initial prenatal care visit		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
2001F	Weight record		M					
2002F	Clin sign vol ovoid assess		M					
2004F	Initial exam involved joints		M					
2010F	Vital signs recorded		M					
2014F	Mental status assess		M					
2018F	Hydration status assess		M					
2019F	Dilated macul exam done		M					
2020F	Dilated fundus eval done		M					
2021F	Dilat macul+ exam done		M					
2022F	Dil retina exam interp rev		M					
2024F	7 field photo interp doc rev		M					
2027F	Eye image valid to dx rev		M					
2027F	Optic nerve head eval done		M					
2028F	Foot exam performed		M					
2029F	Complete phys skin exam done		M					
2030F	H2o stat docd, normal		M					
2031F	H2o stat docd, dehydrated		M					
2035F	Tymp memb motion examd		M					
2040F	Bk pn xm on init visit date		M					
2044F	Doc mnt lst b/4 bk trmnt		M					
2050F	Wound char size etc docd		E					
2060F	Pt talk eval hlt/wktr re mdd	NI	E					
3006F	Cxr doc rev		M					
3008F	Body mass index docd	NI	E					
3011F	Lipid panel doc rev		M					
3014F	Screen mammo doc rev		M					
3015F	Cerv cancer screen docd	NI	E					
3018F	Pt smrd unhlthy OH use		M					
3017F	Colorectal ca screen doc rev		M					
3018F	Pre-prxd rsk et al docd		E					
3020F	Lvf assess		M					
3021F	Lvel mod/sever deprs syst		M					
3022F	Lvel >=40% systolic		M					
3023F	Spirom doc rev		M					
3025F	Spirom lev/fvc<70% w copd		M					
3027F	Spirom lev/fvc>=70%/w/o copd		M					
3028F	O2 saturation doc rev		M					
3035F	O2 saturation<=88% /paO<=55		M					
3037F	O2 saturation> 88% /paO>55		M					
3038F	Pulm fx win 12 mon b/4 surg	NI	E					
3040F	Fev<40% predicted value		M					
3042F	Fev>=40% predicted value		M					
3044F	Hg a1c level lt 7.0%		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
1055F	Visual funct status assess		M					
1060F	Doc perm/cont/parox air fib		M					
1061F	Doc lack perm-cont+parox fib		M					
1065F	Iscim stroke symp i3 hrsb/4		M					
1066F	Iscim stroke symp ge3 hrsb/4		M					
1070F	Alarm symp assessed-absent		M					
1071F	Alarm symp assessed-1+ prsnt		M					
1090F	Pres/absn urine incoon assess		M					
1091F	Urine incoon characterized		M					
1100F	Pt falls assess-docd ge2+/yr		M					
1101F	Pt falls assess-docd le1/yr		M					
1110F	Pt lift/lgt fac w/in 60 days		M					
1111F	Dschrg med/current med merge		M					
1116F	Autic/peri pain assessed		M					
1116F	GERD symps assessed 12 month		M					
1118F	Init eval for condition		M					
1121F	Subs eval for condition		M					
1123F	Acq discuss/dscn mkr docd		M					
1124F	Acq discuss-no dscnmkr docd		M					
1125F	Amnt pain noted pain prsnt		M					
1126F	Amnt pain noted none prsnt		M					
1127F	New episode for condition	CH	D					
1128F	Subs episode for condition	CH	D					
1130F	Bk pain + fxn assessed		M					
1134F	Epsd bk pain for <= 6 wks		M					
1135F	Epsd bk pain for > 6 wks		M					
1136F	Epsd bk pain for <= 12 wks		M					
1137F	Epsd bk pain for > 12 wks		M					
1150F	Doc pt rsk death w/in 1yr		E					
1151F	Doc no pt rsk death w/in 1yr		E					
1152F	Doc advncd dis comfort 1st		E					
1153F	Doc advncd dis cmfrt not 1st		E					
1157F	Advnc care plan in rcd		E					
1158F	Advnc care plan lk docd		M					
1159F	Med list docd in rcd		E					
1160F	Fvw meds by rx/dr in rcd		E					
1170F	Fxn status assessed		M					
1180F	Thromboemb risk assessed		E					
1200F	Seizure type(s)+ frq docd	NI	E					
1205F	EPI eliol synd hvwd and docd	NI	E					
1220F	Pt screened for depression		M					
2000F	Blood pressure measure		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
3200F	Barium swallow test not req		M					
3210F	Grp a strep test performed		M					
3215F	Pt immunity to hep a docd		M					
3218F	Pt immunity to hep b docd		M					
3218F	Rna tsmg hep c docd-done		M					
3220F	Hep c quant ma tsmg docd		M					
3230F	Note trng ist w/in 6 mon		M					
3250F	Nonprim loc anat bx site tum		M					
3260F	Pt cat/pn cat/hist grd docd		M					
3265F	Rna tsmg hepc vlr ord/doct		M					
3268F	Hepe gn tsmg docd b/4bxmrt		M					
3268F	Psat/qdisc docd b/4 bxmrt		M					
3269F	Bone scn b/4 bxmrt/vaftr Dx		M					
3270F	No bone scn b/4 bxmrt/vaftrDx		M					
3271F	Low risk prostate cancer		M					
3272F	Med risk prostate cancer		M					
3273F	High risk prostate cancer		M					
3274F	Prost Cnrc risk not tvr/mf/hgh		M					
3278F	Serum lvis CA199/tpd ord		M					
3279F	Hgb lv >= 13 g/dl		M					
3280F	Hgb lv < 11 g/dl		M					
3281F	Hgb lv < 11 g/dl		M					
3284F	IOP down >= 15% of pre-svc lv		M					
3285F	IOP down < 15% of pre-svc lv		M					
3288F	Fall risk assessment docd		M					
3290F	Pt-D(RH)- and unsensitized		M					
3291F	Pt-d(h)- or sensitized		M					
3292F	Hiv tsmg asked/doct/revwd		M					
3293F	Abo rh blood typing docd	NI	E					
3294F	Grp b strep screening docd	NI	E					
3300F	AJCC stage docd b/4 thxpy		M					
3301F	Cancer stage docd metast		M					
3315F	Er+ or pr+ breast cancer		M					
3316F	ER- or PR- breast cancer		M					
3317F	Path rpt malig cancer docd		M					
3318F	Path rpt malig cancer docd		M					
3319F	X-ray/ct/ultrnsnd et al ord		M					
3320F	No xray/ct/ et al ord		M					
3321F	AJCC cnrc 0/IA melan docd		E					
3322F	Melan >AJCC stage 0 or IA		E					
3323F	Clin node stng doct/b/4 surg	NI	M					
3324F	Mri ct scan ord hvwd rgstd	NI	E					
3325F	Preop asses 4 cataract surg		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
3045F	Hg a tc level 7.0-9.0%		M					
3046F	Hemoglobin a tc level > 9.0%		M					
3048F	Ldt-c < 100 mg/dl		M					
3049F	Ldt-c 100-129 mg/dl		M					
3050F	Ldt-c >= 130 mg/dl		M					
3060F	Pos microalbuminuria rev		M					
3061F	Neg microalbuminuria rev		M					
3062F	Pos macroalbuminuria rev		M					
3066F	Nephropathy doc tx		M					
3072F	Low risk for retinopathy		M					
3073F	Pre-surg eye measures docd		M					
3074F	Syst bp lt 130 mm hg		M					
3075F	Syst bp ge 130 - 139mm hg		M					
3077F	Syst bp >= 140 mm hg6 it		M					
3078F	Diastr bp < 80 mm hg		M					
3079F	Diastr bp 80-89 mm hg		M					
3080F	Diastr bp >= 90 mm hg		M					
3082F	K/v lt1.2		M					
3083F	K/v ge 1.2 and <1.7		M					
3084F	K/v ge 1.7		M					
3085F	Suicide risk assessed		M					
3088F	MDD, mild		M					
3089F	MDD, moderate		M					
3090F	MDD, severe; w/o psych		M					
3091F	Mdd, severe; w/ psych		M					
3092F	MDD, in remission		M					
3093F	Doc new diag 1st/addl mdd		M					
3095F	Central dexa results docd		M					
3096F	Central dexa ordered		M					
3100F	Image test ref carot diam		M					
3110F	Doc pres/absn hmrtg/hesion		M					
3111F	Ct/mri brain done w/in 24hrs		M					
3112F	Ct/Mri brain done gt 24 hrs		M					
3120F	12-lead ecg performed		M					
3130F	Upper gi endoscopy performed		M					
3132F	Doc ref upper gi endoscopy		M					
3140F	Upper gi endo shows barrits		M					
3141F	Upper gi endo not barrits		M					
3142F	Barium swallow test ordered		M					
3150F	Forcaps esoph biopsy done		M					
3155F	Cyogen test marrow b/4 tx		M					
3160F	Doc fe+ stores b/4 epo thx		M					
3170F	Flow cyto done b/4 tx		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
3500F	Cd4-cell cnt/% docd as done		M					
3502F	HIV rna vrl id <lims quantif		M					
3503F	HIV rna vrl idnot<lims quantif		M					
3510F	Doc to scrng rsts interpd		E					
3511F	Chimyl/gonrh tsds docd done	CH	M					
3512F	Syph scrng docd as done	CH	M					
3513F	Hep B scrng docd as done		E					
3514F	Hep C scrng docd as done		E					
3515F	Pt has docd immun to hep C		E					
3550F	Low risk thromboembolism		E					
3551F	Intrmed risk thromboembolism		E					
3552F	High risk for thromboembolism		E					
3553F	Pt inr measurement performed		E					
3570F	Rprt bone scint xraf w xray		M					
3572F	Pt consid poss risk tx		E					
3573F	Pt not consid poss risk tx		E					
3650F	Eeg ordered rwd reqstd	NI	E					
4000F	Tobacco use brnmt counseling		M					
4001F	Tobacco use brnmt, pharmacol		M					
4002F	Statin therapy, rx		M					
4003F	Pt ed write/oral, pls w/ ht		M					
4004F	Pt tobacco use done rwd tik	NI	E					
4005F	Pharm tnx for op rxd		M					
4006F	Beta-blocker therapy rx		M					
4009F	Ace/arb inhibitor therapy rx		M					
4011F	Oral antiplatelet therapy rx		M					
4012F	Warfarin therapy rx		M					
4014F	Written discharge instr pvtd		M					
4015F	Persist asthma medicine ctrl		M					
4016F	Anti-inflm/antigsc agent rx		M					
4017F	GI prophylaxis for nsaid rx		M					
4018F	Therapy exercise joint rx		M					
4019F	Doc rept counsl vit d/calc+		M					
4025F	Inhaled bronchodilator rx		M					
4030F	Oxygen therapy rx		M					
4033F	Pulmonary rehab rec		M					
4035F	Influenza imm rec		M					
4037F	Influenza imm order/admn		M					
4040F	Pneumoc vac/admn/cvtd		M					
4041F	Doc order cetazoln/ceturox		M					
4042F	Doc antibiotic not given		M					
4043F	Doc order given stop antibio		M					
4044F	Doc order given vte prophylx		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
3328F	Primmc docd 2 wks b4 surg	NI	E					
3330F	Imaging study ordered (bkg)		M					
3331F	Bk imaging lst not ordered		M					
3340F	Mammo assess inc xray docd		M					
3341F	Mammo assess negative docd		M					
3342F	Mammo assess benign docd		M					
3343F	Mammo probably benign docd		M					
3344F	Mammo assess susp. docd		M					
3345F	Mammo assess highlymalig doc		M					
3350F	Mammo bx proven malig docd		M					
3351F	Neg scm dep symp by deptool		E					
3352F	No sig dep symp by dep tool		E					
3353F	Mild-mtd dep symp by deptool		E					
3354F	Clin sig dep sym by dep tool		E					
3370F	AJCC brst cncr stage 0 docd		M					
3372F	Ajcc brst cncr stage 1+docd		M					
3374F	Ajcc brst cncr stage 1+docd		M					
3376F	AJCC brstcncr stage 2 docd		M					
3378F	AJCC brstcncr stage 3 docd		M					
3380F	AJCC brstcncr stage 4 docd		M					
3382F	AJCC cin cncr stage 0 docd		M					
3384F	AJCC cin cncr stage 1 docd		M					
3386F	AJCC cin cncr stage 2 docd		M					
3388F	AJCC cin cncr stage 3 docd		M					
3390F	AJCC cin cncr stage 4 docd		M					
3450F	Dyspnea scnd, no-mild dysp		E					
3451F	Dyspnea scnd mod-high dysp		E					
3452F	Dyspnea not screened		E					
3455F	TB scrng done-interpd 6mon		M					
3470F	RA disease activity, low		M					
3471F	RA disease activity, mod		M					
3472F	RA disease activity, high		M					
3475F	Disease progrn RA poor docd		M					
3476F	Disease progrn RA good docd		M					
3490F	History - AIDS-defining cond		M					
3491F	HIV unsure baby of HIV+moims		E					
3492F	History cd4+ cell count <350		M					
3493F	No hist cd4+cell cnt<350		M					
3494F	CD4+cell count <200cells/mm3		M					
3495F	Cd4+cell cnt 200-499 cells		M					
3496F	Cd4+ cell count =500 cells		M					
3497F	CD4+ cell percentage <15%		E					
3498F	CD4+ cell percentage >=15%		E					

APPENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
4151F	Pt not recving anitv hep c		M					
4151F	Combo peginnif/rib rx		M					
4155F	Hep A vac series prev recvd		M					
4157F	Hep B vac series prev recvd		M					
4158F	Pt edu re alcoh drinking done		M					
4159F	Controp talk b/4 anitv bmrnt		M					
4163F	Pt couns 4 bmrnt opti prost		M					
4164F	Adlv hrmnl thxpy rxd		M					
4166F	3d-cr/(fmr) received		M					
4167F	Hd bed tilted 1st day vent		M					
4168F	Pt care ICU/vent w/in 24hrs		M					
4169F	No pt care ICU/vent in 24hrs		M					
4171F	Pt recvg esa thxpy		M					
4172F	Pt not recvg esa thxpy		M					
4174F	Couns potent glauc impact		M					
4175F	Vis of >= 20/40 w/in 90 days		M					
4176F	Talk re uv light pt/crov		M					
4177F	Talk pt/crov re areds prev		M					
4178F	Antid gbln rovd w/in 26wks		M					
4179F	Tamoxifen/AI prescribed		M					
4180F	Adlv thxpyrxd/rovd sig3a-c		M					
4181F	Conformal rathn thxpy rovd		M					
4182F	No conformal rathn thxpy		M					
4185F	Continuous ppl or h2ra rovd		M					
4186F	No cont ppl or h2ra rovd		M					
4187F	Anti rheum drughthxpyrxd/gvn		M					
4188F	Approp ACE/ARB tstag done		M					
4189F	Approp digoxin tstag done		M					
4190F	Approp diuretic tstag done		M					
4191F	Approp anticonvuls tstag		M					
4192F	Pt not recvg glucooco thxpy		M					
4193F	Pt recvg<10mg daily predniso		M					
4194F	Pt rec=>10mg prednison qd		M					
4196F	Pt recvg anti-rheum thxpy RA		M					
4196F	Pt not recvg anti-rhm thxpyRA		M					
4200F	External beam to prost only		M					
4201F	Extml beam other than prost		M					
4210F	ACE/ARB thxpy for >= 6 mons		M					
4221F	Diuretic thxpy for >= 6 mons		M					
4230F	Anticonv thxpy for >= 6 mons		M					
4240F	Inst xrcz 4bk pn >12 weeks		M					
4242F	Spnvsd xrcz bk pn >12 weeks		M					

APPENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
4045F	Empiric antibiotic rx		M					
4046F	Doc antibio given b/4 surg		M					
4047F	Doc antibio given b/4 surg		M					
4048F	Doc antibio given b/4 surg		M					
4049F	Doc order given stop antibio		M					
4050F	Ht care plan doc		M					
4051F	Referred for an AV fistula		M					
4052F	Hemodialysis via AV fistula		M					
4053F	Hemodialysis via AV graft		M					
4054F	Hemodialysis via catheter		M					
4055F	Pt recvg periton dialysis		M					
4056F	Approp oral rehyd reccomm		M					
4058F	Ped gastro ed given categvr		M					
4060F	Psych svcs provided		M					
4062F	Pt referral psych docd		M					
4063F	Anidepres rxthxpy not rxd	NI	E					
4064F	Anidepressant rx		M					
4065F	Antipsychotic rx		M					
4066F	ECT provided		M					
4067F	Pt referral for ect docd		M					
4070F	Dvt prophylx recvd day 2		M					
4075F	Anticoag thx rx at dischrng		M					
4077F	Oral antiplat thx rx dischrng		M					
4079F	Doc t-pa admin considered		M					
4084F	Doc rehab svcs considered		M					
4090F	Aspirin recvd w/in 24 hrs		M					
4095F	Pt not recvg epo thxpy		M					
4100F	Biphos thxpy vein ord/recvd		M					
4110F	Beta blkcr admin w/in 24 hrs		M					
4115F	Int mam art used for cabg		M					
4120F	Antibiot rxd/given		M					
4124F	Antibiot not rxd/given		M					
4130F	Topical prep rx aoe		M					
4131F	Syst antimicrobial thx rx		M					
4132F	No syst antimicrobial thx rx		M					
4133F	Antihist/decong rx/recom		M					
4134F	No antihist/decong rx/recom		M					
4136F	Systemic corticosteroids rx		M					
4136F	Syst corticosteroids not rx		M					
4148F	Hep A vac injxn admin/recvd		M					
4149F	Hep B vac injxn admin/recvd		M					
4150F	Pt recvg antivir bmrnt hepc		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
6040F	Appro rad ds dvcx techs dooc		M					
6045F	Radpks in end rpt4fluro pxd		M					
6070F	Pt asked/insid aed effects	NI	E					
7010F	Pt info into recall system		M					
7020F	Mammo assess cat in dbase		M					
7025F	Pt infows alarm 4 nxt mammo		M					
A0021	Outside state ambulance serv		E					
A0080	Noninterest escort in non er		E					
A0090	Interest escort in non er		E					
A0100	Nonemergency transport taxi		E					
A0110	Nonemergency transport bus		E					
A0120	Noner transport mini-bus		E					
A0130	Noner transport wheelch van		E					
A0140	Nonemergency transport air		E					
A0160	Noner transport case worker		E					
A0170	Transport parking fees/tolls		E					
A0180	Noner transport lodging recip		E					
A0190	Noner transport meals recip		E					
A0200	Noner transport lodging escort		E					
A0210	Noner transport meals escort		E					
A0225	Neonatal emergency transport		E					
A0380	Basic life support mileage		E					
A0382	Basic support routine supplis		A					
A0384	Bls defibrillation supplis		A					
A0390	Advanced life support mileag		E					
A0392	Als defibrillation supplis		A					
A0394	Als IV drug therapy supplis		A					
A0396	Als esophageal intub supplis		A					
A0398	Als routine disposable supplis		A					
A0420	Ambulance waiting 1/2 hr		A					
A0422	Ambulance 02 life sustaining		A					
A0424	Extra ambulance attendant		A					
A0425	Ground mileage		A					
A0426	Als 1		A					
A0427	ALS1-emergency		A					
A0428	bis		A					
A0429	BL S-emergency		A					
A0430	Fixed wing air transport		A					
A0431	Rotary wing air transport		A					
A0432	Pt volunteer ambulance co		A					
A0433	als 2		A					
A0434	Specialty care transport		A					
A0435	Fixed wing air mileage		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
4245F	Pt instr norm lifest		M					
4248F	Pt instr-no bd rest>= 4 days		M					
4250F	Wring 4 surg - normothermia		M					
4255F	Anesth >= 60 min as dooc	NI	M					
4256F	Anesth < 60 min as dooc	NI	M					
4260F	Wound srfc culturetech used		E					
4261F	Tech other than surfc cultur		E					
4265F	Wet-dry dressings Rx-recmd		E					
4266F	No wet-dry dressings Rx-recmd		E					
4267F	Compression thapy prescribed		M					
4268F	Pt ed re comp thapy recvd		E					
4269F	Appropos mthd offloading Rxd		E					
4270F	Pt revng anti r-viral thapy		E					
4271F	Pt revng anti r-viral thapy		M					
4274F	Flu immuno admind rovd		E					
4275F	Hep b vac inj admind/ rovd		E					
4276F	Potent antivir thapy Rxd		M					
4279F	PCP prophylaxis Rxd		E					
4280F	PCP prophylax Rxd 3mon low %		E					
4290F	Pt scndd for inj drug use	CH	M					
4293F	Pt scndd - high-rsk sex behav	CH	M					
4300F	Pt revng wart thapy		E					
4301F	Pt not revng wart thapy		E					
4305F	Pt ed re ft care inspct rovd		E					
4306F	Pt lik psych & Rx opd addic		E					
4320F	Pt talk psychsoc-rx oh dprnd		E					
4330F	Cnsing ept spec sfty issues	NI	E					
4340F	Cnsing chldbrng+ women ept	NI	E					
5005F	Pt counseld on exam for moles		M					
5010F	Macul+ findings to dr rmg dm		M					
5015F	Doc fx & test/bmnt for op		M					
5020F	T xmnis 2 main Dr by 1 mon		E					
5050F	Plan 2 main dr by 1 month		M					
5060F	Findngs mammo zpt win 3 days		M					
5062F	Doc f2mammo findng in 5 days		M					
5100F	Rsk fx ref win 24 hrs x-ray		E					
5200F	Eval appros surg thapy epi	NI	E					
6005F	Care level rationale doc		M					
6010F	Dysphag test dome b/4 eating		M					
6015F	Dysphag test dome b/4 eating		M					
6020F	Npo (nothing-mouth) ordered		M					
6030F	Max sterile barriers follwd		M					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A4264	Intrabul occlusion device	NI	E					
A4265	Paraffin		Y					
A4266	Diaphragm		E					
A4267	Male condom		E					
A4268	Female condom		E					
A4269	Spermicide		E					
A4270	Disposable endoscope sheath		N					
A4280	Bst prsths adnsv attachmt		A					
A4281	Replacement breastpump tube		E					
A4282	Replacement breastpump adpt		E					
A4283	Replacement breastpump cap		E					
A4284	Replcmnt breast pump shield		E					
A4285	Replcmnt breast pump bottle		E					
A4286	Replcmnt breastpump lok ring		E					
A4290	Sacral nerve stim test lead		B					
A4300	Cath impi vasc access portal		N					
A4301	Implantable access syst portc		N					
A4306	Drug delivery system >=50 ML		N					
A4310	Insert tray w/o bag/cath		A					
A4311	Catheter w/o bag 2-way latex		A					
A4312	Cath w/o bag 2-way silicone		A					
A4313	Catheter w/bag 3-way		A					
A4314	Cath w/drainage 2-way latex		A					
A4315	Cath w/drainage 2-way silcne		A					
A4316	Cath w/drainage 3-way		A					
A4320	Irrigation tray		A					
A4321	Cath therapeutic irrig agent		A					
A4322	Irrigation syringe		A					
A4326	Male external catheter		A					
A4327	Fem urinary collect dev cup		A					
A4328	Fem urinary collect pouch		A					
A4330	Stool collection pouch		A					
A4331	Extension drainage tubing		A					
A4332	Lube sterile packet		A					
A4333	Urinary cath anchor device		A					
A4334	Urinary cath leg strap		A					
A4335	Incontinence supply		A					
A4336	Urethral insert	NI	A					
A4338	Indwelling catheter latex		A					
A4340	Indwelling catheter special		A					
A4344	Cath indw foley 2 way sillicn		A					
A4346	Cath indw foley 3 way		A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A0436	Rotary wing air mileage		A					
A0888	Noncovered ambulance mileage		E					
A0998	Ambulance response/treatment		E					
A0999	Unlisted ambulance service		A					
A4206	1 CC sterile syringe&needle		E					
A4207	2 CC sterile syringe&needle		E					
A4208	3 CC sterile syringe&needle		E					
A4209	5+ CC sterile syringe&needle		E					
A4210	Nonneedle injection device		E					
A4211	Supp for self-adm injections		E					
A4212	Non-coring needle or syrlt		B					
A4213	20+ CC syringe only		E					
A4215	Sterile needle		E					
A4216	Sterile water/saline, 10 ml		A					
A4217	Sterile water/saline, 500 ml		A					
A4218	Sterile saline or water		N					
A4220	Infusion pump refill kit		N					
A4221	Maint drug infus cath per wk		Y					
A4222	Infusion supplies with pump		Y					
A4223	Infusion supplies w/o pump		E					
A4230	Infus insulin pump non needl		N					
A4231	Infusion insulin pump needle		N					
A4232	Syringe w/needle insulin 3cc		E					
A4233	Alkaliin batt for glucose mon		Y					
A4234	J-cell batt for glucose mon		Y					
A4235	Lithium batt for glucose mon		Y					
A4236	Silvr oxide batt glucose mon		Y					
A4244	Alcohol or peroxide per pint		E					
A4245	Alcohol wipes per box		E					
A4246	Betadine/phisothex solution		E					
A4247	Betadine/iodine swabs/wipes		E					
A4248	Chlorhexidine antisept		E					
A4250	Urine reagent strips/tablets		E					
A4252	Blood ketone test or strip		E					
A4253	Blood glucose/resagent strips		Y					
A4255	Glucose monitor platforms		Y					
A4256	Calibrator solution/chips		Y					
A4257	Replace Lensshield Cartridge		Y					
A4258	Lancet device each		Y					
A4259	Lancets per box		Y					
A4261	Cervical cap contraceptive		E					
A4262	Temporary tear duct plug		N					
A4263	Permanent tear duct plug		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A4397	Irrigation supply sleeve		A					
A4398	Ostomy irrigation bag		A					
A4399	Ostomy irrig cone/cath w brs		A					
A4400	Ostomy irrigation set		A					
A4402	Lubricant per ounce		A					
A4404	Ostomy ring each		A					
A4405	Nonpectin based ostomy paste		A					
A4406	Pectin based ostomy paste		A					
A4407	Ext wear ost skn barr <=4sq"		A					
A4408	Ext wear ost skn barr >4sq"		A					
A4409	Ost skn barr convex <=4 sq i		A					
A4410	Ost skn barr extnd >4 sq		A					
A4411	Ost skn barr extnd =4sq		A					
A4412	Ost pouch drain high output		A					
A4413	2 pc drainable ost pouch		A					
A4414	Ost sknbar w/o conv<=4 sq in		A					
A4415	Ost skn barr w/o conv >4 sqj		A					
A4416	Ost pouch elsd w barrier/filtr		A					
A4417	Ost pouch w barr/blinconv/filtr		A					
A4418	Ost pouch for bar w flange/filtr		A					
A4419	Ost pouch elsd w/o bar w fltr		A					
A4420	Ost pouch elsd for bar w lk fl		A					
A4421	Ostomy supply misc		E					
A4422	Ost pouch absorbent material		A					
A4423	Ost pouch for bar w lk l/filtr		A					
A4424	Ost pouch drain w bar & filter		A					
A4425	Ost pouch drain for barrier fl		A					
A4426	Ost pouch drain 2 piece system		A					
A4427	Ost pouch drain/barr lk flng/f		A					
A4428	Urine ost pouch w faucet/tap		A					
A4429	Urine ost pouch w blinconv		A					
A4430	Ost urine pouch w b/blin conv		A					
A4431	Ost pouch urine w barrier/tapv		A					
A4432	Os pouch urine w barr/fangeltap		A					
A4433	Urine ost pouch bar w lock fln		A					
A4434	Ost pouch urine w lock flng/flt		A					
A4450	Non-waterproof tape		A					
A4452	Waterproof tape		A					
A4455	Adhesive remover per ounce		A					
A4456	Adhesive remover, wipes	NI	A					
A4458	Reusable enema bag		E					
A4461	Surgical dress hold non-reuse		A					
A4463	Surgical dress holder reuse		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A4349	Disposable male external cat		A					
A4351	Straight tip urine catheter		A					
A4352	Coarde tip urinary catheter		A					
A4353	Intermittent urinary cath		A					
A4354	Cath insertion tray w/bag		A					
A4355	Bladder irrigation tubing		A					
A4356	Ext ureth clamp or compr dvc		A					
A4357	Beadside drainage bag		A					
A4358	Urinary leg or abdomen bag		A					
A4360	Disposable ext urethral dev	NI	A					
A4361	Ostomy face plate		A					
A4362	Solid skin barrier		A					
A4363	Ostomy clamp, replacement		A					
A4364	Adhesive, liquid or equal		A					
A4365	Adhesive remover wipes	CH	D					
A4366	Ostomy vent		A					
A4367	Ostomy belt		A					
A4368	Ostomy filter		A					
A4369	Skin barrier liquid per oz		A					
A4371	Skin barrier powder per oz		A					
A4372	Skin barrier solid 4x4 equiv		A					
A4373	Skin barrier with flange		A					
A4375	Drainable plastic pouch w lcpj		A					
A4376	Drainable rubber pouch w lcpjt		A					
A4377	Drainable plastic pouch w/o fp		A					
A4378	Drainable rubber pouch w/o fp		A					
A4379	Urinary plastic pouch w lcpj		A					
A4380	Urinary rubber pouch w lcpjt		A					
A4381	Urinary plastic pouch w/o fp		A					
A4382	Urinary hvy plstc pouch w/o fp		A					
A4383	Urinary rubber pouch w/o fp		A					
A4385	Ostomy facepill/silicone ring		A					
A4387	Ost elsd pouch w att st barr		A					
A4388	Drainable pouch w ex wear barr		A					
A4389	Drainable pouch w st wear barr		A					
A4390	Drainable pouch ex wear convex		A					
A4391	Urinary pouch w ex wear barr		A					
A4392	Urinary pouch w st wear barr		A					
A4393	Urine pouch w ex wear bar conv		A					
A4394	Ostomy pouch liq deodorant		A					
A4395	Ostomy pouch solid deodorant		A					
A4396	Peristomal hernia supprt bit		A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A4625	Trach care kit for new trach.		A					
A4626	Tracheostomy cleaning brush		A					
A4627	Spacer bag/reservoir		E					
A4628	Oropharyngeal suction cath		Y					
A4629	Tracheostomy care kit		A					
A4630	Repl bat t a.n.s. own by pt		Y					
A4633	Uvl replacement bulb		Y					
A4634	Replacement bulb th lighbox		A					
A4635	Underarm crutch pad		Y					
A4636	Handgrip for cane etc		Y					
A4637	Repl tip caner/crutch/walker		Y					
A4638	Repl batt pulse gen sys		Y					
A4639	Infrared ht sys replcmt pad		Y					
A4640	Alternating pressure pad		Y					
A4641	Radiofarm dx agent noc		N					
A4642	In111 satumomab		N					
A4648	Implantable tissue marker		N					
A4649	Surgical supplies		N					
A4650	Implant radiation dosimeter		N					
A4651	Calibrated microcap tube		A					
A4652	Microcapillary tube sealant		A					
A4653	PD catheter anchor belt		A					
A4657	Syringe w/wo needle		N					
A4660	Sphyg/bp app w cuff and stet		N					
A4663	Dialysis blood pressure cuff		N					
A4670	Automatic bp monitor, dial		E					
A4671	Disposable cyclor set		B					
A4672	Drainage ext line, dialysis		B					
A4673	Ext line w easy lock connect		B					
A4674	Chem/antisept solution, 8oz		B					
A4680	Activated carbon filter, ea		N					
A4706	Dialyzer, each		N					
A4707	Bicarbonate conc sol per gal		N					
A4708	Acetate conc sol per gallon		N					
A4709	Acid conc sol per gallon		N					
A4714	Treated water per gallon		N					
A4719	"Y set" tubing		N					
A4720	Dialysat sol fld vol > 249cc		N					
A4721	Dialysat sol fld vol > 999cc		N					
A4722	Dialys sol fld vol > 1999cc		N					
A4723	Dialys sol fld vol > 2999cc		N					
A4724	Dialys sol fld vol > 3999cc		N					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A4465	Non-elastic extremity binder		N					
A4466	Elastic garment/covering	NI	E					
A4470	Gravitee jet washer		N					
A4480	Vabra aspirator		N					
A4481	Tracheostoma filter		A					
A4483	Moisture exchanger		A					
A4490	Above knee surgical stocking		E					
A4495	Thigh length surg stocking		E					
A4500	Below knee surgical stocking		E					
A4510	Full length surg stocking		E					
A4520	Incontinence garment, any type		E					
A4550	Surgical trays		B					
A4554	Disposable underpads		E					
A4556	Electrodes, pair		Y					
A4557	Lead wires, pair		Y					
A4558	Conductive gel or paste		Y					
A4559	Coupling gel or paste		Y					
A4561	Pessary rubber, any type		N					
A4562	Pessary, non rubber, any type		N					
A4565	Slings		N					
A4570	Splint		E					
A4575	Hyperbaric o2 chamber disp		E					
A4580	Cast supplies (plaster)		E					
A4590	Special casting material		E					
A4595	TENS suppl, 2 lead per month		Y					
A4600	Sleeve, inter limb comp dev		Y					
A4601	Lith ion batt, non-pros use		Y					
A4604	Tubing with heating element		Y					
A4605	Trach suction cath close sys		Y					
A4606	Oxygen probe used w oximeter		A					
A4608	Transitracheal oxygen cath		Y					
A4611	Heavy duty battery		Y					
A4612	Battery cables		Y					
A4613	Battery charger		Y					
A4614	Hand-held PEFR meter		Y					
A4615	Cannula nasal		Y					
A4616	Tubing (oxygen) per foot		Y					
A4617	Mouth piece		Y					
A4618	Breathing circuits		Y					
A4619	Face tent		Y					
A4620	Variable concentration mask		Y					
A4623	Tracheostomy inner cannula		A					
A4624	Tracheal suction tube		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A5083	Stoma absorptive cover		A					
A5093	Ostomy accessory convex inse		A					
A5102	Bedside drain bil w/wo tube		A					
A5105	Urinary suspensory		A					
A5112	Urinary leg bag		A					
A5113	Latex leg strap		A					
A5114	Foam fabric leg strap		A					
A5120	Skin barrier, wipe or swab		A					
A5121	Solid skin barrier 6x6		A					
A5122	Solid skin barrier 8x8		A					
A5126	Disk/foam pad +/- adhesive		A					
A5131	Appliance cleaner		A					
A5200	Percutaneous catheter anchor		A					
A5500	Diab shoe for density insert		Y					
A5501	Diabetic custom molded shoe		Y					
A5503	Diabetic shoe w/roller/rocker		Y					
A5504	Diabetic shoe with wedge		Y					
A5505	Diab shoe w/metatarsal bar		Y					
A5506	Diabetic shoe w/off set heel		Y					
A5507	Modification diabetic shoe		Y					
A5508	Diabetic deluxe shoe		Y					
A5510	Compression form shoe insert		E					
A5512	Multi den insert direct form		Y					
A5513	Multi den insert custom mold		Y					
A6000	Wound warming wound cover		E					
A6010	Collagen based wound filler		A					
A6011	Collagen gel/paste wound fil		A					
A6021	Collagen dressing <=16 sq in		A					
A6022	Collagen drsg >16<=48 sq in		A					
A6023	Collagen dressing >48 sq in		A					
A6024	Collagen drsg wound filler		A					
A6025	Silicone gel sheet, each		E					
A6154	Wound pouch each		A					
A6196	Alginate dressing <=16 sq in		A					
A6197	Alginate drsg >16 <=48 sq in		A					
A6198	alginate dressing > 48 sq in		A					
A6199	Alginate drsg wound filler		A					
A6200	Compos drsg <=16 no border		CH					
A6201	Compos drsg >16<=48 no bdr		CH					
A6202	Compos drsg >48 no border		CH					
A6203	Composite drsg <= 16 sq in		A					
A6204	Composite drsg >16<=48 sq in		A					
A6205	Composite drsg > 48 sq in		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A4725	Dialys sol fld vol > 4999cc		N					
A4726	Dialys sol fld vol > 4999cc		N					
A4728	Dialysate solution, non-dex		B					
A4730	Fistula cannulation set, ea		N					
A4736	Topical anesthetic, per gram		N					
A4737	Ini, anesthetic per 10 ml		N					
A4740	Shunt accessory		N					
A4750	Art or venous blood tubing		N					
A4755	Comb art/venous blood tubing		N					
A4760	Dialysate sol test kit, each		N					
A4765	Dialysate conc pow per pack		N					
A4766	Dialysate conc sol add 10 ml		N					
A4770	Blood collection tube/vacuum		N					
A4771	Serum clotting time tube		N					
A4772	Blood glucose test strips		N					
A4773	Occult blood test strips		N					
A4774	Ammonia test strips		N					
A4802	Protamine sulfate per 50 mg		N					
A4860	Disposable catheter tips		N					
A4870	Plumb/elec wk hm hemo equip		N					
A4890	Repair/maint cont hemo equip		N					
A4911	Drain bag/bottle		N					
A4913	Misc dialysis supplies noc		N					
A4918	Venous pressure clamp		N					
A4927	Non-sterile gloves		N					
A4928	Surgical mask		N					
A4929	Touriquet for dialysis, ea		N					
A4930	Sterile, gloves per pair		N					
A4931	Reusable oral thermometer		N					
A4932	Reusable rectal thermometer		E					
A5051	Pouch drsd w barr attached		A					
A5052	Cisd ostomy pouch w/o barr		A					
A5053	Cisd ostomy pouch faceplate		A					
A5054	Cisd ostomy pouch w/riange		A					
A5055	Stoma cap		A					
A5061	Pouch drainable w barrier at		A					
A5062	Drinble ostomy pouch w/o barr		A					
A5063	Drain ostomy pouch w/riange		A					
A5071	Urinary pouch w/barrier		A					
A5072	Urinary pouch w/o barrier		A					
A5073	Urinary pouch on barr w/riang		A					
A5081	Continent stoma plug		A					
A5082	Continent stoma catheter		A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A6253	Absorpt drg > 48 sq in w/o b	A	A					
A6254	Absorpt drg <= 16 sq in w/bdr	A	A					
A6255	Absorpt drg > 16<=48 in w/bdr	A	A					
A6256	Absorpt drg > 48 sq in w/bdr	A	A					
A6257	Transparent film <= 16 sq in	A	A					
A6258	Transparent film > 16<=48 in	A	A					
A6259	Transparent film > 48 sq in	A	A					
A6260	Wound cleanser, any type/size	A	A					
A6261	Wound filler gel/paste /oz	A	A					
A6262	Wound filler dry form / gram	A	A					
A6266	Impreg gauze no h2O/sal/yard	A	A					
A6402	Sterile gauze <= 16 sq in	A	A					
A6403	Sterile gauze > 16 <= 48 sq in	A	A					
A6404	Sterile gauze > 48 sq in	A	A					
A6407	Packing strips, non-impreg	A	A					
A6410	Sterile eye pad	A	A					
A6411	Non-sterile eye pad	A	A					
A6412	Occlusive eye patch	E						
A6413	Adhesive bandage, first-aid	E						
A6441	Pad band w>=3' <5'/yd	A	A					
A6442	Conform band n/s w<3'/yd	A	A					
A6443	Conform band n/s w>=3' <5'/yd	A	A					
A6444	Conform band n/s w>=5'/yd	A	A					
A6445	Conform band s w <3'/yd	A	A					
A6446	Conform band s w>=3' <5'/yd	A	A					
A6447	Conform band s w >=5'/yd	A	A					
A6448	Lt compres band <3'/yd	A	A					
A6449	Lt compres band >=3' <5'/yd	A	A					
A6450	Lt compres band >=5'/yd	A	A					
A6451	Mod compres band w>=3' <5'/yd	A	A					
A6452	High compres band w>=3' <5'/yd	A	A					
A6453	Self-adher band w<3'/yd	A	A					
A6454	Self-adher band w>=3' <5'/yd	A	A					
A6455	Self-adher band >=5'/yd	A	A					
A6456	Zinc paste band w >=3' <5'/yd	A	A					
A6457	Tubular dressing	A	A					
A6501	Compres burngarment bodysuit	A	A					
A6502	Compres burngarment chinstrip	A	A					
A6503	Compres burngarment facehood	A	A					
A6504	Compresburngarment glove-wrist	A	A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A6206	Contact layer <= 16 sq in	A	A					
A6207	Contact layer > 16<= 48 sq in	A	A					
A6208	Contact layer > 48 sq in	A	A					
A6209	Foam drg <= 16 sq in w/o bdr	A	A					
A6210	Foam drg > 16<=48 sq in w/o b	A	A					
A6211	Foam drg > 48 sq in w/o bdr	A	A					
A6212	Foam drg <= 16 sq in w/border	A	A					
A6213	Foam drg > 16<=48 sq in w/bdr	A	A					
A6214	Foam drg > 48 sq in w/border	A	A					
A6215	Foam dressing wound filler	A	A					
A6216	Non-sterile gauze<=16 sq in	A	A					
A6217	Non-sterile gauze> 16<=48 sq	A	A					
A6218	Non-sterile gauze > 48 sq in	A	A					
A6219	Gauze <= 16 sq in w/border	A	A					
A6220	Gauze > 16 <=48 sq in w/bdr	A	A					
A6221	Gauze > 48 sq in w/border	A	A					
A6222	Gauze <= 16 in no w/sal w/o b	A	A					
A6223	Gauze > 16<=48 no w/sal w/o b	A	A					
A6224	Gauze > 48 in no w/sal w/o b	A	A					
A6228	Gauze <= 16 sq in water/sal	A	A					
A6229	Gauze > 16<=48 sq in wat/sal	A	A					
A6230	Gauze > 48 sq in water/saline	A	A					
A6231	Hydrogel drsg<=16 sq in	A	A					
A6232	Hydrogel drg > 16<=48 sq in	A	A					
A6233	Hydrogel dressing >48 sq in	A	A					
A6234	Hydrocolloid drg <= 16 w/o bdr	A	A					
A6235	Hydrocolloid drg > 16<=48 w/o b	A	A					
A6236	Hydrocolloid drg > 48 in w/o b	A	A					
A6237	Hydrocolloid drg <= 16 in w/bdr	A	A					
A6238	Hydrocolloid drg > 16<=48 w/bdr	A	A					
A6239	Hydrocolloid drg > 48 in w/bdr	A	A					
A6240	Hydrocolloid drg filler paste	A	A					
A6241	Hydrocolloid drg filler dry	A	A					
A6242	Hydrogel drg <= 16 in w/o bdr	A	A					
A6243	Hydrogel drg > 16<=48 w/o bdr	A	A					
A6244	Hydrogel drg > 48 in w/o bdr	A	A					
A6245	Hydrogel drg <= 16 in w/bdr	A	A					
A6246	Hydrogel drg > 16<=48 in w/b	A	A					
A6247	Hydrogel drg > 48 sq in w/b	A	A					
A6248	Hydrogel drsg gel filler	A	A					
A6250	Skin seal protect moisturizer	A	A					
A6251	Absorpt drg <= 16 sq in w/o b	A	A					
A6252	Absorpt drg > 16 <=48 w/o bdr	A	A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A7012	Nebulizer water colic device		Y					
A7013	Disposable compressor filter		Y					
A7014	Compressor nondispos filter		Y					
A7015	Aerosol mask used w nebulize		Y					
A7016	Nebulizer dome & mouthpiece		Y					
A7017	Nebulizer not used w oxygen		Y					
A7018	Water distilled w/nebulizer		Y					
A7025	Replace chest compress vest		Y					
A7026	Replace chst empress sys hose		Y					
A7027	Combination oral/nasal mask		Y					
A7028	Repl oral cushion comb mask		Y					
A7029	CPAP full face mask		Y					
A7030	Replacement facemask interfa		Y					
A7032	Replacement nasal cushion		Y					
A7033	Replacement nasal pillows		Y					
A7034	Nasal applicator device		Y					
A7035	Pos airway press headgear		Y					
A7036	Pos airway press chinstrap		Y					
A7037	Pos airway pressure tubing		Y					
A7038	Pos airway pressure filter		Y					
A7039	Filter, non disposable w pap		Y					
A7040	One way chest drain valve		A					
A7041	Water seal drain container		A					
A7042	Implanted pleural catheter		N					
A7043	Vacuum drainagebottle/tubing		A					
A7044	PAP oral interface		Y					
A7045	Repl exhalation port for PAP		Y					
A7046	Repl water chamber, PAP dev		Y					
A7501	Tracheostoma valve w diaphra		A					
A7502	Replacement diaphragm/plate		A					
A7503	HMES filter holder or cap		A					
A7504	Tracheostoma HMES filter		A					
A7505	HMES or trach valve housing		A					
A7506	HMES/trachvalve adhesivedisk		A					
A7507	Integrated filter & holder		A					
A7508	Housing & Integrated Adhesiv		A					
A7509	Heat & moisture exchange sys		A					
A7520	Trach/aryn tube non-cuffed		A					
A7521	Trach/aryn tube cuffed		A					
A7522	Trach/aryn tube stainless		A					
A7523	Tracheostomy shower protect		A					
A7524	Tracheostoma stent/stud/btin		A					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A6505	Compriburngarment glove- elbow		A					
A6506	Compriburngarment glove-axilla		A					
A6507	Compriburngarment foot-knee		A					
A6508	Compriburngarment foot- thigh		A					
A6509	Compres burn garment jacket		A					
A6510	Compres burn garment leotard		A					
A6511	Compres burn garment panty		A					
A6512	Compres burn garment, noc face/neck		A					
A6513	Compression stocking BK18-30		B					
A6530	Compression stocking BK30-40		E					
A6531	Compression stocking BK40-50		A					
A6532	Compression stocking BK40-50		A					
A6533	Gc stocking thighlength 18-30		E					
A6534	Gc stocking thighlength 30-40		E					
A6535	Gc stocking thighlength 40-50		E					
A6536	Gc stocking full length 18-30		E					
A6537	Gc stocking full length 30-40		E					
A6538	Gc stocking full length 40-50		E					
A6539	Gc stocking waistlength 18-30		E					
A6540	Gc stocking waistlength 30-40		E					
A6541	Gc stocking waistlength 40-50		E					
A6542	Gc stocking custom made	CH	D					
A6543	Gc stocking lymphedema	CH	D					
A6544	Gc stocking garter belt		E					
A6545	Grad comp non-elastic BK		A					
A6549	G compression stocking		E					
A6550	Neg pres wound ther drsg set		Y					
A7000	Disposable canister for pump		Y					
A7001	Nondisposable pump canister		Y					
A7002	Tubing used w suction pump		Y					
A7003	Nebulizer administration set		Y					
A7004	Disposable nebulizer smi vol		Y					
A7005	Nondisposable nebulizer set		Y					
A7006	Filtered nebulizer admin set		Y					
A7007	Lg vol nebulizer disposable		Y					
A7008	Disposable nebulizer refill		Y					
A7009	Nebulizer reservoir bottle		Y					
A7010	Disposable corrugated tubing		Y					
A7011	Nondispos corrugated tubing		Y					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A9528	Iodine I-131 iodide cap. dx		N					
A9529	I131 iodide sol. dx		N					
A9530	I131 iodide sol. rx	CH	K	1150		\$11.02		\$2.21
A9531	I131 max 100uCi		N					
A9532	I125 serum albumin, dx		N					
A9535	Injection, methylene blue	CH	D					
A9536	Tc99m depreotide		N					
A9537	Tc99m mebrofenin		N					
A9538	Tc99m pyrophosphate		N					
A9539	Tc99m pentetate		N					
A9540	Tc99m MAA		N					
A9541	Tc99m sulfur colloid		N					
A9542	In111 ibritumomab, dx		N					
A9543	Y90 ibritumomab, rx	CH	K	1643		\$15,532.15		\$3,106.43
A9544	I131 tositumomab, dx		N					
A9545	I131 tositumomab, rx	CH	K	1645		\$9,929.05		\$1,985.81
A9546	C657/68		N					
A9547	In111 oxyquinoline		N					
A9548	In111 pentetate		N					
A9550	Tc99m gluceptate		N					
A9551	Tc99m succimer		N					
A9552	F18 fdg		N					
A9553	Cr51 chromate		N					
A9554	I125 iothalamate, dx		N					
A9555	Rb82 rubidium		N					
A9556	Ga67 gallium		N					
A9557	Tc99m bicisate		N					
A9558	Xe133 xenon 10mci		N					
A9559	Ce67 cyano		N					
A9560	Tc99m labeled rbc		N					
A9561	Tc99m oxidronate		N					
A9562	Tc99m merlitalde		N					
A9563	P32 Na phosphate	CH	K	1675		\$196.49		\$39.30
A9564	P32 chromic phosphate	CH	K	1676		\$113.44		\$22.69
A9566	Tc99m fanlesomab		N					
A9567	Technetium Tc-99m aerosol		N					
A9568	Technetium tc99m arctumomab		N					
A9569	Technetium Tc-99m auto WBC		N					
A9570	Indium In-111 auto WBC		N					
A9571	Indium In-111 auto platelet		N					
A9572	Indium In-111 pentioretide		N					
A9576	Inj prochance multipack		N					
A9577	Inj multihance		N					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A7525	Tracheostomy mask		A					
A7526	Tracheostomy tube collar		A					
A7527	Trachealryn tube plug/stop		A					
A8000	Soft protect helmet prefab		Y					
A8001	Hard protect helmet prefab		Y					
A8002	Soft protect helmet custom		Y					
A8003	Hard protect helmet custom		Y					
A8004	Repl soft interface, helmet		Y					
A9150	Misc/expir non-prescript dru		B					
A9152	Single vitamin nos		E					
A9153	Multi-vitamin nos		E					
A9155	Artificial saliva		B					
A9180	Lice treatment, topical		E					
A9270	Non-covered item or service		E					
A9274	Ext amp insulin delivery sys		E					
A9275	Disp home glucose monitor		E					
A9276	Disposable sensor, CGM sys		E					
A9277	External transmitter, CGM		E					
A9278	External receiver, CGM sys		E					
A9279	Monitoring feature/device/NOC		E					
A9280	Alert device, noc		E					
A9281	Reaching/grabbing device		E					
A9282	Wig any type		E					
A9283	Foot press off board supp dev		E					
A9284	Non-electronic spirometer		E					
A9300	Exercise equipment		E					
A9500	Tc99m sestamibi		N					
A9501	Technetium Tc-99m tetrofosmin		N					
A9502	Tc99m tetrofosmin		N					
A9503	Tc99m medronate		N					
A9504	Tc99m apcitide		N					
A9505	TL201 thallium		N					
A9507	In111 capromab		N					
A9508	I131 iodobenguante, dx		N					
A9509	Iodine I-123 sod iodide mil		N					
A9510	Tc99m disofenin		N					
A9512	Tc99m partechnetate		N					
A9516	Iodine I-123 sod iodide mic		N					
A9517	I131 iodide cap. rx	CH	K	1064		\$16.51		\$3.31
A9521	Tc99m exametazime		N					
A9524	I131 serum albumin, dx		N					
A9526	Nitrogen N-13 ammonia		N					
A9527	Iodine I-125 sodium iodide		U	2632	0.5626	\$37.92		\$7.59

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
B4178	Parenteral sol amino acid >		Y					
B4180	Parenteral sol catb > 50%		Y					
B4185	Parenteral sol 10 gm lipids		B					
B4189	Parenteral sol amino acid &		Y					
B4193	Parenteral sol 52-73 gm prot		Y					
B4197	Parenteral sol 74-100 gm pro		Y					
B4199	Parenteral sol > 100gm prote		Y					
B4216	Parenteral nutrition additiv		Y					
B4220	Parenteral supply kit premix		Y					
B4224	Parenteral administration ki		Y					
B5000	Parenteral sol renal-amiroxy		Y					
B5100	Parenteral sol hepatic-freem		Y					
B5200	Parenteral sol stress-brnch c		Y					
B9000	Enter infusion pump w/o airm		Y					
B9002	Enter infusion pump w/ ala		Y					
B9004	Parenteral infus pump portab		Y					
B9006	Parenteral infus pump statio		Y					
B9998	Enterl supp not otherwise c		Y					
B9999	Parenteral supp not othwrs c		Y					
C1300	HYPERBARIC Oxygen		S	0659	1.588	\$107.04		\$21.41
C1713	Anchor/screw brn/bn,ls/bn		N					
C1714	Cath, trans atherectomy, dir		N					
C1715	Brachytherapy headle		N					
C1716	Brachytx, non-str, Gold-198		U	1716	0.6357	\$42.85		\$6.57
C1717	Brachytx, non-str, HDR Ir-192		U	1717	3.4326	\$231.38		\$46.28
C1719	Brachytx, NS, Non-HDR Ir-192		U	1719	0.9497	\$64.02		\$12.81
C1721	AICD, dual chamber		N					
C1722	AICD, single chamber		N					
C1724	Cath, trans atheroc, rotation		N					
C1725	Cath, transillum non-laser		N					
C1726	Cath, bal dil, non-vascular		N					
C1727	Cath, bal tis dis, non-vas		N					
C1728	Cath, brachytx seed adm		N					
C1729	Cath, drainage		N					
C1730	Cath, EP, 19 or few elect		N					
C1731	Cath, EP, 20 or more elec		N					
C1732	Cath, EP, diag/lab, 3D/vect		N					
C1733	Cath, EP, othr than cool-hip		N					
C1750	Cath, hemodialysis, long-term		N					
C1751	Cath, inf, per/cent/inidline		N					
C1752	Cath, hemodialysis, short-term		N					
C1753	Cath, intravas ultrasound		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
A9578	Inj multihance multipack		N					
A9579	Gad-base MR contrast		N					
A9580	NOS, 1ml		N					
A9580	Sodium fluoride F-18		N					
A9581	Gadoxetate disodium inj	NI	G	9246		\$13.50		
A9582	Iodine I-123 iobenguane	NI	G	9247		\$2,329.83		\$465.97
A9583	Gadofosvesat trisodium inj	NI	G	1299		\$1.29		
A9600	Sr89 strontium	CH	K	0701		\$647.67		\$129.54
A9604	Sm 153 lexidronam	NI	K	1295		\$4,671.66		\$934.34
A9605	Sm 153 lexidronam	CH	D					
A9698	Non-rad contrast material/MOC		N					
A9699	Radiopharm rx agent noc		N					
A9700	Echocardiography Contrast		B					
A9900	Supply/accessory/service		Y					
A9901	Delivery/set up/dispensing		A					
A9999	DME supply or accessory, nos		Y					
B4034	Enter feed supkit syr by day		Y					
B4035	Enterl feed supp pump per d		Y					
B4036	Enterl feed sup kit grav by		Y					
B4081	Enterl ng tubing w/ stylet		Y					
B4082	Enterl ng tubing w/o stylet		Y					
B4083	Enterl stomach tube levine		Y					
B4087	Gastro/jejuno tube, std		A					
B4088	Gastro/jejuno tube, low-pro		A					
B4100	Food thickener oral		E					
B4102	EF adult fluids and electro		Y					
B4103	EF ped fluid and electrolyte		Y					
B4104	Additive for enteral formula		E					
B4149	EF blenderized foods		Y					
B4150	EF complete w/intact nutrient		Y					
B4152	EF calorie dense>=1.5Kcal		Y					
B4153	EF hydrolyzed/amino acids		Y					
B4154	EF spec metabolic noninherit		Y					
B4155	EF incomplete/modular		Y					
B4157	EF special metabolic inherit		Y					
B4158	EF ped complete intact nut		Y					
B4159	EF ped complete soy based		Y					
B4160	EF ped caloric dense>=0.7kc		Y					
B4161	EF ped hydrolyzed/amino acid		Y					
B4162	EF ped specmetabolic inherit		Y					
B4164	Parenteral 50% dextrose solu		Y					
B4168	Parenteral sol amino acid 3		Y					
B4172	Parenteral sol amino acid 5		Y					
B4176	Parenteral sol amino acid 7		Y					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
C1875	Stent, coated/cov w/o del sy	N	N					
C1876	Stent, non-coat/non-cov w/del	N	N					
C1877	Stent, non-coat/cov w/o del	N	N					
C1878	Matr for vocal cord	N	N					
C1879	Tissue marker, implantable	N	N					
C1880	Vena cava filter	N	N					
C1881	Dialysis access system	N	N					
C1882	AICD, other than sing/dual	N	N					
C1883	Adapt/ext, pacing/neuro lead	N	N					
C1884	Embolization Protect syst	N	N					
C1885	Cath, translumin angio laser	N	N					
C1887	Catheter, guiding	N	N					
C1888	Endovas non-cardiac abi cath	N	N					
C1891	Infusion pump non-prog, perm	N	N					
C1892	Intro/sheath, fixed, peel-away	N	N					
C1893	Intro/sheath, fixed, non-peel	N	N					
C1894	Intro/sheath, non-laser	N	N					
C1895	Lead, AICD, endo dual coil	N	N					
C1896	Lead, AICD, non sing/dual	N	N					
C1897	Lead, neurostim test kit	N	N					
C1898	Lead, pmkr, other than trans	N	N					
C1899	Lead, pmkr/AICD combination	N	N					
C1900	Lead, coronary venous	N	N					
C2614	Probe, perc lumb disc	N	N					
C2615	Sealant, pulmonary, liquid	N	N					
C2616	Brachytx, non-str, Yttrium-90	U	2616		234.0942	\$15,779.35		\$3,155.87
C2617	Stent, non-cor, tem w/o del	N	N					
C2618	Probe, cryoablation	N	N					
C2619	Pmkr, dual, non rate-resp	N	N					
C2620	Pmkr, single, non rate-resp	N	N					
C2621	Pmkr, other than sing/dual	N	N					
C2622	Prosthesis, penile, non-inf	N	N					
C2625	Stent, non-cor, tem w/del sy	N	N					
C2626	Infusion pump, non-prog, temp	N	N					
C2627	Cath, suprapubic/cystoscopic	N	N					
C2628	Catheter, occlusion	N	N					
C2629	Intro/sheath, laser	N	N					
C2630	Cath, EP, cool-hip	N	N					
C2631	Rep dev, urinary, w/o sling	N	N					
C2634	Brachytx, non-str, HA, I-125	U	2634		0.8871	\$59.80		\$11.96
C2635	Brachytx, non-str, HA, P-103	U	2635		0.4242	\$28.59		\$5.72
C2636	Brachy linear, non-str, P-103	U	2636		0.2873	\$19.37		\$3.88
C2637	Brachy, non-str, Yttrium-169	B	B					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
C1754	Catheter, intradiscal	N	N					
C1755	Catheter, intraspinal	N	N					
C1756	Cath, pacing, transepiph	N	N					
C1757	Cath, thrombectomy/embolact	N	N					
C1758	Catheter, ureteral	N	N					
C1759	Cath, intra echocardiography	N	N					
C1760	Closure dev, vasc	N	N					
C1762	Conn tiss, human(tric fascia)	N	N					
C1763	Conn tiss, non-human	N	N					
C1764	Event recorder, cardiac	N	N					
C1765	Adhesion barrier	N	N					
C1766	Intro/sheath, stifle, non-peel	N	N					
C1767	Generator, neuro non-recharge	N	N					
C1768	Graft, vascular	N	N					
C1769	Guide wire	N	N					
C1770	Imaging coil, MR, insertable	N	N					
C1771	Rep dev, urinary, w/sling	N	N					
C1772	Infusion pump, programmable	N	N					
C1776	Joint device (implantable)	N	N					
C1777	Lead, AICD, endo single coil	N	N					
C1778	Lead, neurostimulator	N	N					
C1779	Lead, pmkr, transvenous VDD	N	N					
C1780	Lens, intraocular (new tech)	N	N					
C1781	Mesh (implantable)	N	N					
C1782	Morcellator	N	N					
C1783	Ocular imp, aqueous drain de	N	N					
C1784	Ocular dev, intracop, det ret	N	N					
C1785	Pmkr, dual, rate-resp	N	N					
C1786	Pmkr, single, rate-resp	N	N					
C1787	Patient progr, neurostim	N	N					
C1788	Port, indwelling, imp	N	N					
C1789	Prosthesis, breast, imp	N	N					
C1813	Prosthesis, penile, inflatable	N	N					
C1814	Retinal lamp, silicone oil	N	N					
C1815	Pros, urinary sph, imp	N	N					
C1816	Receiver/transmitter, neuro	N	N					
C1817	Septal defect imp, sys	N	N					
C1818	Integrated keratoprosthesis	N	N					
C1819	Tissue localization-excision	N	N					
C1820	Generator neuro rechg bat sys	N	N					
C1821	Interspinous implant	N	N					
C1874	Stent, coated/cov w/del sys	N	N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
C9249	Inj, certolizumab pegol	CH	D					\$27.12
C9250	Artiss fibrin sealant	CH	D	9250		\$138.20		
C9251	Inj, C1 esterase inhibitor	CH	D					
C9252	Injection, plerixator	CH	D					
C9253	Injection, temozolomide	CH	D					
C9254	Injection, lacosamide	NI	K	9254		\$0.18		\$0.04
C9255	Paliperidone palmitate inj	NI	G	9255		\$6.71		\$1.35
C9256	Dexamethasone intravitreal	NI	G	9256		\$196.10		\$38.48
C9257	Bevacizumab injection	NI	K	1281		\$1.41		\$0.29
C9352	Neuragen nerve guide, per cm		N					
C9353	Neurawrap nerve protector,cm		N					
C9354	Veritas collagen matrix, cm2	CH	N					
C9355	Neuromatrix nerve cuff, cm	CH	N					
C9356	TenoGlide tendon prot, cm2		G	9356		\$24.86		
C9358	SurqMend, fetal		G	9358		\$10.76		\$2.11
C9359	Impinj,bon void filler-putty		G	9359		\$63.54		
C9360	SurqMend, neonatal		G	9360		\$10.67		\$2.09
C9361	NeuroMend nerve wrap		G	9361		\$247.29		
C9362	Impinj,bon void filler-strip		G	9362		\$83.60		
C9363	Integra Meshed Bil Wound Mat		G	9363		\$25.62		\$5.03
C9364	Porcine Implant, Permacol		G	9364		\$17.21		
C9399	Unclassified drugs or biolog		A					
C9716	Radiofrequency energy to anu		T	0150	32.1812	\$2,189.21	\$437.12	\$439.85
C9724	EPS gast cardia plc		T	0422	24.2703	\$1,635.96	\$437.96	\$327.20
C9725	Place endorectal app		T	0148	5.365	\$361.63		\$72.33
C9726	Rxt breast appl piece/remov		T	0028	24.7516	\$1,668.41		\$339.69
C9727	Insert palate implants		T	0252	7.6196	\$513.61	\$109.16	\$102.73
C9728	Place device/marker, non pros		X	0310	13.7576	\$927.34		\$185.47
C9898	Implt stay radiolabeled item		N					
C9899	Implt implant pros dev,no cov		A					
D0120	Periodic oral evaluation		E					
D0140	Limit oral eval problem focus		E					
D0145	Oral evaluation, pt < 3yrs		E					
D0150	Comprehensive oral evaluation		S	0330	10.3227	\$695.81		\$139.17
D0160	Extensv oral eval prob focus		E					
D0170	Re-eval,est pt,problem focus		E					
D0180	Comp periodontal evaluation		E					
D0210	Intraoral complete film series		E					
D0220	Intraoral periapical first f		E					
D0230	Intraoral periapical ea add		E					
D0240	Intraoral occlusal film		S	0330	10.3227	\$695.81		\$139.17
D0250	Extraoral first film		S	0330	10.3227	\$695.81		\$139.17
D0260	Extraoral ea additional film		S	0330	10.3227	\$695.81		\$139.17

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
C2638	Brachyx, stranded, I-125	U		2638	0.6302	\$42.48		\$8.50
C2639	Brachytx, non-stranded,I-125	U		2639	0.5368	\$36.18		\$7.24
C2640	Brachytx, stranded, P-103	U		2640	0.8955	\$60.36		\$12.08
C2641	Brachytx, non-stranded,P-103	U		2641	0.8474	\$57.12		\$11.43
C2642	Brachytx, stranded, C-131	U		2642	1.6295	\$109.84		\$21.97
C2643	Brachytx, non-stranded,C-131	U		2643	0.9805	\$66.09		\$13.22
C2698	Brachytx, stranded, NOS	U		2698	0.6302	\$42.48		\$8.50
C2699	Brachytx, non-stranded, NOS	U		2699	0.4242	\$28.59		\$5.72
C8900	MRA w/cont, abd	O3	0284	6.2901	\$423.99	\$147.21	\$147.21	\$64.80
C8901	MRA w/cont, abd	O3	0336	5.1855	\$349.53	\$137.40	\$137.40	\$69.91
C8902	MRA w/cont, lwr ext	O3	0337	7.9432	\$535.42	\$198.32	\$198.32	\$107.09
C8903	MRI w/cont, breast, uni	O3	0284	6.2901	\$423.99	\$147.21	\$147.21	\$64.80
C8904	MRI w/cont, breast, uni	O3	0336	5.1855	\$349.53	\$137.40	\$137.40	\$69.91
C8905	MRI w/cont, breast, uni	O3	0337	7.9432	\$535.42	\$198.32	\$198.32	\$107.09
C8906	MRI w/cont, breast, bi	O3	0284	6.2901	\$423.99	\$147.21	\$147.21	\$64.80
C8907	MRI w/cont, breast, bi	O3	0336	5.1855	\$349.53	\$137.40	\$137.40	\$69.91
C8908	MRI w/cont, breast, uni	O3	0337	7.9432	\$535.42	\$198.32	\$198.32	\$107.09
C8909	MRA w/cont, chest	O3	0284	6.2901	\$423.99	\$147.21	\$147.21	\$64.80
C8910	MRA w/cont, chest	O3	0336	5.1855	\$349.53	\$137.40	\$137.40	\$69.91
C8911	MRA w/cont, lwr ext	O3	0284	6.2901	\$423.99	\$147.21	\$147.21	\$64.80
C8912	MRA w/cont, lwr ext	O3	0337	7.9432	\$535.42	\$198.32	\$198.32	\$107.09
C8913	MRA w/cont, lwr ext	O3	0336	5.1855	\$349.53	\$137.40	\$137.40	\$69.91
C8914	MRA w/cont, pelvis	O3	0284	6.2901	\$423.99	\$147.21	\$147.21	\$64.80
C8915	MRA w/cont, pelvis	O3	0336	5.1855	\$349.53	\$137.40	\$137.40	\$69.91
C8916	MRA w/cont, pelvis	O3	0337	7.9432	\$535.42	\$198.32	\$198.32	\$107.09
C8917	TTE w or w/o fol w/cont, com	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8918	TTE w or w/o fol w/cont, flu	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8919	2D TTE w or w/o fol w/cont,co	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8920	2D TTE w or w/o fol w/cont,flu	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8921	2D TTE w or w/o fol w/cont,in	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8922	TEE w or w/o fol w/cont,mon	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8923	TEE w or w/o fol w/cont,mon	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8924	TTE w or w/o fol w/cont,sitres	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8925	TTE w or w/o fol w/cont, Doppler	S	0128	9.6604	\$651.17	\$216.29	\$216.29	\$130.24
C8926	Prolonged IV inf, req pump	S	0440	3.2632	\$219.96			\$44.00
C9113	Inj pentiprazole sodium, via		N					
C9121	Injection, argatroban		K	9121		\$18.10		\$3.62
C9245	Injection, romiplostim	CH	D					
C9246	Inj, gadoxlate disodium	CH	D					
C9247	Inj, lobenguan, I-123, dx	CH	D					
C9248	Inj, clevitidine butyrate		G	9248		\$3.39		\$0.67

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D1120	Dental prophylaxis child		E					
D1203	Topical app fluoride child		E					
D1204	Topical app fluoride adult		E					
D1206	Topical fluoride varnish		E					
D1310	Nutri counsel-control caries		E					
D1320	Tobacco counseling		E					
D1330	Oral hygiene instruction		E					
D1351	Dental sealant per tooth		E					
D1510	Space maintainer fxd unilat		S	0330	10.3227	\$695.81		\$139.17
D1515	Fixed bilat space maintainer		S	0330	10.3227	\$695.81		\$139.17
D1520	Remove unilat space maintain		S	0330	10.3227	\$695.81		\$139.17
D1525	Remove bilat space maintain		S	0330	10.3227	\$695.81		\$139.17
D1550	Recement space maintainer		S	0330	10.3227	\$695.81		\$139.17
D1555	Remove fix space maintainer		E					
D2140	Amalgam one surface permanent		E					
D2150	Amalgam two surfaces perm		E					
D2160	Amalgam three surfaces perma		E					
D2161	Amalgam 4 or > surfaces perm		E					
D2300	Resin one surface-anterior		E					
D2331	Resin two surfaces-anterior		E					
D2332	Resin three surfaces-anterior		E					
D2335	Resin 4/2 surf or w incis an		E					
D2390	Ant resin-based cmpst crown		E					
D2391	Post 1 srfc resinbased cmpst		E					
D2392	Post 2 srfc resinbased cmpst		E					
D2393	Post 3 srfc resinbased cmpst		E					
D2394	Post >=4srfc resinbase cmpst		E					
D2410	Dental gold foil one surface		E					
D2420	Dental gold foil two surface		E					
D2430	Dental gold foil three surfa		E					
D2510	Dental inlay metallic 1 surf		E					
D2520	Dental inlay metallic 2 surf		E					
D2530	Dental inlay metall 3/more sur		E					
D2542	Dental onlay metallic 2 surf		E					
D2543	Dental onlay metallic 3 surf		E					
D2544	Dental onlay metall 4/more sur		E					
D2610	Inlay porcelain/ceramic 1 su		E					
D2620	Inlay porcelain/ceramic 2 su		E					
D2630	Dental onlay porc 3/more sur		E					
D2642	Dental onlay porcelain 2 surf		E					
D2643	Dental onlay porcelain 3 surf		E					
D2644	Dental onlay porc 4/more sur		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D0270	Dental bitewing single film		S	0330	10.3227	\$695.81		\$139.17
D0272	Dental bitewings two films		S	0330	10.3227	\$695.81		\$139.17
D0273	Bitewings - three films		E					
D0274	Dental bitewings four films		S	0330	10.3227	\$695.81		\$139.17
D0277	Vert bitewings-sev to eight		S	0330	10.3227	\$695.81		\$139.17
D0290	Dental film skull/facial bon		E					
D0310	Dental radiography		E					
D0320	Dental tmj arthrogram incl i		E					
D0321	Dental other tmj films		E					
D0322	Dental tomographic survey		E					
D0330	Dental panoramic film		E					
D0340	Dental cephalometric film		E					
D0350	Oral/facial photo images		E					
D0360	Cone beam ct		E					
D0362	Cone beam, two dimensional		E					
D0363	Cone beam, three dimensional		E					
D0415	Collection of microorganisms		E					
D0416	Viral culture		B					
D0417	Collect & prep saliva sample		E					
D0418	Analysis of saliva sample		E					
D0421	Gen tst suscept oral disease		B					
D0425	Caries susceptibility test		E					
D0431	Diag tst detect mucos abnorm		B					
D0460	Pulp vitality test		S	0330	10.3227	\$695.81		\$139.17
D0470	Diagnostic casts		E					
D0472	Gross exam, prep & report		B					
D0473	Micro exam, prep & report		B					
D0474	Micro w exam of surg margins		B					
D0475	Decalcification procedure		B					
D0476	Spec stains for microorganis		B					
D0477	Spec stains not for microorg		B					
D0478	Immunohistochemical stains		B					
D0479	Tissue in-situ hybridization		B					
D0480	Cytopath smear prep & report		B					
D0481	Electron microscopy diagnost		B					
D0482	Direct immunofluorescence		B					
D0483	Indirect immunofluorescence		B					
D0484	Consult slides prep elsewhere		B					
D0485	Consult inc prep of slides		B					
D0486	Accession of brush biopsy		E					
D0502	Other oral pathology procedu		B					
D0999	Unspecified diagnostic proce		B					
D1110	Dental prophylaxis adult		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D2970	Temp crown (fractured tooth)	E	E					
D2971	Add proc construct new crown	E	E					
D2975	Coping	E	E					
D2980	Crown repair	E	E					
D2989	Dental unspec restorative pr	S	S	0330	10.3227	\$695.81		\$139.17
D3110	Pulp cap direct	E	E					
D3120	Pulp cap indirect	E	E					
D3220	Therapeutic pulpotomy	E	E					
D3221	Gross pulpal debridement	E	E					
D3222	Part pulp for apexogenesis	E	E					
D3230	Pulpal therapy anterior prim	E	E					
D3240	Pulpal therapy posterior pri	E	E					
D3310	End thxpy, anterior tooth	E	E					
D3320	End thxpy, bicuspid tooth	E	E					
D3330	End thxpy, molar	E	E					
D3331	Non-surg b root canal obs	E	E					
D3332	Incomplete endodontic tx	E	E					
D3333	Internal root repair	E	E					
D3346	Retreat root canal anterior	E	E					
D3347	Retreat root canal bicuspid	E	E					
D3348	Retreat root canal molar	E	E					
D3351	Apexification/recalc initial	E	E					
D3352	Apexification/recalc interm	E	E					
D3353	Apexification/recalc final	E	E					
D3410	Apicoect/perirad surg anter	E	E					
D3421	Root surgery bicuspid	E	E					
D3425	Root surgery molar	E	E					
D3426	Root surgery ea add root	E	E					
D3430	Retrograde filling	E	E					
D3450	Root amputation	E	E					
D3460	Endodontic endosseous implan	S	S	0330	10.3227	\$695.81		\$139.17
D3470	Intentional replantation	E	E					
D3910	Isolation- tooth w rubb dam	E	E					
D3920	Tooth splitting	E	E					
D3950	Canal prep/fitting of dowel	E	E					
D3999	Endodontic procedure	S	S	0330	10.3227	\$695.81		\$139.17
D4210	Gingivectomy/plasty per quad	E	E					
D4211	Gingivectomy/plasty per root	E	E					
D4230	Ana crown exp 4 or> per quad	E	E					
D4231	Ana crown exp 1-3 per quad	E	E					
D4240	Gingival flap proc w/ plann	E	E					
D4241	Gingiv flap w rootplan 1-3 th	E	E					
D4245	Apically positioned flap	E	E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D2650	Inlay composite/resin one su	E	E					
D2651	Inlay composite/resin two su	E	E					
D2652	Dental inlay resin 3/mre sur	E	E					
D2662	Dental onlay resin 2 surface	E	E					
D2663	Dental onlay resin 3 surface	E	E					
D2664	Dental onlay resin 4/mre sur	E	E					
D2710	Crown resin-based indirect	E	E					
D2712	Crown 3/4 resin-based compos	E	E					
D2720	Crown resin w/ high noble me	E	E					
D2721	Crown resin w/ base metal	E	E					
D2722	Crown resin w/ noble metal	E	E					
D2740	Crown porcelain/ceramic subs	E	E					
D2750	Crown porcelain w/ h noble m	E	E					
D2751	Crown porcelain fused base m	E	E					
D2752	Crown porcelain w/ noble met	E	E					
D2780	Crown 3/4 cast h/ noble met	E	E					
D2781	Crown 3/4 cast base metal	E	E					
D2782	Crown 3/4 cast noble metal	E	E					
D2783	Crown 3/4 porcelain/ceramic	E	E					
D2790	Crown full cast high noble m	E	E					
D2791	Crown full cast base metal	E	E					
D2792	Crown full cast noble metal	E	E					
D2794	Crown-titanium	E	E					
D2799	Provisional crown	E	E					
D2910	Recement inlay onlay or part	E	E					
D2915	Recement cast or prefab post	E	E					
D2920	Dental recement crown	E	E					
D2930	Prefab smless steel crown pri	E	E					
D2931	Prefab smless steel crown pe	E	E					
D2932	Prefabricated resin crown	E	E					
D2933	Prefab stainless steel crown	E	E					
D2934	Prefab steel crown primary	E	E					
D2940	Dental sedative filling	E	E					
D2950	Core build-up incl any pins	E	E					
D2951	Tooth pin retention	E	E					
D2952	Post and core cast + crown	E	E					
D2953	Each addtl cast post	E	E					
D2954	Prefab post/core + crown	E	E					
D2955	Post removal	E	E					
D2957	Each addtl prefab post	E	E					
D2960	Laminate labial veneer	E	E					
D2961	Lab labial veneer resin	E	E					
D2962	Lab labial veneer porcelain	E	E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D5630	Rep partial denture clasp	E	E					
D5640	Replace part denture teeth	E	E					
D5650	Add tooth to partial denture	E	E					
D5660	Add clasp to partial denture	E	E					
D5670	Repic thâ on mtl firmwk	E	E					
D5671	Repic thâ mandibular	E	E					
D5710	Dentures rebase cmplt maxil	E	E					
D5711	Dentures rebase cmplt mand	E	E					
D5720	Dentures rebase part maxil	E	E					
D5721	Dentures rebase part mand	E	E					
D5730	Denture rein cmplt maxil chr	E	E					
D5731	Denture rein cmplt mand chr	E	E					
D5740	Denture rein part mand chr	E	E					
D5741	Denture rein part mand chr	E	E					
D5750	Denture rein cmplt max lab	E	E					
D5751	Denture rein cmplt mand lab	E	E					
D5760	Denture rein part maxil lab	E	E					
D5761	Denture rein part mand lab	E	E					
D5810	Denture interim cmplt maxil	E	E					
D5811	Denture interim cmplt mand	E	E					
D5820	Denture interim part maxil	E	E					
D5821	Denture interim part mand	E	E					
D5850	Denture liss conditn maxil	E	E					
D5851	Denture liss conditn mand	E	E					
D5860	Overdenture complete	E	E					
D5861	Overdenture partial	E	E					
D5862	Precision attachment	E	E					
D5867	Replacement of precision att	E	E					
D5875	Prosthesis modification	E	E					
D5899	Removable prosthodontic proc	E	E					
D5911	Facial moulage sectional	S	S	0330	10.3227	\$695.81		\$139.17
D5912	Facial moulage complete	S	S	0330	10.3227	\$695.81		\$139.17
D5913	Nasal prosthesis	E	E					
D5914	Auricular prosthesis	E	E					
D5915	Orbital prosthesis	E	E					
D5916	Ocular prosthesis	E	E					
D5919	Facial prosthesis	E	E					
D5922	Nasal septal prosthesis	E	E					
D5923	Ocular prosthesis interim	E	E					
D5924	Cranial prosthesis	E	E					
D5925	Facial augmentation implant	E	E					
D5926	Replacement nasal prosthesis	E	E					
D5927	Auricular replacement	E	E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D4249	Crown lengthen hard tissue	E	E					
D4260	Osseous surgi-3teethperquad	S	S	0330	10.3227	\$695.81		\$139.17
D4261	Osseous surgi-3teethperquad	E	E					
D4263	Bone replce graft first site	S	S	0330	10.3227	\$695.81		\$139.17
D4264	Bone replce graft each add	S	S	0330	10.3227	\$695.81		\$139.17
D4265	Blo mifils to aid soft/os reg	E	E					
D4266	Guided liss regen resorb	E	E					
D4267	Guided tiss regen nonresorb	E	E					
D4268	Surgical revision procedure	S	S	0330	10.3227	\$695.81		\$139.17
D4270	Pericle soft tissue graft pr	S	S	0330	10.3227	\$695.81		\$139.17
D4271	Free soft tissue graft proc	S	S	0330	10.3227	\$695.81		\$139.17
D4273	Subepithelial tissue graft	S	S	0330	10.3227	\$695.81		\$139.17
D4274	Distal/proximal wedge proc	E	E					
D4275	Soft tissue allograft	E	E					
D4276	Con tissue w dble ped graft	E	E					
D4320	Provision splint intracoronal	E	E					
D4321	Provisional splint extracor	E	E					
D4341	Periodontal scaling & root	E	E					
D4342	Periodontal scaling 1-3teeth	E	E					
D4355	Full mouth debridement	S	S	0330	10.3227	\$695.81		\$139.17
D4381	Localized delivery antimicro	S	S	0330	10.3227	\$695.81		\$139.17
D4910	Periodontal maint procedures	E	E					
D4920	Unscheduled dressing change	E	E					
D4999	Unspecified periodontal proc	E	E					
D5110	Dentures complete maxillary	E	E					
D5120	Dentures complete mandible	E	E					
D5130	Dentures immediat maxillary	E	E					
D5140	Dentures immediat mandible	E	E					
D5211	Dentures maxil part resin	E	E					
D5212	Dentures mand part resin	E	E					
D5213	Dentures maxil part metal	E	E					
D5214	Dentures mandib part metal	E	E					
D5225	Maxillary part denture flex	E	E					
D5226	Mandibular part denture flex	E	E					
D5281	Removable partial denture	E	E					
D5410	Dentures adjust cmplt maxil	E	E					
D5411	Dentures adjust cmplt mand	E	E					
D5421	Dentures adjust part maxil	E	E					
D5422	Dentures adjust part mand	E	E					
D5510	Dentur repr broken compl bas	E	E					
D5520	Replace denture teeth cmplt	E	E					
D5610	Dentures repair resin base	E	E					
D5620	Rep part denture cast frame	E	E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D6066	Implant supported mtl crown	E	E					
D6067	Implant supported mtl crown	E	E					
D6068	Abutment supported retainer	E	E					
D6069	Abutment supported retainer	E	E					
D6070	Abutment supported retainer	E	E					
D6071	Abutment supported retainer	E	E					
D6072	Abutment supported retainer	E	E					
D6073	Abutment supported retainer	E	E					
D6074	Abutment supported retainer	E	E					
D6075	Implant supported retainer	E	E					
D6076	Implant supported retainer	E	E					
D6077	Implant supported retainer	E	E					
D6078	Implant/abut supprt fixd dent	E	E					
D6079	Implnt/abut supprt fixd dent	E	E					
D6080	Implant maintenance	E	E					
D6090	Repair implant	E	E					
D6091	Repl semi/precision attach	E	E					
D6092	Recement supp crown	E	E					
D6093	Recement supp part denture	E	E					
D6094	Abut support crown titanium	E	E					
D6095	Odontics repr abutment	E	E					
D6100	Removal of implant	E	E					
D6190	Radio/surgical implant index	E	E					
D6194	Abut support retainer titani	E	E					
D6199	Implant procedure	E	E					
D6205	Pontic-indirect resin based	E	E					
D6210	Prosthodont high noble metal	E	E					
D6211	Bridge base metal cast	E	E					
D6212	Bridge noble metal cast	E	E					
D6214	Pontic titanium	E	E					
D6240	Bridge porcelain high noble	E	E					
D6241	Bridge porcelain base metal	E	E					
D6242	Bridge porcelain nobel metal	E	E					
D6245	Bridge porcelain/ceramic	E	E					
D6250	Bridge resin w/high noble	E	E					
D6251	Bridge resin base metal	E	E					
D6252	Bridge resin w/noble metal	E	E					
D6253	Provisional pontic	E	E					
D6545	Dental retainr cast metl	E	E					
D6548	Porcelain/ceramic retainer	E	E					
D6600	Porcelain/ceramic inlay 2srf	E	E					
D6601	Porc/ceram inlay >= 3 surfac	E	E					
D6602	Cst high nble mtl inlay 2 srf	E	E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D5928	Orbital replacement	E	E					
D5929	Facial replacement	E	E					
D5931	Surgical obturator	E	E					
D5932	Postsurgical obturator	E	E					
D5933	Refitting of obturator	E	E					
D5934	Mandibular flange prosthesis	E	E					
D5935	Mandibular denture prosth	E	E					
D5936	Temp obturator prosthesis	E	E					
D5937	Trismus appliance	E	E					
D5951	Feeding aid	E	E					
D5952	Pediatric speech aid	E	E					
D5953	Adult speech aid	E	E					
D5954	Superimposed prosthesis	E	E					
D5955	Palatal lift prosthesis	E	E					
D5958	Intraoral con def rtrr plit	E	E					
D5959	Intraoral con def mod palat	E	E					
D5960	Modify speech aid prosthesis	E	E					
D5982	Surgical stent	E	E					
D5983	Radiation applicator	S	S	0330	10.3227	\$695.81		\$139.17
D5984	Radiation shield	S	S	0330	10.3227	\$695.81		\$139.17
D5985	Radiation cone locator	S	S	0330	10.3227	\$695.81		\$139.17
D5986	Fluoride applicator	E	E					
D5987	Commissionure splint	S	S	0330	10.3227	\$695.81		\$139.17
D5988	Surgical splint	E	E					
D5991	Topical medicament carrier	E	E					
D5999	Maxillofacial prosthesis	E	E					
D6010	Odontics endosteal implant	E	E					
D6012	Endosteal implant	E	E					
D6040	Odontics eposteal implant	E	E					
D6050	Odontics transosteal impint	E	E					
D6053	Implnt/abtmnt spprt remv dnt	E	E					
D6054	Implnt/abtmnt spprt remyvrnl	E	E					
D6055	Implant connecting bar	E	E					
D6056	Prefabricated abutment	E	E					
D6057	Custom abutment	E	E					
D6058	Abutment supported crown	E	E					
D6059	Abutment supported mtl crown	E	E					
D6060	Abutment supported mtl crown	E	E					
D6061	Abutment supported mtl crown	E	E					
D6062	Abutment supported mtl crown	E	E					
D6063	Abutment supported mtl crown	E	E					
D6064	Abutment supported mtl crown	E	E					
D6065	Implant supported crown	E	E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D6985	Pediatric partial denture fx	E	E					
D6989	Fixed prosthodontic proc	E	E					
D7111	Extraction coronal remnants	S	S	0330	10.3227	\$695.81		\$139.17
D7140	Extraction erupted tooth/lext	S	S	0330	10.3227	\$695.81		\$139.17
D7210	Rem imp tooth w mucoper flap	S	S	0330	10.3227	\$695.81		\$139.17
D7220	Impact tooth remov soft tiss	S	S	0330	10.3227	\$695.81		\$139.17
D7230	Impact tooth remov part bony	S	S	0330	10.3227	\$695.81		\$139.17
D7240	Impact tooth remov comp bony	S	S	0330	10.3227	\$695.81		\$139.17
D7241	Impact tooth rem bony w/comp	S	S	0330	10.3227	\$695.81		\$139.17
D7250	Tooth root removal	S	S	0330	10.3227	\$695.81		\$139.17
D7260	Oral antral fistula closure	S	S	0330	10.3227	\$695.81		\$139.17
D7261	Primary closure sinus perf	S	S	0330	10.3227	\$695.81		\$139.17
D7270	Tooth reimplantation	E	E					
D7272	Tooth transplantation	E	E					
D7280	Exposure impact tooth onftod	E	E					
D7282	Mobilize erupted/malpos toot	E	E					
D7283	Place device impacted tooth	B	B					
D7285	Biopsy of oral tissue hard	E	E					
D7286	Biopsy of oral tissue soft	E	E					
D7287	Exfoliative cytolog collect	E	E					
D7288	Brush biopsy	B	B					
D7290	Repositioning of teeth	E	E					
D7291	Transseptal fiberotomy	S	S	0330	10.3227	\$695.81		\$139.17
D7292	Screw retained plate	E	E					
D7294	Temp anchorage dev w/o flap	E	E					
D7310	Alveoplasty w/ extraction	E	E					
D7311	Alveoplasty w/extract 1-3	E	E					
D7321	Alveoplasty not w/extracts	B	B					
D7340	Vestibuloplasty ridge extens	E	E					
D7410	Rad exc lesion up to 1.25 cm	E	E					
D7411	Excision benign lesion>1.25c	E	E					
D7412	Excision benign lesion compl	E	E					
D7413	Excision malign lesion<=1.25c	E	E					
D7414	Excision malign lesion>1.25cm	E	E					
D7415	Excision malign les complicat	E	E					
D7440	Malign tumor exc to 1.25 cm	E	E					
D7441	Malign tumor > 1.25 cm	E	E					
D7450	Rem odontogen cyst to 1.25cm	E	E					
D7451	Rem odontogen cyst > 1.25 cm	E	E					
D7460	Rem nonodontic cyst to 1.25cm	E	E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D6603	Cst high noble mtl inlay >=3sr	E	E					
D6604	Cst base mtl inlay 2 surfaces	E	E					
D6605	Cst base mtl inlay >= 3 surfa	E	E					
D6606	Cast noble metal inlay 2 sur	E	E					
D6607	Cst noble mtl inlay >=3 surf	E	E					
D6608	Onlay poro/crmc 2 surfaces	E	E					
D6609	Onlay poro/crmc >=3 surfaces	E	E					
D6610	Onlay cst high nbl mtl 2 surfc	E	E					
D6611	Onlay cst high nbl mtl >=3srfl	E	E					
D6612	Onlay cst base mtl 2 surface	E	E					
D6613	Onlay cst base mtl >=3 surfa	E	E					
D6615	Onlay cst nbl mtl 2 surfaces	E	E					
D6616	Onlay cst nbl mtl >=3 surfac	E	E					
D6624	Inlay titanium	E	E					
D6624	Inlay titanium	E	E					
D6710	Crown-indirect resin based	E	E					
D6720	Retain crown resin w hi nble	E	E					
D6721	Crown resin w/base metal	E	E					
D6722	Crown resin w/noble metal	E	E					
D6740	Crown porcelain/ceramic	E	E					
D6750	Crown porcelain high noble	E	E					
D6751	Crown porcelain base metal	E	E					
D6752	Crown porcelain noble metal	E	E					
D6780	Crown 3/4 high noble metal	E	E					
D6781	Crown 3/4 cast based metal	E	E					
D6782	Crown 3/4 cast noble metal	E	E					
D6783	Crown 3/4 porcelain/ceramic	E	E					
D6790	Crown full high noble metal	E	E					
D6791	Crown full base metal cast	E	E					
D6792	Crown full noble metal cast	E	E					
D6793	Provisional retainer crown	E	E					
D6794	Crown titanium	E	E					
D6820	Dental connector bar	S	S	0330	10.3227	\$695.81		\$139.17
D6930	Dental cement bridge	E	E					
D6940	Stress breaker	E	E					
D6950	Precision attachment	E	E					
D6970	Post & core plus retainer	E	E					
D6972	Prefab post & core plus reta	E	E					
D6973	Core build up for retainer	E	E					
D6975	Coping metal	E	E					
D6976	Each addtl cast post	E	E					
D6977	Each addtl prefab post	E	E					
D6980	Bridge repair	E	E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D7865	Tmj reshaping components		E					
D7870	Tmj aspiration joint fluid		E					
D7871	Lysis + lavage w catheters		E					
D7872	Tmj diagnostic arthroscopy		E					
D7873	Tmj arthroscopy lysis adhes		E					
D7874	Tmj arthroscopy disc reposit		E					
D7875	Tmj arthroscopy synovectomy		E					
D7876	Tmj arthroscopy discectomy		E					
D7877	Tmj arthroscopy debridement		E					
D7880	Occlusal orthotic appliance		E					
D7899	Tmj unspecified therapy		E					
D7910	Dent suture recent wnd to 5cm		E					
D7911	Dental suture wound to 5 cm		E					
D7912	Suture complicate wnd > 5 cm		E					
D7920	Dental skin graft		E					
D7940	Reshaping bone orthognathic		S	0330	10.3227	\$695.81		\$139.17
D7941	Bone cutting ramus closed		E					
D7943	Cutting ramus open w/graft		E					
D7944	Bone cutting segmented		E					
D7945	Bone cutting body mandible		E					
D7946	Reconstruct maxilla total		E					
D7947	Reconstruct maxilla segment		E					
D7948	Reconstruct midface no graft		E					
D7949	Reconstruct midface w/graft		E					
D7950	Mandible graft		E					
D7951	Sinus aug w bone/bone sup		E					
D7953	Bone replacement graft		E					
D7955	Repair maxillofacial defects		E					
D7960	Frenulectomy/frenulotomy		E					
D7963	Frenuloplasty		E					
D7970	Excision hyperplastic tissue		E					
D7971	Excision pericoronal gingiva		E					
D7972	Surg reduct fibrous tuberosit		E					
D7980	Sialolithomy		E					
D7981	Excision of salivary gland		E					
D7982	Sialodochoplasty		E					
D7983	Closure of salivary fistula		E					
D7990	Emergency tracheotomy		E					
D7991	Dental coronodectomy		E					
D7995	Synthetic graft facial bones		E					
D7996	Implant mandible for augment		E					
D7997	Appliance removal		E					
D7998	Intraoral place of fix dev		E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D7461	Rem nonodonto cyst > 1.25 cm		E					
D7465	Lesion destruction		E					
D7471	Rem exostosis any site		E					
D7472	Removal of torus palatinus		E					
D7473	Remove torus mandibularis		E					
D7485	Surg reduct osseoustuberosit		E					
D7490	Maxilla or mandible resectio		E					
D7510	I&d abscc intraoral soft tiss		E					
D7511	Incision/drain abscess intra		B					
D7520	I&d abscess extraoral		E					
D7521	Incision/drain abscess extra		B					
D7530	Removal fb skin/areolar tiss		E					
D7540	Removal of fb reaction		E					
D7550	Removal of sloughed off bone		E					
D7560	Maxillary sinusotomy		E					
D7610	Maxilla open reduct simple		E					
D7620	Cisd reduct simpl maxilla fx		E					
D7630	Open red simpl mandible fx		E					
D7640	Cisd red simpl mandible fx		E					
D7650	Open red simp malar/zygom fx		E					
D7660	Cisd red simp malar/zygom fx		E					
D7670	Closed reduct splint alveolus		E					
D7671	Alveolus open reduction		E					
D7680	Reduct simple facial bone fx		E					
D7710	Maxilla open reduct compound		E					
D7720	Cisd reduct compd maxilla fx		E					
D7730	Open reduct compd mandible fx		E					
D7740	Cisd reduct compd mandible fx		E					
D7750	Open red comp malar/zygma fx		E					
D7760	Cisd red comp malar/zygma fx		E					
D7770	Open reduct compd alveolus fx		E					
D7771	Alveolus cisd reduct sribz te		E					
D7780	Reduct compnd facial bone fx		E					
D7810	Tmj open reduct-dislocation		E					
D7820	Closed tmp manipulation		E					
D7830	Tmj manipulation under anest		E					
D7840	Removal of tmj condyle		E					
D7850	Tmj meniscectomy		E					
D7852	Tmj repair of joint disc		E					
D7854	Tmj excision of joint membrane		E					
D7856	Tmj cutting of a muscle		E					
D7858	Tmj reconstruction		E					
D7860	Tmj cutting into joint		E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D9920	Behavior management	E	S	0330	10.3227	\$695.81		\$139.17
D9930	Treatment of complications	E	S	0330	10.3227	\$695.81		\$139.17
D9940	Dental occlusal guard	E	S	0330	10.3227	\$695.81		\$139.17
D9941	Fabrication athletic guard	E	S	0330	10.3227	\$695.81		\$139.17
D9942	Repair/reline occlusal guard	E	S	0330	10.3227	\$695.81		\$139.17
D9950	Occlusion analysis	E	S	0330	10.3227	\$695.81		\$139.17
D9951	Limited occlusal adjustment	E	S	0330	10.3227	\$695.81		\$139.17
D9952	Complete occlusal adjustment	E	S	0330	10.3227	\$695.81		\$139.17
D9970	Enamel microabrasion	E	E					
D9971	Odontoplasty 1-2 teeth	E	E					
D9972	Extrnl bleaching per arch	E	E					
D9973	Extrnl bleaching per tooth	E	E					
D9974	Intrnl bleaching per tooth	E	E					
D9999	Adjunctive procedure	E	E					
E0100	Cane adjust/fixd with tip		Y					
E0105	Cane adjust/fixd quad/3 pro		Y					
E0110	Crutch forearm pair		Y					
E0111	Crutch forearm each		Y					
E0112	Crutch underarm pair wood		Y					
E0113	Crutch underarm each wood		Y					
E0114	Crutch underarm pair no wood		Y					
E0116	Crutch underarm each no wood		Y					
E0117	Underarm springassist crutch		Y					
E0118	Crutch substitute		E					
E0130	Walker rigid adjust/fixd ht		Y					
E0135	Walker folding adjust/fixd		Y					
E0140	Walker w trunk support		Y					
E0141	Rigid wheeled walker adj/fix		Y					
E0143	Walker folding wheeled w/o s		Y					
E0144	Enclosed walker w rear seat		Y					
E0147	Walker variable wheel resist		Y					
E0148	Heavyduty walker no wheels		Y					
E0149	Heavy duty wheeled walker		Y					
E0153	Forearm crutch platform atta		Y					
E0154	Walker platform attachment		Y					
E0155	Walker wheel attachment,pair		Y					
E0156	Walker seat attachment		Y					
E0157	Walker crutch attachment		Y					
E0158	Walker leg extenders set of4		Y					
E0159	Brake for wheeled walker		Y					
E0160	Sitz type bath or equipment		Y					
E0161	Sitz bath/equipment w/faucet		Y					
E0162	Sitz bath chair		Y					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
D7999	Oral surgery procedure	E	E					
D8010	Limited dental tx primary	E	E					
D8020	Limited dental tx transition	E	E					
D8030	Limited dental tx adolescent	E	E					
D8040	Limited dental tx adult	E	E					
D8050	Intercep dental tx primary	E	E					
D8060	Intercep dental tx transitn	E	E					
D8070	Compre dental tx transition	E	E					
D8080	Compre dental tx adolescent	E	E					
D8090	Compre dental tx adult	E	E					
D8210	Orthodontic rem appliance tx	E	E					
D8220	Fixed appliance therapy habtt	E	E					
D8660	Preorthodontic tx visit	E	E					
D8670	Periodic orthodontic tx visit	E	E					
D8680	Orthodontic retention	E	E					
D8690	Orthodontic treatment	E	E					
D8691	Repair ortho appliance	E	E					
D8692	Replacement retainer	E	E					
D8693	Rebond/cement/repair retain	E	E					
D8999	Orthodontic procedure	E	E					
D9110	Tx dental pain minor proc	N	N					
D9120	Fix partial denture section	E	E					
D9210	Dent anesthesia w/o surgery	E	E					
D9211	Regional block anesthesia	E	E					
D9212	Trigeminal block anesthesia	E	E					
D9215	Local anesthesia	E	E					
D9220	General anesthesia	E	E					
D9221	General anesthesia ea ad 15m	E	E					
D9230	Analgesia	N	N					
D9241	Intravenous sedation	E	E					
D9242	IV sedation ea ad 30 m	E	E					
D9248	Sedation (non-iv)	N	N					
D9310	Dental consultation	E	E					
D9410	Dental house call	E	E					
D9420	Hospital call	E	E					
D9430	Office visit during hours	E	E					
D9440	Office visit after hours	E	E					
D9450	Case presentation tx plan	E	E					
D9610	Dent therapeutic drug inject	E	E					
D9612	Thera par drugs 2 or > admin	E	E					
D9630	Other drugs/medications	S	S	0330	10.3227	\$695.81		\$139.17
D9910	Dent appl desensitizing med	E	E					
D9911	Appl desensitizing resin	E	E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0240	Bath/shower chair		E					
E0241	Bath tub wall rail		E					
E0242	Bath tub, rail floor		E					
E0243	Toilet rail		E					
E0244	Toilet seat raised		E					
E0245	Tub stool or bench		E					
E0246	Transfer tub rail attachment		E					
E0247	Trans bench w/w/o comm open HD/trans bench w/w/o comm open		E					
E0248	open		E					
E0249	Pad water circulating heat u		Y					
E0250	Hosp bed fixed ht w/ mattress		Y					
E0251	Hosp bed fixed ht w/o mattress		Y					
E0255	Hospital bed var ht w/ matr		Y					
E0256	Hospital bed var ht w/o matr		Y					
E0260	Hosp bed semi-elect w/ matr		Y					
E0261	Hosp bed semi-elect w/o matr		Y					
E0265	Hosp bed total elect w/ mat		Y					
E0266	Hosp bed total elec w/o matr		Y					
E0270	Hospital bed institutional t		E					
E0271	Mattress innerspring		Y					
E0272	Mattress foam rubber		Y					
E0273	Bed board		E					
E0274	Over-bed table		E					
E0275	Bed pan standard		Y					
E0276	Bed pan fracture		Y					
E0277	Powered pres-redu air mattress		Y					
E0280	Bed cradle		Y					
E0290	Hosp bed fx ht w/o rails w/m		Y					
E0291	Hosp bed fx ht w/o rail w/o		Y					
E0292	Hosp bed var ht w/o rail w/o		Y					
E0293	Hosp bed var ht w/o rail w/		Y					
E0294	Hosp bed semi-elect w/ matr		Y					
E0295	Hosp bed semi-elect w/o matr		Y					
E0296	Hosp bed total elect w/ matr		Y					
E0297	Hosp bed total elect w/o matr		Y					
E0300	Enclosed ped crib hosp grade		Y					
E0301	HD hosp bed, 350-600 lbs		Y					
E0302	Ex hd hosp bed > 600 lbs		Y					
E0303	Hosp bed hvy dty extra wide		Y					
E0304	Hosp bed extra hvy dty x wide		Y					
E0305	Rails bed side half length		Y					
E0310	Rails bed side full length		Y					
E0315	Bed accessory brd/tbl/supprt		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0163	Commode chair with fixed arm		Y					
E0165	Commode chair with		Y					
E0167	Commode chair, pail or pan detacharm		Y					
E0168	Heavyduty/wide commode chair		Y					
E0170	Commode chair electric		Y					
E0171	Commode chair non-electric		Y					
E0172	Seat lift mechanism toilet		E					
E0175	Commode chair foot rest		Y					
E0181	Press pad alternating w/ pum		Y					
E0182	Replace pump, alt press pad		Y					
E0184	Dry pressure mattress		Y					
E0185	Gel pressure mattress pad		Y					
E0186	Air pressure mattress		Y					
E0187	Water pressure mattress		Y					
E0188	Synthetic sheepskin pad		Y					
E0189	Lambswool sheepskin pad		Y					
E0190	Positioning cushion		E					
E0191	Protector heel or elbow		Y					
E0193	Powered air flotation bed		Y					
E0194	Air fluidized bed		Y					
E0196	Gel pressure mattress		Y					
E0197	Air pressure pad for mattress		Y					
E0198	Water pressure pad for matr		Y					
E0199	Dry pressure pad for mattress		Y					
E0200	Heat lamp without stand		Y					
E0202	Phototherapy light w/ photom		Y					
E0203	Therapeutic lightbox tabletp		E					
E0205	Heat lamp with stand		Y					
E0210	Electric heat pad standard		Y					
E0215	Electric heat pad moist		Y					
E0217	Water circ heat pad w pump		Y					
E0218	Water circ cold pad w pump		Y					
E0220	Hot water bottle		Y					
E0221	Infrared heating pad system		Y					
E0225	Hydrocollator unit		Y					
E0230	Ice cap or collar		Y					
E0231	Wound warming device		E					
E0232	Warming card for NWT		E					
E0235	Parafin bath unit portable		Y					
E0236	Pump for water circulating p		Y					
E0238	Heat pad non-electric moist		Y					
E0239	Hydrocollator unit portable		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0486	Oral device/appliance cusfab		Y					
E0487	Electronic spirometer		N					
E0500	Ippb, all types		Y					
E0500	Ippb, all types		Y					
E0500	Ippb, all types		Y					
E0550	Humidif extens supplie w ippb		Y					
E0555	Humidifier for use w/ regula		Y					
E0560	Humidifier supplemental w/ i		Y					
E0561	Humidifier nonheated w PAP		Y					
E0562	Humidifier heated used w PAP		Y					
E0565	Compressor air power source		Y					
E0566	Compressor with compression		Y					
E0571	Aerosol compressor for seneb		Y					
E0572	Aerosol compressor adjust pr		Y					
E0574	Ultrasonic generator w svneb		Y					
E0575	Nebulizer ultrasonic		Y					
E0580	Nebulizer for use w/ regulat		Y					
E0585	Nebulizer w/ compressor & he		Y					
E0600	Suction pump portab horn modl		Y					
E0601	Cont airway pressure device		Y					
E0602	Manual breast pump		Y					
E0603	Electric breast pump		N					
E0604	Hosp grade elec breast pump		A					
E0605	Vaporizer room type		Y					
E0606	Drainage board postural		Y					
E0607	Blood glucose monitor home		Y					
E0610	Pacemaker monitr audible/vis		Y					
E0615	Pacemaker monitr digital/vis		Y					
E0616	Cardiac event recorder		N					
E0617	Automatic ext defibrillator		Y					
E0618	Apnea monitor		Y					
E0619	Apnea monitor w recorder		Y					
E0620	Cap bid skin piercing laser		Y					
E0621	Patient lift sling or seat		Y					
E0625	Patient lift bathroom or toi		E					
E0627	Seat lift incorp lift-chair		Y					
E0628	Seat lift for pt turn-electr		Y					
E0629	Seat lift for pt turn-non-el		Y					
E0630	Patient lift hydraulic		Y					
E0635	Patient lift electric		Y					
E0636	PT support & positioning sys		Y					
E0637	Combination sit to stand sys		E					
E0638	Standing frame Sys		E					
E0639	Moveable patient lift system		E					
E0640	Fixed patient lift system		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0316	Bed safety enclosure		Y					
E0325	Urinal male lug-type		Y					
E0326	Urinal female lug-type		Y					
E0328	Ped hospital bed, manual		Y					
E0329	Ped hospital bed semi/elect		Y					
E0350	Control unit bowel system		E					
E0352	Disposable pack w/bowel syst		E					
E0370	Air elevator for heel		E					
E0371	Nonpower mattress overlay		Y					
E0372	Powered air mattress overlay		Y					
E0373	Nonpowered pressure mattress		Y					
E0424	Stationary compressed gas O2		Y					
E0425	Gas system stationary compre		E					
E0430	Oxygen system, gas portable		E					
E0431	Portable gaseous O2		Y					
E0433	Portable liquid oxygen sys	NI	Y					
E0434	Portable liquid oxygen sys		Y					
E0435	Oxygen system, liquid portabl		Y					
E0439	Stationary liquid O2		Y					
E0440	Oxygen system liquid station		E					
E0441	Stationary O2 contents, gas		Y					
E0442	Stationary O2 contents, liq		Y					
E0443	Portable O2 contents, gas		Y					
E0444	Portable O2 contents, liquid		Y					
E0445	Oximeter non-invasive		N					
E0450	Vol control vent invasiv int		Y					
E0455	Oxygen tent excl crouched t		Y					
E0457	Chest shell		Y					
E0459	Chest wrap		Y					
E0460	Neg press vent portabl/stati		Y					
E0461	Vol control vent noninv int		Y					
E0462	Rocking bed w/ or w/o side r		Y					
E0463	Press supp vent invasive int		Y					
E0464	Press supp vent noninv int		Y					
E0470	RAD w/o backup non-inv intrfc		Y					
E0471	RAD w/backup non inv intrfc		Y					
E0472	RAD w backup invasive intrfc		Y					
E0480	Percussor elec/pneum home		Y					
E0481	Intrplumtry percuss vent sys		E					
E0482	Cough stimulating device		Y					
E0483	Chest compression gen system		Y					
E0484	Non-elec oscillatory pep dvc		Y					
E0485	Oral device/appliance prefab		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0769	Electric wound treatment dev		B					
E0770	Functional electric stim NOS		Y					
E0776	Iv pole		Y					
E0779	Amb infusion pump mechanical		Y					
E0780	Mech amb infusion pump <8hrs		Y					
E0781	External ambulatory infus pu		Y					
E0782	Non-programmable infusion pump		N					
E0783	Programmable infusion pump		Y					
E0784	Ext amb infus pump insulin		N					
E0785	Replacement impl pump cathet		N					
E0786	Implantable pump replacement		N					
E0791	Parenteral infusion pump sta		Y					
E0830	Ambulatory traction device		N					
E0840	Tract frame attach headboard		Y					
E0849	Cervical pneum trac equip		Y					
E0850	Traction stand free standing		Y					
E0855	Cervical traction equipment		Y					
E0856	Cervic collar w air bladder		Y					
E0860	Tract equip cervical tract		Y					
E0870	Tract frame attach toothboard		Y					
E0880	Trac stand free stand extrem		Y					
E0890	Traction frame attach pelvic		Y					
E0900	Trac stand free stand pelvic		Y					
E0910	Trapeze bar attached to bed		Y					
E0911	HD trapeze bar attach to bed		Y					
E0912	HD trapeze bar free standing		Y					
E0920	Fracture frame attached to b		Y					
E0930	Fracture frame free standing		Y					
E0935	Cont pas motion exercise dev		Y					
E0936	CPM device, other than knee		E					
E0940	Trapeze bar free standing		Y					
E0941	Gravity assisted traction de		Y					
E0942	Cervical head harness/halter		Y					
E0944	Pelvic belt/harness/boot		Y					
E0945	Belt/harness extremity		Y					
E0946	Fracture frame dual w cross		Y					
E0947	Fracture frame attachments pe		Y					
E0948	Fracture frame attachments ce		Y					
E0950	Tray		Y					
E0951	Loop heel		Y					
E0952	Toe loop/holder, each		Y					
E0955	Cushioned headrest		Y					
E0956	W/c lateral trunk/hip suppor		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0641	Multi-position stand fram sys		E					
E0642	Dynamic standing frame		E					
E0650	Pneuma compressor non-segment		Y					
E0651	Pneum compressor segmental		Y					
E0652	Pneum compres wical pressure		Y					
E0655	Pneumatic appliance half arm		Y					
E0656	Segmental pneumatic trunk		Y					
E0657	Segmental pneumatic chest		Y					
E0660	Pneumatic appliance full leg		Y					
E0665	Pneumatic appliance full arm		Y					
E0666	Pneumatic appliance half leg		Y					
E0667	Seg pneumatic appl full leg		Y					
E0668	Seg pneumatic appl full arm		Y					
E0669	Seg pneumatic appl half leg		Y					
E0671	Pressure pneum appl full leg		Y					
E0672	Pressure pneum appl full arm		Y					
E0673	Pressure pneum appl half leg		Y					
E0675	Pneumatic compression device		Y					
E0676	Inter limb compress dev NOS		Y					
E0691	Uvi prt 2 sq ft or less		Y					
E0692	Uvi sys panel 4 ft		Y					
E0693	Uvi sys panel 6 ft		Y					
E0694	Uvi md cabinet sys 6 ft		Y					
E0700	Safety equipment		E					
E0705	Transfer device		B					
E0710	Restraints any type		E					
E0720	Tens two lead		Y					
E0730	Tens four lead		Y					
E0731	Conductive garment for lens/		Y					
E0740	Incontinence treatment system		Y					
E0744	Neuromuscular stim for scoli		Y					
E0745	Neuromuscular stim for shock		Y					
E0746	Electromyograph biofeedback		N					
E0747	Elec osteogen stim not spine		Y					
E0748	Elec osteogen stim spinal		Y					
E0749	Elec osteogen stim implanted		N					
E0755	Electronic salivary reflex s		E					
E0760	Osteogen ultrasound stimtor		Y					
E0761	Nontherm electromagnetic device		E					
E0762	Trans elec if stim dev sys		B					
E0764	Functional neuromuscular stim		Y					
E0765	Nerve stimulator for tx n&v		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E1030	W/c vent tray gimbal		Y					
E1031	Rollabout chair with casters		Y					
E1035	Patient transfer system <300		Y					
E1036	Patient transfer system >300	NI	Y					
E1037	Transport chair, ped size		Y					
E1038	Transport chair pt wt<=300lb		Y					
E1039	Transport chair pt wt >300lb		Y					
E1050	Wheelch fxd full length arms		Y					
E1060	Wheelchair detachable arms		Y					
E1070	Wheelchair detachable foot r		Y					
E1083	Hemi-wheelchair fixed arms		Y					
E1084	Hemi-wheelchair detachable a		Y					
E1085	Hemi-wheelchair fixed arms		E					
E1086	Hemi-wheelchair detachable a		E					
E1087	Wheelchair lghtwt fixed arm		Y					
E1088	Wheelchair lghtweight det a		Y					
E1089	Wheelchair lghtwt fixed arm		E					
E1090	Wheelchair lghtweight det a		E					
E1092	Wheelchair wide w/ leg rests		Y					
E1093	Wheelchair wide w/ foot rest		Y					
E1100	Wheelch s-recl fxd arm leg res		Y					
E1110	Wheelchair semi-recl detach		Y					
E1130	Wheelch stand fxd arm ft rest		E					
E1140	Wheelchair standard detach a		E					
E1150	Wheelchair standard w/ leg r		Y					
E1160	Wheelchair fixed arms		Y					
E1161	Manual adult wc w tiltingpac		Y					
E1170	Wheelch ampu fxd arm leg rest		Y					
E1171	Wheelchair amputee w/o leg r		Y					
E1172	Wheelchair amputee detach ar		Y					
E1180	Wheelchair amputee w/ foot r		Y					
E1190	Wheelchair amputee w/ leg re		Y					
E1195	Wheelchair amputee heavy dut		Y					
E1200	Wheelchair amputee fixed arm		Y					
E1220	Wheelch special size/constrc		Y					
E1221	Wheelchair spec size w/ leg		Y					
E1223	Wheelchair spec size w/ foot		Y					
E1224	Wheelchair spec size w/ leg		Y					
E1225	Manual semi-reclining back		Y					
E1226	Manual fully reclining back		B					
E1227	Wheelchair spec sz spec ht a		Y					
E1228	Wheelchair spec sz spec ht b		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E0957	W/c medial thigh support		Y					
E0958	Wheelch att-conv 1 arm drive		Y					
E0959	Amputee adapter		B					
E0960	W/c shoulder harness/straps		Y					
E0961	Wheelchair brake extension		B					
E0966	Wheelchair head rest extensi		Y					
E0967	Manual wc hand rim w project		Y					
E0968	Wheelchair commode seat		Y					
E0969	Wheelchair narrowing device		E					
E0970	Wheelchair no. 2 footplates		Y					
E0971	Wheelchair anti-tipping devi		B					
E0973	W/Ch access det adj armrest		B					
E0974	W/Ch access anti-rollback		B					
E0978	W/C acc. saf belt pelv strap		B					
E0980	Wheelchair safety vest		Y					
E0981	Seat upholstery, replacement		Y					
E0982	Back upholstery, replacement		Y					
E0983	Add pwr joystick		Y					
E0984	Add pwr tiller		Y					
E0985	W/c seat lift mechanism		Y					
E0986	Man w/c push-rim pow assist		Y					
E0990	Wheelchair elevating leg res		B					
E0992	Wheelchair solid seat insert		B					
E0994	Wheelchair arm rest		Y					
E0995	Wheelchair calf rest		B					
E1002	Pwr seat tilt		Y					
E1003	Pwr seat recline		Y					
E1004	Pwr seat recline mech		Y					
E1005	Pwr seat recline pwr		Y					
E1006	Pwr seat combo w/o shear		Y					
E1007	Pwr seat combo w/shear		Y					
E1008	Pwr seat combo pwr shear		Y					
E1009	Add mech leg elevation		Y					
E1010	Add pwr leg elevation		Y					
E1011	Ped wc modify width adjustm		Y					
E1014	Reclining back add ped w/c		Y					
E1015	Shock absorber for man w/c		Y					
E1016	Shock absorber for power w/c		Y					
E1017	HD shck absbr for hd man wc		Y					
E1018	HD shck absbr for hd powwc		Y					
E1020	Residual limb support system		Y					
E1028	W/c manual swingaway		Y					
E1029	W/c vent tray fixed		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E1550	Bath conductivity meter		A					
E1560	Replace blood leak detector		A					
E1570	Adjustable chair for sead pt		A					
E1575	Transducer protect/rid bar		A					
E1580	Unipuncture control system		A					
E1590	Hemodialysis machine		A					
E1592	Auto intern peritoneal dialy		A					
E1594	Cycler dialysis machine		A					
E1600	De/in/install chng hemo equip		A					
E1610	Reverse osmosis h2o puri sys		A					
E1615	Deionizer h2o puri system		A					
E1620	Replacement blood pump		A					
E1625	Water softening system		A					
E1630	Reciprocating peritoneal dia		A					
E1632	Wearable artificial kidney		A					
E1634	Peritoneal dialysis clamp		B					
E1635	Compact travel hemodialyzer		A					
E1636	Sorbent cartridges per 10		A					
E1637	Hemostats for dialysis, each		A					
E1639	Dialysis scale		A					
E1699	Dialysis equipment noc		A					
E1700	Jaw motion rehab system		Y					
E1701	Rept cushions for jaw motion		Y					
E1800	Adjust elbow ext/flex device		Y					
E1801	SPS elbow device		Y					
E1802	Adjust forearm prof/sup device		Y					
E1805	Adjust wrist ext/flex device		Y					
E1806	SPS wrist device		Y					
E1810	Adjust knee ext/flex device		Y					
E1811	SPS knee device		Y					
E1812	Knee ext/flex w act res ctrl		Y					
E1815	Adjust ankle ext/flex device		Y					
E1816	SPS ankle device		Y					
E1818	SPS forearm device		Y					
E1820	Soft interface material		Y					
E1821	Replacement interface SPSD		Y					
E1825	Adjust finger ext/flex devc		Y					
E1830	Adjust toe ext/flex device		Y					
E1840	Adj shoulder ext/flex device		Y					
E1841	Static str shldr dev rom adj		Y					
E1902	AAC non-electronic board		Y					
E2000	Gastric suction pump hme mdl		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E1229	Pediatric wheelchair NOS		Y					
E1230	Power operated vehicle		Y					
E1231	Rigid ped w/c tilt-in-space		Y					
E1232	Folding ped w/c tilt-in-space		Y					
E1233	Rig ped w/c tilnspc w/o seat		Y					
E1234	Fld ped w/c tilnspc w/o seat		Y					
E1235	Rigid ped w/c adjustable		Y					
E1236	Folding ped w/c adjustable		Y					
E1237	Rigd ped w/c adjustabl w/o seat		Y					
E1238	Fld ped w/c adjustabl w/o seat		Y					
E1239	Ped power wheelchair NOS		Y					
E1240	Whchr fltr det arm leg rest		Y					
E1250	Wheelchair lghtwt fixed arm	E						
E1260	Wheelchair lghtwt foot rest	E						
E1270	Wheelchair lghtweight leg r	Y						
E1280	Whchr h-duty det arm leg res	Y						
E1285	Wheelchair heavy duty fixed	E						
E1290	Wheelchair hvy duty detach a	E						
E1295	Wheelchair heavy duty fixed	Y						
E1296	Wheelchair special seat heig	Y						
E1297	Wheelchair special seat dept	Y						
E1298	Wheelchair spec seat depth/w	Y						
E1300	Whirpool portable	E						
E1310	Whirpool non-portable	Y						
E1340	Repair for DME, per 15 min	CH	D					
E1353	Oxygen supplies regulator		Y					
E1354	Wheelied cart, port cyl/conc		Y					
E1355	Oxygen supplies stand/rack		Y					
E1356	Batt pack/cart, port conc		Y					
E1357	Battery charger, port conc		Y					
E1358	DC power adapter, port conc		Y					
E1372	Oxy suppl heater for nebuliz		Y					
E1390	Oxygen concentrator		Y					
E1391	Oxygen concentrator, dual		Y					
E1392	Portable oxygen concentrator		Y					
E1399	Durable medical equipment mi		Y					
E1405	O2/water vapor enrich wheat		Y					
E1406	O2/water vapor enrich w/o he		Y					
E1500	Centrifuge		A					
E1510	Kidney dialysate delivery sys		A					
E1520	Heparin infusion pump		A					
E1530	Replacement air bubble detec		A					
E1540	Replacement pressure alarm		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E2313	PWC harness, expand control		Y					
E2321	Hand interface joystick		Y					
E2322	Multi mech switches		Y					
E2323	Special joystick handle		Y					
E2324	Chin cup interface		Y					
E2325	Sip and puff interface		Y					
E2326	Breath tube kit		Y					
E2327	Head control interface mech		Y					
E2328	Head/extremity control inter		Y					
E2329	Head control nonproportional		Y					
E2330	Head control proximity switc		Y					
E2331	Attendant control		Y					
E2340	W/c width 20-23 in seat frame		Y					
E2341	W/c width 24-27 in seat frame		Y					
E2342	W/c depth 20-21 in seat frame		Y					
E2343	W/c depth 22-25 in seat frame		Y					
E2351	Electronic SGD interface		Y					
E2360	22H nonsealed leadacid		Y					
E2361	22H sealed leadacid battery		Y					
E2362	Gr24 nonsealed leadacid		Y					
E2363	Gr24 sealed leadacid battery		Y					
E2364	U1 nonsealed leadacid battery		Y					
E2365	U1 sealed leadacid battery		Y					
E2366	Battery charger, single mode		Y					
E2367	Battery charger, dual mode		Y					
E2368	Power, w/ motor replacement		Y					
E2369	Pwr w/ gear box replacement		Y					
E2370	Pwr w/ motor/gear box combo		Y					
E2372	Gr27 sealed leadacid battery		Y					
E2373	Hand/chin ctrl spec joystick		Y					
E2374	Hand/chin ctrl std joystick		Y					
E2375	Non-expandable controller		Y					
E2376	Expandable controller, repl		Y					
E2377	Expandable controller, inifl		Y					
E2381	Pneum drive wheel tire		Y					
E2382	Tube, pneum wheel drive tire		Y					
E2383	Insert, pneum wheel drive		Y					
E2384	Pneumatic caster tire		Y					
E2385	Tube, pneumatic caster tire		Y					
E2386	Foam filled drive wheel tire		Y					
E2387	Foam filled caster tire		Y					
E2388	Foam drive wheel tire		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E2100	Bld glucose monitor, w voice		Y					
E2101	Bld glucose monitor, w lance		Y					
E2120	Pulse, gen sys tx endolprop fl		Y					
E2201	Men w/ch acc seat w/s=20' <24"		Y					
E2202	Seat width, 24-27 in		Y					
E2203	Frame depth, less than 22 in		Y					
E2204	Frame depth, 22 to 25 in		Y					
E2205	Manual w/ accessory, handrim		Y					
E2206	Complete wheel lock assembly		Y					
E2207	Crutch and cane holder		Y					
E2208	Cylinder tank carrier		Y					
E2209	Arm trough each		Y					
E2210	Wheelchair bearings		Y					
E2211	Pneumatic propulsion tire		Y					
E2212	Pneumatic prop tire tube		Y					
E2213	Pneumatic prop tire insert		Y					
E2214	Pneumatic caster tire each		Y					
E2215	Pneumatic caster tire tube		Y					
E2216	Foam filled propulsion tire		Y					
E2217	Foam filled caster tire each		Y					
E2218	Foam propulsion tire each		Y					
E2219	Foam caster tire any size ea		Y					
E2220	Solid propulsion tire each		Y					
E2221	Solid caster tire each		Y					
E2222	Solid caster integrated whl		Y					
E2223	Valve replacement only each	CH	D					
E2224	Propulsion wtl excludes tire		Y					
E2225	Caster wheel excludes tire		Y					
E2226	Caster fork replacement only		Y					
E2227	Gear reduction drive wheel		Y					
E2228	M/wc acc, wheelchair brake		Y					
E2230	Manual standing system		E					
E2231	Solid seat support base		Y					
E2291	Planar back for ped size wc		Y					
E2292	Planar seat for ped size wc		Y					
E2293	Contour back for ped size wc		Y					
E2294	Contour seat for ped size wc		Y					
E2295	Ped dynamic seating frame		Y					
E2300	Pwr seat elevation sys		Y					
E2301	Pwr standing		Y					
E2310	Electro connect b/w control		Y					
E2311	Electro connect btw 2 sys		Y					
E2312	Mini-prop remote joystick		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E2621	WC planar back cush wds>=22in		Y					
E8000	Posterior gait trainer		E					
E8001	Upright gait trainer		E					
E8002	Anterior gait trainer		E					
G0008	Admin influenza virus vac		S	0350	0.3809	\$25.67		
G0009	Admin pneumococcal vaccine		S	0350	0.3809	\$25.67		
G0010	Admin hepatitis b vaccine		B					
G0027	Semen analysis		A					
G0101	CA screen;pelvic/breast exam		V	0604	0.8593	\$57.92		\$11.59
G0102	Prostate ca screening; drc		N					
G0103	PSA screening		A					
G0104	CA screen;flexi sigmoidoscope		S	0159	3.7227	\$250.93		\$62.73
G0105	Colorectal scm; hi risk ind		T	0158	8.0912	\$545.40		\$136.35
G0106	Colon CA screen;barium		S	0157	1.2511	\$84.33		\$16.87
G0108	enema		A					
G0108	Diab manage trn per indiv		A					
G0109	Diab manage trn ind/group		A					
G0117	Glaucoma scm high risk direc		S	0698	0.9553	\$64.39		\$12.88
G0118	Glaucoma scm high risk direc		S	0230	0.5945	\$40.07		\$6.02
G0120	Colon ca scm; barium enema		S	0157	1.2511	\$84.33		\$16.87
G0121	Colon ca scm not hi risk ind		T	0158	8.0912	\$545.40		\$136.35
G0122	Colon ca scm; barium enema		E					
G0123	Screen cerv/vag thin layer		A					
G0124	Screen c/v thin layer by MD		B					
G0127	Trim nail(s)	CH	T	0012	0.4436	\$29.90		\$5.98
G0128	CORF skilled nursing service		B					
G0129	Partial hosp prog service		P					
G0130	Single energy x-ray study		X	0260	0.6661	\$44.90		\$6.98
G0141	Scr c/v cyto,autosys and imd		B					
G0143	Scr c/v cyto,thinlayer,rescr		A					
G0144	Scr c/v cyto,thinlayer,rescr		A					
G0145	Scr c/v cyto,thinlayer,rescr		A					
G0147	Scr c/v cyto,automated sys		A					
G0148	Scr c/v cyto, autosys, rescr		A					
G0151	HHCP-serv of pt,ea 15 min		B					
G0152	HHCP-serv of ot,ea 15 min		B					
G0153	HHCP-svs of s/l path,ea 15min		B					
G0154	HHCP-svs of rm,ea 15 min		B					
G0155	HHCP-svs of csw,ea 15 min		B					
G0156	HHCP-svs of aid,ea 15 min		B					
G0166	Extrl counterpulse, per tx		T	0678	1.5463	\$104.23		\$20.85
G0168	Wound closure by adhesive		B					
G0173	Linear acc stereo radcur room		S	0067	52.9891	\$3,571.78		\$714.36

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
E2389	Foam caster tire		Y					
E2390	Solid drive wheel tire		Y					
E2391	Solid caster tire		Y					
E2392	Solid caster tire, integrate		Y					
E2393	Valve, pneumatic tire tube	CH	D					
E2394	Drive wheel excludes tire		Y					
E2395	Caster wheel excludes tire		Y					
E2396	Caster fork		Y					
E2397	Pwc acc, lift-based battery		Y					
E2399	Noc interface	CH	D					
E2402	Neg press wound therapy pump		Y					
E2500	SGD digitized pre-rec <=8min		Y					
E2502	SGD prerec msg >8min <=20min		Y					
E2504	SGD prerec msg>20min <=40min		Y					
E2506	SGD prerec msg > 40 min		Y					
E2508	SGD spelling phys contact		Y					
E2510	SGD w multi methods msg/accs		Y					
E2511	SGD shvre prog for PC/PDA		Y					
E2512	SGD accessory, mounting sys		Y					
E2599	SGD accessory noc		Y					
E2601	Gen w/c cushion width < 22 in		Y					
E2602	Gen w/c cushion width >=22 in		Y					
E2603	Skin protect wc cus wd <22in		Y					
E2604	Skin protect wc cus wds>=22in		Y					
E2605	Position wc cush width <22 in		Y					
E2606	Position wc cush width>=22 in		Y					
E2607	Skin prot/pos wc cus wd <22in		Y					
E2608	Skin prot/pos wc cus wds>=22in		Y					
E2609	Custom fabricate w/c cushion		Y					
E2610	Powered w/c cushion		B					
E2611	Gen use back cush width <22in		Y					
E2612	Gen use back cush wds>=22in		Y					
E2613	Position back cush wd <22in		Y					
E2614	Position back cush wds>=22in		Y					
E2615	Pos back post/lat width <22in		Y					
E2616	Pos back post/lat width>=22in		Y					
E2617	Custom fab w/c back cushion		Y					
E2619	Replace cover w/c seat cush		Y					
E2620	WC planar back cush wd <22in		Y					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G0293	Non-cov surg proc,clin trial	X	X	0340	0.6693	\$45.11		\$9.03
G0294	Non-cov proc, clinical trial	X	X	0340	0.6693	\$45.11		\$9.03
G0295	Electromagnetic therapy onc	E						
G0302	Pre-op service LVRS complete	S	0209	0209	11.4315	\$770.55	\$268.73	\$154.11
G0303	Pre-op service LVRS 10-15dos	S	0209	0209	11.4315	\$770.55	\$268.73	\$154.11
G0304	Pre-op service LVRS 1-9 dos	S	0213	0213	2.4043	\$162.06	\$53.58	\$32.42
G0305	Post op service LVRS min 6	S	0213	0213	2.4043	\$162.06	\$53.58	\$32.42
G0306	CBC/diffwbc w/o platelet	A						
G0307	CBC without platelet	A						
G0328	Fecal blood sern immunoassay	A						
G0329	Electromagnetic tx for ulcers	A						
G0333	Dispense fee initial 30 day	M						
G0337	Hospice evaluation prelecti	B						
G0339	Robot lin-radsurg com, first	S	0067	0067	52.9891	\$3,571.78		\$714.36
G0340	Robot lin-radsurg track 2-5	S	0066	0066	36.9119	\$2,488.08		\$497.62
G0341	Percutaneous islet celltrans	C						
G0342	Laparoscopy islet cell trans	C						
G0343	Laparotomy islet cell trans	C						
G0364	Bone marrow aspirate biopsy	X	0340	0340	0.6693	\$45.11		\$9.03
G0365	Vessel mapping hemo access	S	0267	0267	2.3005	\$155.07	\$60.50	\$31.02
G0372	MD service required for PMD	M						
G0378	Hospital observation per hr	N						
G0379	Direct refer hospital observ	Q3	0604	0604	0.8593	\$57.92		\$11.59
G0380	Lev 1 hosp type B ED visit	V	0626	0626	0.6796	\$45.81		\$9.17
G0381	Lev 2 hosp type B ED visit	V	0627	0627	0.9229	\$62.21		\$12.45
G0382	Lev 3 hosp type B ED visit	V	0628	0628	1.4572	\$98.22		\$19.65
G0383	Lev 4 hosp type B ED visit	V	0629	0629	2.1041	\$141.83		\$28.37
G0384	Lev 5 hosp type B ED visit	Q3	0630	0630	3.4466	\$232.32	\$52.26	\$46.47
G0389	Ultrasound exam AAA screen	S	0266	0266	1.4434	\$97.29	\$37.61	\$19.46
G0390	Trauma Resposns w/hosp criti	S	0618	0618	12.3717	\$833.93		\$166.79
G0392	AV fistula or graft arterial	CH	D					
G0393	AV fistula or graft venous	CH	D					
G0396	Alcohol/subs intery 15-30mn	S	0432	0432	0.6013	\$40.53		\$8.11
G0397	Alcohol/subs intery >30 min	S	0432	0432	0.6013	\$40.53		\$8.11
G0398	Home sleep test/type 2 Porta	S	0213	0213	2.4043	\$162.06	\$53.58	\$32.42
G0399	Home sleep test/type 3 Porta	S	0213	0213	2.4043	\$162.06	\$53.58	\$32.42
G0400	Home sleep test/type 4 Porta	S	0213	0213	2.4043	\$162.06	\$53.58	\$32.42
G0402	Initial preventive exam	CH	V	0607	1.683	\$113.44		\$22.69
G0403	EKG for initial preventive exam	M						
G0404	EKG tracing for initial prev	S	0099	0099	0.394	\$26.56		\$5.32
G0405	EKG interpret & report prev	B						
G0406	Telhealth inpt consult 15min	CH	C					
G0407	Telhealth inpt consult 25min	CH	C					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G0175	OPPS Service,sched team conf	CH	V	0607	1.683	\$113.44		\$22.69
G0176	OPPS/PHP:activity therapy	P						
G0177	OPPS/PHP: train & educ serv	N						
G0179	MD recertification HHA pt	M						
G0180	MD certification HHA patient	M						
G0181	Home health care supervision	M						
G0182	Hospice care supervision	M						
G0186	Dsily eye lesn,ldr vssl tech	T	0235	0235	5.8498	\$394.31		\$78.97
G0202	Screeningmammothographydigital	A						
G0204	Diagnosicmammothographydigital	A						
G0206	Diagnosicmammothographydigital	A						
G0219	PET img wholebod melano	E						
G0235	PET not otherwise specified	E						
G0237	Therapeutic proced strg endure	S	0077	0077	0.4058	\$27.35	\$7.74	\$5.47
G0238	Oh resp proc, indiv	S	0077	0077	0.4058	\$27.35	\$7.74	\$5.47
G0239	Oh resp proc, group	S	0077	0077	0.4058	\$27.35	\$7.74	\$5.47
G0245	Initial foot exam pt tops	V	0604	0604	\$57.92			\$11.59
G0246	Followup eval of foot pt top	V	0605	0605	\$69.68			\$13.94
G0247	Routine footcare pt w tops	T	0013	0013	0.8789	\$59.24		\$11.85
G0248	Demonstrate use home fir mon	V	0607	0607	1.683	\$113.44		\$22.69
G0249	Provide INR test mater/equip	V	0607	0607	1.683	\$113.44		\$22.69
G0250	MD INR test revie inter night	M						
G0251	Linear acc based stereo radio	S	0065	0065	14.2808	\$982.61		\$192.53
G0252	PET imaging initial dx	E						
G0255	Current percep threshold tst	E						
G0257	Unsched dialysis ESRD pt hos	S	0170	0170	6.8212	\$459.79		\$91.96
G0259	Injct for sacroiliac joint	N						
G0260	Inj for sacroiliac it anesth	T	0207	0207	7.2002	\$465.34		\$97.07
G0268	Removal of impacted wax ind	N						
G0269	Occlusive device in vein art	N						
G0270	MNT subs tx for change dx	A						
G0271	Group MNT 2 or more 30 mins	A						
G0275	Renal angio, cardiac cath	N						
G0278	lliac art angio,cardiac cath	N						
G0281	Elec stfm unattend for press	A						
G0282	Elect stfm wound care not pd	E						
G0283	Elec stfm other than wound	A						
G0288	Recon, CTA for surg plan	N						
G0289	Arthro, loose body + chondro	N						
G0290	Drug-eluting stents, single	T	0656	0656	110.787	\$7,467.71		\$1,493.55
G0291	Drug-eluting stents,each add	T	0656	0656	110.787	\$7,467.71		\$1,493.55

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G0408	Telhealth inpt consult 35mins	CH	C					
G0409	CORF related serv 15 mins ea	M	M					
G0410	Grp psych partial hosp 45-50	P	P					
G0411	Inter active grp psych part	P	P					
G0412	Open tx iliac spine unibil	C	C	0050	31.7717	\$2,141.60		\$428.32
G0413	Pelvic ring fracture unibil	C	C					
G0414	Pelvic ring fx treat int fix	C	C					
G0415	Open tx post pelvic fracture	C	C					
G0416	Sat biopsy prostate 1-20 spc	S	S	1505		\$350.00		\$70.00
G0417	Sat biopsy prostate 21-40	S	S	1507		\$550.00		\$110.00
G0418	Sat biopsy prostate 41-60	S	S	1511		\$950.00		\$190.00
G0419	Sat biopsy prostate: >60	S	S	1513		\$1,150.00		\$230.00
G0420	Ed svc CKD ind per session	NI	A					
G0421	Ed svc CKD grp per session	NI	A					
G0422	Intens cardiac rehab w/exerc	NI	S	0095	0.5691	\$38.36	\$13.86	\$7.68
G0423	Intens cardiac rehab no exer	NI	S	0095	0.5691	\$38.36	\$13.86	\$7.68
G0424	Pulmonary rehab w exer	NI	S	0102	0.7486	\$50.46		\$10.10
G0425	Inpt telehealth consult 30m	NI	C					
G0426	Inpt telehealth consult 50m	NI	C					
G0427	Inpt telehealth con 70->rn	NI	C					
G0430	Drug screen multi class	NI	A					
G0431	Drug screen single class	NI	A					
G3001	Admin + supply, tositumomab	NI	S	0442	25.7674	\$1,736.88		\$347.38
G8006	AMI pt recd aspirin at arriv	M	M					
G8007	AMI pt did not recv aspiri	M	M					
G8008	AMI pt ineligible for aspiri	M	M					
G8009	AMI pt recd Bbloc at arr	M	M					
G8010	AMI pt did not rec bblock	M	M					
G8011	AMI pt inelig Bbloc at arriv	M	M					
G8012	Pneum pt recv antibiotic 4 h	M	M					
G8013	Pneum pt w/o antibiotic 4 hr	M	M					
G8014	Pneum pt not elig antibiotic	M	M					
G8015	Diabetic pt w/HbA1c<9%	M	M					
G8016	Diabetic pt w/HbA1c=9%	M	M					
G8017	Dm pt inelig for HbA1c measu	M	M					
G8018	Care not provided for HbA1c	M	M					
G8019	Diabetic pt w/LDL<= 100mg/dl	M	M					
G8020	Diab pt w/LDL< 100mg/dl	M	M					
G8021	Diab pt inelig for LDL meas	M	M					
G8022	Care not provided for LDL	M	M					
G8023	DM pt w BP<=140/80	M	M					
G8024	Diabetic pt wBP<140/80	M	M					
G8025	Diabetic pt inelig for BP me	M	M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8026	Diabet pt w no care re BP me	M	M					
G8027	HF p w/LVSD on ACE-I/ARB	M	M					
G8028	HF pt w/LVSD not on ACE-	M	M					
G8028	I/AR	M	M					
G8029	HF pt not elig for ACE-I/ARB	M	M					
G8030	HF pt w/LVSD on Bblocker	M	M					
G8031	HF pt w/LVSD not on Bblocker	M	M					
G8032	HF pt not elig for Bblocker	M	M					
G8033	PMI-CAD pt on Bblocker	M	M					
G8034	PMI-CAD pt not on Bblocker	M	M					
G8035	PMI-CAD pt inelig Bblocker	M	M					
G8036	AMI-CAD pt doc on antiplate	M	M					
G8037	AMI-CAD pt not docu on antipl	M	M					
G8038	AMI-CAD inelig antiplate mea	M	M					
G8039	CAD pt w/LDL>100mg/dl	M	M					
G8040	CAD pt w/LDL<=100mg/dl	M	M					
G8041	CAD pt not eligible for LDL	M	M					
G8051	Osteoporosis assess	M	M					
G8052	Osteopor pt not assess	M	M					
G8053	Pt inelig for osteopor meas	M	M					
G8054	Falls assess not docum 12 mo	M	M					
G8055	Falls assess w/ 12 mon	M	M					
G8056	Not elig for falls assessment	M	M					
G8057	Hearing assess receive	M	M					
G8058	Pt w/o hearing assess	M	M					
G8059	Pt inelig for hearing assess	M	M					
G8060	Urinary incont pt assess	M	M					
G8061	Pt not assess for urinary in	M	M					
G8062	ESRD pt w/ urinary inco	M	M					
G8075	ESRD pt w/ dialy of URB>=65%	M	M					
G8076	ESRD pt w dialy of URR<65%	M	M					
G8077	ESRD pt not elig for URR/KV	M	M					
G8078	ESRD pt w/Hct<or=33	M	M					
G8079	ESRD pt w/Hct<33	M	M					
G8080	ESRD pt inelig for HCT/Hgb	M	M					
G8081	ESRD pt w/ auto AV fistula	M	M					
G8082	ESRD pt w other fistula	M	M					
G8085	ESRD PT inelig auto AV FISTU	M	M					
G8093	COPD pt rec smoking cessat	M	M					
G8094	COPD pt w/o smoke cessat int	M	M					
G8099	Osteopo pt given Ca+VitD sup	M	M					
G8100	Osteop pt inelig for Ca+VitD	M	M					
G8103	New dx osteo pt w/antiresorp	M	M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8209	Clinician did not doc		M					
G8214	Clin not doc order VTE		M					
G8217	Pt not received DVT proph		M					
G8219	Received DVT proph day 2		M					
G8220	Pt not rec DVT proph day 2		M					
G8221	Pt inelig for DVT proph		M					
G8223	Pt not doc for presc antipla		M					
G8226	Pt no presc anticoag at D/C		M					
G8234	Pt not doc for admin t-PA		M					
G8234	Pt not doc dysphagia screen		M					
G8238	Pt not doc to rec rehab serv		M					
G8240	Infer carotid stenosis30-99%		M					
G8243	Pt not doc MRI/CT w/o lesion		M					
G8246	Pt inelig hx w new/chg mole		M					
G8248	Pt w/one alarm symp not doc		M					
G8251	Pt not doc w/Barretts, endo		M					
G8254	Pt w/no doc order for barium		M					
G8257	Pt not doc rev meds D/C		M					
G8260	Pt not doc to have dec maker		M					
G8263	Pt not doc assess urinary in		M					
G8266	Pt not doc charc urtn incon		M					
G8268	Pt not doc rec care urtn inc		M					
G8271	Pt no doc screen fall		M					
G8274	Clin not doc press/abs alarm		M					
G8276	Pt not doc mole change		M					
G8279	Pt not doc rec PE		M					
G8282	Pt not doc to rec couns		M					
G8285	Pt did not rec pres osteo		M					
G8289	Pt not doc rec Ca/Vit D		M					
G8293	COPD pt w/o spir results		M					
G8296	COPD pt not doc bronch ther		M					
G8299	Pt not doc optic nerv eval		M					
G8302	Pt doc w/ target IOP		M					
G8303	Pt not doc w/ IOP		M					
G8304	Clin doc pt inelig IOP		M					
G8305	Clin not prov care POAG		M					
G8306	POAG w/ IOP rec care plan		M					
G8307	POAG w/ IOP no care plan		M					
G8308	POAG w/ IOP not doc plan		M					
G8310	Pt not doc rec antiobx		M					
G8314	Pt not doc to rec mac exam		M					
G8318	Pt not doc to have visual func		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8104	Osteo pt inelig for antireso		M					
G8106	Bone dens meas test perf		M					
G8107	Bone dens meas test inelig		M					
G8108	Pt receiv influenza vacc		M					
G8109	Pt w/o influenza vacc		M					
G8110	Pt inelig for influenza vacc		M					
G8111	Pt receiv mammogram		M					
G8112	Pt not doc mammogram		M					
G8113	Pt ineligble mammography		M					
G8114	Care not provided for mamogr		M					
G8115	Pt receiv pneumo vacc		M					
G8116	Pt did not rec pneumo vacc		M					
G8117	Pt was inelig for pneumo vac		M					
G8126	Pt treat w/antidepress12wks		M					
G8127	Pt not treat w/antidepress12w		M					
G8128	Pt inelig for antidepress med		M					
G8129	Pt treat w/antidepress for 6m		M					
G8130	Pt not treat w/antidepress 6m		M					
G8131	Pt inelig for antidepress med		M					
G8152	Pt w/AB 1 hr prior to incisi		M					
G8154	Pt inelig for AB therapy		M					
G8155	Pt recd thromboemb prophylax		M					
G8156	Pt did not rec thromboemb		M					
G8157	Pt inelig for thrombolism		M					
G8159	Pt w/CABG w/o IMA		M					
G8162	Iso CABG pt w/o preop Bblock		M					
G8164	Iso CABG pt w/prolong intub		M					
G8165	Iso CABG pt w/o prolong intub		M					
G8166	Iso CABG req surg expo		M					
G8167	Iso CABG w/o surg expo		M					
G8170	CEA/ext bypass pt on aspirin		M					
G8171	Pt w/carot endart/ext bypas		M					
G8172	CEA/ext bypass pt not on asp		M					
G8182	CAD pt care not prov LDL		M					
G8183	HF/atrial fib pt on warfarin		M					
G8184	HF/atrial fib pt inelig warf		M					
G8185	Osteoarth pt w/ assess pain		M					
G8186	Osteoarth pt inelig assess		M					
G8193	Antibio not doc prior surg		M					
G8196	Antibio not docum prior surg		M					
G8200	Cefazolin not docum prophy		M					
G8204	MD not doc order to d/c anti		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8400	Pt w/DXA no document or order		M					
G8401	Pt inelig osteo screen measu		M					
G8402	Smoke preven interven counse		M					
G8403	Smoke preven nocounsel		M					
G8404	Low extremity neur exam docum		M					
G8405	Low extremity neur not perfor		M					
G8406	Pt inelig lower extrem neuro		M					
G8407	ABI documented		M					
G8408	ABI not documented		M					
G8409	Pt inelig for ABI measure		M					
G8410	Eval on foot documented		M					
G8415	Eval on foot not performed		M					
G8417	Pt inelig footwear evaluation		M					
G8418	Calc BMI abv up param f/u		M					
G8419	Calc BMI b/w low param f/u		M					
G8420	Calc BMI out nrm param not/fu		M					
G8421	Calc BMI norm parameters		M					
G8422	BMI not calculated		M					
G8422	Pt inelig BMI calculation		M					
G8423	Pt screen flu vac & counsel		M					
G8424	Flu vaccine not screen		M					
G8425	Flu vaccine screen not curre		M					
G8426	Pt not approp screen & couns		M					
G8427	Doc medis verified w/pt or rep		M					
G8428	Medis document w/o verifica		M					
G8429	Incomplete doc pt on medis		M					
G8430	Pt inelig med check		M					
G8431	Pos clin depres scrn f/u doc		M					
G8432	Clin depression screen not d		M					
G8433	Pt inelig, scrn clin dep		M					
G8434	Cognitive impairment screen		M					
G8435	Cognitive screen not document		M					
G8436	Pt inelig for cognitive impairm		M					
G8437	Care plan develop & document		M					
G8438	Pt inelig for devlop care pin		M					
G8439	Care plan develop & not docum		M					
G8440	Pain assess f/u pin document		M					
G8441	No document of pain assess		M					
G8442	Pt inelig pain assessment		M					
G8443	Prescrip by E-Prescrib system		M					
G8445	Prescrip not gen at encounter		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8322	Pt not doc pre axial leng		M					
G8326	Pt not doc rec fundus exam		M					
G8330	Pt not doc rec dilated mac		M					
G8334	Doc of macular not giv MD		M					
G8338	Clin not doc pt test osteo		M					
G8341	Pt not doc for DEXA		M					
G8345	Pt not doc have DEXA		M					
G8351	Pt not doc ECG		M					
G8354	Pt not doc aspirin prior ER		M					
G8357	Pt not doc to have ECG		M					
G8360	Pt not doc vital signs recor		M					
G8362	Pt not doc O2 SAT assess		M					
G8365	Pt not doc mental status		M					
G8367	Pt not doc have empiric AB		M					
G8370	Asthma pt w survey not docum		M					
G8371	Chemother not rec sig3 colon		M					
G8372	Chemother rec sig3 colon ca		M					
G8373	Chemo plan document prior che		M					
G8374	Chemo plan not doc prior che		M					
G8375	CLL pt w/o doc flow cytometr		M					
G8376	Brst ca pt inelig tamoxifen		M					
G8377	MD doc colon ca pt inelig ch		M					
G8378	MD doc pt inelig radiation		M					
G8379	Doc radiat bx recom 12mo ov		M					
G8380	Pt w sigIC-3Brst ca not rec		M					
G8381	Pt w sigIC-3Brst ca rec tam		M					
G8382	MM pt w/o doc TV bisphosphon		M					
G8383	No doc radiation rec 12mo ov		M					
G8384	Base cytogen test MDS notper		M					
G8385	Diabet pt no do Hgb A1c 12m		M					
G8386	Diabet pt not doc LDL/protei		M					
G8387	ESRD pt w Hct/Hgb not docum		M					
G8388	ESRD pt w URR/Ktv not doc eli		M					
G8389	MDS pt no doc FE st prio EPO		M					
G8390	Diabetic w/o document BP 12m		M					
G8391	Pt w asthma no doc med or tx LVEF>=40% doc normal or mild		M					
G8395	LVEF not performed		M					
G8396	Dil macular/fundus exam/w doc		M					
G8397	Dil macular/fundus exam/w doc performe		M					
G8398	Pt w/DXA document or order		M					
G8399			M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8489	CAD measures grp		M					
G8490	RA measures grp		M					
G8491	HIV/AIDS measures grp		M					
G8492	Periop Care measures grp		M					
G8493	Back pain measures grp		M					
G8494	DM meas qual act perform		M					
G8495	CKD meas qual act perform		M					
G8496	Prev Care MG qual act perform		M					
G8497	CABG meas qual act perform		M					
G8498	CAD meas qual act perform		M					
G8499	RA meas qual act perform		M					
G8500	HIV meas qual act perform		M					
G8501	Perio meas qual act perform		M					
G8502	Back Pain MG qual act perform		M					
G8503	Doc proph antibx w/in 1 hr	CH	D					
G8504	Doc ord pro antibx w/in 1 hr	CH	D					
G8505	No doc proph antibx w/in 1hr	CH	D					
G8506	Pt rec ACE/ARB		M					
G8507	Pt inelig pt verif meds		M					
G8508	Pt inelig; pain asses no f/u		M					
G8509	Pain assess no f/u/pln doc		M					
G8510	Pt inelig; neg scrm depre		M					
G8511	Clin depre scrm no f/u doc		M					
G8512	Pain sav quant present	CH	D					
G8513	ABI meas & doc	CH	D					
G8514	PT inelig; ABI measure	CH	D					
G8515	No ABI measurement	CH	D					
G8516	Scrn fal risk >2 fal or w/inj	CH	D					
G8517	Scrn fal risk; <2 falls	CH	D					
G8518	Clin stig b/f lun/eso ca surg		M					
G8519	Pt in; clin ca stig b/f surg		M					
G8520	Clin stig b/f surg not doc		M					
G8521	Antpt recd 48 hrs & disch	CH	D					
G8522	Pt inelig; antipit therapy	CH	D					
G8523	Antpt not recd reas no spec	CH	D					
G8524	Patch closure conv CEA		M					
G8525	No patch closure CEA		M					
G8526	No patch closure conv CEA		M					
G8527	Doc ord anlimic proph	CH	D					
G8528	Pt inelig; proph antibot	CH	D					
G8529	No doc ord anlimic proph	CH	D					
G8530	Auto AV fistula recd		M					
G8531	Pt inelig; auto AV fistula		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8446	Some prescrib print or call		M					
G8447	Pt vis doc use CCHIT cer EHR		M					
G8448	Pt vis doc w/non-CCHIT EHR		M					
G8449	Pt not doc w/EHR due to syst		M					
G8450	Beta-bloc rx pt w/abn lvef		M					
G8451	Pt w/abn lvef inelig b-bloc		M					
G8452	Pt w/abn lvef b-bloc no rx		M					
G8453	Tob use cess int counsel		M					
G8454	Tob use cess int no counsel		M					
G8455	Current tobacco smoker		M					
G8456	Current smkless tobacco user		M					
G8457	Cur tobacco non-user		M					
G8458	Pt inelig geno no antivir tx		M					
G8459	Doc pt rec antivir treat		M					
G8460	Pt inelig RNA no antivir tx		M					
G8461	Pt rec antivir treat hep c		M					
G8462	Pt inelig couns no antivir tx		M					
G8463	Pt rec antiviral treat doc		M					
G8464	Pt inelig; lb to no oter risk		M					
G8465	High risk recurrence pro ca		M					
G8466	Pt inelig suic; MDD remis		M					
G8467	New dx ini/rec episode MDD		M					
G8468	ACE/ARB rx pt w/abn lvef		M					
G8469	Pt w/abn lvef inelig ACE/ARB		M					
G8470	Pt w/ normal lvef		M					
G8471	LVEF not performed/doc		M					
G8472	ACE/ARB no rx pt w/abn lvef		M					
G8473	ACE/ARB thxpy rx'd		M					
G8474	ACE/ARB not rx'd; doc reas		M					
G8475	ACE/ARB thxpy not rx'd		M					
G8476	BP sys <130 and dias <80		M					
G8477	BP sys=130 and/or dias >=80		M					
G8478	BP not performed/doc		M					
G8479	MD rx'd ACE/ARB thxpy		M					
G8480	Pt inelig ACE/ARB thxpy		M					
G8481	MD not rx'd ACE/ARB thxpy		M					
G8482	Flu immunize order/admin		M					
G8483	Flu imm no ord/admin doc rea		M					
G8484	Flu immunize no order/admin		M					
G8485	Report, Diabetes measures		M					
G8486	Report, Prev Care Measures		M					
G8487	Report CKD Measures		M					
G8488	Report ESRD Measures		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8577	Recp req bid grft oth	NI	M					
G8578	No recp req bid grft oth	NI	M					
G8579	Antipit med disch	NI	M					
G8580	Antipit med contraind	NI	M					
G8581	no antipit med disch	NI	M					
G8582	Bblock disch	NI	M					
G8583	Bblock contraind	NI	M					
G8584	No bblock disch	NI	M					
G8585	Antilipid treat disch	NI	M					
G8586	Antilipid disch contra	NI	M					
G8587	No antilipid treat disch	NI	M					
G8588	Sys BP <140	NI	M					
G8589	Sys BP >= 140	NI	M					
G8590	Dia BP < 90	NI	M					
G8591	Dia BP >= 90	NI	M					
G8592	No BP measure	NI	M					
G8593	Lipid pn results	NI	M					
G8594	No lipid prof perf	NI	M					
G8595	Ldl < 100	NI	M					
G8596	No LDL perf	NI	M					
G8597	Ldl >= 100	NI	M					
G8598	Asp therp used	NI	M					
G8599	No asp therp used	NI	M					
G8600	IPA init w/in 3 hrs	NI	M					
G8601	No elig IPA init w/in 3 hrs	NI	M					
G8602	No IPA init w/in 3 hrs	NI	M					
G8603	Spok lang comp score	NI	M					
G8604	No high score spok lang	NI	M					
G8605	No spok lang comp score	NI	M					
G8606	Attention score	NI	M					
G8607	No high score attention	NI	M					
G8608	No attention score	NI	M					
G8609	Memory score	NI	M					
G8610	No high score memory	NI	M					
G8611	No memory score	NI	M					
G8612	Moto speech score	NI	M					
G8613	No high score moto speech	NI	M					
G8614	No moto speech score	NI	M					
G8615	Reading score	NI	M					
G8616	No high score reading	NI	M					
G8617	No reading score	NI	M					
G8618	Spok lang exp score	NI	M					
G8619	No high score spok lang exp	NI	M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8532	No auto AV fistula; no reas	M	D					
G8533	Panic in clin data base reg	CH	D					
G8534	Doc elder mal scrn f/u plan	M	M					
G8535	Pt inelig no eid mal scrn	M	M					
G8536	No doc elder mal scrn	M	M					
G8537	Pt inelig eidmal scrn no f/u	M	M					
G8538	Eld mal scrn no f/u pln	M	M					
G8539	Cur funct assess & care pln	M	M					
G8540	Pt inelig funct assess	M	M					
G8541	No doc cur funct assess	M	M					
G8542	Pt inelig func asses no pln	M	M					
G8543	Cur funct asses; no care pln	M	M					
G8544	CABG measures grp	M	M					
G8545	HepC measures grp	NI	M					
G8546	CAP measures grp	NI	M					
G8547	IVD measures grp	NI	M					
G8548	HF measures grp	NI	M					
G8549	HepC MG qual act perform	NI	M					
G8550	CAP MG qual act perform	NI	M					
G8551	HF MG qual act perform	NI	M					
G8552	IVD MG qual act perform	NI	M					
G8553	1 Rx via qualified eRx sys	NI	M					
G8556	Ref to doc otolog eval	NI	M					
G8557	Pt inelig ref otolog eval	NI	M					
G8558	No ref to doc otolog eval	NI	M					
G8559	Pt ref doc oto eval	NI	M					
G8560	Pt hx act drain prev 90 days	NI	M					
G8561	Pt inelig for ref oto eval	NI	M					
G8562	Pt no hx act drain 90 d	NI	M					
G8563	Pt no ref oto reas no spec	NI	M					
G8564	Pt ref oto eval	NI	M					
G8565	Ver doc hear loss	NI	M					
G8566	Pt inelig ref oto eval	NI	M					
G8567	Pt no doc hear loss	NI	M					
G8568	Pt no ref otob no spec	NI	M					
G8569	Prol intubation req	NI	M					
G8570	No prol intub req	NI	M					
G8571	Ster wd fix 30 d postop	NI	M					
G8572	No ster wd fix	NI	M					
G8573	Slk/CVA CABG	NI	M					
G8574	No strk/CVA CABG	NI	M					
G8575	Postop ren insuf	NI	M					
G8576	No postop ren insuf	NI	M					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G9057	onc pract mgmt differs trial		E					
G9058	onc prac mgmt disagree w/gui		E					
G9059	onc prac mgmt pt opt alterna		E					
G9060	onc prac mgmt dif pt comorb		E					
G9061	onc prac cond noaid by guide		E					
G9062	onc prac guide differs nos		E					
G9063	onc dx nscic stgl no progres		M					
G9064	onc dx nscic stg2 no progres		M					
G9065	onc dx nscic stg3A no progres		M					
G9066	onc dx nscic stg3B-4 melasta		M					
G9067	onc dx nscic dx unknown nos		M					
G9068	onc dx sclc/nscic limited		M					
G9069	onc dx sclc/nscic ext at dx		M					
G9070	onc dx sclc/nscic ext unkwn		M					
G9071	onc dx brst sig1-2B HR,nopro		M					
G9072	onc dx brst sig1-2 noprogres		M					
G9073	onc dx brst sig3-HR, no pro		M					
G9074	onc dx brst sig3-noprogres		M					
G9075	onc dx brst melastat/ recur		M					
G9077	onc dx prostate T1no progres		M					
G9078	onc dx prostate T2no progres		M					
G9079	onc dx prostate T3b-T4noprog		M					
G9080	onc dx prostate w/ris PSA		M					
G9083	onc dx prostate unkwn nos		M					
G9084	onc dx colon t1-3,n1-2,no pr		M					
G9085	onc dx colon T4, N0,w/o prog		M					
G9086	onc dx colon T1-4 no dx prog		M					
G9087	onc dx colon metas evid dx		M					
G9088	onc dx colon metas noevid dx		M					
G9089	onc dx colon extent unknown		M					
G9090	onc dx rectal T1-2 no progr		M					
G9091	onc dx rectal T3 N0 no prog		M					
G9092	onc dx rectal T1-3,N1-2noprog		M					
G9093	onc dx rectal T4,N,M0 no prog		M					
G9094	onc dx rectal M1 w/mets prog		M					
G9095	onc dx rectal extent unkwn		M					
G9096	onc dx esophag T1-T3 noprog		M					
G9097	onc dx esophageal T4 no prog		M					
G9098	onc dx esophageal mets recur		M					
G9099	onc dx esophageal unknown		M					
G9100	onc dx gastric no recurrence		M					
G9101	onc dx gastric p R1-R2noprog		M					
G9102	onc dx gastric unresectable		M					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G8620	No spok lang exp score	NI	M					
G8621	Writing score	NI	M					
G8622	No high score writing	NI	M					
G8623	No writing score	NI	M					
G8624	Swallowing score	NI	M					
G8625	No high score swallowing	NI	M					
G8626	No swallowing score	NI	M					
G8627	Surg proc w/in 30 days	NI	M					
G8628	No surg proc w/in 30 days	NI	M					
G9001	MCCD, initial rate		B					
G9002	MCCD,maintenance rate		B					
G9003	MCCD, risk adj hi, initial		B					
G9004	MCCD, risk adj lo, initial		B					
G9005	MCCD, risk adj, maintenance		B					
G9006	MCCD, Home monitoring		B					
G9007	MCCD, sch team conf		B					
G9008	Mccd,phys coor-care oversght		B					
G9009	MCCD, risk adj, level 3		B					
G9010	MCCD, risk adj, level 4		B					
G9011	MCCD, risk adj, level 5		B					
G9012	Other Specified Case Mgmt		B					
G9013	ESRD demo bundle level I		E					
G9014	ESRD demo bundle-level II		E					
G9016	Demo-smoking cessation coun		E					
G9017	Amantadine HCL, 100mg oral		A					
G9018	Zanamivir,inhalation pwd 10mr		A					
G9019	Oseltamivir, phosphate 75mg		A					
G9020	Rimantadine HCL, 100mg oral		A					
G9033	Amantadine HCL, oral brand		A					
G9034	Zanamivir, inh pwdtr, brand		A					
G9035	Oseltamivir, phosp, brand		A					
G9036	Rimantadine HCL, brand		A					
G9041	Low vision rehab occupationala		A					
G9042	Low vision rehab orient/mobi		A					
G9043	Low vision lowvision therapl		A					
G9044	Low vision rehabilitate teache		A					
G9050	Oncology work-up evaluation		E					
G9051	Oncology tx decision-mgmt		E					
G9052	onc surveillance for disease		E					
G9053	onc expectant management pt		E					
G9054	onc supervision palliative		E					
G9055	onc visit unspecified NOS		E					
G9056	onc prac mgmt adheres guide		E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J0135	Adalimumab injection	K	1083			\$357.53		\$71.51
J0152	Adenosine injection	K	0379			\$9.50		\$1.90
J0170	Adrenalin epinephrin inject	K	0917			\$76.42		\$15.29
J0180	Aqalisidase beta injection	N						
J0190	Inj biperiden lactate/5 mg	K	9208			\$133.69		\$26.74
J0200	Alatroloxacin mesylate	N						
J0205	Algucerase injection	K	0900			\$41.19		\$8.24
J0207	Amifostine	K	7000			\$350.07		\$70.02
J0210	Methyldopate hcl injection	K	2210			\$27.64		\$5.53
J0215	Aliefacet	K	1633			\$30.02		\$6.01
J0220	Algucosidase alfa injection	K	9234			\$124.69		\$24.94
J0256	Alpha 1 proteinase inhibitor	K	0901			\$3.63		\$0.73
J0270	Alprostadi for injection	B						
J0275	Alprostadi urethral suppos	B						
J0278	Amikacin sulfate injection	N						
J0280	Aminophyllin 250 MG inj	N						
J0282	Amiodarone HCl	N						
J0285	Amphotericin B	N						
J0287	Amphoterin b lipid complex	K	9024			\$9.66		\$1.94
J0288	Ampho b cholesteryl sulfate	K	0735			\$13.74		\$2.75
J0289	Amphotericin b liposome inj	K	0736			\$14.96		\$3.00
J0290	Ampicillin 500 MG inj	N						
J0295	Ampicillin sodium per 1.5 gm	N						
J0300	Amobarbital 125 MG inj	N						
J0330	Succinylcholine chloride inj	N						
J0348	Anidulaturgin injection	CH						
J0350	Injection amstreplase 30 u	K	0760			\$1.21		\$0.25
J0360	Hydratazine hcl injection	N						
J0364	Apomorphine hydrochloride	N						
J0365	Aprotinin, 10,000 kiu	K	1682			\$2.60		\$0.52
J0380	Inj metaraminol bitartrate	N						
J0390	Chloroquine injection	N						
J0395	Arbutamine hcl injection	CH						
J0400	Aripiprazole injection	N						
J0456	Azithromycin	N						
J0460	Atropine sulfate injection	CH						
J0461	Atropine sulfate injection	NI						
J0470	Dimecaprol injection	CH	1273			\$26.81		\$5.37
J0475	Baclofen 10 MG injection	K	9032			\$195.31		\$39.07
J0476	Baclofen intrathecal trial	K	1631			\$71.24		\$14.25
J0480	Basiliximab	K	1683			\$1,624.44		\$324.89
J0500	Dicyclomine injection	N						

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
G9103	Onc dx gastric recurrent	M						
G9104	Onc dx gastric unknown NOS	M						
G9105	Onc dx pancreatc p R0 res no	M						
G9106	Onc dx pancreatc p R1/R2 no	M						
G9107	Onc dx pancreatc unresectab	M						
G9108	Onc dx pancreatc unknown NOS	M						
G9109	Onc dx head/neck T1-T2no pig	M						
G9110	Onc dx head/neck T3-4 noprog	M						
G9111	Onc dx head/neck M1 mets rec	M						
G9112	Onc dx head/neck ext unknown	M						
G9113	Onc dx ovarian stg1A-B no pr	M						
G9114	Onc dx ovarian stg1A-B or 2	M						
G9115	Onc dx ovarian stg3/4 noprog	M						
G9116	Onc dx ovarian recurrence	M						
G9117	Onc dx ovarian unknown NOS	M						
G9123	Onc dx CML chronic phase	M						
G9124	Onc dx CML accelr phase	M						
G9125	Onc dx CML blast phase	M						
G9126	Onc dx CML remission	M						
G9128	Onc dx multi myeloma stage I	M						
G9129	Onc dx multi myeloma stg2 hig	M						
G9130	Onc dx multi myeloma unknown	M						
G9131	Onc dx bst unknown NOS	M						
G9132	Onc dx prostate mets no cast	M						
G9133	Onc dx prostate clinical met	M						
G9134	Onc NHL stg 1-2 no relap no	M						
G9135	Onc dx NHL stg 3-4 not relap	M						
G9136	Onc dx NHL trans to Ig Bcell	M						
G9137	Onc dx NHL relapse/refractor	M						
G9138	Onc dx NHL stg unknown	M						
G9139	Onc dx CML dx status unknown	M						
G9140	Frontier extended stay demo	A						
G9141	Influenza A H1N1 admin w cou	S	0350		0.3809	\$25.67		
G9142	Influenza A H1N1, vaccine	E						
G9143	Warfarin respon genetic test	NI						
J0120	Tetracyclin injection	N						
J0128	Abarelix injection	CH						
J0129	Abatacept injection	K	9230			\$18.98		\$3.80
J0130	Abaximab injection	K	1605			\$459.36		\$91.88
J0132	Acetylcysteine injection	CH	1272			\$2.29		\$0.46
J0133	Acyclovir injection	N						

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J0740	Cidofovir injection		K	9033		\$746.46		\$149.30
J0743	Cilastatin sodium injection		N					
J0744	Ciprofloxacin iv		N					
J0745	Inj cidofovir phosphate 730 MG		N					
J0760	Colchicine injection		N					
J0770	Colistimethate sodium inj		N					
J0780	Prochlorperazine injection		N					
J0795	Corticorelin ovine triflutal		K	1684		\$4.24		\$0.85
J0800	Corticotropin injection		K	1280		\$2,394.93		\$478.99
J0833	Cosyntropin injection NOS	NI	K	0835		\$91.84		\$18.37
J0834	Cosyntropin injection	NI	K	1298		\$91.84		\$18.37
J0835	Inj cosyntropin per 0.25 MG	CH	D					
J0850	Cytomegalovirus imm IV /vial		K	0903		\$862.24		\$172.45
J0878	Daptomycin injection		K	9124		\$0.40		\$0.08
J0881	Darbepoetin alfa, non-esrd		K	1685		\$2.76		\$0.56
J0882	Darbepoetin alfa, esrd use		A					
J0885	Epoetin alfa, non-esrd		K	1686		\$9.40		\$1.88
J0886	Epoetin alfa 1000 units ESFD		A					
J0894	Epoetin alfa 1000 units ESFD		K	9231		\$28.42		\$5.69
J0895	Deferoxamine injection		N					
J0895	Deferoxamine mesylate inj		N					
J0900	Testosterone enanthate inj		N					
J0945	Brompheniramine maleate inj	CH	K	1256		\$0.75		\$0.15
J0970	Estradiol valerate injection		N					
J1000	Depo-estradiol cypionate inj		N					
J1020	Methylprednisolone 20 MG inj		N					
J1030	Methylprednisolone 40 MG inj		N					
J1040	Methylprednisolone 80 MG inj		N					
J1051	Mecroxprogesterone inj		N					
J1055	Mecroxprogester acetate inj		E					
J1056	MAEC contraceptiveinjection		E					
J1060	Testosterone cypionate 1 ML		N					
J1070	Testosterone cypionat 100 MG		N					
J1080	Testosterone cypionat 200 MG		N					
J1084	Inj dexamethasone acetate		N					
J1100	Dexamethasone sodium phos		N					
J1110	Inj dhydroergotamine mesylt		N					
J1120	Acetazolamid sodium injectio		N					
J1160	Digoxin injection		N					
J1162	Digoxin immune fab (ovine)		K	1687		\$474.73		\$94.95
J1165	Phenytoin sodium injection		N					
J1170	Hydromorphone injection		N					
J1180	Dyphylline injection		N					
J1190	Dexrazoxane HCl injection		K	0726		\$340.03		\$68.01

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J0515	Inj benzotropine mesylate		N					
J0520	Bethanechol chloride inject		N					
J0530	Penicillin g benzathine inj	CH	D					
J0540	Penicillin g benzathine inj	CH	D					
J0550	Penicillin g benzathine inj	CH	D					
J0559	PenG benzathine/procaine inj	NI	N					
J0560	Penicillin g benzathine inj		N					
J0570	Penicillin g benzathine inj		N					
J0580	Penicillin g benzathine inj		N					
J0583	Bivalirudin		K	3041		\$2.40		\$0.48
J0585	Injection, onabotulinumtoxinA		K	0902		\$5.40		\$1.08
J0586	AbobotulinumtoxinA	NI	K	1289		\$9.23		\$1.65
J0587	Inj, rimabotulinumtoxinB		K	9018		\$10.38		\$2.08
J0592	Buprenorphine hydrochloride		N					
J0594	Busulfan injection		K	1178		\$14.18		\$2.84
J0595	Butorphanol tartrate 1 mg		N					
J0598	C1 esterase inhibitor inj	NI	G	9251		\$41.34		\$8.11
J0600	Ezetate calcium disodium inj	CH	K	1274		\$78.86		\$15.78
J0610	Calcium gluconate injection		N					
J0620	Calcium glycer & lact/10 ML		N					
J0630	Calcitonin salmon injection		K	1220		\$48.37		\$9.68
J0636	Inj calcitriol per 0.1 mcg		N					
J0637	Caspofungin acetate		K	9019		\$11.52		\$2.31
J0640	Leucovorin calcium injection		N					
J0641	Levoleucovorin injection		G	1236		\$0.99		\$0.19
J0670	Inj imipivacaine HCL/10 ml		N					
J0690	Cefazolin sodium injection		N					
J0692	Cefepime HCl for injection		N					
J0694	Cefoxitin sodium injection		N					
J0696	Ceftaxone sodium injection		N					
J0697	Sterile ceturoxime injection		N					
J0698	Cetotaxime sodium injection		N					
J0702	Betamethasone acet&sod phosp		N					
J0704	Betamethasone sod phosp/4 MG		N					
J0706	Caffeine citrate injection		N					
J0710	Cephapirin sodium injection		N					
J0713	Inj ceftazidime per 500 mg		N					
J0715	Ceftiozime sodium / 500 MG		N					
J0718	Certolizumab pegol inj	NI	G	9249		\$3.80		\$0.75
J0720	Chloramphenicol sodium injec		N					
J0725	Chorionic gonadotropin/1000u		N					
J0735	Cionidine hydrochloride		K	0935		\$109.75		\$21.95

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J1530	Gamma globulin 8 CC inj	K	K	0922		\$120.40		\$24.08
J1540	Gamma globulin 9 CC inj	K	K	0923		\$150.50		\$30.10
J1550	Gamma globulin 10 CC inj	K	K	0924		\$150.50		\$30.10
J1560	Gamma globulin > 10 CC inj	K	K	0933		\$150.50		\$30.10
J1561	Gamma globulin > 10 CC inj	K	K	0948		\$36.71		\$7.35
J1562	Vivaglobin, inj	CH	K	1275		\$7.05		\$1.41
J1565	IMVig	CH	K					
J1566	Immune globulin, powder	D	K	2731		\$29.83		\$5.97
J1568	Octagam injection	K	K	0943		\$37.03		\$7.41
J1569	Gammaagard liquid injection	K	K	0944		\$37.85		\$7.57
J1570	Ganciclovir sodium injection	N	N					
J1571	Hepagam b im injection	G	G	0946		\$50.04		\$9.82
J1572	Flebogamma injection	K	K	0947		\$36.51		\$7.31
J1573	Hepagam b intravenous, inj	G	G	1138		\$50.04		\$9.82
J1580	Garantycin gentamicin inj	N	N					
J1590	Gatifloxacin injection	N	N					
J1595	Injection glatiramer acetate	K	K	1015		\$81.23		\$16.25
J1600	Gold sodium thiomaleate inj	N	N					
J1610	Glucagon hydrochloride/1 MG	K	K	9042		\$79.20		\$15.84
J1620	Gonadorelin hydrocl/ 100 mcg	K	K	7005		\$176.89		\$35.38
J1626	Granisetron hcl injection	CH	N					
J1630	Haloperidol injection	N	N					
J1631	Haloperidol decanoate inj	N	N					
J1640	Hemlin, 1 mg	K	K	1690		\$7.73		\$1.55
J1642	Inj heparin sodium per 10 u	N	N					
J1644	Inj heparin sodium per 1000u	N	N					
J1645	Dalteparin sodium	N	N					
J1650	Inj enoxaparin sodium	N	N					
J1652	Fondaparinux sodium	CH	K	1276		\$5.98		\$1.20
J1655	Tinzaparin sodium injection	N	N					
J1670	Tetanus immune globulin inj	K	K	1670		\$199.91		\$39.99
J1675	Histrelin acetate	B	B					
J1680	Human fibrinogen conc inj	NI	G	1290		\$96.46		\$18.93
J1700	Hydrocortisone acetate inj	N	N					
J1710	Hydrocortisone sodium ph inj	N	N					
J1720	Hydrocortisone sodium succ i	N	N					
J1730	Diazoxide injection	K	K	1740		\$112.16		\$22.44
J1740	Ibandronate sodium injection	K	K	9229		\$139.22		\$27.85
J1742	Ibutilide fumarate injection	K	K	9044		\$404.01		\$80.81
J1743	Ictursulfase injection	K	K	9232		\$446.44		\$89.29
J1745	Infliximab injection	K	K	7043		\$57.60		\$11.52
J1750	Inj iron dextran	K	K	1237		\$14.11		\$2.83
J1756	Iron sucrose injection	K	K	9046		\$0.37		\$0.08

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J1200	Diphenhydramine hcl injectio	N	N					
J1205	Chlorothiazide sodium inj	K	K	0747		\$292.02		\$58.41
J1212	Dimethyl sulfoxide 50% 50 ML	K	K	1221		\$67.46		\$13.50
J1230	Methadone injection	N	N					
J1240	Dimenhydrinate injection	N	N					
J1245	Dipyridamole injection	N	N					
J1250	Inj dobutamine HCL/250 mg	N	N					
J1260	Dolasetron mesylate	CH	N					
J1265	Dopamine injection	N	N					
J1267	Doxipenem injection	G	G	9241		\$0.57		\$0.11
J1270	Injection, doxercalciferol	N	N					
J1300	Eculizumab injection	CH	K	9236		\$177.57		\$35.52
J1320	Amiripryline injection	N	N					
J1324	Entuvitride injection	CH	K	1257		\$0.47		\$0.10
J1325	Epoprostenol injection	N	N					
J1327	Eprifibatid injection	K	K	1607		\$18.57		\$3.72
J1330	Eronovine maleate injection	N	N					
J1335	Ertapenem injection	N	N					
J1364	Erythro lactobionate /500 MG	N	N					
J1390	Estradiol valerate 20 MG inj	N	N					
J1390	Estradiol valerate 20 MG inj	N	N					
J1410	Inj estrogen conjugate 25 MG	K	K	9038		\$83.21		\$16.65
J1430	Ethanolamine oleate 100 mg	K	K	1688		\$147.14		\$29.43
J1435	Injection estone per 1 MG	N	N					
J1436	Eltroxate disodium inj	K	K	1436		\$70.06		\$14.02
J1438	Etanercept injection	K	K	1608		\$183.61		\$36.73
J1440	Filgrastim 300 mcg injection	K	K	0728		\$208.54		\$41.71
J1441	Filgrastim 480 mcg injection	K	K	7049		\$324.44		\$64.89
J1450	Fluconazole	N	N					
J1451	Fomepizole, 15 mg	K	K	1689		\$7.99		\$1.60
J1452	Intraocular Formivisen na	CH	E					
J1453	Fosaprepitant injection	G	G	9242		\$1.58		\$0.31
J1455	Foscarnet sodium injection	CH	N					
J1457	Gallium nitrate injection	K	K	0878		\$1.71		\$0.35
J1458	Galsulfase injection	K	K	9224		\$339.04		\$67.81
J1459	Inj IVIG priven 500 mg	G	G	1214		\$35.05		\$6.88
J1460	Gamma globulin 1 CC inj	K	K	3043		\$15.05		\$3.01
J1470	Gamma globulin 2 CC inj	CH	K	1282		\$30.10		\$6.02
J1480	Gamma globulin 3 CC inj	CH	K	1283		\$45.14		\$9.03
J1490	Gamma globulin 4 CC inj	K	K	0904		\$60.20		\$12.04
J1500	Gamma globulin 5 CC inj	CH	K	1284		\$75.26		\$15.06
J1510	Gamma globulin 6 CC inj	K	K	0920		\$90.35		\$18.07
J1520	Gamma globulin 7 CC inj	K	K	0921		\$105.27		\$21.06

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J2310	Inj naloxone hydrochloride		N					
J2315	Naltrexone, depot form		K	0759		\$2.14		\$0.43
J2320	Nandrolone decanoate 50 MG	CH	K	1285		\$7.00		\$1.40
J2321	Nandrolone decanoate 100 MG	CH	K	1286		\$7.00		\$1.40
J2322	Nandrolone decanoate 200 MG	CH	K	1286		\$14.74		\$2.95
J2323	Natalizumab injection		K	9126		\$8.32		\$1.67
J2325	Nesifiridine injection		K	1695		\$36.07		\$7.22
J2353	Ocreotide injection, depot		K	1207		\$105.27		\$21.06
J2354	Ocreotide inj, non-depot		N					
J2355	Oprelvekin injection		K	7011		\$242.16		\$48.44
J2357	Omalizumab injection		K	9300		\$18.86		\$3.78
J2360	Orphenadrine injection		N					
J2370	Phenylephrine hcl injection		N					
J2400	Chlorprocaine hcl injection		N					
J2405	Ondansetron hcl injection		N					
J2410	Oxymorphone hcl injection	CH	N					
J2425	Palferrin injection		K	1696		\$11.06		\$2.22
J2430	Pamidronate disodium /30 MG		K	0730		\$18.42		\$3.69
J2440	Papaverin hcl injection		N					
J2460	Oxytetracycline injection		E					
J2469	Palonosetron hcl		K	9210		\$17.19		\$3.44
J2501	Paricalcitol		N					
J2503	Pegaptanib sodium injection		K	1697		\$1,014.11		\$202.83
J2504	Pegademase bovine, 25 lu		K	1739		\$242.67		\$48.54
J2505	Injection, pegfilgrastim 6mg		K	9119		\$2,222.07		\$444.42
J2510	Penicillin g procaine inj		N					
J2513	Pentastarch 10% solution		K	1222		\$1,270.88		\$254.18
J2515	Pentobarbital sodium inj		N					
J2540	Penicillin g potassium inj	CH	N					
J2543	Piperacillin/tazobactam		N					
J2545	Pentamidine non-comp unit		B					
J2560	Promethazine hcl injection		N					
J2562	Plerixator injection	NI	G	9252		\$268.51		\$52.69
J2590	Oxytocin injection		N					
J2597	Inj desmopressin acetate		N					
J2650	Prechisalone acetate inj		N					
J2670	Totazoline hcl injection		N					
J2675	Inj progesterone per 50 MG Fluphenazine decanoate 25 MG		N					
J2680	Procainamide hcl injection		N					
J2700	Oxacillin sodium injection		N					
J2710	Neostigmine methylsulfate inj		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J1785	Injection imiglucerase /unit		K	0916		\$4.12		\$0.83
J1790	Droperidol injection		N					
J1800	Propranolol injection		N					
J1810	Droperidol/fentanyl inj		E					
J1815	Insulin injection		N					
J1817	Insulin for insulin pump use	CH	K	1277		\$3.34		\$0.67
J1825	Interferon beta-1a		E					
J1830	Interferon beta-1b / 25 MG		K	0910		\$168.90		\$33.78
J1835	Itraconazole injection	CH	N					
J1840	Kanamycin sulfate 500 MG inj		N					
J1850	Kanamycin sulfate 75 MG inj		N					
J1885	Keorotec tromethamine inj		N					
J1890	Cephalothin sodium injection		N					
J1930	Lamotrigine injection		K	9237		\$28.65		\$5.73
J1931	Laromedase injection		K	9209		\$25.08		\$5.02
J1940	Furosemide injection		N					
J1945	Lepirudin		K	1693		\$174.51		\$34.91
J1950	Leuprolide acetate (3.75 MG		K	0800		\$480.20		\$96.04
J1953	Levetiracetam injection		G	9238		\$0.75		\$0.15
J1955	Inj levocarnitine per 1 gm		B					
J1956	Levofloxacin injection		N					
J1960	Levorphanol tartrate inj		N					
J1980	Hyoscyamine sulfate inj		N					
J1990	Chloridazepoxide injection		N					
J2001	Lidocaine injection		N					
J2010	Lincocmycin injection		N					
J2020	Lincosolid injection		K	9001		\$29.37		\$5.88
J2060	Lorazepam injection		N					
J2150	Mannitol injection		N					
J2170	Mecasermin injection		N					
J2175	Meperidine hydrochloride /100 MG		N					
J2180	Meperidine/promethazine inj		N					
J2185	Meropenem		N					
J2210	Methylergonovin maleate inj		N					
J2248	Micafungin sodium injection		K	9227		\$1.08		\$0.22
J2250	Inj midazolam hydrochloride		N					
J2260	Inj milrinone lactate / 5 MG		N					
J2270	Morphine sulfate injection		N					
J2271	Morphine sulfate injection 100mg		N					
J2275	Morphine sulfate injection		N					
J2278	Ziconotide injection		K	1694		\$6.65		\$1.33
J2280	Inj, moxifloxacin 100 mg		N					
J2300	Inj nalbuphine hydrochloride		N					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J3140	Testosterone suspension inj	N	N					
J3150	Testosterone propionate inj	N	N					
J3230	Chlorpromazine hcl injection	N	N					
J3240	Thyrotropin injection	K	9108			\$948.38		\$189.68
J3243	Tigecycline injection	K	9228			\$1.15		\$0.23
J3246	Trofiban HCl	K	7041			\$7.83		\$1.57
J3250	Trimethoprim-sulfamethoxazole hcl inj	N	N					
J3260	Tobramycin sulfate injection	N	N					
J3280	Injection torsemide 10 mg/ml	N	N					
J3280	Thiethylperazine maleate inj	N	N					
J3285	Treprostinil injection	K	1701			\$54.83		\$10.97
J3300	Triamcinolone A inj PFS-free	K	1253			\$3.20		\$0.64
J3301	Triamcinolone acet inj NOS	N	N					
J3302	Triamcinolone diacetate inj	N	N					
J3303	Triamcinolone hexacetate inj	N	N					
J3305	Inj trimetrexate glucuronate	K	7045			\$124.80		\$24.96
J3310	Perphenazine injection	N	N					
J3315	Triptorelin pamoate	K	9122			\$160.83		\$32.17
J3320	Spectinomycin di-hcl inj	N	N					
J3350	Urea Injection	CH	N					
J3355	Urofollitropin, 75 iu	K	1741			\$59.26		\$11.86
J3360	Diazepam injection	N	N					
J3364	Urokinase 5000 IU injection	N	N					
J3365	Urokinase 250,000 IU inj	K	7036			\$449.09		\$89.82
J3370	Vancocycin hcl injection	N	N					
J3396	Verteporfin injection	K	1203			\$9.31		\$1.87
J3400	Triflupromazine hcl inj	CH	N					
J3410	Hydroxyzine hcl injection	N	N					
J3411	Thiamine hcl 100 mg	N	N					
J3415	Pyridoxine hcl 100 mg	N	N					
J3420	Vitamin B12 injection	N	N					
J3465	Injection, voriconazole	K	1052			\$5.26		\$1.06
J3470	Hyaluronidase injection	N	N					
J3471	Ovine, up to 999 USP units	N	N					
J3472	Ovine, 1000 USP units	CH	N					
J3473	Hyaluronidase recombinant	CH	N					
J3475	Inj magnesium sulfate	N	N					
J3480	Inj potassium chloride	N	N					
J3485	Zidovudine	N	N					
J3486	Ziprasidone mesylate	N	N					
J3487	Zoledronic acid	K	9115			\$214.94		\$42.99
J3488	Rheciast injection	CH	K	0951		\$218.59		\$43.72

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J2720	Inj progamine sulfate/10 MG	N	N			\$11.96		\$2.40
J2724	Protein c concentrate	K	1139					
J2725	Inj protirelin per 250 mcg	N	N			\$85.83		\$17.17
J2730	Pralidoxime chloride inj	K	1023					
J2760	Phentolamine mesylate inj	N	N					
J2765	Metoclopramide hcl injection	N	N					
J2770	Curumipristin/dalofopristin	K	2770			\$144.08		\$28.82
J2778	Ranitidine injection	K	9233			\$398.11		\$79.63
J2780	Ranitidine hydrochloride inj	N	N					
J2783	Rasburicase	K	0738			\$164.00		\$32.80
J2785	Regadenoson injection	G	9244			\$50.78		\$9.96
J2788	Rho d immune globulin 50 mcg	K	9023			\$25.76		\$5.16
J2790	Rho d immune globulin inj	K	0884			\$84.39		\$16.88
J2791	Rhophylac injection	K	0945			\$5.13		\$1.03
J2792	Rho(D) immune globulin h, sd	K	1609			\$18.39		\$3.68
J2793	Rilonacept injection	NI	K	1291		\$23.64		\$4.73
J2794	Risperidone, long acting	K	9125			\$4.93		\$0.99
J2795	Ropivacaine HCl injection	N	N					
J2796	Romiplostin injection	NI	G	9245		\$43.75		\$8.58
J2800	Methocarbamol injection	N	N					
J2805	Sinacalide injection	CH	N					
J2810	Inj theophylline per 40 MG	N	N					
J2820	Sargramostim injection	K	0731			\$23.31		\$4.67
J2850	Inj secretin synthetic human	K	1700			\$19.93		\$3.99
J2910	Aurothioglucose injection	N	N					
J2916	Na ferric gluconate complex	N	N					
J2920	Methylprednisolone injection	N	N					
J2930	Methylprednisolone injection	N	N					
J2940	Somatrem injection	K	1225			\$43.99		\$8.80
J2941	Somatropin injection	K	7034			\$59.47		\$10.70
J2950	Promazine hcl injection	N	N					
J2993	Retepase injection	K	9005			\$1,230.80		\$246.16
J2995	Inj streptokinase (2500000 IU	K	1226			\$78.00		\$15.60
J2997	Alteplase recombinant	K	7048			\$35.03		\$7.01
J3000	Streptomycin injection	N	N					
J3010	Fentanyl citrate injection	N	N					
J3030	Sumatriptan succinate / 6 MG	K	3030			\$55.49		\$11.10
J3070	Peniazocine injection	N	N					
J3101	Tenecteplase injection	K	9002			\$40.10		\$8.02
J3105	Terbutaline sulfate inj	N	N					
J3110	Teniparotide injection	B	N					
J3120	Testosterone enanthate inj	N	N					
J3130	Testosterone enanthate inj	N	N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J7330	Cultured chondrocytes implant		B					
J7500	Azathioprine oral 50mg		N					
J7501	Azathioprine parenteral		K	0887		\$90.64		\$18.13
J7502	Cyclosporine oral 100 mg	CH	K	1292		\$3.22		\$0.65
J7504	Lymphocyte immune globulin		K	0690		\$453.67		\$90.74
J7505	Monoclonal antibodies		K	7038		\$1,109.45		\$221.89
J7506	Prednisone oral		N					
J7507	Tacrolimus oral per 1 MG		K	0891		\$3.96		\$0.80
J7509	Methylprednisolone oral		N					
J7510	Prednisolone oral per 5 mg		N					
J7511	Antithymocyte globulin rabbit		K	9104		\$414.44		\$82.89
J7513	Dacizumab, parenteral		K	1612		\$378.20		\$75.64
J7515	Cyclosporine oral 25 mg	CH	K	1294		\$0.82		\$0.17
J7516	Cyclosporin parenteral 250mg		K	1204		\$21.24		\$4.25
J7517	Mycophenolate mofetil oral		K	9015		\$2.45		\$0.49
J7518	Mycophenolic acid	CH	N					
J7520	Stimulus, oral		K	9020		\$9.44		\$1.89
J7525	Tacrolimus injection		K	9006		\$136.82		\$27.37
J7599	Immunosuppressive drug noc		N					
J7604	Acetylcysteine comp unit		M					
J7605	Arformoterol non-comp unit		M					
J7606	Formoterol fumarate, inh		M					
J7607	Levalbuterol comp con		M					
J7608	Acetylcysteine non-comp unit		M					
J7609	Albuterol comp unit		M					
J7610	Albuterol comp con		M					
J7611	Albuterol non-comp con		M					
J7612	Levalbuterol non-comp con		M					
J7613	Albuterol non-comp unit		M					
J7614	Levalbuterol non-comp unit		M					
J7615	Levalbuterol comp unit		M					
J7620	Albuterol ipratrop non-comp		M					
J7622	Beclomethasone comp unit		M					
J7624	Betamethasone comp unit		M					
J7626	Budesonide non-comp unit		M					
J7627	Budesonide comp unit		M					
J7628	Biotin mesylate comp con		M					
J7629	Biotin mesylate comp unit		M					
J7631	Cromolyn sodium noncomp unit		M					
J7632	Cromolyn sodium comp unit		M					
J7633	Budesonide non-comp con		M					
J7634	Budesonide comp con		M					
J7635	Atropine comp con		M					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J3490	Drugs unclassified injection		N					
J3520	Edate discoidum per 150 mg		E					
J3530	Nasal vaccine inhalation		N					
J3535	Metered dose inhaler drug		E					
J3570	Laetile amygdalin vit B17		E					
J3590	Unclassified biologics		N					
J7030	Normal saline solution infus		N					
J7040	Normal saline solution infus		N					
J7042	5% dextrose/normal saline		N					
J7050	Normal saline solution infus		N					
J7060	5% dextrose/water		N					
J7070	D5w infusion		N					
J7100	Dextran 40 infusion		N					
J7110	Dextran 75 infusion		N					
J7120	Fingers lactate infusion		N					
J7130	Hypertonic saline solution		N					
J7185	Xyntha in	NI	K	1268		\$1.06		\$0.22
J7186	Antihemophilic viii/vwf comp		K	1213		\$0.84		\$0.17
J7187	Humate-P, inj		K	1704		\$0.87		\$0.18
J7189	Factor vlla		K	1705		\$1.29		\$0.26
J7190	Factor viii		K	0925		\$0.84		\$0.17
J7191	Factor VIII (porcine)	CH	K	1279		\$2.00		\$0.40
J7192	Factor viii recombinant NOS		K	0927		\$1.08		\$0.22
J7193	Factor IX non-recombinant		K	0931		\$0.88		\$0.18
J7194	Factor ix complex		K	0928		\$0.85		\$0.17
J7195	Factor IX recombinant		K	0932		\$1.06		\$0.22
J7197	Antithrombin iii injection	CH	K	1263		\$2.28		\$0.46
J7198	Anti-inhibitor		K	0929		\$1.53		\$0.31
J7199	Hemophilia clot factor noc		B					
J7300	Intraut copper contraceptive		E					
J7302	Levonorgestrel tu contraceptive		E					
J7304	Contraceptive vaginal ring		E					
J7306	Contraceptive hormone patch		E					
J7307	Levonorgestrel implant sys		E					
J7308	Etonogestrel implant system		E					
J7310	Aminolevulinic acid hel top		K	7308		\$127.60		\$25.52
J7311	Ganciclovir long act implant		K	0913		\$16,640.00		\$3,328.00
J7321	Fluocinolone acetonide impit		K	9225		\$18,980.00		\$3,796.00
J7322	Hyalgan/supartz inj per dose		K	0873		\$91.87		\$18.38
J7323	Synvisc inj per dose	CH	D					
J7324	Euflexxa inj per dose		K	0875		\$113.96		\$22.80
J7325	Orthovisc inj per dose		K	0877		\$177.68		\$35.54
J7325	Synvisc or Synvisc-One	NI	K	0874		\$11.47		\$2.30

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J8597	Antiemetic drug oral NOS		N					
J8600	Melphalan oral 2 MG		N					
J8610	Mefloquine oral 2.5 MG		N					
J8650	Nabilone oral	CH	N					
J8700	Temozolomide		K	1086		\$8.59		\$1.72
J8705	Topotecan oral		G	1238		\$71.35		\$14.00
J8999	Oral prescription drug chemo		B					
J9000	Doxorubicin hcl injection		N					
J9001	Doxorubicin hcl liposome inj		K	7046		\$450.51		\$90.11
J9010	Alendronate injection		K	9110		\$559.46		\$111.90
J9015	Alendronate injection		K	0807		\$631.49		\$166.30
J9017	Arsenic trioxide injection		K	9012		\$36.73		\$7.35
J9020	Asparaginase injection		K	0814		\$56.92		\$11.39
J9025	Azacioline injection		K	1709		\$4.78		\$0.96
J9027	Clofarabine injection		K	1710		\$114.21		\$22.85
J9031	Ecig live intravesical vac		K	0809		\$111.08		\$22.22
J9033	Bendamustine injection		G	9243		\$18.53		\$3.64
J9035	Bevacizumab injection		K	9214		\$56.39		\$11.28
J9040	Bleomycin sulfate injection		N					
J9041	Bortezomib injection		K	9207		\$36.54		\$7.31
J9045	Carboplatin injection		N					
J9050	Carmustine injection		K	0812		\$173.73		\$34.75
J9055	Cetuximab injection		K	9215		\$48.79		\$9.76
J9060	Cisplatin 10 MG injection		N					
J9062	Cisplatin 50 MG injection		N					
J9065	Inj cladribine per 1 MG		K	0858		\$25.15		\$5.03
J9070	Cyclophosphamide 100 MG inj		N					
J9080	Cyclophosphamide 200 MG inj		N					
J9090	Cyclophosphamide 500 MG inj		N					
J9091	Cyclophosphamide 1.0 grm inj		N					
J9092	Cyclophosphamide 2.0 grm inj		N					
J9093	Cyclophosphamide lyophilized		N					
J9094	Cyclophosphamide lyophilized		N					
J9095	Cyclophosphamide lyophilized		N					
J9096	Cyclophosphamide lyophilized		N					
J9097	Cyclophosphamide lyophilized		N					
J9098	Cytarabine liposome inj		K	1166		\$480.19		\$96.04
J9100	Cytarabine hcl 100 MG inj		N					
J9110	Cytarabine hcl 500 MG inj		N					
J9120	Dactinomycin injection		K	0752		\$533.21		\$106.65
J9130	Dacarbazine 100 mg inj		N					
J9140	Dacarbazine 200 MG inj		N					
J9150	Daunorubicin injection		K	0820		\$14.95		\$2.99

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J7636	Atropine comp unit		M					
J7637	Dexamethasone comp con		M					
J7638	Dexamethasone comp unit		M					
J7639	Dornase alfa non-comp unit		M					
J7640	Formoterol comp unit		E					
J7641	Flunisolide comp unit		M					
J7642	Glycopyrrolate comp con		M					
J7643	Glycopyrrolate comp unit		M					
J7644	Ipratropium bromide non-comp		M					
J7645	Ipratropium bromide comp		M					
J7647	Isoetharine comp con		M					
J7648	Isoetharine non-comp con		M					
J7649	Isoetharine non-comp unit		M					
J7650	Isoetharine comp unit		M					
J7657	Isoproterenol comp con		M					
J7658	Isoproterenol non-comp con		M					
J7659	Isoproterenol non-comp unit		M					
J7660	Isoproterenol comp unit		M					
J7667	Metaproterenol comp con		M					
J7668	Metaproterenol non-comp con		M					
J7669	Metaproterenol non-comp unit		M					
J7670	Metaproterenol comp unit		M					
J7674	Methacholine chloride, neb		N					
J7676	Pentamidine comp unit dose		M					
J7680	Terbutaline suif comp con		M					
J7681	Terbutaline suif comp unit		M					
J7682	Tobramycin non-comp unit		M					
J7683	Triamcinolone comp con		M					
J7684	Triamcinolone comp unit		M					
J7685	Tobramycin comp unit		M					
J7699	Inhalation solution for DME		M					
J7739	Non-inhalation drug for DME		N					
J8498	Antiemetic, rectal/supp NOS		B					
J8499	Oral prescrip drug non chemo		E					
J8501	Oral aprepitant		K	0868		\$5.42		\$1.09
J8510	Oral busulfan	CH	N					
J8515	Cabergoline, oral 0.25mg		E					
J8520	Capecitabine, oral, 150 mg		K	7042		\$5.68		\$1.14
J8521	Capecitabine, oral, 500 mg		K	0934		\$18.73		\$3.75
J8530	Cyclophosphamide oral 25 MG		N					
J8540	Oral dexamethasone		N					
J8560	Etoposide oral 50 MG		K	0802		\$0.45		\$0.09
J8565	Gefitinib oral		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J9293	Mitoxantrone hydrochl / 5 MG		K	0864		\$65.51		\$13.11
J9300	Gemtuzumab ozogamicin inj		K	9004		\$2,572.82		\$514.57
J9303	Panitumumab injection		K	9235		\$85.21		\$17.05
J9305	Pemetrexed injection		K	9213		\$48.50		\$9.70
J9310	Rituximab injection		K	0849		\$552.70		\$110.54
J9320	Streptozocin injection		K	0850		\$278.35		\$55.67
J9328	Temozolomide injection	NI	G	9253		\$4.90		\$0.96
J9330	Temsrolimus injection	CH	K	1168		\$47.93		\$9.59
J9340	Thiotepa injection		K	0851		\$97.69		\$19.54
J9350	Topotecan injection		K	0852		\$988.88		\$197.78
J9355	Trastuzumab injection		K	1613		\$63.51		\$12.71
J9357	Valrubicin injection		K	1235		\$953.16		\$190.64
J9360	Vinblastine sulfate inj		N					
J9370	Vincristine sulfate 1 MG inj		N					
J9375	Vincristine sulfate 2 MG inj		N					
J9380	Vincristine sulfate 5 MG inj		N					
J9390	Vincristine tartrate inj	CH	N					
J9395	Vincristine tartrate inj		K	9120		\$80.63		\$16.13
J9600	Portimer sodium injection		K	0856		\$2,745.46		\$549.10
J9999	Chemotherapy drug		N					
K0001	Standard wheelchair		N					
K0002	Stnd hemi (low seat) wheelch		Y					
K0003	Lightweight wheelchair		Y					
K0004	High strength lwt wheelch		Y					
K0005	Ultralightweight wheelchair		Y					
K0006	Heavy duty wheelchair		Y					
K0007	Extra heavy duty wheelchair		Y					
K0009	Other manual wheelchair/base		Y					
K0010	Stnd wt frame power wheelch		Y					
K0011	Stnd wt pwr wheelch w control		Y					
K0012	Lvrt portbl power wheelch		Y					
K0014	Other power wheelch base		Y					
K0015	Detach non-adjust height armst		Y					
K0017	Detach adjust armrest base		Y					
K0018	Detach adjust armst upper		Y					
K0019	Arm pad each		Y					
K0020	Fixed adjust armrest pair		Y					
K0037	High mount flip-up footrest		Y					
K0038	Leg strap each		Y					
K0039	Leg strap h style each		Y					
K0040	Adjustable angle footplate		Y					
K0041	Large size footplate each		Y					
K0042	Standard size footplate each		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
J9151	Dauunonubcin citrate inj		K	0821		\$55.27		\$11.06
J9155	Degarelix injection	NI	G	1296		\$2.23		\$0.44
J9160	Demileukin diftiox inj		K	1084		\$1,448.32		\$289.67
J9165	Diethylstilbestrol Injection		K	1209		\$1,257.36		\$251.48
J9170	Docetaxel injection	CH	D					
J9171	Docetaxel injection	NI	K	0823		\$16.95		\$3.39
J9175	Elliotts b solution per ml		N					
J9178	inj, epirubicin hcl, 2 mg		K	1167		\$2.55		\$0.51
J9181	Etoposide injection		N					
J9185	Fludarabine phosphate inj		K	0842		\$151.36		\$30.28
J9190	Fluorouracil injection		N					
J9200	Floxuridine injection		K	0827		\$46.60		\$9.32
J9201	Gemcitabine hcl injection		K	0828		\$139.10		\$27.82
J9202	Goserelin acetate implant		K	0810		\$193.02		\$38.61
J9206	Irinotecan injection		K	0830		\$13.18		\$2.64
J9207	kabepalone injection		G	9240		\$63.74		\$12.51
J9208	fosfomide injection		K	0831		\$29.39		\$5.88
J9209	Mesna injection		K	0732		\$4.34		\$0.87
J9211	Idarubicin hcl injection		K	0832		\$19.34		\$3.94
J9212	Interferon alfacon-1 inj	CH	K	1266		\$6.75		\$1.35
J9213	Interferon alfa-2a inj		K	0834		\$10.60		\$2.12
J9214	Interferon alfa-2b inj		K	0836		\$15.54		\$3.11
J9215	Interferon alfa-n3 inj		K	0865		\$17.89		\$3.58
J9216	Interferon gamma 1-b inj		K	0838		\$294.03		\$58.81
J9217	Leuprolide acetate suspension		K	9217		\$210.52		\$42.11
J9218	Leuprolide acetate injection		K	0861		\$5.29		\$1.06
J9219	Leuprolide acetate implant		K	7051		\$4,728.88		\$945.78
J9225	Vantas implant		G	1711		\$1,473.60		\$289.16
J9226	Supprelin LA implant		G	1142		\$14,875.43		\$2,918.95
J9230	Mechlorethamine hcl inj		K	0751		\$144.56		\$28.92
J9245	inj melphalan hydrochl 50 MG		K	0840		\$1,622.81		\$324.57
J9250	Methotrexate sodium inj		N					
J9260	Methotrexate sodium inj		N					
J9261	Nelarabine injection	CH	K	0825		\$101.28		\$20.26
J9263	Oxaliplatin		K	1738		\$9.55		\$1.91
J9264	Paclitaxel protein bound		K	1712		\$9.09		\$1.82
J9265	Paclitaxel injection	CH	N					
J9266	Pegaspargase injection		K	0843		\$2,695.67		\$539.14
J9268	Pentostatin injection		K	0844		\$1,399.56		\$279.92
J9270	Plicamycin (mithramycin) inj	CH	N					
J9280	Mitomycin 5 MG inj		K	1232		\$17.74		\$3.55
J9290	Mitomycin 20 MG inj		K	1233		\$70.98		\$14.20
J9291	Mitomycin 40 MG inj		K	1234		\$141.95		\$28.39

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
K0739	Repair/svc DME non-oxygen eq		Y					
K0740	Repair/svc oxygen equipment		E					
K0800	POV group 1 std up to 300lbs		Y					
K0801	POV group 1 vhd 301-450 lbs		Y					
K0802	POV group 1 vhd 451-600 lbs		Y					
K0806	POV group 2 std up to 300lbs		Y					
K0807	POV group 2 hd 301-450 lbs		Y					
K0808	POV group 2 vhd 451-600 lbs		Y					
K0812	Power operated vehicle NOC		Y					
K0813	PWC gp 1 std pont seat/back		Y					
K0814	PWC gp 1 std pont cap chair		Y					
K0815	PWC gp 1 std seat/back		Y					
K0816	PWC gp 1 std cap chair		Y					
K0820	PWC gp 2 std pont seat/back		Y					
K0821	PWC gp 2 std pont cap chair		Y					
K0822	PWC gp 2 std seat/back		Y					
K0823	PWC gp 2 std cap chair		Y					
K0824	PWC gp 2 hd seat/back		Y					
K0825	PWC gp 2 hd cap chair		Y					
K0826	PWC gp 2 vhd seat/back		Y					
K0827	PWC gp vhd cap chair		Y					
K0828	PWC gp 2 xtra hd seat/back		Y					
K0829	PWC gp 2 xtra hd cap chair		Y					
K0830	PWC gp2 std seat elevate s/b		Y					
K0831	PWC gp2 std seat elevate cap		Y					
K0835	PWC gp2 std sing pow opt s/b		Y					
K0836	PWC gp2 std sing pow opt cap		Y					
K0837	PWC gp 2 hd sing pow opt s/b		Y					
K0838	PWC gp 2 hd sing pow opt cap		Y					
K0839	PWC gp2 vhd sing pow opt s/b		Y					
K0840	PWC gp2 vhd sing pow opt s/b		Y					
K0841	PWC gp2 std mult pow opt s/b		Y					
K0842	PWC gp2 std mult pow opt cap		Y					
K0843	PWC gp2 hd mult pow opt s/b		Y					
K0848	PWC gp 3 std seat/back		Y					
K0849	PWC gp 3 std cap chair		Y					
K0850	PWC gp 3 hd seat/back		Y					
K0851	PWC gp 3 hd cap chair		Y					
K0852	PWC gp 3 vhd seat/back		Y					
K0853	PWC gp 3 vhd cap chair		Y					
K0854	PWC gp 3 xhd seat/back		Y					
K0855	PWC gp 3 xhd cap chair		Y					
K0856	PWC gp3 std sing pow opt s/b		Y					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
K0043	First lower extension tube		Y					
K0044	First upper hanger bracket		Y					
K0045	Footrest complete assembly		Y					
K0046	Elevat legrst low extension		Y					
K0047	Elevat legrst up hangr brack		Y					
K0050	Ratchet assembly		Y					
K0051	Cam release assem frst/legrst		Y					
K0052	Swingaway detach footrest		Y					
K0053	Elevate footrest articulate		Y					
K0056	Seat ht <17 or >=21 lwt wc		Y					
K0065	Spoke protectors		Y					
K0069	Rear whl complete solid tire		Y					
K0070	Rear whl compl pneu tire		Y					
K0071	Front castr compl pneu tire		Y					
K0072	Frrnt castr compl semi-pneu tir		Y					
K0073	Caster pin lock each		Y					
K0077	Front caster assem complete		Y					
K0098	Drive belt power wheelchair		Y					
K0105	Iv hanger		Y					
K0108	W/c component-accessory NOS		Y					
K0195	Elevating wheelchair leg rests		Y					
K0455	Pump uninterrupted infusion		Y					
K0462	Temporary replacement eqpmnt		Y					
K0552	Supply/ext inf pump syr bype		Y					
K0601	Repl batt silver oxide 1.5 v		Y					
K0602	Repl batt silver oxide 3 v		Y					
K0603	Repl batt alkaline 1.5 v		Y					
K0604	Repl batt lithium 3.6 v		Y					
K0605	Repl batt lithium 4.5 v		Y					
K0606	AED garment w elec analysis		Y					
K0607	Repl batt for AED		Y					
K0608	Repl garment for AED		Y					
K0609	Repl electrode for AED		Y					
K0669	Seat/back cus no sedmerc ver		Y					
K0672	Removable soft interface LE		A					
K0730	Ctrl dose inh drug deliv sys		Y					
K0733	12-24hr sealed lead acid		Y					
K0734	Adj skin pro w/c cus wd=22in		Y					
K0735	Adj skin pro wc cus wd>=22in		Y					
K0736	Adj skin pro/pos wc cus=22in		Y					
K0737	Adj skin pro/pos wc cus>=22"		Y					
K0738	Portable gas oxygen system		Y					

APPENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L0458	TLSO 2Mod symphis-xipho pre		A					
L0460	TLSO2Mod symphis-sterm pre		A					
L0462	TLSO 3Mod sacro-scap pre		A					
L0464	TLSO 4Mod sacro-scap pre		A					
L0466	TLSO rigid frame pre soft ap		A					
L0468	TLSO rigid frame prefab pelv		A					
L0470	TLSO rigid frame pre subclav		A					
L0472	TLSO rigid frame hyperex pre		A					
L0480	TLSO rigid plastic custom fa		A					
L0482	TLSO rigid lined custom tab		A					
L0484	TLSO rigid lined cust tab		A					
L0486	TLSO rigid plastic cust tab		A					
L0488	TLSO rigid lined pre one pie		A					
L0490	TLSO rigid plastic pre one		A					
L0491	TLSO 2 piece rigid shell		A					
L0492	TLSO 3 piece rigid shell		A					
L0621	SIO flex pelvisacral prefab		A					
L0622	SIO flex pelvisacral custom		A					
L0623	SIO panel prefab		A					
L0624	SIO panel custom		A					
L0625	LO flexibl L1-below L5 pre		A					
L0626	LO sag stays/panels pre-fab		A					
L0627	LO sagitt rigid panel prefab		A					
L0628	LO flex w/o rigid stays pre		A					
L0629	LSO flex w/rigid stays cust		A					
L0630	LSO post rigid panel pre		A					
L0631	LSO sag-coro rigid frame pre		A					
L0632	LSO sag rigid frame cust		A					
L0633	LSO flexion control prefab		A					
L0634	LSO flexion control custom		A					
L0635	LSO sagitt rigid panel prefab		A					
L0636	LSO sagittal rigid panel cus		A					
L0637	LSO sag-coronal panel prefab		A					
L0638	LSO sag-coronal panel custom		A					
L0639	LSO s/c shell/panel prefab		A					
L0640	LSO s/c shell/panel custom		A					
L0700	Cliso a-p-i control molded		A					
L0710	Cliso a-p-i control w/ inter		A					
L0810	Halo cervical into jckt vest		A					
L0820	Halo cervical into body jckt		A					
L0830	Halo cerv into milwaukee typ		A					
L0859	MFL compatible system		A					
L0861	Halo repl liner/interface		A					

APPENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
K0857	PWC gp3 std sing pow opt cap		Y					
K0858	PWC gp3 hd sing pow opt s/b		Y					
K0859	PWC gp3 hd sing pow opt cap		Y					
K0860	PWC gp3 vhd sing pow opt s/b		Y					
K0861	PWC gp3 std mult pow opt s/b		Y					
K0862	PWC gp3 hd mult pow opt s/b		Y					
K0863	PWC gp3 vhd mult pow opt s/b		Y					
K0864	PWC gp3 xhd mult pow opt s/b		Y					
K0868	PWC gp 4 std seat/back		Y					
K0869	PWC gp 4 std cap chair		Y					
K0870	PWC gp 4 hd seat/back		Y					
K0871	PWC gp 4 vhd seat/back		Y					
K0877	PWC gp4 std sing pow opt s/b		Y					
K0878	PWC gp4 std sing pow opt cap		Y					
K0879	PWC gp4 hd sing pow opt s/b		Y					
K0880	PWC gp4 vhd sing pow opt s/b		Y					
K0884	PWC gp4 std mult pow opt s/b		Y					
K0885	PWC gp4 std mult pow opt cap		Y					
K0886	PWC gp4 hd mult pow s/b		Y					
K0890	PWC gp5 ped sing pow opt s/b		Y					
K0891	PWC gp5 ped mult pow opt s/b		Y					
K0898	Power wheelchair NOC		Y					
K0899	Pow mobil dev no SADMERC		Y					
L0112	Cranial cervical orthosis		A					
L0113	Cranial cervical torticollis		A					
L0120	Cerv flexible non-adjustable		A					
L0130	Flex thermoplastic collar mo		A					
L0140	Cervical semi-rigid adjustab		A					
L0150	Cerv semi-rg adj molded chrn		A					
L0160	Cerv semi-rg wire occ/mand		A					
L0170	Cervical collar molded to pt		A					
L0172	Cerv col thermplas foam 2 pi		A					
L0174	Cerv col foam 2 piece w thor		A					
L0180	Cerv post col occ/main sup adj		A					
L0190	Cerv collar supp adj cerv ba		A					
L0200	Cerv col supp adj bar & thor		A					
L0210	Thoracic fib belt	CH	D					
L0220	Thor rib belt custom fabrica		A					
L0430	Dewall posture protector		A					
L0450	TLSO flex prefab thoracic		A					
L0452	tlso flex custom fab thoraci		A					
L0454	TLSO flex prefab sacroccc-t9		A					
L0456	TLSO flex prefab		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L1610	Abduct hip flex frejka covr	A	A					
L1620	Abduct hip flex pavlik harn	A	A					
L1630	Abduct control hip semi-flex	A	A					
L1640	Pelv band/spread bar thigh c	A	A					
L1650	HO abduction hip adjustable	A	A					
L1660	HO bi thighcuffs w sprd bar	A	A					
L1680	HO abduction static plastic	A	A					
L1685	Pelvic & hip control thigh c	A	A					
L1686	Post-op hip abduct custom fa	A	A					
L1688	HO post-op hip abduction	A	A					
L1690	Combination bilateral HO	A	A					
L1700	Leg perthes orth toronto typ	A	A					
L1710	Leg perthes orth newington	A	A					
L1720	Legg perthes orthos triat	A	A					
L1730	Legg perthes orth scottish r	A	A					
L1755	Legg perthes patten bottom t	A	A					
L1800	Knee orthoses elas w stays	CH	D					
L1810	Ko elastic with joints	A	A					
L1815	Elastic with condylar pads	CH	D					
L1820	Ko elas w/ condyle pads & jo	CH	D					
L1825	Ko elastic knee cap	A	A					
L1830	Ko immobilizer canvas longit	A	A					
L1831	Knee orth pos locking joint	A	A					
L1832	KO adj jnt pos rigid support	A	A					
L1834	Ko w/O joint rigid molded to	A	A					
L1836	Rigid KO w/o joints	A	A					
L1840	Ko derot ant cruciate custom	A	A					
L1843	KO single upright custom fit	A	A					
L1844	Ko w/adj jt rot cntrl molded	A	A					
L1845	Ko w/ adj flex/ext rotat cus	A	A					
L1846	Ko w adj flex/ext rotat mold	A	A					
L1847	KO adjustable w air chambers	A	A					
L1850	Ko swedish type	A	A					
L1860	Ko supracondylar socket mold	A	A					
L1900	Afo sprng wr drstlx call bd	CH	D					
L1901	Prefab ankle orthosis	A	A					
L1902	Afo ankle gauntlet	A	A					
L1904	Afo molded ankle gauntlet	A	A					
L1906	Afo multiligamentus ankle su	A	A					
L1907	AFO supramalleolar custom	A	A					
L1910	Afo sing bar clasp attach sh	A	A					
L1920	Afo sing upright w/ adjust s	A	A					
L1930	Afo plastic	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L0970	Tiso corset front	A	A					
L0972	Lso corset front	A	A					
L0974	Tiso full corset	A	A					
L0976	Lso full corset	A	A					
L0978	Axillary crutch extension	A	A					
L0980	Peroneal straps pair	A	A					
L0982	Stocking supp grips set of f	A	A					
L0984	Protective body sock each	A	A					
L0999	Add to spinal orthosis NOS	A	A					
L1000	Ctiso miwaukee initial model	A	A					
L1001	CTLSO infant immobilizer	A	A					
L1005	Tension based scoliosis orth	A	A					
L1010	Ctiso axilla sling	A	A					
L1020	Kyphosis pad	A	A					
L1025	Kyphosis pad floating	A	A					
L1030	Lumbar bolster pad	A	A					
L1040	Lumbar or lumbar rib pad	A	A					
L1050	Sternal pad	A	A					
L1060	Thoracic pad	A	A					
L1070	Trapezius sling	A	A					
L1080	Outrigger	A	A					
L1085	Outrigger bil w/ vert extens	A	A					
L1090	Lumbar sling	A	A					
L1100	Ring flange plastic/leather	A	A					
L1110	Ring flange plas/leather mol	A	A					
L1120	Covers for upright each	A	A					
L1200	Furnsh initial orthosis only	A	A					
L1210	Lateral thoracic extension	A	A					
L1220	Anterior thoracic extension	A	A					
L1230	Milwaukee type superstructur	A	A					
L1240	Lumbar derotation pad	A	A					
L1250	Anterior asis pad	A	A					
L1260	Anterior thoracic derotation	A	A					
L1270	Abdominal pad	A	A					
L1280	Rib gusset (elastic) each	A	A					
L1290	Lateral trochanteric pad	A	A					
L1300	Body jacket mold to patient	A	A					
L1310	Post-operative body jacket	A	A					
L1499	Spinal orthosis NOS	A	A					
L1500	Thkao mobility frame	A	A					
L1510	Thkao standing frame	A	A					
L1520	Thkao swivel walker	A	A					
L1600	Abduct hip flex frejka w cov	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L2200	Limited ankle motion ea jnt		A					
L2210	Dorsiflexion assist each jnt		A					
L2220	Dorsi & plantar flex ass/res		A					
L2230	Split flat caliper strir & p		A					
L2232	Rocker bottom, contact AFO		A					
L2240	Round caliper and plate attn		A					
L2250	Foot plate molded stirrup at		A					
L2260	Reinforced solid stirrup		A					
L2265	Long tongue stirrup		A					
L2270	Vanus/valgus strap padded/i		A					
L2275	Plastic mod low ext pad/line		A					
L2280	Molded inner boot		A					
L2290	Abduction bar jointed adjust		A					
L2310	Abduction bar-straight		A					
L2320	Non-molded lacer		A					
L2330	Lacer molded to patient molde		A					
L2335	Anterior swing band		A					
L2340	Pre-tibial shell molded to p		A					
L2350	Prosthetic type socket molde		A					
L2360	Extended steel shank		A					
L2370	Patten bottom		A					
L2375	Torsion ank & half solid sti		A					
L2380	Torsion straight knee joint		A					
L2385	Straight knee joint heavy du		A					
L2390	Add LE poly knee custom		A					
L2387	KAFO		A					
L2390	Offset knee joint each		A					
L2395	Offset knee joint heavy duty		A					
L2397	Suspension sleeve lower ext		A					
L2405	Knee joint drop lock ea jnt		A					
L2415	Knee joint cam lock each joi		A					
L2425	Knee disc/dial lock/adj flex		A					
L2430	Knee jnt ratchet lock ea jnt		A					
L2492	Knee lift loop drop lock rin		A					
L2500	Thigh/ischia wgt bearing		A					
L2510	Thigh/bear quad-lat brim m		A					
L2520	Thigh/bear quad-lat brim c		A					
L2525	Thigh/bear near m-1 brim mo		A					
L2528	Thigh/bear near m-1 brim cu		A					
L2530	Thigh/wght bear lacer non-mo		A					
L2540	Thigh/wght bear lacer molded		A					
L2550	Thigh/wght bear high roll cu		A					
L2570	Hip clevis type 2 posit jnt		A					
L2580	Pelvic control pelvic sling		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L1932	Ato rig ant tib prefab TCF/=		A					
L1940	Ato molded to patient: plasti		A					
L1945	Ato molded plas rig ant tib		A					
L1950	Ato spiral molded to pt plas		A					
L1951	AFO spiral prefabricated		A					
L1960	Ato pos solid ank plastic mo		A					
L1970	Ato plastic molded w/ankle j		A					
L1971	AFO w/ankle joint, prefab		A					
L1980	Ato sing solid stirrup calf		A					
L1990	Ato doub solid stirrup calf		A					
L2000	Kato sing tre strir th/calf		A					
L2005	KAFO sing/dbi mechanical act		A					
L2010	Kato sng solid stirrup w/o j		A					
L2020	Kato dbi solid stirrup band/		A					
L2030	Kato dbi solid stirrup w/o j		A					
L2034	KAFO pla sin up w/w/o k/a cus		A					
L2035	KAFO plastic pediatric size		A					
L2036	Kato plas doub free knee mol		A					
L2037	Ato plas sing free knee mol		A					
L2038	Kato w/o joint multi-axis an		A					
L2040	Hkato torsion bil rot straps		A					
L2050	Hkato torsion cable hip pelv		A					
L2060	Hkato torsion ball bearing j		A					
L2070	Hkato torsion unilat rot str		A					
L2080	Hkato unilat torsion cable		A					
L2090	Hkato unilat torsion ball br		A					
L2106	Ato tib fx cast plaster mold		A					
L2108	Ato tib fx cast molded to pt		A					
L2112	Ato tibial fracture soft		A					
L2114	Ato tib fx semi-rigid		A					
L2116	Ato tibial fracture rigid		A					
L2126	Kato fem fx cast thermoplas		A					
L2128	Kato fem fx cast molded to p		A					
L2132	Kato femoral fx cast soft		A					
L2134	Kato fem fx cast semi-rigid		A					
L2136	Kato femoral fx cast rigid		A					
L2180	Plas shoe insert w ank joint		A					
L2182	Drop lock knee		A					
L2184	Limited motion knee joint		A					
L2186	Adj motion knee jnt lerman t		A					
L2188	Quadrilateral brim		A					
L2190	Waist belt		A					
L2192	Pelvic band & belt thigh fla		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L3100	Hallux-valgus right dynamic s		A					
L3140	Abduction rotation bar shoe		A					
L3150	Abduct rotation bar w/o shoe		A					
L3160	Shoe styler positioning dev		A					
L3170	Foot plastic heel stabilizer		A					
L3201	Oxford w supinat/pronat inf		A					
L3202	Oxford w supinat/pronator c		A					
L3203	Oxford w supinator/pronator		A					
L3204	Hightop w/ sup/pronator inf		A					
L3206	Hightop w/ sup/pronator chi		A					
L3207	Hightop w/ sup/pronator jun		A					
L3208	Surgical boot each infant		A					
L3209	Surgical boot each child		A					
L3211	Surgical boot each junior		A					
L3212	Benesch boot pair infant		A					
L3213	Benesch boot pair child		A					
L3214	Benesch boot pair junior		A					
L3215	Orthopedic fwear ladies oxf		E					
L3216	Orthopedic ladies shoes depth i		E					
L3217	Ladies shoes hightop depth i		E					
L3219	Orthopedic mens shoes oxford		E					
L3221	Orthopedic mens shoes depth i		E					
L3222	Mens shoes hightop depth inf		E					
L3224	Woman's shoe oxford brace		A					
L3225	Man's shoe oxford brace		A					
L3230	Custom shoes depth inlay		A					
L3250	Custom mold shoe remov prost		A					
L3251	Shoe molded to pt silicone s		A					
L3252	Shoe molded plastazote cust		A					
L3253	Shoe molded plastazote cust		A					
L3254	Orth foot non-standard size/w		A					
L3257	Orth foot add charge split s		A					
L3260	Ambulatory surgical boot eac		E					
L3265	Plastazote sandal each		A					
L3300	Sho lift taper to metatarsal		A					
L3310	Shoe lift elev heel/sole neo		A					
L3320	Shoe lift elev heel/sole cor		A					
L3330	Lifts elevation metal extens		A					
L3332	Shoe lifts tapered to one-ha		A					
L3334	Shoe lifts elevation heel /i		A					
L3340	Shoe wedge each		A					
L3350	Shoe heel wedge		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L2600	Hip clevis/thrust bearing fr		A					
L2610	Hip clevis/thrust bearing lo		A					
L2620	Pelvic control hip heavy dut		A					
L2622	Hip joint adjustable flexion		A					
L2624	Hip adj flex ext abduct cont		A					
L2627	Plastic mold recipro hip & c		A					
L2628	Metal frame recipro hip & ca		A					
L2630	Pelvic control band & belt u		A					
L2640	Pelvic control band & belt b		A					
L2650	Paiv & thor control gluteal		A					
L2660	Thoracic control thoracic ba		A					
L2670	Thorac cont paraspinat uprig		A					
L2680	Thorac cont lat support upri		A					
L2750	Plating chrome/nickel pr bar		A					
L2755	Carbon graphite lamination		A					
L2760	Extension per extension per		A					
L2768	Ortho sidebar disconnect		A					
L2770	Low ext orthosis per bar/jnt	CH	D					
L2780	Non-corrosive finish		A					
L2785	Drop lock retainer each		A					
L2795	Knee control full kneecap		A					
L2800	Knee cap medial or lateral p		A					
L2810	Knee control condylar pad		A					
L2820	Soft interface below knee se		A					
L2830	Soft interface above knee se		A					
L2840	Tibial length sock tx or equ		A					
L2850	Femoral lgh sock fx or equ		A					
L2861	Torsion mechanism knee/ankle	NI	E					
L2999	Lower extremity orthosis NOS		A					
L3000	Ft insert ucb berkeley shell		A					
L3001	Foot insert remov molded spe		A					
L3002	Foot insert plastazote or eq		A					
L3003	Foot insert silicone gel eac		A					
L3010	Foot longitudinal arch suppo		A					
L3020	Foot longitud/metatarsal sup		A					
L3030	Foot arch support remov prem		A					
L3031	Foot lamini/prepreg composite		A					
L3040	Ft arch suprt premold longit		A					
L3050	Foot arch supp premold metat		A					
L3060	Foot arch supp longitud/meta		A					
L3070	Arch suprt att to sho longit		A					
L3080	Arch supp att to shoe metata		A					
L3090	Arch supp att to shoe long/m		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010										
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment		
L3700	Elbow orthosis elas w stays	CH	D							
L3701	Prefab elbow orthosis	CH	D							
L3702	EO w/o joints CF		A							
L3710	Elbow elastic with metal loi		A							
L3720	Forearm/arm cuffs free motio		A							
L3730	Forearm/arm cuffs ext/flex a		A							
L3740	Cuffs adj lock w active con		A							
L3760	EO withjoint, Prefabricated		A							
L3762	Rigid EO w/o joints		A							
L3763	EWHO rigid w/o jnts CF		A							
L3764	EWHO w/joint(s) CF		A							
L3765	EMHFO rigid w/o jnts CF		A							
L3766	EMHFO w/joint(s) CF		A							
L3806	WHFO w/joint(s) custom fab		A							
L3807	WHFO no joint, prefabricated		A							
L3808	WHFO, rigid w/o joints		A							
L3891	Torsion mechanism wrist/elbo	NI	E							
L3900	Hinge extension/flex wrist/f		A							
L3901	Hinge ext/flex wrist finger		A							
L3904	Wrist electric custom fitted		A							
L3905	WHO w/intorsion jnt(s) CF		A							
L3906	WHO w/o joints CF		A							
L3908	Wrist cock-up non-molded		A							
L3909	Prefab wrist orthosis	CH	D							
L3911	Prefab hand finger orthosis	CH	D							
L3912	Flex glove w/elastic finger		A							
L3913	HFO w/o joints CF		A							
L3915	WHO w nontor jnts) prefab		A							
L3917	Prefab metacarp i x orthosis		A							
L3919	HO w/o joints CF		A							
L3921	HFO w/joint(s) CF		A							
L3923	HFO w/o joints PF		A							
L3925	FO pld/dip with joint/spring		A							
L3927	FO pld/dip w/o joint/spring		A							
L3929	HFO nontorsion joint, prefab		A							
L3931	WHFO nontorsion joint prefab		A							
L3933	FO w/o joints CF		A							
L3935	FO nontorsion joint CF		A							
L3956	Add joint upper ext orthosis		A							
L3960	Sewho airplan desig abdu pos		A							
L3961	SEWHO cap design w/o jnts CF		A							
L3962	Sewho erbs palsey design abd		A							
L3964	Sewo mobile arm sup att to wc		Y							

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010										
HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment		
L3360	Shoe sole wedge outside sole		A							
L3370	Shoe sole wedge between sole		A							
L3380	Shoe clubfoot wedge		A							
L3390	Shoe outflare wedge		A							
L3400	Shoe metatarsal bar wedge ro		A							
L3410	Shoe metatarsal bar between		A							
L3420	Full sole/heel wedge between		A							
L3430	Sho heel count plast reinfor		A							
L3440	Heel leather reinforced		A							
L3450	Shoe heel sach cushion type		A							
L3455	Shoe heel new leather standrd		A							
L3460	Shoe heel new rubber standrd		A							
L3465	Shoe heel thomas with wedge		A							
L3470	Shoe heel thomas extend to b		A							
L3480	Shoe heel pad & depress for		A							
L3485	Shoe heel pad removable for		A							
L3500	Ortho shoe add leather insol		A							
L3510	Orthopedic shoe add tub insl		A							
L3520	O shoe add felt w leath insl		A							
L3530	Ortho shoe add half sole		A							
L3540	Ortho shoe add full sole		A							
L3550	O shoe add standard toe tap		A							
L3560	O shoe add horseshoe toe tap		A							
L3570	O shoe add instep extension		A							
L3580	O shoe add instep velcro clo		A							
L3590	O shoe convert to sof counte		A							
L3595	Ortho shoe add march bar		A							
L3600	Trans shoe call plate exist		A							
L3610	Trans shoe caliper plate new		A							
L3620	Trans shoe solid stirrup ext		A							
L3630	Trans shoe solid stirrup new		A							
L3640	Shoe dennis browne splint bo		A							
L3649	Orthopedic shoe modifca NOS		A							
L3650	Slider fig 8 abduct restrain		A							
L3651	Prefab shoulder orthosis	CH	D							
L3652	Prefab dbl shoulder orthosis	CH	D							
L3660	Abduct restrainer canvas&web		A							
L3670	Acromioclavicular canvas&web		A							
L3671	SO cap design w/o jnts CF		A							
L3672	SO airplane w/o jnts CF		A							
L3673	SO airplane w/joint CF		A							
L3675	Canvas vest SO		A							
L3677	SO hard plastic stabilizer		E							

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L4380	Pneumatic knee splint		A					
L4386	Non-pneum walk boot prefab		A					
L4392	Replace AFO soft interface		A					
L4394	Replace foot drop splint		A					
L4396	Static AFO		A					
L4398	Foot drop splint recumbent		A					
L5000	Sho insert w arch toe filler		A					
L5010	Mold socket ank hgt w/ toe f		A					
L5020	Tibial tubercle hgt w/ toe f		A					
L5050	Ank symes mold sock sach ft		A					
L5080	Symes met fr leath socket ar		A					
L5100	Molded socket shin sach foot		A					
L5105	Plast socket lts/thgh lacer		A					
L5150	Mold sock ext knee shin sach		A					
L5160	Mold socket bent knee shin s		A					
L5200	Kne sing axis fric shin sach		A					
L5210	No knee/ankle joints w/ ft b		A					
L5220	No knee joint with artic ali		A					
L5230	Fem focal detic constant fri		A					
L5250	Hip caned sing axi cons fric		A					
L5270	Tilt table locking hip sing		A					
L5280	Hemipelvect canad sing axis		A					
L5301	BK mold socket SACH ft endo		A					
L5311	Knee disart. SACH ft. endo		A					
L5321	AK open end SACH		A					
L5331	Hip disart canadian SACH ft		A					
L5341	Hemipelvectomy canadian SACH		A					
L5400	Postop dress & 1 cast chg bk		A					
L5410	Postop dsg bk ea add cast ch		A					
L5420	Postop dsg & 1 cast chg ak/d		A					
L5430	Postop dsg ak ea add cast ch		A					
L5450	Postop app non-wgt bear dsg		A					
L5480	Postop app non-wgt bear dsg		A					
L5500	Init bk ptb plaster direct		A					
L5505	Init ak ischial plstr direct		A					
L5520	Prep BK ptb thermopls direct		A					
L5530	Prep BK ptb plaster molded		A					
L5535	Prep BK ptb thermopls molded		A					
L5540	Prep BK ptb open end socket		A					
L5580	Prep AK ischial plast molded		A					
L5570	Prep AK ischial direct form		A					
L5580	Prep AK ischial thermo mold		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L3965	Arm supp att to wc rancho ty		Y					
L3966	Mobile arm supports reclini		Y					
L3967	SEWHO airplane w/o jnts CF		A					
L3968	Friction dampening arm supp		Y					
L3969	Monosuspension arm/hand supp		Y					
L3970	Elevat proximal arm support		Y					
L3971	SEWHO cap design w/jnt(s)		A					
L3972	Offset/lat rocker arm w/ ela		Y					
L3973	SEWHO airplane w/jnt(s) CF		A					
L3974	Mobile arm support supinator		Y					
L3975	SEWHO cap design w/o jnt		A					
L3976	CF		A					
L3977	SEWHO airplane w/o jnts CF		A					
L3978	SEWHO cap desgn w/jnt(s)		A					
L3978	SEWHO airplane w/jnt(s) CF		A					
L3980	Upp ext fx orthosis humeral		A					
L3982	Upp ext fx orthosis rad/ul		A					
L3984	Upp ext fx orthosis wrist		A					
L3995	Sock fracture or equal each		A					
L3999	Upper limb orthosis NOS		A					
L4000	Repl girdle milwaukee orth		A					
L4002	Replace strap, any orthosis		A					
L4010	Replace ilialateral socket br		A					
L4020	Replace quadrat socket brim		A					
L4030	Replace socket brim cust fit		A					
L4040	Replace molded thigh lacer		A					
L4045	Replace non-molded thigh lac		A					
L4050	Replace molded calf lacer		A					
L4055	Replace non-molded calf lace		A					
L4060	Replace high roll cuff		A					
L4070	Replace prox. & dist upright		A					
L4080	Repl met band kato-ato prox		A					
L4090	Repl met band kato-ato calf/		A					
L4100	Repl leath cuff kato-ato prox th		A					
L4110	Repl leath cuff kato-ato calf		A					
L4130	Replace prefribial shell		A					
L4205	Ortho dvc repair per 15 min		A					
L4210	Orth dev repair/repl minor p		A					
L4350	Ankle control orthosi prefab		A					
L4360	Pneumati walking boot prefab		A					
L4370	Pneumatic full leg splint		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L5685	Multi-durometer below knee	A	A					
L5686	Below knee cuff suspension	A	A					
L5688	Socket insert w/o lock tower	A	A					
L5670	Bk molded supracondylar susp	A	A					
L5671	BK/AK locking mechanism	A	A					
L5672	Bk removable medial brim sus	A	A					
L5673	Socket insert w lock mech	A	A					
L5676	Bk knee joints single axis p	A	A					
L5677	Bk knee joints polycentric p	A	A					
L5678	Bk joint covers pair	A	A					
L5679	Socket insert w/o lock mech	A	A					
L5680	Bk thigh lacer non-molded	A	A					
L5681	Intl custom cong/latyp insert	A	A					
L5682	Bk thigh lacer glut/schia m	A	A					
L5683	Initial custom socket insert	A	A					
L5684	Bk fork strap	A	A					
L5685	Below knee sus/seal sleeve	A	A					
L5686	Bk back check	A	A					
L5688	Bk waist belt webbing	A	A					
L5690	Bk waist belt padded and lin	A	A					
L5692	Bk pelvic control belt light	A	A					
L5694	Bk pelvic control belt pad/	A	A					
L5695	Bk sleeve susp neoprene/lequa	A	A					
L5696	Bk/knee disartic pelvic join	A	A					
L5697	Bk/knee disartic pelvic band	A	A					
L5698	Bk/knee disartic silesian ba	A	A					
L5699	Shoulder harness	A	A					
L5700	Replace socket below knee	A	A					
L5701	Replace socket above knee	A	A					
L5702	Replace socket hip	A	A					
L5703	Symes ankle w/o (SACH) foot	A	A					
L5704	Custom shape cover BK	A	A					
L5705	Custom shape cover AK	A	A					
L5706	Custom shape cvr knee disart	A	A					
L5707	Custom shape cvr hip disart	A	A					
L5710	Knee-shin exo sng axi mnl loc	A	A					
L5711	Knee-shin exo mnl lock ultra	A	A					
L5712	Knee-shin exo frict swg & st	A	A					
L5714	Knee-shin exo variable frict	A	A					
L5716	Knee-shin exo mech stance ph	A	A					
L5718	Knee-shin exo frct swg & sta	A	A					
L5722	Knee-shin pneurom swg frct exo	A	A					
L5724	Knee-shin exo fluid swing ph	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L5585	Prep AK ischial open end	A	A					
L5590	Prep AK ischial laminated	A	A					
L5595	Hip disartic sach thermopls	A	A					
L5600	Hip disart sach laminat mold	A	A					
L5610	Above knee hydracadece	A	A					
L5611	AK 4 bar link w/fric swing	A	A					
L5613	AK 4 bar link w/hydraul swig	A	A					
L5614	4-bar link above knee w/swing	A	A					
L5616	AK univ multiplex sys frict	A	A					
L5617	AK/BK self-aligning unit ea	A	A					
L5618	Test socket symes	A	A					
L5620	Test socket below knee	A	A					
L5622	Test socket knee disarticula	A	A					
L5624	Test socket above knee	A	A					
L5626	Test socket hip disarticulat	A	A					
L5628	Test socket hemipelvectomy	A	A					
L5629	Below knee acrylic socket	A	A					
L5630	Syme typ expandabl wall sock	A	A					
L5631	Bk/knee disartic acrylic soc	A	A					
L5632	Symes type pfb brim design s	A	A					
L5634	Symes type poster opening so	A	A					
L5636	Symes type medial opening so	A	A					
L5637	Below knee total contact	A	A					
L5638	Above knee leather socket	A	A					
L5639	Below knee wood socket	A	A					
L5640	Knee disarticulat leather so	A	A					
L5642	Above knee leather socket	A	A					
L5643	Hip flex inner socket ext fr	A	A					
L5644	Above knee wood socket	A	A					
L5645	Bk flex inner socket ext fra	A	A					
L5646	Below knee cushion socket	A	A					
L5647	Below knee suction socket	A	A					
L5648	Above knee cushion socket	A	A					
L5649	Isch containm/narrow m-i so	A	A					
L5650	Tot contact ak/knee disart s	A	A					
L5651	AK flex inner socket ext fra	A	A					
L5652	Suction susp ak/knee disart	A	A					
L5653	Knee disart expand wall sock	A	A					
L5654	Socket insert symes	A	A					
L5655	Socket insert below knee	A	A					
L5656	Socket insert knee articulat	A	A					
L5658	Socket insert above knee	A	A					
L5661	Multi-durometer symes	A	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L5974	Foot single axis ankle/foot		A					
L5975	Combo ankle/foot prosthesis		A					
L5976	Energy storing foot		A					
L5978	Fl prosth multiaxial ank/ft		A					
L5979	Multi-axial ankle/ft prosth		A					
L5980	Flex foot system		A					
L5981	Flex-walk sys low ext prosth		A					
L5982	Exoskeletal axial rotation u		A					
L5984	Exoskeletal axial rotation		A					
L5985	Lwr ext dynamic prosth pylon		A					
L5986	Multi-axial rotation unit		A					
L5987	Shank ft w vert load pylon		A					
L5988	Vertical shock reducing pylon		A					
L5990	User adjustable heel height		A					
L5999	Low extremity prosthesis NOS		A					
L6000	Par hand robin-aids thumb rem		A					
L6010	Hand robin-aids little/ring		A					
L6020	Part hand robin-aids no fing		A					
L6025	Part hand disart myoelectric		A					
L6050	Wrst MLD sock fix ring tri pad		A					
L6055	Wrst mold sock wiexp interfa		A					
L6100	Eib mold sock flex hinge pad		A					
L6110	Elbow mold sock suspension t		A					
L6120	Elbow mold doub split soc ste		A					
L6130	Elbow stump activated lock h		A					
L6205	Elbow mold outside lock hinge		A					
L6250	Elbow inter loc elbow forearm		A					
L6300	Shldr disart int lock elbow		A					
L6310	Shoulder passive restor comp		A					
L6320	Shoulder passive restor cap		A					
L6350	Thoracic intern lock elbow		A					
L6360	Thoracic passive restor comp		A					
L6370	Thoracic passive restor cap		A					
L6382	Postop dsg cast chg wrist/elb		A					
L6384	Postop dsg cast chg elb/dis/		A					
L6386	Postop ea cast chg & realign		A					
L6388	Postop applicat rigid dsg on		A					
L6400	Below elbow prosth liss shap		A					
L6450	Eib disart prosth tiss shap		A					
L6500	Above elbow prosth tiss shap		A					
L6550	Shldr disart prosth tiss shap		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L5726	Knee-shin ext lntis fld swg e		A					
L5728	Knee-shin fluid swg & stance		A					
L5780	Knee-shin pneu/hydra pneu		A					
L5781	Lower limb pros vacuum pump		A					
L5782	HD low limb pros vacuum pump		A					
L5785	Exoskeletal bk ultrait mater		A					
L5790	Exoskeletal ak ultra-light m		A					
L5795	Exoskel hip ultra-light mate		A					
L5810	Endoskel knee-shin mnl lock		A					
L5811	Endo knee-shin mnl lock ultra		A					
L5812	Endo knee-shin frct swg & st		A					
L5814	Endo knee-shin hydral swg ph		A					
L5818	Endo knee-shin polyc mch sta		A					
L5822	Endo knee-shin pneu swg frc		A					
L5824	Endo knee-shin fluid swing p		A					
L5826	Miniature knee joint		A					
L5828	Endo knee-shin fluid swg/sta		A					
L5830	Endo knee-shin pneu/swg		A					
L5840	Multi-axial knee/shin system		A					
L5845	Knee-shin sys stance flexion		A					
L5848	Knee-shin sys hydral stance		A					
L5850	Endo ak/hip knee extens assi		A					
L5855	Mech hip extension assist		A					
L5856	Elec knee-shin swing/stance		A					
L5857	Elec knee-shin swing only		A					
L5858	Stance phase only		A					
L5910	Endo below knee alignable sy		A					
L5920	Endo ak/hip alignable system		A					
L5925	Above knee manual lock		A					
L5930	High activity knee frame		A					
L5940	Endo bk ultra-light material		A					
L5950	Endo ak ultra-light material		A					
L5960	Endo hip ultra-light materia		A					
L5962	Below knee flex cover system		A					
L5964	Above knee flex cover system		A					
L5966	Hip flexible cover system		A					
L5968	Multiaxial ankle w dorsiflex		A					
L5970	Foot external keel sach foot		A					
L5971	SACH foot replacement		A					
L5972	Flexible keel foot		A					
L5973	Ank-foot sys dors-plant flex	NI	A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L6682	Test sock elbw disart/above		A					
L6684	Test sockel shldr disart/tho		A					
L6686	Suction socket		A					
L6687	Frame typ socket bel elbow/w		A					
L6688	Frame typ sock above elb/dis		A					
L6689	Frame typ socket shoulder di		A					
L6690	Frame typ sock interscap-tho		A					
L6691	Removable insert each		A					
L6692	Silicone gel insert or equal		A					
L6693	Lockingelbow forearm cntrlbal		A					
L6694	Elbow socket ins use w/lock		A					
L6695	Elbow socket ins use w/o lock		A					
L6696	Cus elbo skt in for con/atyp		A					
L6697	Cus elbo skt in not con/atyp		A					
L6698	Below/above elbow lock mech		A					
L6703	Term dev, passive hand mitt		A					
L6704	Term dev, sport/rec/work att		A					
L6706	Term dev mech hook vol open		A					
L6707	Term dev mech hook vol close		A					
L6708	Term dev mech hand vol open		A					
L6709	Term dev mech hand vol close		A					
L6711	Ped term dev, hook, vol open		A					
L6712	Ped term dev, hook, vol clos		A					
L6713	Ped term dev, hand, vol open		A					
L6714	Ped term dev, hand, vol clos		A					
L6721	Hook/hand, hvy dty, vol open		A					
L6722	Hook/hand, hvy dty, vol clos		A					
L6805	Term dev modifier wrist unit		A					
L6810	Term dev precision pinch dev		A					
L6881	Term dev auto grasp feature		A					
L6882	Microprocessor control uplmb		A					
L6883	Repic sockt below e/w disa		A					
L6884	Repic sockt above elbow disa		A					
L6885	Repic sockt shldr dis/interc		A					
L6890	Prefab glove for term device		A					
L6895	Custom glove for term device		A					
L6900	Hand restorat thumb/1 finger		A					
L6905	Hand restoration multiple fi		A					
L6910	Hand restoration no fingers		A					
L6915	Hand restoration replacmnt g		A					
L6920	Wrist disarticul switch ctrl		A					
L6925	Wrist disart myoelectronic c		A					
L6930	Below elbow switch control		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L6570	Scap thorac prosth tiss shap		A					
L6580	Wrist/elbow bowden cable mol		A					
L6582	Wrist/elbow bowden cbl dir f		A					
L6584	Elbow fair lead cable molded		A					
L6586	Elbow fair lead cable dir fo		A					
L6588	Shdr fair lead cable molded		A					
L6590	Shdr fair lead cable direct		A					
L6600	Polycentric hinge pair		A					
L6605	Single pivot hinge pair		A					
L6610	Flexible metal hinge pair		A					
L6611	Additional switch, ext power		A					
L6615	Disconnect locking wrist uni		A					
L6616	Disconnect insert locking wr		A					
L6620	Flexion/extension wrist unit		A					
L6621	Flex/ext wrist w/w friction		A					
L6623	Spring-ass rot wrist w/ latch		A					
L6624	Flex/ext/rotation wrist unit		A					
L6625	Rotation wrst w/ cable lock		A					
L6628	Quick disconn hook adapter o		A					
L6629	Lamination collar w/ couplin		A					
L6630	Stainless steel any wrist		A					
L6632	Latex suspension sleeve each		A					
L6635	Lift assist for elbow		A					
L6637	Nudge control elbow lock		A					
L6638	Elec lock on manual pw elbow		A					
L6639	Heavy duty elbow feature	CH	A					
L6640	Shoulder abduction joint pai		A					
L6641	Excursion amplifier pulley t		A					
L6642	Excursion amplifier lever ty		A					
L6645	Shoulder flexion-abduction j		A					
L6646	Multiplo locking shoulder jnt		A					
L6647	Shoulder lock actuator		A					
L6648	Ext pwrd shldr lock/unlock		A					
L6650	Shoulder universal joint		A					
L6655	Standard control cable extra		A					
L6660	Heavy duty control cable		A					
L6665	Teflon or equal cable lining		A					
L6670	Hook to hand cable adapter		A					
L6672	Harness chest/shldr saddle		A					
L6675	Harness figure of 8 sing con		A					
L6676	Harness figure of 8 dual con		A					
L6677	UE triple control harness		A					
L6680	Test sock wrist disart/bel e		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L7900	Male vacuum erection system		A					
L8000	Mastectomy bra		A					
L8001	Breast prosthesis bra & form		A					
L8002	Breast prosthesis bra & form		A					
L8010	Mastectomy sleeve		A					
L8015	Ext breast prosthesis garment		A					
L8020	Mastectomy form		A					
L8030	Breast prosthesis w/o adhesive		A					
L8031	Breast prosthesis w adhesive	NI	A					
L8032	Reusable nipple prosthesis	NI	A					
L8035	Custom breast prosthesis		A					
L8039	Breast prosthesis NOS		A					
L8040	Nasal prosthesis		A					
L8041	Midfacial prosthesis		A					
L8042	Orbital prosthesis		A					
L8043	Upper facial prosthesis		A					
L8044	Hemi-facial prosthesis		A					
L8045	Auricular prosthesis		A					
L8046	Partial facial prosthesis		A					
L8047	Nasal septal prosthesis		A					
L8048	Unspec maxillofacial prosth		A					
L8049	Repair maxillofacial prosth		A					
L8300	Truss single w/ standard pad		A					
L8310	Truss double w/ standard pad		A					
L8320	Truss addition to std pad wa		A					
L8330	Truss add to std pad scrotal		A					
L8400	Sheath below knee		A					
L8410	Sheath above knee		A					
L8415	Sheath upper limb		A					
L8417	Pros sheath/sock w gel cushn		A					
L8420	Prosthetic sock multi ply BK		A					
L8430	Prosthetic sock multi ply AK		A					
L8435	Pros sock multi ply upper lm		A					
L8440	Shrinker below knee		A					
L8460	Shrinker above knee		A					
L8465	Shrinker upper limb		A					
L8470	Pros sock single ply BK		A					
L8480	Pros sock single ply AK		A					
L8485	Pros sock single ply upper l		A					
L8499	Unlisted misc prosthetic ser		A					
L8500	Artificial larynx		A					
L8501	Tracheostomy speaking valve		A					
L8505	Artificial larynx, accessory		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L6935	Below elbow myoelectric ct		A					
L6940	Elbow disarticulation switch		A					
L6945	Elbow disart myoelectric c		A					
L6950	Above elbow switch control		A					
L6955	Above elbow myoelectric ct		A					
L6960	Shldr disartic switch contro		A					
L6965	Shldr disartic myoelectric		A					
L6970	Interscapular-thor switch ct		A					
L6975	Interscap-thor myoelectric		A					
L7007	Adult electric hand		A					
L7008	Pediatric electric hand		A					
L7009	Adult electric hook		A					
L7040	Prehensile actuator		A					
L7045	Pediatric electric hook		A					
L7170	Electronic elbow hosmer swit		A					
L7180	Electronic elbow sequential		A					
L7181	Electronic elbo simultaneous		A					
L7185	Electron elbow adoescent sw		A					
L7186	Electron elbow child switch		A					
L7190	Elbow adolescent myoelectron		A					
L7191	Elbow child myoelectric ct		A					
L7260	Electron wrist rotator otto		A					
L7261	Electron wrist rotator utiah		A					
L7266	Servo control steeper or equ		A					
L7272	Analogous control unb or equa		A					
L7274	Proportional ctt 12 volt utia		A					
L7360	Six volt bat otto book/eq ea		A					
L7362	Battery chgr six volt otto		A					
L7364	Twelve volt battery utah/egu		A					
L7366	Battery chgr 12 volt utiah/e		A					
L7367	Replacemnt lithium ionbatter		A					
L7368	Lithium ion battery charger		A					
L7400	Add UE prost be/wtd, utilite		A					
L7401	Add UE prost a/e utilite mat		A					
L7402	Add UE prost s/d utilite mat		A					
L7403	Add UE prost b/e acrylic		A					
L7404	Add UE prost a/e acrylic		A					
L7405	Add UE prost s/d acrylic		A					
L7499	Upper extremity prosthes NOS		A					
L7500	Prosthetic dvc repair hourly		A					
L7510	Prosthetic device repair rep		A					
L7520	Repair prosthesis per 15 min		A					
L7600	Prosthetic donning sleeve		E					

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L8687	Impit nrostrm pls gen dua rec		N					
L8688	Impit nrostrm pls gen dua non		N					
L8689	External recharg sys intern		A					
L8690	Aud osseo dev, int/ext comp		N					
L8691	Csseintegrated snd proc rpl		A					
L8692	Non-osseointegrated snd proc	NI	E					
L8695	External recharg sys extern		A					
L8699	Prosthetic implant NOS		N					
L9000	O&P supply/accessory/service		N					\$22.69
M0064	Visit for drug monitoring	CH	O3	0607	1.683	\$113.44		
M0075	Cellular therapy		E					
M0076	Prothrotherapy		E					
M0100	Intragastric hypothermia		E					
M0300	IV chelationtherapy		E					
M0301	Fabric wrapping of aneurysm		E					
P2028	Cephalin flocculation test		A					
P2029	Congo red blood test		A					
P2031	Hair analysis		E					
P2033	Blood thymol turbidity		A					
P2038	Blood mucoprotein		A					
P3000	Screen pap by tech w md supv		A					
P3001	Screening pap smear by phys		B					
P7001	Culture bacterial urine		E					
P9010	Whole blood for transfusion		R	0950	3.0598	\$206.25		\$41.25
P9011	Blood split unit		R	0967	1.2965	\$87.39		\$17.48
P9012	Cryoprecipitate each unit		R	0952	0.691	\$46.98		\$9.32
P9016	RBC leukocytes reduced		R	0954	2.7703	\$186.73		\$37.35
P9017	Plasma, I donor frz w/m 8 hr		R	9508	1.1278	\$76.02		\$15.21
P9019	Platelets, each unit		R	0957	0.9882	\$66.61		\$13.33
P9020	Platelet rich plasma unit		R	0958	2.0294	\$136.79		\$27.36
P9021	Washed red blood cells unit		R	0959	2.1027	\$141.73		\$28.35
P9022	Washed red blood cells unit		R	0960	3.6495	\$246.00		\$49.20
P9023	Frozen plasma, pooled, sd		R	0949	0.7589	\$51.15		\$10.23
P9031	Platelets leukocytes reduced		R	1013	1.5541	\$104.76		\$20.96
P9032	Platelets, irradiated		R	9500	2.232	\$150.45		\$30.09
P9033	Platelets leukoreduced irr ad		R	0968	1.9576	\$131.95		\$26.39
P9034	Platelets, pheresis		R	9507	6.9594	\$469.11		\$93.83
P9035	Platelet pheres leukoreduced		R	9501	7.5974	\$512.11		\$102.43
P9036	Platelet pheresis irradiated		R	9502	5.3105	\$357.96		\$71.60
P9037	Plate pheres leukoredu irr ad		R	1019	10.0373	\$676.57		\$135.32
P9038	RBC irradiated		R	9505	3.3499	\$225.80		\$45.16
P9039	RBC deglycerolized		R	9504	5.3988	\$363.91		\$72.79
P9040	RBC leukoreduced irradiated		R	0969	3.635	\$245.02		\$49.01

ADDENDUM B.—FINAL OPPS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
L8507	Trach-esoph voice pros pt in		A					
L8509	Trach-esoph voice pros rmd in		A					
L8510	Voice amplifier		A					
L8511	Indwelling trach insert		A					
L8512	Gel cap for trach voice pros		A					
L8513	Trach pros cleaning device		A					
L8514	Repl trach puncture dilator		A					
L8515	Gel cap app device for trach		A					
L8600	Implant breast silicone/eq		N					
L8603	Collagen imp urinary 2.5 ml		N					
L8604	Dextranomer/hyaluronic acid		N					
L8606	Synthetic implant urinary 1ml		N					
L8609	Artificial cornea		N					
L8610	Ocular implant		N					
L8612	Aqueous shunt prosthesis		N					
L8613	Ossicular implant		N					
L8614	Cochlear device		N					
L8615	Coch implant headset replace		A					
L8616	Coch implant microphone repl		A					
L8617	Coch implant trans coil repl		A					
L8618	Coch implant tran cable repl		A					
L8619	Coch imp ext proc/contr rpic		A					
L8621	Repl zinc air battery		A					
L8622	Repl alkaline battery		A					
L8623	Lith ion batt CID non-eat/d		A					
L8624	Lith ion batt CID, ear level		A					
L8627	CID ext speech process repl	NI	A					
L8628	CID ext controller repl	NI	A					
L8629	CID transmit coil and cable	NI	A					
L8630	Metacarpophalangeal implant		N					
L8631	MCP joint repl 2 pc or more		N					
L8641	Metatarsal joint implant		N					
L8642	Hallux implant		N					
L8658	Interphalangeal joint spacer		N					
L8659	Interphalangeal joint repl		N					
L8670	Vascular graft, synthetic		N					
L8680	Impit neurostim elctr each	CH	N					
L8681	Pt prgm for impit neurostim		A					
L8682	Impit neurostim radiofreq rec		N					
L8683	Radiofreq tsmtr for impit neu		A					
L8684	Radiof tsmtr impit scrf neu		A					
L8685	Impit nrostrm pls gen sng rec		N					
L8686	Impit nrostrm pls gen sng non		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
Q0169	Promethazine HCl 12.5mg oral		N					
Q0170	Promethazine HCl 25 mg oral		N					
Q0171	Chlorpromazine HCl 10mg oral		N					
Q0172	Chlorpromazine HCl 25mg oral		N					
Q0173	Trimethoprimazole HCl 250mg		N					
Q0174	Thiethylperazine maleate 10mg		N					
Q0175	Perphenazine 4mg oral		N					
Q0176	Perphenazine 8mg oral		N					
Q0177	Hydroxyzine pamoate 25mg		N					
Q0178	Hydroxyzine pamoate 50mg		N					
Q0179	Ondansetron hcl 8 mg oral	CH	N					
Q0180	Dolasetron mesylate oral	CH	N					
Q0181	Unspecified oral anti-emetic		E					
Q0480	Driver pneumatic vial, rep		A					
Q0481	Microprscr cu elec vial, rep		A					
Q0482	Microprscr cu combo vial, rep		A					
Q0483	Monitor elec vial, rep		A					
Q0484	Monitor elec or comb vial, rep		A					
Q0485	Monitor cable elec vial, rep		A					
Q0486	Mon cable elec/pneum vial, rep		A					
Q0487	Leads any type vial, rep only		A					
Q0488	Pwr pack base elec vial, rep		A					
Q0489	Pwr pck base elec vial, rep		A					
Q0490	Emr pwr source elec vial, rep		A					
Q0491	Emr pwr source combo vial, rep		A					
Q0492	Emr pwr cbl elec vial, rep		A					
Q0493	Emr pwr cbl combo vial, rep		A					
Q0494	Emr hd bmp elec/combo, rep		A					
Q0495	Charger elec/combo vial, rep		A					
Q0496	Battery elec/combo vial, rep		A					
Q0497	Bat clips elec/combo vial, rep		A					
Q0498	Hoister elec/combo vial, rep		A					
Q0499	Belt/vest elec/combo vial, rep		A					
Q0500	Filters elec/combo vial, rep		A					
Q0501	Shwr cov elec/combo vial, rep		A					
Q0502	Mobility cart pneum vial, rep		A					
Q0503	Battery pneum vial, rep		A					
Q0504	Pwr adpt pneum vial, rep, veh		A					
Q0505	Misc supply/accessory vial		A					
Q0506	Lith-ion bat elec/pneum VAD	NI	A					
Q0510	Dispens fee immunosuppressive		B					
Q0511	Sup fee antiem, antica, immuno		B					
Q0512	Px sup fee anti-can, sub, pres		B					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
P9041	Albumin (human), 5%, 50ml		K	0961		\$16.89		\$3.38
P9043	Plasma, protein fract, 5%, 50ml		R	0956	0.9754	\$65.75		\$13.15
P9044	Cryoprecipitate, reduced plasma		R	1009	1.4035	\$94.60		\$18.92
P9045	Albumin (human), 5%, 250 ml		K	0963		\$60.58		\$12.12
P9046	Albumin (human), 25%, 20 ml		K	0964		\$25.67		\$5.14
P9047	Albumin (human), 25%, 50ml		K	0965		\$62.05		\$12.41
P9048	Plasmaprotein fract, 5%, 250ml		R	0966	1.6017	\$107.96		\$21.60
P9050	Granulocytes, pheresis unit		R	9506	0.6664	\$44.92		\$8.99
P9051	Blood, /fr, cmv-neg		R	1010	2.0076	\$135.32		\$27.07
P9052	Platelets, hla-m, /fr, unit		R	1011	10.929	\$736.68		\$147.34
P9053	Plt, pher, /fr cmv-neg, /fr		R	1020	9.7428	\$656.72		\$131.35
P9054	Blood, /fr, froz/deg/wash		R	1016	1.5373	\$103.62		\$20.73
P9055	Plt, apha/pher, /fr, cmv-neg		R	1017	6.2195	\$419.23		\$83.65
P9056	Blood, /fr, irradiated		R	1018	2.4502	\$165.16		\$33.04
P9057	RBC, frz/deg/wsh, /fr, /frad		R	1021	5.3858	\$363.04		\$72.61
P9058	RBC, /fr, cmv-neg, /frad		R	1022	4.3604	\$293.92		\$58.79
P9059	Plasma, frz between 8-24hour		R	0955	1.1491	\$77.46		\$15.50
P9060	Fr /fr, plasma donor retested		R	9503	1.0664	\$71.88		\$14.38
P9603	One-way allow prorated milles		A					
P9604	One-way allow prorated trip		A					
P9612	Catheterize for urine spec		A					
P9615	Urine specimen collect mult		N					
G0035	Cardiokymography		X	0100	2.6136	\$176.17	\$41.44	\$35.24
G0081	Infusion ther other than che		B					
G0083	Chemo by other than infusion		B					
G0084	Chemotherapy by infusion		B					
G0085	Chemo by both infusion and o		B					
G0091	Obtaining screen pap smear		T	0191	0.132	\$8.90	\$2.09	\$1.78
G0092	Set up port xray equipment		N					
G0111	Wet mounts/ w preparations		A					
G0112	Potassium hydroxide preps		A					
G0113	Pinworm examinations		A					
G0114	Fern test		A					
G0115	Post-coital mucous exam		A					
G0138	Ferumoxylol, non-esrd	NI	G	1297		\$0.82		\$0.16
G0139	Ferumoxylol, esrd use	NI	A					
G0144	Azithromycin dihydrate, oral		E					
G0163	Diphenhydramine HCl 50mg		N					
G0164	Prochlorperazine maleate 5mg		N					
G0165	Prochlorperazine maleate 10mg		N					
G0166	Granisetron hcl 1 mg oral	CH	N					
G0167	Dronabinol 2.5mg oral		N					
G0168	Dronabinol 5mg oral		N					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
Q4028	Cast sup hip spica ped fibril	B	B					
Q4029	Cast sup long leg plaster	B	B					
Q4030	Cast sup long leg fiberglass	B	B					
Q4031	Cast sup lng leg ped plaster	B	B					
Q4032	Cast sup lng leg ped fibril	B	B					
Q4033	Cast sup lng leg cylinder pl	B	B					
Q4034	Cast sup lng leg cylinder fb	B	B					
Q4035	Cast sup lngleg cyindr ped p	B	B					
Q4036	Cast sup lngleg cyindr ped f	B	B					
Q4037	Cast sup shrt leg plaster	B	B					
Q4038	Cast sup shrt leg fiberglass	B	B					
Q4039	Cast sup shrt leg ped plaster	B	B					
Q4040	Cast sup shrt leg ped fibril	B	B					
Q4041	Cast sup lng leg splint plstr	B	B					
Q4042	Cast sup lng leg splint fibril	B	B					
Q4043	Cast sup lng leg splint ped p	B	B					
Q4044	Cast sup lng leg splint ped f	B	B					
Q4045	Cast sup shrt leg splint plstr	B	B					
Q4046	Cast sup shrt leg splint fibril	B	B					
Q4047	Cast sup shrt leg splint ped p	B	B					
Q4048	Cast sup shrt leg splint ped f	B	B					
Q4049	Finger splint, static	B	B					
Q4050	Cast supplies unlisted	B	B					
Q4051	Splint supplies misc	B	B					
Q4074	Ilprost non-comp unit dose	NI	Y					
Q4080	Ilprost non-comp unit dose	CH	D					
Q4081	Epoetin alfa, 100 units ESRD	A	A					
Q4082	Drug/bio NOC part B drug CAP	B	B					
Q4100	Skin substitute, NOS	N	N					
Q4101	Abigrat skin sub	K	K	1240		\$32.16		\$6.44
Q4102	Oasis wound matrix skin sub	K	K	1241		\$4.12		\$0.83
Q4103	Oasis burn matrix skin sub	K	K	1242		\$4.12		\$0.83
Q4104	Integra BMWD skin sub	K	K	1243		\$11.77		\$2.36
Q4105	Integra DRT skin sub	K	K	1244		\$11.77		\$2.36
Q4106	Dermagraft skin sub	K	K	1245		\$39.25		\$7.85
Q4107	Graftjacket skin sub	K	K	1246		\$69.23		\$17.85
Q4108	Integra matrix skin sub	K	K	1247		\$17.98		\$3.60
Q4109	Tissuemend skin sub	N	N					
Q4110	Primatrix skin sub	K	K	1248		\$33.99		\$6.80
Q4111	Gammagraft skin sub	K	K	1252		\$7.13		\$1.43
Q4112	Cymetra allograft	K	K	1249		\$327.47		\$65.50
Q4113	Graftjacket express allograf	K	K	1250		\$327.47		\$65.50
Q4114	Integra flowable wound matri	G	G	1251		\$907.96		\$178.05

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
Q0513	Disp fee inhal drugs/30 days	B	B					
Q0514	Disp fee inhal drugs/90 days	B	B					
Q0515	Sermonelin acetate injection	K	K	3050		\$1.77		\$0.36
Q1003	Nitro category 3	N	E					
Q1004	Nitro category 4	E	E					
Q1005	Nitro category 5	E	E					
Q2004	Bladder calculi irrig sol	CH	K	1293		\$29.28		\$5.86
Q2009	Fosphenytoin inj PE	N	N					
Q2017	Teniposide, 50 mg	K	K	7035		\$319.43		\$63.89
Q2023	Xyntba, inj	CH	D					
Q2024	Bevacizumab injection	CH	D					
Q3001	Brachytherapy Radioelements	B	B					
Q3014	Telehealth facility fee	A	A					
Q3025	IM inj interferon beta 1-a	K	K	9022		\$187.24		\$37.45
Q3026	Subc inj interferon beta-1a	E	E					
Q3031	Collagen skin test	N	N					
Q4001	Cast sup body cast plaster	B	B					
Q4002	Cast sup body cast fiberglass	B	B					
Q4003	Cast sup shoulder cast plstr	B	B					
Q4004	Cast sup shoulder cast fibril	B	B					
Q4005	Cast sup long arm adult plst	B	B					
Q4006	Cast sup long arm adult fibril	B	B					
Q4007	Cast sup long arm ped plaster	B	B					
Q4008	Cast sup long arm ped fibril	B	B					
Q4009	Cast sup shrt arm adult plstr	B	B					
Q4010	Cast sup shrt arm adult fibril	B	B					
Q4011	Cast sup shrt arm ped plaster	B	B					
Q4012	Cast sup shrt arm ped fibril	B	B					
Q4013	Cast sup gauntlet plaster	B	B					
Q4014	Cast sup gauntlet fiberglass	B	B					
Q4015	Cast sup gauntlet ped plaster	B	B					
Q4016	Cast sup gauntlet ped fibril	B	B					
Q4017	Cast sup lng arm splint plst	B	B					
Q4018	Cast sup lng arm splint fibril	B	B					
Q4019	Cast sup lng arm splint ped p	B	B					
Q4020	Cast sup lng arm splint ped f	B	B					
Q4021	Cast sup shrt arm splint plst	B	B					
Q4022	Cast sup shrt arm splint fibril	B	B					
Q4023	Cast sup shrt arm splint ped p	B	B					
Q4024	Cast sup shrt arm splint ped f	B	B					
Q4025	Cast sup hip spica plaster	B	B					
Q4026	Cast sup hip spica fiberglass	B	B					
Q4027	Cast sup hip spica ped plstr	B	B					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
V2102	Singl vlsn sphere 7.12-20.00		A					
V2103	Sphero cylindr 4.00d/12-2.00d		A					
V2104	Sphero cylindr 4.00d/2.12-4d		A					
V2105	Sphero cylindr 4.00d/4.25-6d		A					
V2106	Sphero cylindr 4.00d/6-6.00d		A					
V2107	Sphero cylindr 4.25d/12-2d		A					
V2108	Sphero cylindr 4.25d/2.12-4d		A					
V2109	Sphero cylindr 4.25d/4.25-6d		A					
V2110	Sphero cylindr 4.25d/over 6d		A					
V2111	Sphero cylindr 7.25d/2.25-2.25		A					
V2112	Sphero cylindr 7.25d/2.25-4d		A					
V2113	Sphero cylindr 7.25d/4.25-6d		A					
V2114	Sphero cylindr over 12.00d		A					
V2115	Lens lenticular bifocal		A					
V2118	Lens aniseikonic single		A					
V2121	Lenticular lens, single		A					
V2199	Lens single vision not oth c		A					
V2200	Lens sphere bifoc plano 4.00d		A					
V2201	Lens sphere bifocal 4.12-7.0		A					
V2202	Lens sphere bifocal 7.12-20.		A					
V2203	Lens sphcyl bifocal 4.00d/1		A					
V2204	Lens sphcyl bifocal 4.00d/2.1		A					
V2205	Lens sphcyl bifocal 4.00d/4.2		A					
V2206	Lens sphcyl bifocal 4.00d/ove		A					
V2207	Lens sphcyl bifocal 4.25-7d/.		A					
V2208	Lens sphcyl bifocal 4.25-7/2.		A					
V2209	Lens sphcyl bifocal 4.25-7/4.		A					
V2210	Lens sphcyl bifocal 4.25-7/ov		A					
V2211	Lens sphcyl bifo 7.25-12/25-		A					
V2212	Lens sphcyl bifo 7.25-12/2.2		A					
V2213	Lens sphcyl bifo 7.25-12/4.2		A					
V2214	Lens sphcyl bifocal over 12.		A					
V2215	Lens lenticular bifocal		A					
V2218	Lens aniseikonic bifocal		A					
V2219	Lens bifocal seg width over		A					
V2220	Lens bifocal add over 3.25d		A					
V2221	Lenticular lens, bifocal		A					
V2299	Lens bifocal speciality		A					
V2300	Lens sphere trifocal 4.00d		A					
V2301	Lens sphere trifocal 4.12-7.		A					
V2302	Lens sphere trifocal 7.12-20		A					
V2303	Lens sphcyl trifocal 4.0/12-		A					
V2304	Lens sphcyl trifocal 4.0/2.25		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
Q4115	Allokin skin sub		K	1287		\$9.36		\$1.88
Q4116	Alloderm skin sub		K	1270		\$31.72		\$6.35
Q5001	Hospice in patient home		B					
Q5002	Hospice in assisted living		B					
Q5003	Hospice in L7/noon-skilled NF		B					
Q5004	Hospice in SNF		B					
Q5005	Hospice, inpatient hospital		B					
Q5006	Hospice in hospice facility		B					
Q5007	Hospice in LTCH		B					
Q5008	Hospice in inpatient psych		B					
Q5009	Hospice care, NDS		B					
Q9951	LOCM >= 400 mg/ml iodine, 1ml		N					
Q9953	Inj Fe-based MR contrast,1ml		N					
Q9954	Oral MR contrast, 100 ml		N					
Q9955	Inj peritexane lip micros/ml		N					
Q9956	Inj octafluoropropane mic/ml		N					
Q9957	Inj perflutren lip micros/ml		N					
Q9957	HOCCM <=149 mg/ml iodine, 1ml		N					
Q9958	HOCCM 150-199mg/ml iodine, 1ml		N					
Q9959	HOCCM 200-249mg/ml iodine, 1ml		N					
Q9960	HOCCM 250-299mg/ml iodine, 1ml		N					
Q9961	HOCCM 300-349mg/ml iodine, 1ml		N					
Q9962	HOCCM 350-399mg/ml iodine, 1ml		N					
Q9963	HOCCM >= 400mg/ml iodine, 1ml		N					
Q9964	LOCM 100-199mg/ml iodine, 1ml		N					
Q9965	LOCM 200-299mg/ml iodine, 1ml		N					
Q9966	LOCM 300-399mg/ml iodine, 1ml		N					
Q9967	LOCM >= 400mg/ml iodine, 1ml		N					
Q9968	Visualization adjunct		N					
R0070	Transport portable x-ray	NI	K	1288		\$4.11		\$0.83
R0075	Transport port x-ray multipl		B					
R0076	Transport portable EKG		B					
V2020	Vision svcs frames purchases		A					
V2025	Eyeglasses delux frames		E					
V2100	Lens spher single plano 4.00		A					
V2101	Single vlsn sphere 4.12-7.00		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
V2629	Prosthetic eye other type		A					
V2630	Anter chamber intraocul lens		N					
V2631	Iris support intraocul lens		N					
V2632	Post chmbr intraocul lens		N					
V2700	Balance lens		A					
V2702	Deluxe lens feature		E					
V2710	Glass/plastic slab off prism		A					
V2715	Prism lenses		A					
V2718	Fresnell prism press-on lens		A					
V2730	Special base curve		A					
V2744	Tint photochromatic lens/és		A					
V2745	Tint, any color/solid/grad		A					
V2750	Anti-reflective coating		A					
V2755	UV lenses		A					
V2756	Eye glass case		E					
V2760	Scratch resistant coating		A					
V2761	Mirror coating		B					
V2762	Polarization, any lens		A					
V2770	Occluder lens/és		A					
V2780	Oversize lens/és		A					
V2781	Progressive lens per lens		B					
V2782	Lens, 1.54-1.65 p/1.60-1.79g		A					
V2783	Lens, >= 1.66 p/>=1.80 g		A					
V2784	Lens polycarb or equal		A					
V2785	Corneal tissue processing		F					
V2786	Occupational multifocal lens		A					
V2787	Astigmatism-correct function		E					
V2788	Presbyopia-correct function		E					
V2790	Amniotic membrane		N					
V2797	Vis item/svc in other code		A					
V2799	Miscellaneous vision service		A					
V5008	Hearing screening		E					
V5010	Assessment for hearing aid		E					
V5011	Hearing aid fitting/checking		E					
V5014	Hearing aid repair/modifying		E					
V5020	Conformity evaluation		E					
V5030	Body-worn hearing aid air		E					
V5040	Body-worn hearing aid bone		E					
V5050	Hearing aid monaural in ear		E					
V5060	Behind ear hearing aid		E					
V5070	Glasses air conduction		E					
V5080	Glasses bone conduction		E					
V5090	Hearing aid dispensing fee		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
V2305	Lens sphcy trifocal 4.0/4.25		A					
V2306	Lens sphcy trifocal 4.00/5-6		A					
V2307	Lens sphcy trifocal 4.25-7/.		A					
V2308	Lens sphc trifocal 4.25-7/2.		A					
V2309	Lens sphc trifocal 4.25-7/4.		A					
V2310	Lens sphc trifocal 4.25-7/5-6		A					
V2311	Lens sphc trifo 7.25-12/25-		A					
V2312	Lens sphc trifo 7.25-12/2.25		A					
V2313	Lens sphc trifo 7.25-12/4.25		A					
V2314	Lens sphcy trifocal over 12		A					
V2315	Lens lenticular trifocal		A					
V2318	Lens aniseikonic trifocal		A					
V2319	Lens trifocal seg width > 28		A					
V2320	Lens trifocal add over 3.25d		A					
V2321	Lenticular lens, trifocal		A					
V2399	Lens trifocal speciality		A					
V2410	Lens variab asphericity sing		A					
V2430	Lens variable asphericity bi		A					
V2499	Variable asphericity lens		A					
V2500	Contact lens pmma spherical		A					
V2501	Contact lens pmma-toric/prism		A					
V2502	Contact lens pmma bifocal		A					
V2503	Contact lens pmma color vision		A					
V2510	Contact gas permeable spherical		A					
V2511	Contact toric prism ballast		A					
V2512	Contact lens gas permibi bifocl		A					
V2513	Contact lens extended wear		A					
V2520	Contact lens hydrophilic		A					
V2521	Contact lens hydrophilic tonic		A					
V2522	Contact lens hydrophilic bifocl		A					
V2523	Contact lens hydrophil extend		A					
V2530	Contact lens gas impermeable		A					
V2531	Contact lens gas permeable		A					
V2599	Contact lenses/és other type		A					
V2600	Hand held low vision aids		A					
V2610	Single lens spectacle mount		A					
V2615	Telescop/othr compound lens		A					
V2623	Plastic eye prosthr custom		A					
V2624	Polishing artificial eye		A					
V2625	Enlargemnt of eye prosthesis		A					
V2626	Reduction of eye prosthesis		A					
V2627	Scleral cover shell		A					
V2628	Fabrication & fitting		A					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
V5268	ALD Telephone Amplifier		E					
V5269	Alerting device, any type		E					
V5270	ALD, TV amplifier, any type		E					
V5271	ALD, TV caption decoder		E					
V5272	Tdd		E					
V5273	ALD for cochlear implant		E					
V5274	ALD unspecified		E					
V5275	Ear Impression		E					
V5298	Hearing aid noc		E					
V5299	Hearing services		B					
V5396	Repair communication device		E					
V5362	Speech screening		E					
V5363	Language screening		E					
V5364	Dysphagia screening		E					

ADDENDUM B.—FINAL OPPTS PAYMENT BY HCPCS CODE FOR CY 2010

HCPCS Code	Short Descriptor	CI	SI	APC	Relative Weight	Payment Rate	National Unadjusted Copayment	Minimum Unadjusted Copayment
V5095	Implant mid ear hearing pros		E					
V5100	Body-worn bilat hearing aid		E					
V5110	Hearing aid dispensing fee		E					
V5120	Body-worn binaur hearing aid		E					
V5130	In ear binaural hearing aid		E					
V5140	Behind ear binaur hearing ai		E					
V5150	Glasses binaural hearing aid		E					
V5160	Dispensing fee binaural		E					
V5170	Within ear cros hearing aid		E					
V5180	Behind ear cros hearing aid		E					
V5190	Glasses cros hearing aid		E					
V5200	Cros hearing aid dispens fee		E					
V5210	In ear bicros hearing aid		E					
V5220	Behind ear bicros hearing ai		E					
V5230	Glasses bicros hearing aid		E					
V5240	Dispensing fee bicros		E					
V5241	Dispensing fee, monaural		E					
V5242	Hearing aid, monaural, cic		E					
V5243	Hearing aid, monaural, itc		E					
V5244	Hearing aid, prog, mon, cic		E					
V5245	Hearing aid, prog, mon, itc		E					
V5246	Hearing aid, prog, mon, ite		E					
V5247	Hearing aid, prog, mon, ble		E					
V5248	Hearing aid, binaural, cic		E					
V5249	Hearing aid, binaural, itc		E					
V5250	Hearing aid, prog, bin, cic		E					
V5251	Hearing aid, prog, bin, itc		E					
V5252	Hearing aid, prog, bin, ite		E					
V5253	Hearing aid, prog, bin, ble		E					
V5254	Hearing aid, digit, mon, cic		E					
V5255	Hearing aid, digit, mon, itc		E					
V5256	Hearing aid, digit, mon, ite		E					
V5257	Hearing aid, digit, mon, ble		E					
V5258	Hearing aid, digit, bin, cic		E					
V5259	Hearing aid, digit, bin, itc		E					
V5260	Hearing aid, digit, bin, ite		E					
V5261	Hearing aid, digit, bin, ble		E					
V5262	Hearing aid, disp, monaural		E					
V5263	Hearing aid, disp, binaural		E					
V5264	Ear mold/insert		E					
V5265	Ear mold/insert, disp		E					
V5266	Battery for hearing device		E					
V5267	Hearing aid supply/accessory		E					

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
70170	X-ray exam of tear duct		N1		
70190	X-ray exam of eye sockets		Z3		\$18.18
70200	X-ray exam of eye sockets	CH	Z3		\$21.87
70210	X-ray exam of sinuses		Z3		\$15.62
70220	X-ray exam of sinuses		Z3		\$19.32
70240	X-ray exam, pituitary saddle		Z3		\$17.61
70250	X-ray exam of skull		Z3		\$22.16
70260	X-ray exam of skull		Z3		\$6.25
70300	X-ray exam of teeth		Z2		\$17.90
70310	X-ray exam of teeth		Z2	0.4275	\$17.90
70320	Full mouth x-ray of teeth		Z2	0.4275	\$17.90
70328	X-ray exam of jaw joint		Z3		\$15.62
70330	X-ray exam of jaw joints	CH	Z3		\$25.57
70332	X-ray exam of jaw joint		N1		
70336	Magnetic image, jaw joint		Z2	4.961	\$207.73
70350	X-ray head for orthodontia		Z3		\$9.52
70355	Panoramic x-ray of jaws		Z3		\$6.24
70360	X-ray exam of neck		Z3		\$13.63
70370	Throat x-ray & fluoroscopy	CH	Z3		\$44.88
70371	Speech evaluation, complex	CH	Z3		\$39.48
70373	Contrast x-ray of larynx		N1		\$21.02
70380	X-ray exam of salivary gland		Z3		
70390	X-ray exam of salivary duct		N1		
70450	Ct head/brain w/o dye		Z2	2.7687	\$115.93
70460	Ct head/brain w/dye	CH	Z3		\$152.26
70470	Ct head/brain w/o & w/dye	CH	Z3		\$187.48
70480	Ct orbit/ear/fossa w/o dye		Z2	2.7687	\$115.93
70481	Ct orbit/ear/fossa w/dye		Z2	4.2158	\$176.53
70482	Ct orbit/ear/fossa w/o&w/dye		Z2	4.7337	\$198.21
70486	Ct maxillofacial w/o dye		Z2	2.7687	\$115.93
70487	Ct maxillofacial w/dye		Z2	4.2158	\$176.53
70488	Ct maxillofacial w/o & w/dye		Z2	4.7337	\$198.21
70490	Ct soft tissue neck w/o dye		Z2	2.7687	\$115.93

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Note 2: Payment indicators for radiology services (Z2, Z3) are based on a comparison of the final rates according to the ASC standard reresetting methodology and the MPFS. Under current law, the MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
0042T	Ct perfusion w/contrast, cbf		N1		
0067T	Ct colonography/dx	CH	D5		\$250.33
0073T	Delivery, comp limit		Z2	5.9784	
0126T	Chd risk int study		N1		
0144T	CT heart w/o dye, liquid calc		D5		
0145T	CT heart w/o dye funct		D5		
0146T	CCTA w/o dye		D5		
0147T	CCTA w/o, quan calcium		D5		
0148T	CCTA w/o, strxr		D5		
0149T	CCTA w/o, strxr quan calc		D5		
0150T	CCTA w/o, disease strxr		D5		
0151T	CT heart funct add-on		D5		
0159T	Card breast mri		N1		
0174T	Card cxr with interp		N1		
0175T	Card cxr remote		N1		
0182T	Hdr elect brachytherapy		Z2	11.0358	\$462.10
0185T	Comptr probability analysis		N1		
70010	Contrast x-ray of brain		N1		
70015	Contrast x-ray of brain		N1		
70030	X-ray eye for foreign body		Z3		\$14.49
70100	X-ray exam of jaw		Z3		\$16.19
70110	X-ray exam of jaw		Z3		\$19.60
70120	X-ray exam of mastoids		Z3		\$18.18
70130	X-ray exam of mastoids		Z2	0.6373	\$26.69
70134	X-ray exam of middle ear		Z3		\$21.30
70140	X-ray exam of facial bones		Z3		\$14.77
70150	X-ray exam of facial bones	CH	Z3		\$21.59
70160	X-ray exam of nasal bones		Z3		\$17.33

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPSC Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
71100	X-ray exam of ribs		Z3	23	\$15.62
71101	X-ray exam of ribs/chest		Z3	23	\$19.03
71110	X-ray exam of ribs		Z3	23	\$19.88
71111	X-ray exam of ribs/chest		Z3	23	\$26.42
71120	X-ray exam of breastbone		Z3	23	\$16.19
71130	X-ray exam of breastbone		Z3	23	\$19.32
71250	Ct thorax w/o dye		Z2	2,7687	\$115.93
71260	Ct thorax w/dye		Z2	4,2158	\$176.53
71270	Ct thorax w/o & w/dye		Z2	4,7337	\$198.21
71275	Ct angiography, chest		Z2	4,8324	\$202.35
71550	Mri chest w/o dye		Z2	4,961	\$207.73
71551	Mri chest w/dye		Z2	6,0177	\$251.98
71552	Mri chest w/o & w/dye		Z2	7,5993	\$318.21
72010	X-ray exam of spine	CH	Z3	23	\$35.79
72020	X-ray exam of neck spine		Z3	23	\$11.65
72040	X-ray exam of neck spine		Z3	23	\$19.03
72050	X-ray exam of neck spine		Z3	23	\$26.70
72052	X-ray exam of neck spine	CH	Z3	23	\$34.94
72069	X-ray exam of trunk spine		Z3	23	\$17.90
72070	X-ray exam of thoracic spine		Z3	23	\$16.48
72072	X-ray exam of thoracic spine		Z3	23	\$19.60
72074	X-ray exam of thoracic spine	CH	Z3	23	\$24.43
72080	X-ray exam of trunk spine		Z3	23	\$17.61
72090	X-ray exam of trunk spine		Z3	23	\$24.15
72100	X-ray exam of lower spine		Z3	23	\$20.45
72110	X-ray exam of lower spine	CH	Z3	23	\$28.41
72120	X-ray exam of lower spine		Z3	23	\$39.48
72125	Ct neck spine w/o dye		Z2	2,7687	\$115.93
72126	Ct neck spine w/dye		Z2	4,2158	\$176.53
72127	Ct neck spine w/o & w/dye		Z2	4,7337	\$198.21
72128	Ct chest spine w/o dye		Z2	2,7687	\$115.93
72129	Ct chest spine w/dye		Z2	4,2158	\$176.53

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPSC Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
70491	Ct soft tissue neck w/dye		Z2	4,2158	\$176.53
70492	Ct soft tissue neck w/o & w/dye		Z2	4,7337	\$198.21
70496	Ct angiography, head		Z2	4,8324	\$202.35
70498	Ct angiography, neck		Z2	4,8324	\$202.35
70540	Mri orbit/face/neck w/o dye		Z2	4,961	\$207.73
70542	Mri orbit/face/neck w/dye		Z2	6,0177	\$251.98
70543	Mri orbit/face/neck w/o & w/dye		Z2	7,5993	\$318.21
70545	Mri angiography head w/dye		Z2	4,961	\$207.73
70546	Mri angiography head w/o & w/dye		Z2	6,0177	\$251.98
70547	Mri angiography neck w/o dye		Z2	4,961	\$207.73
70549	Mri angiography neck w/dye		Z2	6,0177	\$251.98
70551	Mri brain w/o dye		Z2	7,5993	\$318.21
70552	Mri brain w/dye		Z2	4,961	\$207.73
70553	Mri brain w/o & w/dye		Z2	6,0177	\$251.98
70554	Fmri brain by tech		Z2	7,5993	\$318.21
70555	Fmri brain by phys/psych		Z2	4,961	\$207.73
70557	Mri brain w/o dye		Z2	4,961	\$207.73
70558	Mri brain w/dye		Z2	6,0177	\$251.98
70559	Mri brain w/o & w/dye		Z2	7,5993	\$318.21
71010	Chest x-ray		Z3	23	\$10.79
71015	Chest x-ray		Z3	23	\$14.20
71020	Chest x-ray		Z3	23	\$14.77
71021	Chest x-ray		Z3	23	\$17.90
71022	Chest x-ray	CH	Z3	23	\$22.44
71030	Chest x-ray and fluoroscopy	CH	Z3	23	\$22.44
71034	Chest x-ray and fluoroscopy	CH	Z3	23	\$35.79
71035	Chest x-ray		Z3	23	\$48.57
71040	Contrast x-ray of bronchi		N1	23	\$18.75
71060	Contrast x-ray of bronchi		N1	23	\$18.75
71090	X-ray & pacemaker insertion		N1	23	\$18.75

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
72295	X-ray of lower spine disk		N1		
73000	X-ray exam of collar bone		Z3		\$14.49
73010	X-ray exam of shoulder blade		Z3		\$14.77
73020	X-ray exam of shoulder		Z3		\$11.65
73030	X-ray exam of shoulder		Z3		\$14.77
73040	Contrast x-ray of shoulder		N1		
73050	X-ray exam of shoulders		Z3		\$19.03
73060	X-ray exam of humerus		Z3		\$14.77
73070	X-ray exam of elbow		Z3		\$14.49
73080	X-ray exam of elbow		Z3		\$19.03
73085	Contrast x-ray of elbow		N1		
73090	X-ray exam of forearm		Z3		\$14.20
73092	X-ray exam of arm, infant		Z3		\$15.06
73100	X-ray exam of wrist		Z3		\$15.34
73110	X-ray exam of wrist		Z3		\$19.32
73115	Contrast x-ray of wrist		N1		
73120	X-ray exam of hand		Z3		\$13.92
73130	X-ray exam of hand		Z3		\$16.76
73140	X-ray exam of finger(s)		Z3		\$17.04
73200	Ct upper extremity w/o dye		Z2	2.7687	\$15.93
73201	Ct upper extremity w/dye		Z2	4.2158	\$76.53
73202	Ct upper extremity w/o&w/dye		Z2	4.7337	\$198.21
73206	Ct angio upr extrem w/o&w/dye		Z2	4.8324	\$202.35
73218	Mri upper extremity w/o dye		Z2	4.961	\$207.73
73219	Mri upper extremity w/dye		Z2	6.0177	\$251.98
73220	Mri upper extremity w/o&w/dye		Z2	7.5993	\$318.21
73221	Mri joint upr extrem w/o dye		Z2	4.961	\$207.73
73222	Mri joint upr extrem w/dye		Z2	6.0177	\$251.98
73223	Mri joint upr extr w/o&w/dye		Z2	7.5993	\$318.21
73500	X-ray exam of hip		Z3		\$12.21
73510	X-ray exam of hip		Z3		\$19.03
73520	X-ray exam of hips		Z3		\$19.32
73525	Contrast x-ray of hip		N1		

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
72130	Ct chest spine w/o & w/dye		Z2	4.7337	\$198.21
72131	Ct lumbar spine w/o dye		Z2	2.7687	\$115.93
72132	Ct lumbar spine w/dye		Z2	4.2158	\$176.53
72133	Ct lumbar spine w/o & w/dye		Z2	4.7337	\$198.21
72141	Mri neck spine w/o dye		Z2	4.961	\$207.73
72142	Mri neck spine w/dye		Z2	6.0177	\$251.98
72147	Mri chest spine w/o dye		Z2	4.961	\$207.73
72148	Mri chest spine w/dye		Z2	6.0177	\$251.98
72149	Mri lumbar spine w/o dye		Z2	4.961	\$207.73
72156	Mri lumbar spine w/dye		Z2	6.0177	\$251.98
72157	Mri neck spine w/o & w/dye		Z2	7.5993	\$318.21
72158	Mri lumbar spine w/o & w/dye		Z2	7.5993	\$318.21
72170	X-ray exam of pelvis		Z3		\$12.78
72190	X-ray exam of pelvis	CH	Z3		\$21.30
72191	Ct angiograph pelv w/o&w/dye		Z2	4.8324	\$202.35
72192	Ct pelvis w/o dye		Z2	2.7687	\$115.93
72193	Ct pelvis w/dye		Z2	4.2158	\$176.53
72194	Ct pelvis w/o & w/dye		Z2	4.7337	\$198.21
72195	Mri pelvis w/o dye		Z2	4.961	\$207.73
72196	Mri pelvis w/dye		Z2	6.0177	\$251.98
72197	Mri pelvis w/o & w/dye		Z2	7.5993	\$318.21
72200	X-ray exam sacroiliac joints		Z3		\$15.06
72202	X-ray exam sacroiliac joints		Z3		\$18.18
72220	X-ray exam of tailbone		Z3		\$15.06
72240	Contrast x-ray of neck spine		N1		
72255	Contrast x-ray, thorax spine		N1		
72270	Contrast x-ray, spine		N1		
72275	Epidurography		N1		
72285	X-ray ct spine disk		N1		
72291	Perq verte/sacrospisty, fluor		N1		
72292	Perq verte/sacrospisty, ct		N1		

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
74160	Ct abdomen w/dye		Z2	4.2158	\$176.53
74170	Ct abdomen w/o & w/dye		Z2	4.7337	\$198.21
74175	Ct angio abdom w/o & w/dye		Z2	4.8324	\$202.35
74181	Mri abdomen w/o dye		Z2	4.961	\$207.73
74182	Mri abdomen w/dye		Z2	6.0177	\$251.98
74183	Mri abdomen w/o dye		Z2	7.5993	\$318.21
74190	X-ray exam of peritoneum		N1		
74210	Contrast x-ray exam of throat	CH	Z3		\$43.18
74220	Contrast x-ray, esophagus	CH	Z3		\$48.01
74230	Cine/vid x-ray, throa/esoph	CH	Z3		\$46.89
74235	Remove esophagus obstruction		N1		
74240	X-ray exam, upper gi tract		Z2	1.2423	\$52.02
74241	X-ray exam, upper gi tract		Z2	1.2423	\$52.02
74245	X-ray exam, upper gi tract		Z2	2.0092	\$84.13
74246	Contrast x-ray uppr gi tract		Z2	1.2423	\$52.02
74247	Contrast x-ray uppr gi tract		Z2	1.2423	\$52.02
74249	Contrast x-ray uppr gi tract		Z2	2.0092	\$84.13
74250	X-ray exam of small bowel		Z2	1.2423	\$52.02
74251	X-ray exam of small bowel		Z2	2.0092	\$84.13
74260	X-ray exam of small bowel		Z2	1.2423	\$52.02
74261	Ct colonography, w/o dye	NI	Z2	2.7687	\$115.93
74262	Ct colonography, w/dye	NI	Z2	4.2158	\$176.53
74270	Contrast x-ray exam of colon		Z2	1.2423	\$52.02
74280	Contrast x-ray exam of colon		Z2	2.0092	\$84.13
74283	Contrast x-ray exam of colon		Z2	1.2423	\$52.02
74290	Contrast x-ray, gallbladder		Z3		\$36.93
74291	Contrast x-rays, gallbladder		Z3		\$36.84
74300	X-ray bile ducts/pancreas		N1		
74301	X-rays at surgery add-on		N1		
74305	X-ray bile ducts/pancreas		N1		
74320	Contrast x-ray of bile ducts		N1		
74327	X-ray bile stone removal		N1		
74328	X-ray bile duct endoscopy		N1		

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73530	X-ray exam of hip		N1		
73540	X-ray exam of pelvis & hips		Z3		\$20.45
73542	X-ray exam, sacroiliac joint		N1		
73550	X-ray exam of thigh		Z3		\$14.20
73560	X-ray exam of knee, 1 or 2		Z3		\$14.77
73562	X-ray exam of knee, 3		Z3		\$18.46
73564	X-ray exam, knee, 4 or more		Z3		\$21.30
73565	X-ray exam of knees		Z3		\$16.48
73580	Contrast x-ray of knee joint		N1		
73590	X-ray exam of lower leg		Z3		\$13.63
73592	X-ray exam of leg, infant		Z3		\$15.34
73600	X-ray exam of ankle		Z3		\$14.20
73610	X-ray exam of ankle		Z3		\$16.76
73615	Contrast x-ray of ankle		N1		
73620	X-ray exam of ankle		Z3		\$13.63
73630	X-ray exam of foot		Z3		\$16.48
73650	X-ray exam of heel		Z3		\$13.92
73660	X-ray exam of toes		Z3		\$15.91
73700	Ct lower extremity w/o dye		Z2	2.7687	\$115.93
73701	Ct lower extremity w/dye		Z2	4.2158	\$176.53
73702	Ct lwr extremity w/o&w/dye		Z2	4.7337	\$198.21
73706	Ct angio lwr extr w/o&w/dye		Z2	4.8324	\$202.35
73718	Mri lower extremity w/o dye		Z2	4.961	\$207.73
73719	Mri lower extremity w/dye		Z2	6.0177	\$251.98
73720	Mri lwr extremity w/o&w/dye		Z2	7.5993	\$318.21
73721	Mri jnt of lwr extre w/o dye		Z2	4.961	\$207.73
73722	Mri joint of lwr extr w/dye		Z2	6.0177	\$251.98
73723	Mri joint lwr extr w/o&w/dye		Z2	7.5993	\$318.21
74000	X-ray exam of abdomen		Z3		\$11.93
74010	X-ray exam of abdomen		Z3		\$19.03
74020	X-ray exam of abdomen		Z3		\$19.32
74022	X-ray exam series, abdomen		Z3		\$23.58
74150	Ct abdomen w/o dye		Z2	2.7687	\$115.93

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
75600	Contrast x-ray exam of aorta		N1		
75605	Contrast x-ray exam of aorta		N1		
75625	Contrast x-ray exam of aorta		N1		
75630	X-ray aorta, leg arteries		N1		
75635	Ct angio abdominal arteries		N1		
75658	Attery x-rays, arm		N1		
75662	Attery x-rays, head & neck		N1		
75665	Attery x-rays, head & neck		N1		
75676	Attery x-rays, head & neck		N1		
75680	Attery x-rays, neck		N1		
75685	Attery x-rays, spine		N1		
75705	Attery x-rays, spine		N1		
75710	Attery x-rays, arm/leg		N1		
75716	Attery x-rays, arms/legs		N1		
75722	Attery x-rays, kidney		N1		
75724	Attery x-rays, kidneys		N1		
75726	Attery x-rays, abdomen		N1		
75731	Attery x-rays, adrenal gland		N1		
75733	Attery x-rays, adrenals		N1		
75736	Attery x-rays, pelvis		N1		
75741	Attery x-rays, lung		N1		
75743	Attery x-rays, lungs		N1		
75746	Attery x-rays, lung		N1		
75756	Attery x-rays, chest		N1		
75774	Attery x-ray, each vessel		N1		
75790	Visualize A-V shunt	CH	D5		
75791	Av dialysis shunt imaging	NI	N1		
75801	Lymph vessel x-ray, arm/leg		N1		
75803	Lymph vessel x-ray, arms/legs		N1		
75805	Lymph vessel x-ray, trunk		N1		

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HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
74329	X-ray for pancreas endoscopy		N1		
74330	X-ray bile/panc endoscopy		N1		
74340	X-ray guide for GI tube		N1		
74355	X-ray guide, intestinal tube		N1		
74360	X-ray guide, GI dilation		N1		
74363	X-ray, bile duct dilation		N1		
74400	Contrst x-ray, urinary tract		Z3		\$62.78
74410	Contrst x-ray, urinary tract		Z3		\$66.19
74415	Contrst x-ray, urinary tract		Z3		\$90.39
74420	Contrst x-ray, urinary tract		Z2	2.4358	\$101.99
74425	Contrst x-ray, urinary tract		N1		
74430	Contrast x-ray, bladder		N1		
74440	X-ray, male genital tract		N1		
74445	X-ray exam of penis		N1		
74450	X-ray, uretra/bladder		N1		
74455	X-ray, uretra/bladder		N1		
74470	X-ray exam of kidney lesion		N1		
74475	X-ray control, cath insert		N1		
74480	X-ray control, cath insert		N1		
74485	X-ray guide, GU dilation		N1		
74710	X-ray measurement of pelvis		Z3		\$17.61
74740	X-ray, female genital tract		N1		
74742	X-ray, fallopian tube		N1		
74775	X-ray exam of perineum		Z2	2.4358	\$101.99
75557	Cardiac mri for morph		Z2	4.961	\$207.73
75559	Cardiac mri w/stress img		Z2	4.961	\$207.73
75561	Cardiac mri for morph w/dye		Z2	7.5993	\$318.21
75563	Card mri w/stress img & dye		Z2	7.5993	\$318.21
75565	Card mri vel flw map add-on	NI	N1		
75571	Ct hrt w/cv w/ca test	NI	Z2	0.6403	\$26.81
75572	Ct hrt w/3d image	NI	Z3		\$40.62
75573	Ct hrt w/3d image, congen	NI	Z3		\$48.57
75574	Ct angio hrt w/3d image	NI	Z2	3.82	\$159.95

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
75968	Repair artery blockage, each		N1		
75970	Vascular biopsy		N1		
75978	Repair venous blockage		N1		
75980	Contrast xray exam bile duct		N1		
75982	Contrast xray exam bile duct		N1		
75984	Xray control catheter change		N1		
75989	Abscess drainage under x-ray		N1		
75992	Atherectomy, x-ray exam		N1		
75993	Atherectomy, x-ray exam		N1		
75994	Atherectomy, x-ray exam		N1		
75995	Atherectomy, x-ray exam		N1		
75996	Atherectomy, x-ray exam		N1		
76000	Fluoroscope examination		N1		
76001	Fluoroscope exam, extensive		N1		\$13.63
76010	X-ray, nose to rectum		Z3		
76080	X-ray exam of fistula		N1		
76098	X-ray exam, breast specimen	CH	N1		
76100	X-ray exam of body section		Z2	1.0678	\$44.71
76101	Complex body section x-ray	CH	Z3		\$111.07
76102	Complex body section x-rays		Z2	2.9641	\$124.95
76120	Cine/video x-rays	CH	Z3		\$42.04
76125	Cine/video x-rays add-on		N1		
76150	X-ray exam, dty process		Z3		\$14.49
76350	Special x-ray contrast study		N1		
76376	3d render w/o postprocess		N1		
76377	3d rendering w/postprocess		N1		
76380	CAT scan follow-up study		Z2	1.5586	\$65.26
76496	Fluoroscopic procedure		Z2	1.2143	\$50.85
76497	CI procedure		Z2	1.5586	\$65.26
76498	Mfi procedure		Z2	4.961	\$207.73
76506	Radiographic procedure		Z2	0.6373	\$26.69
76509	Echo exam of head		Z2	0.8866	\$37.12
76510	Ophthalm us, b & quant a		Z3		\$52.55

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for Which Payment is Packaged)

HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
75907	Lymph vessel x-ray, trunk		N1		
75909	Nonvascular shunt, x-ray		N1		
75810	Vein x-ray, spleen/liver		N1		
75820	Vein x-ray, arm/leg		N1		
75822	Vein x-ray, arms/legs		N1		
75825	Vein x-ray, trunk		N1		
75827	Vein x-ray, chest		N1		
75831	Vein x-ray, kidney		N1		
75833	Vein x-ray, kidneys		N1		
75840	Vein x-ray, adrenal gland		N1		
75842	Vein x-ray, adrenal glands		N1		
75860	Vein x-ray, neck		N1		
75870	Vein x-ray, skull		N1		
75872	Vein x-ray, skull		N1		
75880	Vein x-ray, eye socket		N1		
75885	Vein x-ray, liver		N1		
75887	Vein x-ray, liver		N1		
75889	Vein x-ray, liver		N1		
75891	Vein x-ray, liver		N1		
75893	Venous sampling by catheter		N1		
75894	X-rays, transcath therapy		N1		
75896	X-rays, transcath therapy		N1		
75898	Follow-up angiography		N1		
75901	Remove cva device obstruct		N1		
75902	Remove cva lumen obstruct		N1		
75940	X-ray placement, vein filter		N1		
75945	Intravascular us		N1		
75946	Intravascular us add-on		N1		
75960	Transcath iv silent rsk		N1		
75961	Retrieval, broken catheter		N1		
75962	Repair arterial blockage		N1		
75964	Repair artery blockage, each		N1		
75966	Repair arterial blockage		N1		

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HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
76827	Echo exam of fetal heart	CH	Z3		\$29.26
76828	Echo exam of fetal heart		Z3	1.3609	\$16.76
76830	Transvaginal us, non-ob		Z3	1.3609	\$57.82
76831	Echo exam, uterus		Z3	1.3609	\$64.20
76856	Us exam, pelvic, complete		Z2	1.3609	\$57.82
76857	Us exam, pelvic, limited		Z2	0.8866	\$37.12
76870	Us exam, scrotum		Z2	1.3609	\$57.82
76872	Us, transrectal		Z2	1.3609	\$57.82
76873	Echograp trans r, pros study		Z2	1.3609	\$57.82
76880	Us exam, extremity		Z2	1.3609	\$57.82
76885	Us exam infant hips, dynamic		Z2	0.8866	\$37.12
76886	Us exam infant hips, static		Z2	0.8866	\$37.12
76930	Echo guide, cardiocentesis		N1		
76932	Echo guide for heart biopsy		N1		
76936	Echo guide for artery repair		Z2	1.5379	\$64.40
76937	Us guide, vascular access		N1		
76940	Us guide, tissue ablation		N1		
76941	Echo guide for transfusion		N1		
76942	Echo guide for biopsy		N1		
76945	Echo guide, villus sampling		N1		
76946	Echo guide for amniocentesis		N1		
76948	Echo guide, ova aspiration		N1		
76950	Echo guidance radiotherapy		N1		
76965	Echo guidance radiotherapy		N1		
76970	Ultrasound exam follow-up		Z2	0.8866	\$37.12
76975	GI endoscopic ultrasound		N1		
76977	Us bone density measure		Z3		\$6.53
76998	Us guide, intraop		N1		
76999	Echo examination procedure		Z2	0.8866	\$37.12
77001	Fluoroguide for vein device		N1		
77002	Needle localization by xray		N1		
77003	Fluoroguide for spine inject		N1		
77011	CI scan for localization		N1		

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HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
76511	Ophth us, quant a only		Z3		\$35.22
76512	Ophth us, b w/non-quant a		Z3		\$29.83
76513	Echo exam of eye, water bath		Z3		\$38.06
76514	Echo exam of eye, thickness		Z3		\$2.84
76516	Echo exam of eye		Z3		\$29.83
76519	Echo exam of eye		Z3		\$33.24
76529	Echo exam of eye		Z3		\$28.97
76536	Us exam of head and neck		Z2	1.3809	\$57.82
76604	Us exam, chest		Z2	0.8866	\$37.12
76645	Us exam, breast(s)		Z2	0.8866	\$37.12
76700	Us exam, abdom, complete		Z2	1.3809	\$57.82
76705	Echo exam of abdomen	CH	Z3		\$55.68
76770	Us exam abdo back wall, comp		Z2	1.3809	\$57.82
76775	Us exam abdo back wall, lim		Z2	1.3809	\$57.82
76776	Us exam k transpl w/doppler		Z2	1.3809	\$57.82
76800	Us exam, spinal canal	CH	Z3		\$53.40
76801	Ob us < 14 wks, single fetus		Z2	1.3809	\$57.82
76802	Ob us < 14 wks, addl fetus		Z3		\$21.59
76805	Ob us >= 14 wks, singl fetus		Z2	1.3809	\$57.82
76810	Ob us >= 14 wks, addl fetus		Z3		\$35.51
76811	Ob us, detailed, singl fetus	CH	Z3		\$74.71
76812	Ob us, detailed, addl fetus		Z2	0.8866	\$37.12
76813	Ob us nuchal meas, 1 gest		Z2	0.8866	\$37.12
76814	Ob us nuchal meas, add-on		Z3		\$23.58
76815	Ob us, limited, fetus(s)		Z2	0.8866	\$37.12
76816	Ob us, follow-up, per fetus		Z2	0.8866	\$37.12
76817	Transvaginal us, obstetric		Z2	0.8866	\$37.12
76818	Fetal biophyis profile wrist	CH	Z3		\$51.13
76819	Fetal biophyis profil w/o inst		Z3		\$39.77
76820	Umbilical artery echo		Z3		\$18.18
76821	Middle cerebral artery echo		Z3	0.8866	\$37.12
76825	Echo exam of fetal heart		Z3		\$96.58
76826	Echo exam of fetal heart		Z3		\$58.80

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HCPSC Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
77321	Special telelex port plan		Z3		\$48.86
77326	BrachyX isodose calc simp		Z2	1.461	\$61.18
77327	BrachyX isodose calc intemp		Z3		\$100.27
77328	BrachyX isodose plan compl	CH	Z3		\$128.96
77331	Special radiation dosimetry		Z3		\$14.20
77332	Radiation treatment aid(s)		Z3		\$39.06
77333	Radiation treatment aid(s)		Z3		\$15.62
77334	Radiation treatment aid(s)		Z3		\$69.59
77336	Radiation physics consult		Z3		\$41.76
77338	Design mic device for imrt	NI	Z2	2.7055	\$113.29
77370	Radiation physics consult		Z2	1.461	\$61.18
77371	Srs, multisource		Z2	104.238	\$4,364.76
77399	External radiation dosimetry		Z2	1.461	\$61.18
77401	Radiation treatment delivery		Z3		\$20.17
77402	Radiation treatment delivery		Z2	1.3168	\$55.14
77403	Radiation treatment delivery		Z2	1.3168	\$55.14
77404	Radiation treatment delivery		Z2	1.3168	\$55.14
77406	Radiation treatment delivery		Z2	2.2033	\$92.26
77407	Radiation treatment delivery		Z2	1.3168	\$55.14
77408	Radiation treatment delivery		Z2	1.3168	\$55.14
77409	Radiation treatment delivery		Z2	1.3168	\$55.14
77411	Radiation treatment delivery		Z2	2.2033	\$92.26
77412	Radiation treatment delivery		Z2	2.2033	\$92.26
77413	Radiation treatment delivery		Z2	2.2033	\$92.26
77414	Radiation treatment delivery		Z2	2.2033	\$92.26
77416	Radiation treatment delivery		Z2	2.2033	\$92.26
77417	Radiology port film(s)		NI		
77418	Radiation x-ray guidance		Z2	5.9784	\$250.33
77421	Stereoscopic x-ray guidance		NI		
77422	Neutron beam tx, simple		Z2	2.2033	\$92.26
77423	Neutron beam tx, complex		Z2	2.2033	\$92.26
77435	Sort management		NI		
77470	Special radiation treatment		Z3		\$86.34

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77012	Ct scan for needle biopsy		NI		
77013	Ct guide for tissue ablation		NI		
77014	Ct scan for therapy guide		NI		
77021	Mri guidance for needle place		NI		
77022	Mri for tissue ablation		NI		
77031	Stereotact guide for trst bx		NI		
77032	Guidance for needle, breast		NI		
77053	X-ray of mammary duct		NI		
77054	X-ray of mammary ducts		NI		
77071	X-ray stress view		Z3		\$19.60
77072	X-rays for bone age		Z3		\$9.94
77073	X-rays, bone length studies		Z3		\$17.33
77074	X-rays, bone survey, limited		Z3		\$34.09
77075	X-rays, bone survey complete		Z2	1.0678	\$44.71
77076	X-rays, bone survey, infant		Z2	1.0678	\$44.71
77077	Joint survey, single view		Z3		\$18.75
77078	Ct bone density, axial		Z2	1.0289	\$43.08
77079	Ct bone density, peripheral		Z3		\$32.10
77080	Dxa bone density, axial		Z3		\$40.34
77081	Dxa bone density/peripheral		Z3		\$13.35
77082	Dxa bone density, vert tx		Z3		\$14.77
77083	Radiographic absorptiometry		Z3		\$11.36
77084	Magnetic image, bone marrow		Z2	4.961	\$207.73
77280	Set radiation therapy field		Z2	1.461	\$61.18
77285	Set radiation therapy field		Z2	3.7799	\$158.28
77290	Set radiation therapy field		Z2	3.7799	\$158.28
77295	Set radiation therapy field		Z3		\$284.91
77299	Radiation therapy planning		Z2	1.461	\$61.18
77300	Radiation therapy dose plan		Z3		\$29.26
77301	Radiotherapy dose plan, imrt		Z2	13.1619	\$651.13
77305	TeleX isodose plan simple		Z3		\$25.28
77310	TeleX isodose plan intermed		Z3		\$32.95
77315	TeleX isodose plan complex		Z3		\$49.71

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78070	Parathyroid nuclear imaging		Z3		\$93.74
78075	Adrenal nuclear imaging		Z3		\$278.66
78099	Endocrine nuclear procedure		Z2	2.066	\$86.51
78102	Bone marrow imaging, lrd		Z3		\$99.42
78103	Bone marrow imaging, mult	CH	Z3		\$132.37
78104	Bone marrow imaging, body		Z2		\$153.60
78111	Plasma volume, multiple		Z3		\$51.13
78120	Red cell mass, single		Z3		\$53.97
78121	Red cell mass, multiple		Z3		\$60.22
78122	Blood volume		Z3		\$71.30
78130	Red cell survival study		Z3		\$99.48
78135	Red cell survival kinetics	CH	Z3		\$216.44
78140	Red cell sequestration		Z3		\$81.24
78185	Spleen imaging		Z3		\$126.69
78190	Platelet survival, kinetics		Z2	2.5656	\$107.43
78191	Platelet survival		Z2	2.5656	\$107.43
78195	Lymph system imaging		Z2	3.6682	\$153.60
78199	Blood/lymph nuclear exam		Z2	3.6682	\$153.60
78201	Liver imaging		Z3		\$115.33
78202	Liver imaging with flow	CH	Z3		\$129.25
78205	Liver imaging (3D)	CH	Z3		\$143.45
78206	Liver image (3d) with flow		Z2	4.1271	\$172.81
78215	Liver and spleen imaging		Z3		\$119.31
78216	Liver & spleen image/flow		Z3		\$77.26
78220	Liver function study		Z3		\$84.37
78223	Hepatobiliary imaging		Z2	4.1271	\$172.81
78230	Salivary gland imaging		Z3		\$101.13
78231	Salivary gland imaging		Z3		\$76.13
78232	Salivary gland function exam		Z3		\$75.28
78258	Esophageal motility study	CH	Z3		\$136.05
78261	Gastric mucosa imaging		Z2	3.5113	\$147.03
78262	Gastroesophageal reflux exam		Z2	3.5113	\$147.03

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
77520	Proton trmt, simple w/o comp		Z2	13.3743	\$560.02
77522	Proton trmt, simple w/comp		Z2	13.3743	\$560.02
77523	Proton trmt, intermediate		Z2	17.4955	\$732.59
77525	Proton treatment, complex		Z2	17.4955	\$732.59
77600	Hyperthermia treatment		Z2	5.4016	\$226.18
77610	Hyperthermia treatment		Z2	5.4016	\$226.18
77615	Hyperthermia treatment		Z2	5.4016	\$226.18
77620	Hyperthermia treatment		Z2	5.4016	\$226.18
77750	Intuse radioactive materials		Z3		\$72.44
77761	Apply intracav radiat simple		Z3		\$126.98
77762	Apply intracav radiat interm		Z3		\$149.13
77763	Apply intracav radiat compl	CH	Z2	4.2904	\$179.65
77776	Apply intersit radiat simpl		Z3		\$136.63
77777	Apply intersit radiat inter		Z3		\$147.43
77778	Apply intersit radiat compl		Z3		\$198.56
77785	Hdr brachytx, 1 channel		Z3		\$86.64
77786	Hdr brachytx, 2-12 channel		Z3		\$279.80
77787	Hdr brachytx over 12 chan	CH	Z3		\$426.09
77789	Apply surface radiation		Z3		\$38.35
77790	Radiation handling		N1		
77799	Radium/radioscope therapy		Z2	4.2904	\$179.65
78001	Thyroid, single uptake		Z3		\$44.31
78003	Thyroid, multiple uptake	CH	Z3		\$55.96
78006	Thyroid suppress/stimul		Z3		\$45.17
78007	Thyroid imaging with uptake		Z2	3.147	\$131.77
78010	Thyroid image, mult uptakes		Z3		\$77.55
78011	Thyroid imaging with flow		Z2	2.066	\$86.51
78015	Thyroid met imaging		Z3		\$128.40
78016	Thyroid met imaging/studies		Z2	4.1171	\$172.40
78018	Thyroid met imaging, body		Z2	4.1171	\$172.40
78020	Thyroid met uptake		N1		

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78472	Gated heart, planar, single	CH	Z3	4.3402	\$150.85
78473	Gated heart, multiple	CH	Z3	4.3402	\$181.74
78478	Heart wall motion add-on	CH	D5		
78480	Heart function add-on	CH	D5		
78481	Heart first pass, single	CH	Z3		\$122.71
78483	Heart first pass, multiple	CH	Z3		\$167.31
78491	Heart image (pet), single	CH	Z2	20.3369	\$851.57
78492	Heart image (pet), multiple	CH	Z2	20.3369	\$851.57
78494	Heart image, spect	CH	Z3		\$157.97
78496	Heart first pass add-on	CH	N1		
78499	Cardiovascular nuclear exam	CH	Z2	4.3402	\$181.74
78560	Lung perfusion imaging	CH	Z2	2.9299	\$122.68
78584	Lung V/Q image single breath	CH	Z3		\$76.13
78585	Lung V/Q imaging	CH	Z2	4.5952	\$192.41
78586	Aerosol lung image, single	CH	Z3		\$103.68
78587	Aerosol lung image, multiple	CH	Z2	2.9299	\$122.68
78598	Perfusion lung image	CH	Z2	4.5952	\$192.41
78591	Vent image, 1 breath, 1 proj	CH	Z3		\$105.10
78594	Vent image, multi proj, gas	CH	Z2	2.9299	\$122.68
78596	Vent image, multi proj, gas	CH	Z2	2.9299	\$122.68
78599	Lung differential function	CH	Z2	4.5952	\$192.41
78600	Brain image < 4 views	CH	Z3		\$112.49
78601	Brain image w/flow < 4 views	CH	Z3		\$134.08
78605	Brain image 4+ views	CH	Z2	2.7917	\$116.90
78606	Brain image w/flow 4+ views	CH	Z3		\$206.80
78607	Brain imaging (3D)	CH	Z3		\$219.01
78608	Brain imaging (PET)	CH	Z2	14.7231	\$616.50
78610	Brain flow imaging only	CH	Z2	2.7917	\$116.90
78630	Cerebrospinal fluid scan	CH	Z3		\$214.75
78635	CSF ventriculography	CH	Z3		\$202.25
78645	CSF shunt evaluation	CH	Z2	2.7917	\$116.90
78647	Cerebrospinal fluid scan	CH	Z3		\$207.36

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78264	Gastric emptying study	CH	Z2	3.5113	\$147.03
78270	Vit B-12 absorption exam	CH	Z3		\$50.85
78271	Vit b-12 abstr exam, int fac	CH	Z3		\$53.40
78272	Vit B-12 absorp, combined	CH	Z3		\$56.24
78278	Acute GI blood loss imaging	CH	Z2	3.5113	\$147.03
78280	GI protein loss exam	CH	Z2	3.5113	\$147.03
78291	Meckels divert exam	CH	Z2	3.5113	\$147.03
78299	Leveen/shunt patency exam	CH	Z2	3.5113	\$147.03
78300	GI nuclear procedure	CH	Z3	3.5113	\$147.03
78300	Bone imaging, limited area	CH	Z3		\$102.83
78305	Bone imaging, multiple areas	CH	Z3		\$136.07
78306	Bone imaging, whole body	CH	Z2	3.5118	\$147.05
78315	Bone imaging, 3 phase	CH	Z2	3.5118	\$147.05
78320	Bone imaging (3D)	CH	Z3		\$143.17
78399	Musculoskeletal nuclear exam	CH	Z2	3.5118	\$147.05
78414	Non-imaging heart function	CH	Z2	4.3402	\$181.74
78428	Cardiac shunt imaging	CH	Z3		\$114.76
78445	Vascular flow imaging	CH	Z3		\$105.10
78451	Ht muscle image spect, sing	CH	Z3		\$236.34
78452	Ht muscle image spect, mult	CH	Z3		\$236.34
78453	Ht muscle image, planar, sing	CH	Z3		\$113.91
78454	Ht muscle image, planar, mult	CH	Z3		\$95.73
78456	Acute venous thrombus image	CH	Z2	2.8394	\$112.49
78457	Venous thrombosis imaging	CH	Z2	2.8394	\$112.49
78458	Ven thrombosis images, bilat	CH	Z3		\$115.61
78459	Heart muscle imaging (PET)	CH	Z2	20.3369	\$851.57
78460	Heart muscle blood, single	CH	D5		
78461	Heart muscle blood, multiple	CH	D5		
78464	Heart image (3D), single	CH	D5		
78465	Heart image (3D), multiple	CH	D5		
78466	Heart infarct image	CH	Z3		\$105.67
78468	Heart infarct image (3D)	CH	Z3		\$132.94
78469	Heart infarct image (3D)	CH	Z3		\$151.69

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79300	Nuclir rx, interstit colloid		Z2	3.0955	\$129.62
79403	Hematopoietic nuclear tx		Z3		\$71.87
79440	Nuclear rx, intra-articular		Z3		\$40.34
79445	Nuclear rx, intra-arterial		Z2	3.0955	\$129.62
79999	Nuclear medicine therapy		Z2	3.0955	\$129.62
90371	Hep b ig, im		K2		\$111.20
90375	Rabies ig, im/sc		K2		\$142.79
90376	Rabies ig, heat treated		K2		\$130.16
90378	Rsv, mab, im, 50mg		K2		\$937.29
90385	Rh ig, minidose, im		N1		
90396	Vaccinia-zoster ig, im		K2		\$130.49
90476	Adenovirus vaccine, type 4	CH	K2		\$72.17
90585	Ecg vaccine, percut		K2		\$111.66
90632	Hep a vaccine, adult im		N1		
90633	Hep a vacc, ped/adol, 2 dose		N1		
90634	Hep a vacc, ped/adol, 3 dose		N1		
90636	Hep a/hcp b vacc, adult im		N1		
90645	Hib vaccine, hibcc, im		N1		
90646	Hib vaccine, prp-d, im		N1		
90647	Hib vaccine, prp-omp, im		N1		
90648	Hib vaccine, prp-t, im		N1		
90655	Flu vaccine no preserv, 6-35m		L1		
90656	Flu vaccine no preserv, 3 & >		L1		
90657	Flu vaccine, 3 yrs, im		L1		
90658	Flu vaccine, 3 yrs & >, im		L1		
90660	Flu vaccine, nasal		L1		\$0.93
90665	Lyme disease vaccine, im		K2		
90669	Pneumococcal vacc, 7 val im		L1		\$151.97
90675	Rabies vaccine, im		K2		\$96.27
90680	Rotavirus vacc, 3 dose, oral	CH	K2		\$72.37
90681	Rotavirus vacc, 2 dose oral		K2		\$106.60
90690	Typhoid vaccine, oral		N1		

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78650	GSF leakage imaging		Z3		\$211.34
78660	Nuclear exam of tear flow	CH	Z3		\$103.40
78699	Nervous system nuclear exam		Z2	2.7917	\$116.90
78700	Kidney imaging, morphol		Z3		\$109.65
78701	Kidney imaging with flow		Z3		\$134.93
78707	K flow/func image w/o drug		Z3		\$138.62
78708	K flow/func image w/drug		Z3		\$91.47
78709	K flow/func image, multiple		Z2	4.6133	\$193.17
78710	Kidney imaging (3D)		Z3		\$142.03
78725	Kidney function study		Z3		\$59.65
78730	Urinary bladder retention		Z3		\$48.96
78740	Ureteral reflux study		Z3		\$134.08
78761	Testicular imaging w/flow		Z3		\$125.55
78799	Genitourinary nuclear exam		Z2	4.6133	\$193.17
78800	Tumor imaging, limited area		Z3		\$111.35
78801	Tumor imaging, mult areas		Z3		\$151.69
78802	Tumor imaging, whole body		Z3		\$204.81
78803	Tumor imaging (3D)		Z3		\$217.59
78804	Tumor imaging, whole body		Z3		\$362.63
78805	Abscess imaging, ltd area		Z3		\$107.09
78806	Abscess imaging, whole body		Z3		\$214.47
78807	Nuclear localization/abscess	CH	Z2	4.1171	\$172.40
78808	Iv inj ra drug dx study		N1		
78811	Pet image, skull-thigh		Z2	14.7231	\$616.50
78812	Pet image, ltd area		Z2	14.7231	\$616.50
78813	Pet image, full body		Z2	14.7231	\$616.50
78814	Pet image w/ct, ltd		Z2	14.7231	\$616.50
78815	Pet image w/ct, skull-thigh		Z2	14.7231	\$616.50
78816	Pet image w/ct, full body		Z2	14.7231	\$616.50
78999	Nuclear diagnostic exam		Z3	1.5972	\$66.88
79005	Nuclear rx, oral admin		Z3		\$42.33
79101	Nuclear rx, iv admin		Z3		\$46.02
79200	Nuclear rx, intracav admin		Z3		\$51.13

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90749	Vaccine toxoid		N1		
A4218	Sterile saline or water		N1		
A4220	Infusion pump refill kit		N1		
A4248	Chlorhexidine antisept		N1		
A4262	Temporary tear duct plug		N1		
A4263	Permanent tear duct plug		N1		
A4270	Disposable endoscope sheath		N1		
A4300	Cath impl vasc access port		N1		
A4301	Implantable access syst perc		N1		
A4305	Drug delivery system >=50 ML		N1		
A4308	Drug delivery system <=50 ml		N1		
A4641	Radiofarm dx sigent noc		N1		
A4642	In111 satumomab		N1		
A4648	Implantable tissue marker		N1		
A4650	Implant radiation dosimeter		N1		
A9500	Tc99m sestambi		N1		
A9501	Technetium Tc-99m leboroxime		N1		
A9502	Tc99m tetrofosmin		N1		
A9503	Tc99m medronate		N1		
A9504	Tc99m apolide		N1		
A9505	Tl201 thallium		N1		
A9507	In111 capromab		N1		
A9508	I131 iodobenguat, dx		N1		
A9509	Iodine I-123 sod iodide ml		N1		
A9510	Tc99m disofenin		N1		
A9512	Tc99m pertechnetate		N1		
A9516	Iodine I-123 sod iodide mic		N1		
A9524	Tc99m exametazime		N1		
A9526	Nitrogen N-13 ammonia		N1		
A9527	Iodine I-125 sodium iodide	CH	H2		
A9528	Iodine I-131 iodide cap, dx		N1		
A9529	I131 iodide sol, dx		N1		\$37.92

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90691	Typhoid vaccine, im		N1		
90692	Typhoid vaccine, h-p, sc/ld		N1		
90696	Dtap-ipv vacc 4-6 yr im	CH	N1		
90698	Dtap-hib-ipv vaccine, im		N1		
90700	Dtap vaccine, < 7 yrs, im		N1		
90701	Dtp vaccine, im		N1		
90702	Dt vaccine < 7, im		N1		
90703	Tetanus vaccine, im		N1		
90704	Mumps vaccine, sc		N1		
90705	Measles vaccine, sc		N1		
90706	Rubella vaccine, sc		N1		
90707	Mmr vaccine, sc		N1		
90708	Measles-rubella vaccine, sc		N1		
90710	Mmr vaccine, sc		N1		
90712	Oral poliovirus vaccine		N1		
90713	Poliovirus, ipv, sc/im		N1		
90714	Td vaccine no prsv >= 7 im		N1		
90715	Tdap vaccine > 7 im		N1		\$0.16
90717	Yellow fever vaccine, sc		N1		
90718	Td vaccine > 7, im		N1		
90719	Diphtheria vaccine, im		N1		\$96.66
90720	Dtp/hib vaccine, im		N1		\$102.46
90721	Dtap/hib vaccine, im		N1		\$100.15
90725	Cholera vaccine, injectable	CH	K2		
90732	Pneumococcal vaccine		L1		
90733	Meningococcal vaccine, sc		K2		
90734	Meningococcal vaccine, im		K2		
90735	Encephalitis vaccine, sc	CH	K2		
90740	Hepb vacc, ill pat 3 dose im		F4		
90743	Hep b vacc, adol, 2 dose, im		F4		
90744	Hepb vacc ped/adol 3 dose im		F4		
90746	Hep b vaccine, adult, im		F4		
90747	Hepb vacc, ill pat 4 dose im		F4		

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A9572	Iridium In-111 pentetreotide		N1		
A9576	Inj prochloraz multipack		N1		
A9577	Inj multithance		N1		
A9578	Inj multithance multipack		N1		
A9579	Gad-base MR contrast NIOS, 1ml		N1		
A9580	Sodium fluoride F-18		N1		
A9581	Gadoxetate disodium inj	NI	K2		\$13,350
A9582	Iodine I-123 iobenguane	NI	K2		\$2,329.83
A9583	Gadofosveset trisodium inj	NI	K2		\$1,29
A9588	Non-rad contrast materialNOC		N1		
C1713	Anchor/screw brn/bn,ls/bn		N1		
C1714	Cath, trans atherectomy, dir		N1		
C1715	Brachytherapy needle		N1		
C1716	Brachytx, non-str, HDR Ir-192	CH	H2		\$42,85
C1717	Brachytx, non-str, HDR Ir-192	CH	H2		\$231,38
C1719	Brachytx, NS, Non-HDRIr-192	CH	H2		\$64,02
C1721	AICD, dual chamber		N1		
C1722	AICD, single chamber		N1		
C1724	Cath, trans atheroc, rotation		N1		
C1725	Cath, trans atheroc, rotation		N1		
C1726	Cath, bal dil, non-vascular		N1		
C1727	Cath, bal tis dis, non-vas		N1		
C1728	Cath, brachytx seed adm		N1		
C1729	Cath, drainage		N1		
C1730	Cath, EP, 19 or few elec		N1		
C1731	Cath, EP, 20 or more elec		N1		
C1732	Cath, EP, diag/abl, 3D/vect		N1		
C1733	Cath, EP, other than cook-tip		N1		
C1750	Cath, hemodialysis, long-term		N1		
C1751	Cath, int, per/cent/midline		N1		
C1752	Cath, hemodialysis, short-term		N1		
C1753	Cath, intravas ultrasound		N1		
C1754	Catheter, intradiscal		N1		

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A9531	I131 max 100uCi		N1		
A9532	I125 serum albumin, dx		N1		
A9535	Injection, methylene blue	CH	D5		
A9536	Tc99m depreotide		N1		
A9537	Tc99m mebrofenin		N1		
A9538	Tc99m methosphate		N1		
A9539	Tc99m penicilate		N1		
A9540	Tc99m MAA		N1		
A9541	Tc99m sulfur colloid		N1		
A9542	In111 ibritumomab, dx		N1		
A9544	I131 besitumomab, dx		N1		
A9546	Co57/58		N1		
A9547	In111 oxyquinoline		N1		
A9548	In111 pentetate		N1		
A9550	Tc99m gluceptate		N1		
A9551	Tc99m succimer		N1		
A9552	F18 fdg		N1		
A9553	Cr51 chromate		N1		
A9554	I125 iohalamate, dx		N1		
A9555	Rb82 rubidium		N1		
A9556	Ga67 gallium		N1		
A9557	Tc99m bisphate		N1		
A9558	Xe133 xenon 10mci		N1		
A9559	Co57 cyano		N1		
A9560	Tc99m labeled fbc		N1		
A9561	Tc99m oxidronate		N1		
A9562	Tc99m mercaptide		N1		
A9566	Tc99m tetrofosminab		N1		
A9567	Technetium Tc-99m aerosol		N1		
A9568	Technetium tc99m arctiumomab		N1		
A9569	Technetium Tc-99m auto WBC		N1		
A9570	Iridium In-111 auto WBC		N1		
A9571	Iridium In-111 auto platelet		N1		

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPSC Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
C1814	Retinal tamp, silicone oil		N1		
C1815	Pros, urinary sph, imp		N1		
C1816	Receiver/transmitter, neuro		N1		
C1817	Septal defect imp sys		N1		
C1818	Integrated heratocprosthesis		N1		
C1819	Tissue focalization-excision		N1		
C1820	Generator neuro rechg bat sy		N1		
C1821	Interspirous implant		N1		
C1874	Stent, coated/cov w/del sys		N1		
C1875	Stent, coated/cov w/del sy		N1		
C1876	Stent, non-coat/non-cov w/del		N1		
C1877	Stent, non-coat/cov w/del		N1		
C1878	Matt for vocal cord		N1		
C1879	Tissue marker, implantable		N1		
C1880	Vena cava filter		N1		
C1881	Dialysis access system		N1		
C1882	AICD, other than sing/dual		N1		
C1883	Adapt/exit, pacing/neuro lead		N1		
C1884	Embolization Protect syst		N1		
C1885	Cath, transilumin angio laser		N1		
C1887	Catheter, guiding		N1		
C1886	Endovas non-cardiac abl cath		N1		
C1891	Infusion pump,non-prog, perm		N1		
C1892	Intro/sheath fixed,peel-away		N1		
C1893	Intro/sheath, fixed,non-peel		N1		
C1894	Intro/sheath, non-laser		N1		
C1895	Lead, AICD, endo dual coil		N1		
C1896	Lead, AICD, non sing/dual		N1		
C1897	Lead, neurostim leat kit		N1		
C1898	Lead, pmkr, other than trans		N1		
C1899	Lead, pmkr/AICD combination		N1		
C1900	Lead, coronary venous		N1		
C2814	Probe, peric lumb disc		N1		

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C1755	Catheter, intraspinal		N1		
C1756	Cath, pacing, transeptoph		N1		
C1757	Cath, thrombectomy/embolact		N1		
C1758	Catheter, ureteral		N1		
C1759	Cath, intra echocardiography		N1		
C1760	Closure dev, vasc		N1		
C1762	Conn tss, human(inc fascia)		N1		
C1763	Conn tss, non-human		N1		
C1764	Event recorder, cardiac		N1		
C1765	Adhesion barrier		N1		
C1766	Intro/sheath,stable,non-peel		N1		
C1767	Generator, neuro non-rechrg		N1		
C1768	Graft, vascular		N1		
C1769	Guide wire		N1		
C1770	Imaging coil, MR, insertable		N1		
C1771	Rep dev, urinary, wsling		N1		
C1772	Infusion pump, programmable		N1		
C1773	Ret dev, insertable		N1		
C1776	Joint device (implantable)		N1		
C1777	Lead, AICD, endo single coil		N1		
C1778	Lead, neurostimulator		N1		
C1779	Lead, pmkr, transvenous VDD		N1		
C1780	Lens, intraocular (new tech)		N1		
C1781	Mesh (implantable)		N1		
C1782	Morcellator		N1		
C1783	Ocular imp, aqueous drain de		N1		
C1784	Ocular dev, intraop, delret		N1		
C1785	Pmkr, dual, rate-resp		N1		
C1786	Pmkr, single, rate-resp		N1		
C1787	Patient progr, neurostim		N1		
C1788	Port, indwelling, imp		N1		
C1789	Prosthesis, breast, imp		N1		
C1813	Prosthesis, penile, inflatable		N1		

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HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
C8907	MRI w/o cont, breast, bi		Z2	4.961	\$207.73
C8908	MRI w/o fol w/cont, breast,		Z2	7.5993	\$318.21
C8909	MRA w/cont, chest		Z2	6.0177	\$251.98
C8910	MRA w/o cont, chest		Z2	4.961	\$207.73
C8911	MRA w/o fol w/cont, chest		Z2	7.5993	\$318.21
C8912	MRA w/cont, lwr ext		Z2	6.0177	\$251.98
C8913	MRA w/o fol w/cont, lwr ext		Z2	4.961	\$207.73
C8914	MRA w/o fol w/cont, lwr ext		Z2	7.5993	\$318.21
C8918	MRA w/cont, pelvis		Z2	6.0177	\$251.98
C8919	MRA w/o cont, pelvis		Z2	4.961	\$207.73
C8920	MRA w/o fol w/cont, pelvis		Z2	7.5993	\$318.21
C9113	Inj pantoprazole sodium, via		N1		\$18.10
C9121	Injection, argatroban	CH	K2		
C9245	Injection, romiplostim	CH	D5		
C9246	Inj, gadoxetate disodium	CH	D5		
C9247	Inj, obenguan, I-123, dx	CH	D5		\$3.39
C9248	Inj, clevodipine butyrate	CH	K2		
C9249	Inj, certolizumab pegol	CH	D5		
C9250	Artiss fibrin sealant	CH	K2		\$138.20
C9251	Inj, C1 esterase inhibitor	CH	D5		
C9252	Injection, plerixator	CH	D5		
C9253	Injection, ferrozolamide	CH	D5		
C9254	Injection, lacosamide	NI	K2		\$0.18
C9255	Paliperidone palmitate inj	NI	K2		\$6.71
C9256	Dexamethasone intravitreal	NI	K2		\$196.10
C9257	Bevacizumab injection	NI	K2		\$1.41
C9352	Neuragen nerve guide, per cm		N1		
C9353	Neurwrap nerve protector,cm		N1		
C9354	Verifas collagen matrix, cm2	CH	N1		
C9355	Neuromatrix nerve cuff, cm	CH	N1		
C9356	TenoGlide tendon prof, cm2		K2		\$24.86
C9358	SurgiMend, fetal		K2		\$10.76
C9359	Impint,bon void filler-putty		K2		\$63.54

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C2615	Sealant, pulmonary, liquid		N1		\$15,779.35
C2616	BrachyX, non-str, Yttrium-90	CH	H2		
C2617	Stent, non-cor, tem w/o del		N1		
C2618	Probe, cryoablation		N1		
C2619	Pmkr, dual, non rate-resp		N1		
C2620	Pmkr, single, non rate-resp		N1		
C2621	Pmkr, other than sing/dual		N1		
C2622	Prosthesis, penile, non-inf		N1		
C2625	Stent, non-cor, tem w/del sv		N1		
C2626	Infusion pump, non-prog,temp		N1		
C2627	Cath, suprapubic/cystoscopic		N1		
C2628	Catheter, occlusion		N1		
C2629	Intro/sheath, laser		N1		
C2630	Cath, EP, cool-hip		N1		
C2631	Rep dev, urinary, w/o sling		N1		\$59.80
C2634	BrachyX, non-str, HA, I-125	CH	H2		\$28.59
C2635	BrachyX, non-str, HA, P-103	CH	H2		\$19.37
C2636	Brachy linear, non-str, P-103	CH	H2		\$42.48
C2638	BrachyX, stranded, I-125	CH	H2		\$36.18
C2639	BrachyX, non-stranded, I-125	CH	H2		\$60.36
C2640	BrachyX, stranded, P-103	CH	H2		\$57.12
C2641	BrachyX, non-stranded, P-103	CH	H2		\$109.84
C2642	BrachyX, stranded, C-131	CH	H2		\$66.09
C2643	BrachyX, non-stranded, C-131	CH	H2		\$42.48
C2698	BrachyX, stranded, NOS	CH	H2		\$28.59
C2699	BrachyX, non-stranded, NOS	CH	H2		\$251.98
C8900	MRA w/cont, abd		Z2	6.0177	\$207.73
C8901	MRA w/o cont, abd		Z2	4.961	\$318.21
C8902	MRA w/o fol w/cont, abd		Z2	7.5993	\$251.98
C8903	MRI w/cont, breast, uni		Z2	6.0177	\$207.73
C8904	MRI w/o cont, breast, uni		Z2	4.961	\$207.73
C8905	MRI w/o fol w/cont, brst, un		Z2	7.5993	\$318.21
C8906	MRI w/cont, breast, bi		Z2	6.0177	\$251.98

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J0215	Alefacept		K2		\$30.02
J0220	Alglucosidase alfa Injection		K2		\$124.69
J0256	Alpha 1 proteinase inhibitor		K2		\$3.63
J0278	Amikacin sulfate injection		N1		
J0280	Aminophyllin 250 MG inj		N1		
J0285	Amiodarone HCl		N1		
J0287	Amphotericin B		N1		
J0288	Ampho b cholesterol sulfate		K2		\$9.66
J0289	Amphotericin b liposome inj		K2		\$13.74
J0290	Ampicillin 500 MG inj		K2		\$14.96
J0295	Ampicillin sodium per 1.5 gm		N1		
J0300	Amobarbital 125 MG inj		N1		
J0330	Succinylcholine chloride inj		N1		
J0348	Anidulafungin injection		K2		\$1.21
J0360	Hydratizine hcl Injection		N1		
J0364	Apomorphine hydrochloride		N1		
J0365	Aprotinin, 10,000 kiu		K2		\$2.60
J0380	Inj metaraminol bitartrate		N1		
J0390	Chloroquine injection		N1		
J0400	Aripiprazole injection		N1		
J0456	Azithromycin		N1		
J0460	Atropine sulfate Injection	CH	D5		
J0461	Atropine sulfate Injection	NI	N1		\$26.81
J0470	Dimecaprol injection		K2		\$195.31
J0475	Baclofen 10 MG injection		K2		\$71.24
J0476	Baclofen intrathecal trial		K2		\$1,624.44
J0500	Dicyclomine injection		N1		
J0515	Inj benzotropine mesylate		N1		
J0520	Bethanechol chloride inject		N1		
J0530	Penicillin g benzathine inj	CH	D5		
J0540	Penicillin g benzathine inj	CH	D5		

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C9360	SurgiMend, neonatal		K2		\$10.67
C9361	NeuroMend nerve wrap		K2		\$247.29
C9362	ImpinjLbon void filler-strip		K2		\$63.60
C9363	Integra Meshed BI Wound Mat		K2		\$25.62
C9364	Porcine implant, Permacol		K2		\$17.21
C9369	Unclassified Drugs or biolog		K7		
E0616	Cardiac event recorder		N1		
E0749	Elec osteogen stim implanted		N1		
E0782	Non-programmable infusion pump		N1		
E0783	Programmable infusion pump		N1		
E0785	Replacement tmpl pump cathet		N1		
E0786	Implantable pump replacement		N1		
G0130	Single energy x-ray study		Z3		\$15.91
G0173	Linear acc stereo radsur com		Z2	50.6947	\$2,122.74
G0251	Linear acc based stereo radio		Z2	13.6824	\$572.09
G0288	Recon, CTA for surg plan		N1		
G0340	Robot lin-radsurg com, first		Z2	50.6947	\$2,122.74
J0120	Tetracyclin injection		Z2	35.3136	\$1,478.69
J0129	Abatacept injection		N1		
J0130	Abacimab injection		K2		\$18.98
J0132	Acetylcysteine injection		K2		\$459.36
J0133	Acyclovir injection		K2		\$2.29
J0135	Adalimumab injection		N1		
J0150	Injection adenosine 6 MG		K2		\$357.53
J0152	Adenosine injection		K2		\$9.50
J0170	Adrenalin epinephrin inject		K2		\$76.42
J0180	Agalsidase beta injection		N1		
J0190	Inj biperiden lactate/5 mg		K2		\$133.69
J0200	Alatrofloxacin mesylate		N1		
J0205	Aglucerase injection		N1		
J0207	Amifostine		K2		\$41.19
J0210	Meflydopate hcl injection		K2		\$50.07
J0210	Meflydopate hcl injection		K2		\$27.64

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J0715	Ceftiozime sodium / 500 MG		N1		
J0718	Cerlitzumab pegol inj	NI	K2		\$3.80
J0720	Chloramphenicol sodium injec		N1		
J0725	Chlorthalidone hydrochloride		N1		
J0735	Clonidine hydrochloride		K2		\$109.75
J0740	Cidofovir injection		K2		\$746.46
J0743	Cilastatin sodium injection		N1		
J0744	Ciprofloxacin iv		N1		
J0745	Ini codoine phosphate /30 MG		N1		
J0760	Colchicine injection		N1		
J0770	Colistimethate sodium inj		N1		
J0780	Prochlorperazine injection		N1		
J0795	Cortocorelin ovine triflutal		K2		\$4.24
J0800	Corticotropin injection		K2		\$2,394.93
J0833	Cosyntropin injection NOS	NI	K2		\$91.84
J0834	Cosyntropin cortrosyn inj	NI	K2		\$91.84
J0835	Ini cosyntropin per 0.25 MG	CH	D5		
J0850	Cytomegalovirus imm IV /vial		K2		\$862.24
J0878	Daptomycin injection		K2		\$0.40
J0881	Darbepoetin alfa, non-ead		K2		\$2.76
J0885	Epoetin alfa, non-ead		K2		\$9.40
J0894	Dextabine injection		K2		\$28.42
J0895	Deferoxamine mesylate inj		N1		
J0900	Testosterone enanthate inj		N1		
J0945	Brompheniramine maleate inj	CH	K2		\$0.75
J0970	Estradiol valerate injection		N1		
J1000	Depo-estradiol cypionate inj		N1		
J1020	Methylprednisolone 20 MG inj		N1		
J1030	Methylprednisolone 40 MG inj		N1		
J1040	Methylprednisolone 80 MG inj		N1		
J1051	Medroxyprogesterone inj		N1		
J1060	Testosterone cypionate 1 ML		N1		
J1070	Testosterone cypionat 100 MG		N1		

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J0550	Penicillin g benzathine inj	CH	D5		
J0559	PeriG benzathine/procaraine inj	NI	N1		
J0560	Penicillin g benzathine inj		N1		
J0570	Penicillin g benzathine inj		N1		
J0580	Penicillin g benzathine inj		K2		\$2.40
J0583	Bivalirudin		K2		\$5.40
J0585	Injection, onabotulinumtoxinA	NI	K2		\$8.23
J0586	Abobotulinumtoxin type A		K2		\$10.38
J0587	Ini, rimabotulinumtoxinB		N1		
J0592	Buprenorphine hydrochloride		N1		
J0594	Buserelin injection		K2		\$14.18
J0595	Butorphanol tartrate 1 mg		N1		
J0598	C1 esterase inhibitor inj	NI	K2		\$41.34
J0600	Edetate calcium disodium inj		K2		\$78.86
J0610	Calcium gluconate injection		N1		
J0620	Calcium glycer & lact 10 ML		N1		
J0630	Calcitonin salmon injection		K2		\$48.37
J0636	Ini calcitriol per 0.1 mcg		N1		
J0637	Casopofungin acetate		K2		\$11.52
J0640	Leucovorin calcium injection		N1		
J0641	Levoleucovorin injection		K2		\$0.99
J0670	Inj meflvacaine HCl/10 ml		N1		
J0690	Cefazolin sodium injection		N1		
J0692	Cefepime HCl for injection		N1		
J0694	Cefoxitin sodium injection		N1		
J0696	Ceftiozime sodium injection		N1		
J0697	Stetit cefuroxime injection		N1		
J0698	Cefotaxime sodium injection		N1		
J0702	Betamethasone acet&sod phosp		N1		
J0704	Betamethasone sod phosp/4 MG		N1		
J0706	Caffeine citrate injection		N1		
J0710	Cefepirin sodium injection		N1		
J0713	Inj ceftazidime per 500 mg		N1		

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J1430	Ethinamine oleate 100 mg		K2		\$147.14
J1435	Injection estrone per 1 MG		N1		
J1436	Elidronate disodium inj		K2		\$70.06
J1438	Etanercept injection		K2		\$183.61
J1440	Fligastin 300 mcg injection		K2		\$208.54
J1441	Fligastin 480 mcg injection		K2		\$24.44
J1450	Fluconazole		N1		
J1451	Fomepizole, 15 mg		K2		\$7.99
J1453	Fosaprepitant injection		K2		\$1.58
J1455	Foscarnet sodium injection	CH	N1		
J1457	Gallium nitrate injection		K2		\$1.71
J1458	Galsulfase injection		K2		\$339.04
J1459	Inj IVIG privenge 500 mg		K2		\$35.05
J1460	Gamma globulin 1 CC inj		K2		\$15.05
J1470	Gamma globulin 2 CC inj		K2		\$30.10
J1480	Gamma globulin 3 CC inj		K2		\$45.14
J1490	Gamma globulin 4 CC inj		K2		\$60.20
J1500	Gamma globulin 5 CC inj		K2		\$75.26
J1510	Gamma globulin 6 CC inj		K2		\$90.35
J1520	Gamma globulin 7 CC inj		K2		\$105.27
J1530	Gamma globulin 8 CC inj		K2		\$120.40
J1540	Gamma globulin 9 CC inj		K2		\$150.50
J1550	Gamma globulin 10 CC inj		K2		\$150.50
J1560	Gamma globulin > 10 CC inj		K2		\$150.50
J1561	Gamunex injection		K2		\$36.71
J1562	Vivaglobin, inj		K2		\$7.05
J1565	RSV-ivig	CH	D5		
J1566	Immune globulin, powder		K2		\$29.83
J1568	Octagam injection		K2		\$37.03
J1569	Gammagard liquid injection		K2		\$37.85
J1570	Ganciclovir sodium injection		N1		
J1571	Hepagam b tm injection		K2		\$50.04
J1572	Flebogamma injection		K2		\$36.51

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Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
J1080	Testosterone cypionate 200 MG		N1		
J1094	Inj dexamethasone acetate		N1		
J1100	Dexamethasone sodium phos		N1		
J1110	Inj dhydroergotamine mesyl		N1		
J1120	Acetazolamid sodium injecto		N1		
J1180	Digoxin injection		K2		\$474.73
J1162	Digoxin immune tab (ovine)		N1		
J1165	Phenofen sodium injection		N1		
J1170	Hydromorphone injection		N1		
J1180	Dipyrilline injection		N1		
J1190	Dezoxazane HCl injection		K2		\$340.03
J1200	Diphenhydramine hcl injectio		N1		
J1205	Chlorothiazide sodium inj		K2		\$292.02
J1212	Dimethyl sulfoxide 50% 50 ML		K2		\$67.46
J1230	Mebarone injection		N1		
J1240	Dimethylhydinate injection		N1		
J1245	Dipyridamide injection		N1		
J1260	Inj dobutamine HCL/250 mg		N1		
J1265	Dolasetron mesylate	CH	N1		
J1267	Dopamine injection		N1		
J1270	Injection, doxercalciferol		K2		\$0.57
J1300	Ecuzumab injection		K2		\$177.57
J1320	Amirtipiline injection		N1		
J1324	Entuvirtide injection	CH	K2		\$0.47
J1325	Epoprostenol injection		N1		
J1327	Epifibatid injection		K2		\$18.57
J1330	Ergonovine maleate injection		N1		
J1335	Erlapenem injection		N1		
J1364	Erythro lactobionate /500 MG		N1		
J1380	Estradiol valerate 10 MG inj		N1		
J1390	Estradiol valerate 20 MG inj		N1		
J1410	Inj estrogen conjugate 25 MG		K2		\$83.21

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HCPSC Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
J1817	Insulin for insulin pump use	CH	K2		\$3.34
J1830	Insulin beta-1b / 25 MG		K2		\$168.90
J1835	Itraconazole injection	CH	N1		
J1840	Kanamycin sulfate 500 MG inj		N1		
J1855	Kanamycin sulfate 75 MG inj		N1		
J1885	Ketorolac tromethamine inj		N1		
J1890	Cephalexin sodium injection		N1		
J1930	Lanreotide injection		K2		\$26.65
J1931	Lanreotide injection		K2		\$25.08
J1940	Furosemide injection		N1		
J1945	Lepruquin		K2		\$174.51
J1950	Leuprolide acetate /3.75 MG		K2		\$480.20
J1953	Levetiracetam injection		K2		\$0.75
J1956	Levofloxacin injection		N1		
J1960	Levoprolol tartrate inj		N1		
J1980	Hyoscyamine sulfate inj		N1		
J1990	Chondrozepoxide injection		N1		
J2001	Lidocaine injection		N1		
J2010	Lincomycin injection		N1		
J2020	Linezolid injection		K2		\$29.37
J2060	Lorazepam injection		N1		
J2150	Mannitol injection		N1		
J2170	Mecasermin injection		N1		
J2175	Meperidine hydrochloride /100 MG		N1		
J2180	Meperidine/promethazine inj		N1		
J2185	Meropenem		N1		
J2210	Methylglucosyl maltate inj		N1		
J2248	Micafungin sodium injection		K2		\$1.08
J2250	Inj midazolam hydrochloride		N1		
J2260	Inj milrinone lactate / 5 MG		N1		
J2270	Morphine sulfate injection		N1		
J2271	Morphine sulfate injection 100mg		N1		
J2275	Morphine sulfate injection		N1		

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HCPSC Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
J1573	Hepapatin b intravenous, inj		K2		\$50.04
J1580	Garamycin gentamicin inj		N1		
J1590	Gatifloxacin injection		N1		\$81.23
J1595	Injection, glatiramer acetate		K2		
J1600	Gold sodium thiomalate inj		N1		\$79.20
J1610	Glucagon hydrochloride/1 MG		K2		\$176.89
J1620	Gonadorelin hydrochloride/1 MG		N1		
J1626	Granisetron hcl injection	CH	N1		
J1630	Haloperidol injection		N1		
J1631	Haloperidol decanoate inj		N1		
J1640	Hemlin, 1 mg		K2		\$7.73
J1642	Inj heparin sodium per 10 u		N1		
J1644	Inj heparin sodium per 1000u		N1		
J1645	Dalteparin sodium		N1		
J1650	Inj enoxaparin sodium		N1		\$5.98
J1652	Fondaparinux sodium		K2		
J1655	Tinzaparin sodium injection		N1		\$199.91
J1670	Tetanus immune globulin inj		K2		\$96.46
J1680	Human fibrinogen conc inj	NI	K2		
J1700	Hydrocortisone acetate inj		N1		
J1710	Hydrocortisone sodium ph inj		N1		
J1720	Hydrocortisone sodium succ i		N1		
J1730	Diazoxide injection		K2		\$112.16
J1740	Ibandronate sodium injection		K2		\$139.22
J1742	Ibutilide fumarate injection		K2		\$404.01
J1743	Intrusulfase injection		K2		\$446.44
J1745	Infliximab injection		K2		\$57.60
J1750	Inj iron dextran		K2		\$14.11
J1756	Iron sucrose injection		K2		\$0.37
J1785	Injection imiglucerase /unit		K2		\$4.12
J1790	Droperidol injection		N1		
J1800	Propranolol injection		N1		
J1815	Insulin injection		N1		

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HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
J2560	Phenobarbital sodium inj		N1		
J2562	Plerixafor injection	NI	K2		\$268.51
J2590	Oxytocin injection		N1		
J2597	Inf desmopressin acetate		N1		
J2650	Prednisolone acetate inj		N1		
J2670	Totaxoline hcl injection		N1		
J2675	Inf progesterone per 50 MG		N1		
J2680	Fluphenazine decanoate 25 MG		N1		
J2690	Procainamide hcl injection		N1		
J2700	Oxacillin sodium injection		N1		
J2710	Neostigmine methylsulfite inj		N1		
J2720	Inf protamine sulfate/10 MG		N1		
J2724	Protein c concentrate		K2		\$11.96
J2725	Inf proflerin per 250 mcg		N1		
J2730	Pralidoxime chloride inj		K2		\$85.83
J2760	Phentolane mesylate inj		N1		
J2765	Metoclopramide hcl injection		N1		
J2770	Gumpristin/daltopristin		K2		\$144.08
J2778	Ranibizumab injection		K2		\$398.11
J2780	Ranitidine hydrochloride inj		N1		
J2783	Rasburicase		K2		\$164.00
J2785	Regadenoson injection		K2		\$50.78
J2788	Rho d immune globulin 50 mcg		K2		\$25.76
J2790	Rho d immune globulin inj		K2		\$84.39
J2791	Rhophylac injection		K2		\$5.13
J2792	Rho(D) immune globulin h_sd		K2		\$18.39
J2793	Ritonacept injection	NI	K2		\$23.64
J2794	Risperidone, long acting		K2		\$41.93
J2795	Ropivacaine HCl injection		N1		
J2796	Romiplostim injection	NI	K2		\$43.75
J2800	Methocarbamol injection		N1		
J2805	Sincalide injection	CH	N1		
J2810	Inf theophylline per 40 MG		N1		

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HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
J2278	Ziconotide injection		K2		\$6.65
J2280	Inf, moxifloxacin 100 mg		N1		
J2300	Inf nalbuphine hydrochloride		N1		
J2310	Inf naloxone hydrochloride		N1		
J2315	Naltrexone, depot form		K2		\$2.14
J2320	Nandrolone decanoate 50 MG	CH	K2		\$7.00
J2321	Nandrolone decanoate 100 MG	CH	K2		\$7.00
J2322	Nandrolone decanoate 200 MG	CH	K2		\$14.74
J2323	Natalizumab injection		K2		\$8.32
J2325	Nasafitide injection		K2		\$36.07
J2353	Ocrotetide injection, depot		K2		\$105.27
J2354	Ocrotetide inj, non-depot		N1		
J2355	Oprelvekin injection		K2		\$242.16
J2357	Omalizumab injection		K2		\$18.86
J2360	Orphenadrine injection		N1		
J2370	Phenyphrine hcl injection		N1		
J2400	Chlorprocaine hcl injection		N1		
J2405	Ondansetron hcl injection	CH	N1		
J2410	Oxymorphone hcl injection		N1		
J2425	Palfarin injection		K2		\$11.06
J2430	Pamidronate disodium 30 MG		K2		\$18.42
J2440	Papaverin hcl injection		N1		
J2469	Palonosetron hcl		K2		\$17.19
J2501	Paricalcitol		N1		
J2503	Pegadotarb sodium injection		K2		\$1,014.11
J2504	Pegademase bovine, 25 iu		K2		\$242.67
J2505	Injection, pegfilgrastim 6mg		K2		\$2,222.07
J2510	Penicillin g procaine inj		N1		
J2513	Penta starch 10% solution		K2		\$1,270.88
J2515	Pentobarbital sodium inj	CH	N1		
J2540	Penicillin g potassium inj		N1		
J2543	Piperacillin/tazobactam		N1		
J2550	Promethazine hcl injection		N1		

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J3302	Triamcinolone diacetate [in]		N1		
J3303	Triamcinolone hexacetonate [in]		N1		
J3305	Inj trimetrexate glucuronate		K2		\$124.80
J3310	Perphenazine injection		N1		
J3315	Triptonelin pamoate		K2		\$160.83
J3320	Spectinomycin d-hcl inj		N1		
J3350	Urea injection	CH	N1		
J3355	Urofollitropin, 75 lu		K2		\$59.26
J3360	Diazepam injection		N1		
J3364	Urokinase 5000 IU injection		N1		
J3365	Urokinase 250,000 IU [in]		K2		\$449.09
J3370	Vancomycin hcl injection		N1		
J3396	Verteporfin injection	CH	K2		\$9.31
J3400	Trifluromazine hcl inj		N1		
J3410	Hydroxyzine hcl injection		N1		
J3411	Thiamine hcl 100 mg		N1		
J3415	Pyridoxine hcl 100 mg		N1		
J3420	Vitamin b12 injection		N1		
J3430	Vitamin k phyttonadione [in]		N1		
J3465	Injection, voriconazole		K2		\$5.26
J3470	Hyaluronidase injection		N1		
J3471	Ovine, up to 999 USP units		N1		
J3472	Ovine, 1000 USP units	CH	N1		
J3473	Hyaluronidase recombinant	CH	N1		
J3475	Inj magnesium sulfate		N1		
J3480	Inj potassium chloride		N1		
J3485	Zidovudine		N1		
J3486	Ziprasidone mesylate		N1		
J3487	Zoledronic acid		N1		
J3488	Reclast injection		K2		\$214.94
J3490	Drugs unclassified injection		K2		\$218.59
J3530	Nasal vaccine inhalation		N1		
J3590	Unclassified biologics		N1		

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J2920	Sargramostim injection		K2		\$23.31
J2950	Inj, secretin synthetic human		K2		\$19.83
J2910	Aurothioglucose injection		N1		
J2916	Na ferric gluconate complex		N1		
J2920	Methylprednisolone injection		N1		
J2930	Methylprednisolone injection		N1		
J2940	Somatropin injection		K2		\$43.99
J2941	Somatropin injection		K2		\$53.47
J2950	Promazine hcl injection		N1		
J2963	Retepase injection		K2		\$1,230.80
J2995	Inj streptokinase /250000 IU		K2		\$78.00
J2997	Alteplase recombinant		K2		\$35.03
J3000	Streptomycin injection		N1		
J3010	Fentanyl citrate injection		N1		
J3030	Sumatriptan succinate / 6 MG		K2		\$55.49
J3070	Pentacocaine injection		N1		
J3101	Tenecteplase injection		K2		\$40.10
J3105	Terbutaline sulfate [in]		N1		
J3120	Testosterone enanthate [in]		N1		
J3130	Testosterone enanthate inj		N1		
J3140	Testosterone suspension [in]		N1		
J3150	Testosterone propionate [in]		N1		
J3230	Chlorpromazine hcl injection		N1		
J3240	Thyrotropin injection		K2		\$948.38
J3243	Tigecycline injection		K2		\$1.15
J3246	Tirofiban HCl		K2		\$7.83
J3250	Trimethoprimamide hcl [in]		N1		
J3260	Tobramycin sulfate injection		N1		
J3265	Injection tobramycin 10 mg/ml		N1		
J3280	Thiethylperazine maleate [in]		N1		
J3285	Treprostinil injection		K2		\$54.83
J3300	Triamcinolone A inj PFS-free		K2		\$3.20
J3301	Triamcinolone aced [in] NCS		N1		

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J7504	Lymphocyte immune globulin		K2		\$453.67
J7505	Monoclonal antibodies		K2		\$1,109.45
J7506	Prednisone oral		N1		
J7507	Tacrolimus oral per 1 MG		K2		\$3.96
J7509	Methylprednisolone oral		N1		
J7510	Prednisolone oral per 5 mg		N1		
J7511	Anti-thymocyte globulin rabbit		K2		\$414.44
J7513	Dacizumab, parenteral		K2		\$378.20
J7515	Cyclosporine oral 25 mg	CH	K2		\$0.82
J7516	Cyclosporin parenteral 250mg		K2		\$21.24
J7517	Mycophenolate mofetil oral		K2		\$2.45
J7518	Mycophenolic acid	CH	N1		
J7520	Sirolimus, oral		K2		\$9.44
J7525	Tacrolimus injection		K2		\$136.82
J7599	Immunosuppressive drug nec		N1		
J7674	Methacholine chloride, nebul		N1		
J7799	Non-inhalation drug for DME		N1		
J8501	Oral aprepitant		K2		\$5.42
J8510	Oral busulfan	CH	N1		
J8520	Capecitabine, oral, 150 mg		K2		\$5.68
J8521	Capecitabine, oral, 500 mg		K2		\$18.73
J8530	Cyclophosphamide oral 25 MG		N1		
J8540	Oral dexamethasone		N1		
J8560	Etoposide oral 50 MG		K2		\$0.45
J8597	Antiemetic drug oral NOS		N1		
J8600	Melphalan oral 2 MG		N1		
J8510	Methotrexate oral 2.5 MG		N1		
J8650	Nabifone oral	CH	N1		
J8700	Topotecan oral		K2		\$8.59
J8705	Topotecan oral		K2		\$71.35
J9000	Doxorubicin hcl injection		N1		
J9001	Doxorubicin hcl liposome inj		K2		\$450.51
J9010	Alemtuzumab injection		K2		\$559.46

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J7030	Normal saline solution infus		N1		
J7040	Normal saline solution infus		N1		
J7042	5% dextrose/normal saline		N1		
J7050	Normal saline solution infus		N1		
J7060	5% dextrose/water		N1		
J7070	D5w infusion		N1		
J7100	Dextran 40 infusion		N1		
J7110	Dextran 75 infusion		N1		
J7120	Fingers lactate infusion		N1		
J7130	Hypertonic saline solution		N1		
J7185	Xyrista inj	NI	K2		\$1.06
J7186	Antihemophilic viii/wf comp		K2		\$0.84
J7187	Humate-P, inj		K2		\$0.87
J7189	Factor vlla		K2		\$1.29
J7190	Factor viii		K2		\$0.84
J7191	Factor VIII (porcine)		K2		\$2.00
J7192	Factor viii recombinant NOS		K2		\$1.08
J7193	Factor IX non-recombinant		K2		\$0.88
J7194	Factor ix complex		K2		\$0.85
J7195	Factor IX recombinant		K2		\$1.06
J7197	Antithrombin iii injection		K2		\$2.28
J7198	Anti-inhibitor		K2		\$1.53
J7308	Aminolevulinic acid hcl top		K2		\$127.60
J7310	Ganciclovir long act implant		K2		\$16,640.00
J7311	Fluocinolone acetonide impit		K2		\$18,960.00
J7321	Hyalgan/supartz inj per dose		K2		\$91.87
J7322	Synvisc inj per dose	CH	D5		
J7323	Euflexa inj per dose		K2		\$113.96
J7324	Orthovisc inj per dose		K2		\$177.68
J7325	Synvisc or Synvisc-One	NI	K2		\$11.47
J7500	Azathioprine oral 50mg		N1		
J7501	Azathioprine parenteral		K2		\$90.64
J7502	Cyclosporine oral 100 mg		K2		\$3.22

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J9151	Daurorubicin citrate inj		K2		\$55.27
J9155	Degarelix injection	NI	K2		\$2.23
J9160	Denileukin difitox inj		K2		\$1,448.32
J9165	Diethylstilbestrol injection		K2		\$1,257.36
J9170	Docetaxel injection	CH	D5		
J9171	Docetaxel injection	NI	K2		\$16.95
J9175	Eliquis b solution per ml		N1		
J9178	Inj. egtrubicin hcl, 2 mg		K2		\$2.55
J9181	Etoposide injection		N1		
J9185	Fludarabine phosphate inj		K2		\$151.36
J9190	Fluorouracil injection		N1		
J9200	Floxuridine injection		K2		\$46.60
J9201	Gemcitabine hcl injection		K2		\$139.10
J9202	Goserelin acetate implant		K2		\$193.02
J9206	Irinotecan injection		K2		\$13.18
J9207	Ixabepilone injection		K2		\$63.74
J9208	Ifostomide injection		K2		\$29.39
J9209	Mesna injection		K2		\$4.34
J9211	Idarubicin hcl injection		K2		\$96.70
J9212	Interferon alfacon-1 inj	CH	K2		\$6.75
J9213	Interferon alfa-2a inj		K2		\$10.60
J9214	Interferon alfa-2b inj		K2		\$15.54
J9215	Interferon alfa-n3 inj		K2		\$17.89
J9216	Interferon gamma 1-b inj		K2		\$294.03
J9217	Leuprolide acetate suspension		K2		\$210.52
J9218	Leuprolide acetate injection		K2		\$5.29
J9219	Leuprorelin acetate implant		K2		\$4,726.88
J9225	Vanas implant		K2		\$1,473.60
J9226	Supprelin LA implant		K2		\$14,875.43
J9230	Mechlorethamine hcl inj		K2		\$144.56
J9245	Inj. melphalan hydrochl 50 MG		K2		\$1,622.81
J9260	Methotrexate sodium inj		N1		
J9260	Methotrexate sodium inj		N1		

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Note 2: Payment indicators for radiology services (Z2, Z3) are based on a comparison of the final rates according to the ASC standard reassigning methodology and the MPFS. Under current law, the MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
J9015	Adesleukin injection		K2		\$831.49
J9017	Arsenic trioxide injection		K2		\$36.73
J9020	Asparaginase injection		K2		\$56.92
J9025	Azacitidine injection		K2		\$4.78
J9027	Cicafibrate injection		K2		\$114.21
J9031	Bag live intravesical vac		K2		\$111.06
J9033	Bendamustine injection		K2		\$18.53
J9035	Bevacizumab injection		K2		\$56.39
J9040	Bleomycin sulfate injection		N1		
J9041	Bortezomib injection		K2		\$36.54
J9045	Carboplatin injection		N1		
J9050	Carmustine injection		K2		\$173.73
J9055	Cetuximab injection		K2		\$48.79
J9062	Cisplatin 10 MG injection		N1		
J9062	Cisplatin 50 MG injection		N1		
J9065	Inj cladribine per 1 MG		K2		\$25.15
J9070	Cyclophosphamide 100 MG inj		N1		
J9080	Cyclophosphamide 200 MG inj		N1		
J9090	Cyclophosphamide 500 MG inj		N1		
J9091	Cyclophosphamide 1.0 grm inj		N1		
J9092	Cyclophosphamide 2.0 grm inj		N1		
J9093	Cyclophosphamide lyophilized		N1		
J9094	Cyclophosphamide lyophilized		N1		
J9095	Cyclophosphamide lyophilized		N1		
J9096	Cyclophosphamide lyophilized		N1		
J9097	Cyclophosphamide lyophilized		N1		
J9098	Cyclophosphamide lyophilized		N1		
J9100	Cyclophosphamide lyophilized		N1		
J9110	Cytarabine hcl 100 MG inj		N1		\$480.19
J9120	Dactinomycin injection		K2		\$533.21
J9130	Dacarbazine 100 mg inj		N1		
J9140	Dacarbazine 200 MG inj		N1		
J9150	Daurorubicin injection		K2		\$14.95

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Note 2: Payment indicators for radiology services (Z2, Z3) are based on a comparison of the final rates according to the ASC standard reassigning methodology and the MPFS. Under current law, the MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
L8606	Synthetic implant urinary 1ml		N1		
L8609	Artificial cornea		N1		
L8610	Ocular implant		N1		
L8612	Aqueous shunt prosthesis		N1		
L8613	Ocular implant		N1		
L8630	Metacarpophalangeal implant		N1		
L8631	MCP joint repl 2 pc or more		N1		
L8641	Metatarsal joint implant		N1		
L8642	Hallux implant		N1		
L8658	Interphalangeal joint spacer		N1		
L8659	Interphalangeal joint repl		N1		
L8670	Vascular graft, synthetic		N1		
L8682	Implt neurostim radiotr rec		N1		
L8690	Aud osseo dev, intext comp		N1		
L8699	Prosthetic implant NOS		N1		
P9041	Albumin (human), 5%, 50ml		K2		\$16.89
P9045	Albumin (human), 5%, 250 ml		K2		\$60.58
P9046	Albumin (human), 25%, 20 ml		K2		\$25.67
P9047	Albumin (human), 25%, 50ml		K2		\$62.05
C0138	Ferumoxytol, non-esrd	NI	K2		\$0.82
C0163	Diphenhydramine HCl 50mg		N1		
C0164	Prochlorperazine maleate 5mg		N1		
C0166	Granisetron hcl 1 mg oral	CH	N1		
C0167	Dronabinol 2.5mg oral		N1		
C0169	Promethazine HCl 12.5mg oral		N1		
C0171	Chlorpromazine HCl 10mg oral		N1		
C0173	Trimethoprimazole HCl 250mg		N1		
C0174	Triethylperazine maleate 10mg		N1		
C0175	Perphenazine 4mg oral		N1		
C0177	Hydroxyzine pamoate 25mg		N1		
C0179	Ondansetron hcl 8 mg oral	CH	N1		
C0180	Dolasetron mesylate oral	CH	N1		

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Note 2: Payment indicators for radiology services (Z2, Z3) are based on a comparison of the final rates according to the ASC standard ratemaking methodology and the MPFS. Under current law, the MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HCPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Weight	CY 2010 Third Year Transition Payment
J9261	Neitarabine injection		K2		\$101.28
J9263	Oxaliplatin		K2		\$9.55
J9264	Paclitaxel protein bound		K2		\$9.09
J9265	Paclitaxel injection	CH	N1		
J9266	Pegaspargase injection		K2		\$2,695.67
J9268	Pentostatin injection		K2		\$1,399.56
J9270	Plicamycin (miframycin) [n]		N1		
J9280	Mitomycin 5 MG inj		K2		\$17.74
J9290	Mitomycin 20 MG inj		K2		\$70.98
J9291	Mitomycin 40 MG inj		K2		\$141.95
J9293	Mitoxantrone hydrochl / 5 MG		K2		\$65.51
J9300	Gemtuzumab ozogamion [n]		K2		\$2,572.82
J9303	Panitumumab injection		K2		\$85.21
J9305	Pemetrexed injection		K2		\$48.50
J9310	Rituximab injection		K2		\$52.70
J9320	Streptozocin injection		K2		\$278.35
J9328	Temozolomide injection		K2		\$4.90
J9340	Thiotepa injection		K2		\$47.93
J9350	Topotecan injection		K2		\$988.88
J9355	Trastuzumab injection		K2		\$63.51
J9357	Vindesine injection		K2		\$953.16
J9360	Vinblastine sulfate [n]		N1		
J9370	Vincristine sulfate 1 MG inj		N1		
J9375	Vincristine sulfate 2 MG inj		N1		
J9380	Vincristine sulfate 5 MG inj		N1		
J9380	Vincristine tartrate inj		N1		
J9395	Injection, Fulvestrant	CH	N1		
J9600	Porfimer sodium injection		K2		\$80.83
J9999	Chemotherapy drug		K2		\$2,745.46
L8600	Implant breast silicone/eq		N1		
L8603	Collagen imp urinary 2.5 ml		N1		
L8604	Dextranomer/hyaluronic acid		N1		

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Note 2: Payment indicators for radiology services (Z2, Z3) are based on a comparison of the final rates according to the ASC standard ratemaking methodology and the MPFS. Under current law, the MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
C9960	HOCM 200-249mg/ml iodine, 1ml		N1		
C9961	HOCM 250-299mg/ml iodine, 1ml		N1		
C9962	HOCM 300-349mg/ml iodine, 1ml		N1		
C9963	HOCM 350-399mg/ml iodine, 1ml		N1		
C9964	HOCM >= 400mg/ml iodine, 1ml		N1		
C9965	LOCM 100-199mg/ml iodine, 1ml		N1		
C9966	LOCM 200-299mg/ml iodine, 1ml		N1		
C9967	LOCM 300-399mg/ml iodine, 1ml		N1		
C9968	Visualization adjunct	NI	K2		\$4.11
V2630	Anter chamber, intraocular lens		N1		
V2631	Iris support, intraocular lens		N1		
V2632	Post chamber, intraocular lens		N1		
V2752	Corneal tissue processing		F4		
V2790	Amniotic membrane		N1		

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Note 2: Payment indicators for radiology services (Z2, Z3) are based on a comparison of the final rates according to the ASC standard assessing methodology and the MPFS. Under current law, the MPFS payment rates will have a negative update for CY 2010. For a discussion of those rates, we refer readers to the CY 2010 MPFS final rule.

Addendum BB -- Final ASC Covered Ancillary Services Integral to Covered Surgical Procedures for CY 2010 (Including Ancillary Services for Which Payment is Packaged)

HPCS Code	Short Descriptor	CY 2010 Comment Indicator	CY 2010 Payment Indicator	CY 2010 Third Year Transition Payment Weight	CY 2010 Third Year Transition Payment
Q0515	Sermorelin acetate injection		K2		\$1.77
Q1003	Nicot category 3		L6		\$50.00
Q2004	Bladder calculus irrig sol	CH	K2		\$29.28
Q2009	Fosphenytoin, inj PE		N1		
Q2017	Teniposide, 50 mg		K2		\$319.43
Q2023	Xyrista, inj	CH	D5		
Q2024	Bevacizumab injection	CH	D5		
C9025	IM inj interferon beta 1-a		K2		\$187.24
Q4100	Skin substitute, NOS		N1		
Q4101	Apilgraf skin sub		K2		\$32.16
Q4102	Oasis wound matrix skin sub		K2		\$4.12
Q4103	Oasis burn matrix skin sub		K2		\$4.12
Q4104	Integra BMWD skin sub		K2		\$11.77
Q4105	Integra DRT skin sub		K2		\$11.77
Q4106	Dermagraft skin sub		K2		\$39.25
Q4107	Graftjacket skin sub		K2		\$89.23
Q4108	Integra matrix skin sub		K2		\$17.99
Q4109	Tissuemend skin sub		N1		
Q4110	Primatrix skin sub		K2		\$33.99
Q4111	Gamma graft skin sub		K2		\$7.13
Q4112	Cymetra allograft		K2		\$327.47
Q4113	Graftjacket express allograft		K2		\$327.47
Q4114	Integra flowable wound matr		K2		\$907.36
Q4115	Alloskin skin sub		K2		\$9.36
Q4116	Alcoderm skin sub		K2		\$31.72
C9951	LOCM >= 400 mg/ml iodine, 1ml		N1		
C9953	Inj Fe-based MR contrast, 1ml		N1		
C9954	Oral MR contrast, 100 ml		N1		
C9955	Inj perflhexane lip micros, ml		N1		
C9956	Inj octafluoropropane mic, ml		N1		
C9957	Inj perfluren lip micros, ml		N1		
C9958	HOCM <= 149 mg/ml iodine, 1ml		N1		
C9959	HOCM 150-199mg/ml iodine, 1ml		N1		

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ADDENDUM D1.—FINAL OPPS PAYMENT STATUS INDICATORS FOR CY 2010

Indicator	Item/Code/Service	OPPS Payment Status
D	Discontinued Codes	Not paid under OPSS or any other Medicare payment system.
E	Items, Codes, and Services: <ul style="list-style-type: none"> • That are not covered by any Medicare outpatient benefit based on statutory exclusion. • That are not covered by any Medicare outpatient benefit for reasons other than statutory exclusion. • That are not recognized by Medicare for outpatient claims but for which an alternate code for the same item or service may be available. • For which separate payment is not provided on outpatient claims. 	Not paid by Medicare when submitted on outpatient claims (any outpatient bill type).
F	Corneal Tissue Acquisition; Certain CRNA Services and Hepatitis B Vaccines	Not paid under OPSS. Paid at reasonable cost.
G	Pass-Through Drugs and Biologicals	Paid under OPSS; separate APC payment.
H	Pass-Through Device Categories	Separate cost-based pass-through payment; not subject to copayment.
K	Nonpass-Through Drugs and Nontimplantable Biologicals, Including Therapeutic Radiopharmaceuticals	Paid under OPSS; separate APC payment.
L	Influenza Vaccine; Pneumococcal Pneumonia Vaccine	Not paid under OPSS. Paid at reasonable cost; not subject to deductible or coinsurance.
M	Items and Services Not Billable to the Fiscal Intermediary/MAC	Not paid under OPSS.
N	Items and Services Packaged into APC Rates	Paid under OPSS; payment is packaged into payment for other services. Therefore, there is no separate APC payment.

ADDENDUM D1.—FINAL OPSS PAYMENT STATUS INDICATORS FOR CY 2010

Indicator	Item/Code/Service	OPSS Payment Status
A	Services furnished to a hospital outpatient that are paid under a fee schedule or payment system other than OPSS, for example: <ul style="list-style-type: none"> • Ambulance Services • Clinical Diagnostic Laboratory Services • Non-Implantable Prosthetic and Orthotic Devices • EPO for ESRD Patients • Physical, Occupational, and Speech Therapy • Routine Dialysis Services for ESRD Patients Provided in a Certified Dialysis Unit of a Hospital • Diagnostic Mammography • Screening Mammography 	Not paid under OPSS. Paid by fiscal intermediaries/MACs under a fee schedule or payment system other than OPSS. Not subject to deductible or coinsurance.
B	Codes that are not recognized by OPSS when submitted on an outpatient hospital Part B bill type (12x and 13x).	Not paid under OPSS. <ul style="list-style-type: none"> • May be paid by fiscal intermediaries/MACs when submitted on a different bill type, for example, 75x (CORF), but not paid under OPSS. • An alternate code that is recognized by OPSS when submitted on an outpatient hospital Part B bill type (12x and 13x) may be available.
C	Inpatient Procedures	Not paid under OPSS. Admit patient. Bill as inpatient.

ADDENDUM D1.—FINAL OPPTS PAYMENT STATUS INDICATORS FOR CY 2010		
Indicator	Item/Code/Service	OPPS Payment Status
S	Significant Procedure, Not Discounted When Multiple	Paid under OPPTS; separate APC payment.
T	Significant Procedure, Multiple Reduction Applies	Paid under OPPTS; separate APC payment.
U	Brachytherapy Sources	Paid under OPPTS; separate APC payment.
V	Clinic or Emergency Department Visit	Paid under OPPTS; separate APC payment.
X	Ancillary Services	Paid under OPPTS; separate APC payment.
Y	Non-Implantable Durable Medical Equipment	Not paid under OPPTS. All institutional providers other than home health agencies bill to DMERC.

ADDENDUM DD1.—FINAL ASC PAYMENT INDICATORS FOR CY 2010	
Indicator	Payment Indicator Definition
A2	Surgical procedure on ASC list in CY 2007; payment based on OPPTS relative payment weight.
D5	Deleted/discontinued code; no payment made.
F4	Corneal tissue acquisition, hepatitis B vaccine; paid at reasonable cost.
G2	Non office-based surgical procedure added in CY 2008 or later; payment based on OPPTS relative payment weight.
H2	Brachytherapy source paid separately when provided integral to a surgical procedure on ASC list; payment based on OPPTS rate.
H8	Device-intensive procedure on ASC list in CY 2007; paid at adjusted rate.
J7	OPPTS pass-through device paid separately when provided integral to a surgical procedure on ASC list; payment contractor-priced.

ADDENDUM D1.—FINAL OPPTS PAYMENT STATUS INDICATORS FOR CY 2010		
Indicator	Item/Code/Service	OPPS Payment Status
P	Partial Hospitalization	Paid under OPPTS; per diem APC payment.
Q1	STVX-Packaged Codes	Paid under OPPTS; Addendum B displays APC assignments when services are separately payable. (1) Packaged APC payment if billed on the same date of service as a HCPCS code assigned status indicator "S," "T," "V," or "X." (2) In all other circumstances, payment is made through a separate APC payment.
Q2	T-Packaged Codes	Paid under OPPTS; Addendum B displays APC assignments when services are separately payable. (1) Packaged APC payment if billed on the same date of service as a HCPCS code assigned status indicator "T." (2) In all other circumstances, payment is made through a separate APC payment.
Q3	Codes That May Be Paid Through a Composite APC	Paid under OPPTS; Addendum B displays APC assignments when services are separately payable. Addendum M displays composite APC assignments when codes are paid through a composite APC. (1) Composite APC payment based on OPPTS composite-specific payment criteria. Payment is packaged into a single payment for specific combinations of service. (2) In all other circumstances, payment is made through a separate APC payment or packaged into payment for other services.
R	Blood and Blood Products	Paid under OPPTS; separate APC payment.

ADDENDUM D2.—FINAL OPPTS COMMENT INDICATORS FOR CY 2010	
Comment Indicator	Descriptor
NI	New code for the next calendar year or existing code with substantial revision to its code descriptor in the next calendar year as compared to current calendar year, interim APC assignment; comments will be accepted on the interim APC assignment for the new code.
CH	Active HCPCS code in current year and next calendar year, status indicator and/or APC assignment has changed; or active HCPCS code that will be discontinued at the end of the current calendar year.

ADDENDUM DD2.—FINAL ASC COMMENT INDICATORS FOR CY 2010	
CI	Comment Indicator Meanings
CH	Active HCPCS code in current year and next calendar year, payment indicator assignment has changed; or active HCPCS code that is newly recognized as payable in ASC; or active HCPCS code that is discontinued at the end of the current calendar year.
NI	New code for the next calendar year or existing code with substantial revision to its code descriptor in the next calendar year, interim payment indicator assignment; comments will be accepted on the interim payment indicator for the new code.

ADDENDUM DD1.—FINAL ASC PAYMENT INDICATORS FOR CY 2010	
Indicator	Payment Indicator Definition
J8	Device-intensive procedure added to ASC list in CY 2008 or later; paid at adjusted rate.
K2	Drugs and biologicals paid separately when provided integral to a surgical procedure on ASC list; payment based on OPSS rate.
K7	Unclassified drugs and biologicals; payment contractor-priced.
L1	Influenza vaccine; pneumococcal vaccine. Packaged item/service; no separate payment made.
L6	New Technology Intraocular Lens (NTIOL); special payment.
N1	Packaged service/item; no separate payment made.
P2	Office-based surgical procedure added to ASC list in CY 2008 or later with MPFS nonfacility PE RVUs; payment based on OPSS relative payment weight.
P3	Office-based surgical procedure added to ASC list in CY 2008 or later with MPFS nonfacility PE RVUs; payment based on MPFS nonfacility PE RVUs.
R2	Office-based surgical procedure added to ASC list in CY 2008 or later without MPFS nonfacility PE RVUs; payment based on OPSS relative payment weight.
Z2	Radiology service paid separately when provided integral to a surgical procedure on ASC list; payment based on OPSS relative payment weight.
Z3	Radiology service paid separately when provided integral to a surgical procedure on ASC list; payment based on MPFS nonfacility PE RVUs.

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
00176	Anesth, pharyngeal surgery	C	
00192	Anesth, facial bone surgery	C	
00211	Anesth, cran surg, hemotoma	C	
00214	Anesth, skull drainage	C	
00215	Anesth, skull repair/fract	C	
00452	Anesth, surgery of shoulder	C	
00474	Anesth, surgery of rib(s)	C	
00524	Anesth, chest drainage	C	
00540	Anesth, chest surgery	C	
00542	Anesth, release of lung	C	
00546	Anesth, lung, chest wall surg	C	
00560	Anesth, heart surg w/o pump	C	
00561	Anesth, heart surg < age 1	C	
00562	Anesth hrt surg w/pmp age 1+	C	
00567	Anesth, cabg w/pump	C	
00580	Anesth, heart/lung transplnt	C	
00604	Anesth, sitting procedure	C	
00622	Anesth, removal of nerves	C	
00632	Anesth, removal of nerves	C	
00670	Anesth, spine, cord surgery	C	
00792	Anesth, hemorr/excise liver	C	
00794	Anesth, pancreas removal	C	
00796	Anesth, for liver transplant	C	
00802	Anesth, fat layer removal	C	
00844	Anesth, pelvis surgery	C	
00846	Anesth, hysterectomy	C	
00848	Anesth, pelvic organ surg	C	
00864	Anesth, removal of bladder	C	
00865	Anesth, removal of prostate	C	
00866	Anesth, removal of adrenal	C	
00868	Anesth, kidney transplant	C	
00882	Anesth, major vein ligation	C	
00904	Anesth, perineal surgery	C	
00908	Anesth, removal of prostate	C	
00932	Anesth, amputation of penis	C	
00934	Anesth, penis, nodes removal	C	
00936	Anesth, penis, nodes removal	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
00944	Anesth, vaginal hysterectomy	C	
01140	Anesth, amputation at pelvis	C	
01150	Anesth, pelvic tumor surgery	C	
01212	Anesth, hip disarticulation	C	
01214	Anesth, hip arthroplasty	C	
01232	Anesth, amputation of femur	C	
01234	Anesth, radical femur surg	C	
01272	Anesth, femoral artery surg	C	
01274	Anesth, femoral embolectomy	C	
01402	Anesth, knee arthroplasty	C	
01404	Anesth, amputation at knee	C	
01442	Anesth, knee artery surg	C	
01444	Anesth, knee artery repair	C	
01486	Anesth, ankle replacement	C	
01502	Anesth, lwr leg embolectomy	C	
01634	Anesth, shoulder joint amput	C	
01636	Anesth, forequarter amput	C	
01638	Anesth, shoulder replacement	C	
01652	Anesth, shoulder vessel surg	C	
01654	Anesth, shoulder vessel surg	C	
01656	Anesth, arm-leg vessel surg	C	
01756	Anesth, radical humerus surg	C	
01990	Support for organ donor	C	
11004	Debride genitalia & perineum	C	
11005	Debride abdom wall	C	
11006	Debride genit/per/abdom wall	C	
11008	Remove mesh from abd wall	C	
15756	Free myo/skin flap microvasc	C	
15757	Free skin flap, microvasc	C	
15758	Free fascial flap, microvasc	C	
16036	Escharotomy; addl incision	C	
19271	Revision of chest wall	C	
19272	Extensive chest wall surgery	C	
19305	Mast, radical	C	
19306	Mast, rad, urban type	C	
19361	Breast reconstr w/lat flap	C	
19364	Breast reconstruction	C	
19367	Breast reconstruction	C	
19368	Breast reconstruction	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
19369	Breast reconstruction	C	
20661	Application of head brace	C	
20664	Halo brace application	C	
20802	Replantation, arm, complete	C	
20805	Replant forearm, complete	C	
20808	Replantation hand, complete	C	
20816	Replantation digit, complete	C	
20824	Replantation thumb, complete	C	
20827	Replantation thumb, complete	C	
20838	Replantation foot, complete	C	
20930	Sp bone agrft morsel add-on	C	
20931	Sp bone agrft struct add-on	C	
20936	Sp bone agrft local add-on	C	
20937	Sp bone agrft morsel add-on	C	
20938	Sp bone agrft struct add-on	C	
20955	Fibula bone graft, microvasc	C	
20956	Iliac bone graft, microvasc	C	
20957	Mt bone graft, microvasc	C	
20962	Other bone graft, microvasc	C	
20969	Bone/skin graft, microvasc	C	
20970	Bone/skin graft, iliac crest	C	
21045	Extensive jaw surgery	C	
21141	Reconstruct midface, lefort	C	
21142	Reconstruct midface, lefort	C	
21143	Reconstruct midface, lefort	C	
21145	Reconstruct midface, lefort	C	
21146	Reconstruct midface, lefort	C	
21147	Reconstruct midface, lefort	C	
21151	Reconstruct midface, lefort	C	
21154	Reconstruct midface, lefort	C	
21155	Reconstruct midface, lefort	C	
21159	Reconstruct midface, lefort	C	
21160	Reconstruct midface, lefort	C	
21179	Reconstruct entire forehead	C	
21180	Reconstruct entire forehead	C	
21182	Reconstruct cranial bone	C	
21183	Reconstruct cranial bone	C	
21184	Reconstruct cranial bone	C	
21188	Reconstruction of midface	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
21193	Reconst lwr jaw w/o graft	C	
21194	Reconst lwr jaw w/graft	C	
21196	Reconst lwr jaw w/fixation	C	
21247	Reconstruct lower jaw bone	C	
21255	Reconstruct lower jaw bone	C	
21268	Revise eye sockets	C	
21343	Treatment of sinus fracture	C	
21344	Treatment of sinus fracture	C	
21346	Treat nose/jaw fracture	C	
21347	Treat nose/jaw fracture	C	
21348	Treat nose/jaw fracture	C	
21366	Treat cheek bone fracture	C	
21395	Treat eye socket fracture	C	
21422	Treat mouth roof fracture	C	
21423	Treat mouth roof fracture	C	
21431	Treat craniofacial fracture	C	
21432	Treat craniofacial fracture	C	
21433	Treat craniofacial fracture	C	
21435	Treat craniofacial fracture	C	
21436	Treat craniofacial fracture	C	
21510	Drainage of bone lesion	C	
21615	Removal of rib	C	
21616	Removal of rib and nerves	C	
21620	Partial removal of sternum	C	
21627	Sternal debridement	C	
21630	Extensive sternum surgery	C	
21632	Extensive sternum surgery	C	
21705	Revision of neck muscle/rib	C	
21740	Reconstruction of sternum	C	
21750	Repair of sternum separation	C	
21810	Treatment of rib fracture(s)	C	
21825	Treat sternum fracture	C	
22010	I&d, p-spine, c/t/cerv-thor	C	
22015	I&d, p-spine, l/s/l	C	
22110	Remove part of neck vertebra	C	
22112	Remove part, thorax vertebra	C	
22114	Remove part, lumbar vertebra	C	
22116	Remove extra spine segment	C	
22206	Cut spine 3 col, thor	C	
22207	Cut spine 3 col, lumb	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
22208	Cut spine 3 col, addl seg	C	
22210	Revision of neck spine	C	
22212	Revision of thorax spine	C	
22214	Revision of lumbar spine	C	
22216	Revise, extra spine segment	C	
22220	Revision of neck spine	C	
22224	Revision of lumbar spine	C	
22226	Revise, extra spine segment	C	
22318	Treat odontoid fx w/o graft	C	
22319	Treat odontoid fx w/graft	C	
22325	Treat spine fracture	C	
22326	Treat neck spine fracture	C	
22327	Treat thorax spine fracture	C	
22328	Treat each add spine fx	C	
22532	Lat thorax spine fusion	C	
22533	Lat lumbar spine fusion	C	
22534	Lat thor/lumb, addl seg	C	
22548	Neck spine fusion	C	
22554	Neck spine fusion	C	
22556	Thorax spine fusion	C	
22558	Lumbar spine fusion	C	
22585	Additional spinal fusion	C	
22590	Spine & skull spinal fusion	C	
22595	Neck spinal fusion	C	
22600	Neck spine fusion	C	
22610	Thorax spine fusion	C	
22630	Lumbar spine fusion	C	
22632	Spine fusion, extra segment	C	
22800	Fusion of spine	C	
22802	Fusion of spine	C	
22804	Fusion of spine	C	
22808	Fusion of spine	C	
22810	Fusion of spine	C	
22812	Fusion of spine	C	
22818	Kyphectomy, 1-2 segments	C	
22819	Kyphectomy, 3 or more	C	
22830	Exploration of spinal fusion	C	
22840	Insert spine fixation device	C	
22841	Insert spine fixation device	C	
22842	Insert spine fixation device	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
22843	Insert spine fixation device	C	
22844	Insert spine fixation device	C	
22845	Insert spine fixation device	C	
22846	Insert spine fixation device	C	
22847	Insert spine fixation device	C	
22848	Insert pelv fixation device	C	
22849	Reinsert spinal fixation	C	
22850	Remove spine fixation device	C	
22852	Remove spine fixation device	C	
22855	Remove spine fixation device	C	
22856	Cerv artific diskectomy	C	
22857	Lumbar artif diskectomy	C	
22861	Revise cerv artific disc	C	
22862	Revise lumbar artif disc	C	
22864	Remove cerv artif disc	C	
22865	Remove lumb artif disc	C	
23200	Resect clavicle tumor	C	
23210	Resect scapula tumor	C	
23220	Resect prox humerus tumor	C	
23332	Remove shoulder foreign body	C	
23472	Reconstruct shoulder joint	C	
23900	Amputation of arm & girdle	C	
23920	Amputation at shoulder joint	C	
24900	Amputation of upper arm	C	
24920	Amputation of upper arm	C	
24930	Amputation follow-up surgery	C	
24931	Amputate upper arm & implant	C	
24940	Revision of upper arm	C	
25900	Amputation of forearm	C	
25905	Amputation of forearm	C	
25909	Amputation follow-up surgery	C	
25915	Amputation of forearm	C	
25920	Amputate hand at wrist	C	
25924	Amputation follow-up surgery	C	
25927	Amputation of hand	C	
26551	Great toe-hand transfer	C	
26553	Single transfer, toe-hand	C	
26554	Double transfer, toe-hand	C	
26556	Toe joint transfer	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
26992	Drainage of bone lesion	C	
27005	Incision of hip tendon	C	
27025	Incision of hip/thigh fascia	C	
27030	Drainage of hip joint	C	
27036	Excision of hip joint/muscle	C	
27054	Removal of hip joint lining	C	
27070	Partial removal of hip bone	C	
27071	Partial removal of hip bone	C	
27075	Resect hip tumor	C	
27076	Resect hip tum incl acetabul	C	
27077	Resect hip tum w/innom bone	C	
27078	Resect hip tum incl femur	C	
27090	Removal of hip prosthesis	C	
27091	Removal of hip prosthesis	C	
27120	Reconstruction of hip socket	C	
27122	Reconstruction of hip socket	C	
27125	Partial hip replacement	C	
27130	Total hip arthroplasty	C	
27132	Total hip arthroplasty	C	
27134	Revise hip joint replacement	C	
27137	Revise hip joint replacement	C	
27138	Revise hip joint replacement	C	
27140	Transplant femur ridge	C	
27146	Incision of hip bone	C	
27147	Revision of hip bone	C	
27151	Incision of hip bones	C	
27156	Revision of hip bones	C	
27158	Revision of pelvis	C	
27161	Incision of neck of femur	C	
27165	Incision/fixation of femur	C	
27170	Repair/graft femur head/neck	C	
27175	Treat slipped epiphysis	C	
27176	Treat slipped epiphysis	C	
27177	Treat slipped epiphysis	C	
27178	Treat slipped epiphysis	C	
27181	Treat slipped epiphysis	C	
27185	Revision of femur epiphysis	C	
27187	Reinforce hip bones	C	
27222	Treat hip socket fracture	C	
27226	Treat hip wall fracture	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
27227	Treat hip fracture(s)	C	
27228	Treat hip fracture(s)	C	
27232	Treat thigh fracture	C	
27236	Treat thigh fracture	C	
27240	Treat thigh fracture	C	
27244	Treat thigh fracture	C	
27245	Treat thigh fracture	C	
27248	Treat thigh fracture	C	
27253	Treat hip dislocation	C	
27254	Treat hip dislocation	C	
27258	Treat hip dislocation	C	
27259	Treat hip dislocation	C	
27268	Citx thigh fx w/mnpj	C	
27269	Optx thigh fx	C	
27280	Fusion of sacroiliac joint	C	
27282	Fusion of pubic bones	C	
27284	Fusion of hip joint	C	
27286	Fusion of hip joint	C	
27290	Amputation of leg at hip	C	
27295	Amputation of leg at hip	C	
27303	Drainage of bone lesion	C	
27365	Resect femur/knee tumor	C	
27445	Revision of knee joint	C	
27447	Total knee arthroplasty	C	
27448	Incision of thigh	C	
27450	Incision of thigh	C	
27454	Realignment of thigh bone	C	
27455	Realignment of knee	C	
27457	Realignment of knee	C	
27465	Shortening of thigh bone	C	
27466	Lengthening of thigh bone	C	
27468	Shorten/lengthen thighs	C	
27470	Repair of thigh	C	
27472	Repair/graft of thigh	C	
27477	Surgery to stop leg growth	C	
27485	Surgery to stop leg growth	C	
27486	Revise/replace knee joint	C	
27487	Revise/replace knee joint	C	
27488	Removal of knee prosthesis	C	
27495	Reinforce thigh	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
27506	Treatment of thigh fracture	C	
27507	Treatment of thigh fracture	C	
27511	Treatment of thigh fracture	C	
27513	Treatment of thigh fracture	C	
27514	Treatment of thigh fracture	C	
27519	Treat thigh fx growth plate	C	
27535	Treat knee fracture	C	
27536	Treat knee fracture	C	
27540	Treat knee fracture	C	
27556	Treat knee dislocation	C	
27557	Treat knee dislocation	C	
27558	Treat knee dislocation	C	
27580	Fusion of knee	C	
27590	Amputate leg at thigh	C	
27591	Amputate leg at thigh	C	
27592	Amputate leg at thigh	C	
27596	Amputation follow-up surgery	C	
27598	Amputate lower leg at knee	C	
27645	Resect tibia tumor	C	
27646	Resect fibula tumor	C	
27702	Reconstruct ankle joint	C	
27703	Reconstruct, ankle joint	C	
27712	Realignment of lower leg	C	
27715	Revision of lower leg	C	
27724	Repair/graft of tibia	C	
27725	Repair of lower leg	C	
27727	Repair of lower leg	C	
27880	Amputation of lower leg	C	
27881	Amputation of lower leg	C	
27882	Amputation of lower leg	C	
27886	Amputation follow-up surgery	C	
27888	Amputation of foot at ankle	C	
28800	Amputation of midfoot	C	
31225	Removal of upper jaw	C	
31230	Removal of upper jaw	C	
31290	Nasal/sinus endoscopy, surg	C	
31291	Nasal/sinus endoscopy, surg	C	
31360	Removal of larynx	C	
31365	Removal of larynx	C	
31367	Partial removal of larynx	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
31368	Partial removal of larynx	C	
31370	Partial removal of larynx	C	
31375	Partial removal of larynx	C	
31380	Partial removal of larynx	C	
31382	Partial removal of larynx	C	
31390	Removal of larynx & pharynx	C	
31395	Reconstruct larynx & pharynx	C	
31584	Treat larynx fracture	C	
31587	Revision of larynx	C	
31725	Clearance of airways	C	
31760	Repair of windpipe	C	
31766	Reconstruction of windpipe	C	
31770	Repair/graft of bronchus	C	
31775	Reconstruct bronchus	C	
31780	Reconstruct windpipe	C	
31781	Reconstruct windpipe	C	
31786	Remove windpipe lesion	C	
31800	Repair of windpipe injury	C	
31805	Repair of windpipe injury	C	
32035	Exploration of chest	C	
32036	Exploration of chest	C	
32095	Biopsy through chest wall	C	
32100	Exploration/biopsy of chest	C	
32110	Explore/repair chest	C	
32120	Re-exploration of chest	C	
32124	Explore chest free adhesions	C	
32140	Removal of lung lesion(s)	C	
32141	Remove/treat lung lesions	C	
32150	Removal of lung lesion(s)	C	
32151	Remove lung foreign body	C	
32160	Open chest heart massage	C	
32200	Drain, open, lung lesion	C	
32215	Treat chest lining	C	
32220	Release of lung	C	
32225	Partial release of lung	C	
32310	Removal of chest lining	C	
32320	Free/remove chest lining	C	
32402	Open biopsy chest lining	C	
32440	Removal of lung	C	
32442	Sleeve pneumonectomy	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
32445	Removal of lung	C		
32480	Partial removal of lung	C		
32482	Bilobectomy	C		
32484	Segmentectomy	C		
32486	Sleeve lobectomy	C		
32488	Completion pneumonectomy	C		
32491	Lung volume reduction	C		
32500	Partial removal of lung	C		
32501	Repair bronchus add-on	C		
32503	Resect apical lung tumor	C		
32504	Resect apical lung tum/chest	C		
32540	Removal of lung lesion	C		
32650	Thoracoscopy, surgical	C		
32651	Thoracoscopy, surgical	C		
32652	Thoracoscopy, surgical	C		
32653	Thoracoscopy, surgical	C		
32654	Thoracoscopy, surgical	C		
32655	Thoracoscopy, surgical	C		
32656	Thoracoscopy, surgical	C		
32657	Thoracoscopy, surgical	C		
32658	Thoracoscopy, surgical	C		
32659	Thoracoscopy, surgical	C		
32660	Thoracoscopy, surgical	C		
32661	Thoracoscopy, surgical	C		
32662	Thoracoscopy, surgical	C		
32663	Thoracoscopy, surgical	C		
32664	Thoracoscopy, surgical	C		
32665	Thoracoscopy, surgical	C		
32800	Repair lung hernia	C		
32810	Close chest after drainage	C		
32815	Close bronchial fistula	C		
32820	Reconstruct injured chest	C		
32850	Donor pneumonectomy	C		
32851	Lung transplant, single	C		
32852	Lung transplant with bypass	C		
32853	Lung transplant, double	C		
32854	Lung transplant with bypass	C		
32855	Prepare donor lung, single	C		
32856	Prepare donor lung, double	C		
32900	Removal of rib(s)	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
32905	Revise & repair chest wall	C		
32906	Revise & repair chest wall	C		
32940	Revision of lung	C		
32997	Total lung lavage	C		
33015	Incision of heart sac	C		
33020	Incision of heart sac	C		
33025	Incision of heart sac	C		
33030	Partial removal of heart sac	C		
33031	Partial removal of heart sac	C		
33050	Removal of heart sac lesion	C		
33120	Removal of heart lesion	C		
33130	Removal of heart lesion	C		
33140	Heart revascularize (tmr)	C		
33141	Heart tmr w/other procedure	C		
33202	Insert epicard eltrd, open	C		
33203	Insert epicard eltrd, endo	C		
33236	Remove electrode/thoracotomy	C		
33237	Remove electrode/thoracotomy	C		
33238	Remove electrode/thoracotomy	C		
33243	Remove eltrd/thoracotomy	C		
33250	Ablate heart dysrhythm focus	C		
33251	Ablate heart dysrhythm focus	C		
33254	Ablate atria, lmtd	C		
33255	Ablate atria w/o bypass, ext	C		
33256	Ablate atria w/bypass, exten	C		
33257	Ablate atria, lmtd, add-on	C		
33258	Ablate atria, x10sv, add-on	C		
33259	Ablate atria w/bypass add-on	C		
33261	Ablate heart dysrhythm focus	C		
33265	Ablate atria, lmtd, endo	C		
33266	Ablate atria, x10sv, endo	C		
33300	Repair of heart wound	C		
33305	Repair of heart wound	C		
33310	Exploratory heart surgery	C		
33315	Exploratory heart surgery	C		
33320	Repair major blood vessel(s)	C		
33321	Repair major blood vessel	C		
33322	Repair major blood vessel(s)	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
33330	Insert major vessel graft	C		
33332	Insert major vessel graft	C		
33335	Insert major vessel graft	C		
33400	Repair of aortic valve	C		
33401	Valvuloplasty, open	C		
33403	Valvuloplasty, w/cp bypass	C		
33404	Prepare heart-aorta conduit	C		
33405	Replacement of aortic valve	C		
33406	Replacement of aortic valve	C		
33410	Replacement of aortic valve	C		
33411	Replacement of aortic valve	C		
33412	Replacement of aortic valve	C		
33413	Replacement of aortic valve	C		
33414	Repair of aortic valve	C		
33415	Revision, subvalvular tissue	C		
33416	Revise ventricle muscle	C		
33417	Repair of aortic valve	C		
33420	Revision of mitral valve	C		
33422	Revision of mitral valve	C		
33425	Repair of mitral valve	C		
33426	Repair of mitral valve	C		
33427	Repair of mitral valve	C		
33430	Replacement of mitral valve	C		
33460	Revision of tricuspid valve	C		
33463	Valvuloplasty, tricuspid	C		
33464	Valvuloplasty, tricuspid	C		
33465	Replace tricuspid valve	C		
33468	Revision of tricuspid valve	C		
33470	Revision of pulmonary valve	C		
33471	Valvotomy, pulmonary valve	C		
33472	Revision of pulmonary valve	C		
33474	Revision of pulmonary valve	C		
33475	Replacement, pulmonary valve	C		
33476	Revision of heart chamber	C		
33478	Revision of heart chamber	C		
33496	Repair, prosth valve clot	C		
33500	Repair heart vessel fistula	C		
33501	Repair heart vessel fistula	C		
33502	Coronary artery correction	C		
33503	Coronary artery graft	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
33504	Coronary artery graft	C		
33505	Repair artery w/tunnel	C		
33506	Repair artery, translocation	C		
33507	Repair art, intramural	C		
33510	CABG, vein, single	C		
33511	CABG, vein, two	C		
33512	CABG, vein, three	C		
33513	CABG, vein, four	C		
33514	CABG, vein, five	C		
33516	Cabg, vein, six or more	C		
33517	CABG, artery-vein, single	C		
33518	CABG, artery-vein, two	C		
33519	CABG, artery-vein, three	C		
33521	CABG, artery-vein, four	C		
33522	CABG, artery-vein, five	C		
33523	Cabg, art-vein, six or more	C		
33530	Coronary artery, bypass/reop	C		
33533	CABG, arterial, single	C		
33534	CABG, arterial, two	C		
33535	CABG, arterial, three	C		
33536	Cabg, arterial, four or more	C		
33542	Removal of heart lesion	C		
33545	Repair of heart damage	C		
33548	Restore/remodel, ventricle	C		
33572	Open coronary endarterectomy	C		
33600	Closure of valve	C		
33602	Closure of valve	C		
33606	Anastomosis/artery-aorta	C		
33608	Repair anomaly w/conduit	C		
33610	Repair by enlargement	C		
33611	Repair double ventricle	C		
33612	Repair double ventricle	C		
33615	Repair, modified fontan	C		
33617	Repair single ventricle	C		
33619	Repair single ventricle	C		
33641	Repair heart septum defect	C		
33645	Revision of heart veins	C		
33647	Repair heart septum defects	C		
33660	Repair of heart defects	C		
33665	Repair of heart defects	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
33670	Repair of heart chambers	C	
33675	Close mult vsd	C	
33676	Close mult vsd w/resection	C	
33677	Cl mult vsd w/rem pul band	C	
33681	Repair heart septum defect	C	
33684	Repair heart septum defect	C	
33688	Repair heart septum defect	C	
33690	Reinforce pulmonary artery	C	
33692	Repair of heart defects	C	
33694	Repair of heart defects	C	
33697	Repair of heart defects	C	
33702	Repair of heart defects	C	
33710	Repair of heart defects	C	
33720	Repair of heart defect	C	
33722	Repair of heart defect	C	
33724	Repair venous anomaly	C	
33726	Repair pul venous stenosis	C	
33730	Repair heart-vein defect(s)	C	
33732	Repair heart-vein defect	C	
33735	Revision of heart chamber	C	
33736	Revision of heart chamber	C	
33737	Revision of heart chamber	C	
33750	Major vessel shunt	C	
33755	Major vessel shunt	C	
33762	Major vessel shunt	C	
33764	Major vessel shunt & graft	C	
33766	Major vessel shunt	C	
33767	Major vessel shunt	C	
33768	Cavopulmonary shunting	C	
33770	Repair great vessels defect	C	
33771	Repair great vessels defect	C	
33774	Repair great vessels defect	C	
33775	Repair great vessels defect	C	
33776	Repair great vessels defect	C	
33777	Repair great vessels defect	C	
33778	Repair great vessels defect	C	
33779	Repair great vessels defect	C	
33780	Repair great vessels defect	C	
33781	Repair great vessels defect	C	
33782	Nikaidoh proc	C	NI

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
33783	Nikaidoh proc w/ostia implit	C	NI
33786	Repair arterial trunk	C	
33788	Revision of pulmonary artery	C	
33800	Aortic suspension	C	
33802	Repair vessel defect	C	
33803	Repair vessel defect	C	
33813	Repair septal defect	C	
33814	Repair septal defect	C	
33820	Revise major vessel	C	
33822	Revise major vessel	C	
33824	Revise major vessel	C	
33840	Remove aorta constriction	C	
33845	Remove aorta constriction	C	
33851	Remove aorta constriction	C	
33852	Repair septal defect	C	
33853	Repair septal defect	C	
33860	Ascending aortic graft	C	
33861	Ascending aortic graft	C	
33863	Ascending aortic graft	C	
33864	Ascending aortic graft	C	
33870	Transverse aortic arch graft	C	
33875	Thoracic aortic graft	C	
33877	Thoracoabdominal graft	C	
33880	Endovasc taa repr incl subcl	C	
33881	Endovasc taa repr w/o subcl	C	
33883	Insert endovasc prosth, taa	C	
33884	Endovasc prosth, taa, add-on	C	
33886	Endovasc prosth, delayed	C	
33889	Artery transpose/endovas taa	C	
33891	Car-car bp grt/endovas taa	C	
33910	Remove lung artery emboli	C	
33915	Remove lung artery emboli	C	
33916	Surgery of great vessel	C	
33917	Repair pulmonary artery	C	
33920	Repair pulmonary atresia	C	
33922	Transect pulmonary artery	C	
33924	Remove pulmonary shunt	C	
33925	Rpr pul art unifocal w/o cpb	C	
33926	Repr pul art, unifocal w/cpb	C	
33930	Removal of donor heart/lung	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
33933	Prepare donor heart/lung	C		
33935	Transplantation, heart/lung	C		
33940	Removal of donor heart	C		
33944	Prepare donor heart	C		
33945	Transplantation of heart	C		
33960	External circulation assist	C		
33961	External circulation assist	C		
33967	Insert ia percut device	C		
33968	Remove aortic assist device	C		
33970	Aortic circulation assist	C		
33971	Aortic circulation assist	C		
33973	Insert balloon device	C		
33974	Remove intra-aortic balloon	C		
33975	Implant ventricular device	C		
33976	Implant ventricular device	C		
33977	Remove ventricular device	C		
33978	Remove ventricular device	C		
33979	Insert intracorporeal device	C		
33980	Remove intracorporeal device	C		
33981	Replace vad pump ext	C	NI	
33982	Replace vad intra w/o bp	C	NI	
33983	Replace vad intra w/bp	C	NI	
34001	Removal of artery clot	C		
34051	Removal of artery clot	C		
34151	Removal of artery clot	C		
34401	Removal of vein clot	C		
34451	Removal of vein clot	C		
34502	Reconstruct vena cava	C		
34800	Endovas aaa repr w/sm tube	C		
34802	Endovas aaa repr w/2-p part	C		
34803	Endovas aaa repr w/3-p part	C		
34804	Endovas aaa repr w/1-p part	C		
34805	Endovas aaa repr w/long tube	C		
34806	Aneurysm press sensor add-on	C		
34808	Endovas iliac a device addon	C		
34812	Xpose for endoprosth, femoral	C		
34813	Femoral endovas graft add-on	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
34820	Xpose for endoprosth, iliac	C		
34825	Endovasc extend prosth, init	C		
34826	Endovasc exten prosth, addl	C		
34830	Open aortic tube prosth repr	C		
34831	Open aortoiliac prosth repr	C		
34832	Open aortofemor prosth repr	C		
34833	Xpose for endoprosth, iliac	C		
34834	Xpose, endoprosth, brachial	C		
34900	Endovasc iliac repr w/graft	C		
35001	Repair defect of artery	C		
35002	Repair artery rupture, neck	C		
35005	Repair defect of artery	C		
35013	Repair artery rupture, arm	C		
35021	Repair defect of artery	C		
35022	Repair artery rupture, chest	C		
35045	Repair defect of arm artery	C		
35081	Repair defect of artery	C		
35082	Repair artery rupture, aorta	C		
35091	Repair defect of artery	C		
35092	Repair artery rupture, aorta	C		
35102	Repair defect of artery	C		
35103	Repair artery rupture, groin	C		
35111	Repair defect of artery	C		
35112	Repair artery rupture, spleen	C		
35121	Repair defect of artery	C		
35122	Repair artery rupture, belly	C		
35131	Repair defect of artery	C		
35132	Repair artery rupture, groin	C		
35141	Repair defect of artery	C		
35142	Repair artery rupture, thigh	C		
35151	Repair defect of artery	C		
35152	Repair artery rupture, knee	C		
35182	Repair blood vessel lesion	C		
35189	Repair blood vessel lesion	C		
35211	Repair blood vessel lesion	C		
35216	Repair blood vessel lesion	C		
35221	Repair blood vessel lesion	C		
35241	Repair blood vessel lesion	C		
35246	Repair blood vessel lesion	C		
35251	Repair blood vessel lesion	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
35271	Repair blood vessel lesion	C	
35276	Repair blood vessel lesion	C	
35281	Repair blood vessel lesion	C	
35301	Rechanneling of artery	C	
35302	Rechanneling of artery	C	
35303	Rechanneling of artery	C	
35304	Rechanneling of artery	C	
35305	Rechanneling of artery	C	
35306	Rechanneling of artery	C	
35311	Rechanneling of artery	C	
35331	Rechanneling of artery	C	
35341	Rechanneling of artery	C	
35351	Rechanneling of artery	C	
35355	Rechanneling of artery	C	
35361	Rechanneling of artery	C	
35363	Rechanneling of artery	C	
35371	Rechanneling of artery	C	
35372	Rechanneling of artery	C	
35390	Reoperation, carotid add-on	C	
35400	Angioscopy	C	
35450	Repair arterial blockage	C	
35452	Repair arterial blockage	C	
35454	Repair arterial blockage	C	
35456	Repair arterial blockage	C	
35480	Atherectomy, open	C	
35481	Atherectomy, open	C	
35482	Atherectomy, open	C	
35483	Atherectomy, open	C	
35501	Artery bypass graft	C	
35506	Artery bypass graft	C	
35508	Artery bypass graft	C	
35509	Artery bypass graft	C	
35510	Artery bypass graft	C	
35511	Artery bypass graft	C	
35512	Artery bypass graft	C	
35515	Artery bypass graft	C	
35516	Artery bypass graft	C	
35518	Artery bypass graft	C	
35521	Artery bypass graft	C	
35522	Artery bypass graft	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
35523	Artery bypass graft	C	
35525	Artery bypass graft	C	
35526	Artery bypass graft	C	
35531	Artery bypass graft	C	
35533	Artery bypass graft	C	
35535	Artery bypass graft	C	
35536	Artery bypass graft	C	
35537	Artery bypass graft	C	
35538	Artery bypass graft	C	
35539	Artery bypass graft	C	
35540	Artery bypass graft	C	
35548	Artery bypass graft	C	
35549	Artery bypass graft	C	
35551	Artery bypass graft	C	
35556	Artery bypass graft	C	
35558	Artery bypass graft	C	
35560	Artery bypass graft	C	
35563	Artery bypass graft	C	
35565	Artery bypass graft	C	
35566	Artery bypass graft	C	
35570	Artery bypass graft	C	
35571	Artery bypass graft	C	
35583	Vein bypass graft	C	
35585	Vein bypass graft	C	
35587	Vein bypass graft	C	
35600	Harvest art for cabg add-on	C	
35601	Artery bypass graft	C	
35606	Artery bypass graft	C	
35612	Artery bypass graft	C	
35616	Artery bypass graft	C	
35621	Artery bypass graft	C	
35623	Bypass graft, not vein	C	
35626	Artery bypass graft	C	
35631	Artery bypass graft	C	
35632	Artery bypass graft	C	
35633	Artery bypass graft	C	
35634	Artery bypass graft	C	
35636	Artery bypass graft	C	
35637	Artery bypass graft	C	
35638	Artery bypass graft	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
35642	Artery bypass graft	C	
35645	Artery bypass graft	C	
35646	Artery bypass graft	C	
35647	Artery bypass graft	C	
35650	Artery bypass graft	C	
35651	Artery bypass graft	C	
35654	Artery bypass graft	C	
35656	Artery bypass graft	C	
35661	Artery bypass graft	C	
35663	Artery bypass graft	C	
35665	Artery bypass graft	C	
35666	Artery bypass graft	C	
35671	Artery bypass graft	C	
35681	Composite bypass graft	C	
35682	Composite bypass graft	C	
35683	Composite bypass graft	C	
35691	Arterial transposition	C	
35693	Arterial transposition	C	
35694	Arterial transposition	C	
35695	Arterial transposition	C	
35697	Reimplant artery each	C	
35700	Reoperation, bypass graft	C	
35701	Exploration, carotid artery	C	
35721	Exploration, femoral artery	C	
35741	Exploration popliteal artery	C	
35800	Explore neck vessels	C	
35820	Explore chest vessels	C	
35840	Explore abdominal vessels	C	
35870	Repair vessel graft defect	C	
35901	Excision, graft, neck	C	
35905	Excision, graft, thorax	C	
35907	Excision, graft, abdomen	C	
36660	Insertion catheter, artery	C	
36822	Insertion of cannula(s)	C	
36823	Insertion of cannula(s)	C	
37140	Revision of circulation	C	
37145	Revision of circulation	C	
37160	Revision of circulation	C	
37180	Revision of circulation	C	
37181	Splice spleen/kidney veins	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
37182	Insert hepatic shunt (tips)	C	
37616	Ligation of chest artery	C	
37617	Ligation of abdomen artery	C	
37618	Ligation of extremity artery	C	
37660	Revision of major vein	C	
37788	Revascularization, penis	C	
38100	Removal of spleen, total	C	
38101	Removal of spleen, partial	C	
38102	Removal of spleen, total	C	
38115	Repair of ruptured spleen	C	
38380	Thoracic duct procedure	C	
38381	Thoracic duct procedure	C	
38382	Thoracic duct procedure	C	
38562	Removal, pelvic lymph nodes	C	
38564	Removal, abdomen lymph nodes	C	
38724	Removal of lymph nodes, neck	C	
38746	Remove thoracic lymph nodes	C	
38747	Remove abdominal lymph nodes	C	
38765	Remove groin lymph nodes	C	
38770	Remove pelvis lymph nodes	C	
38780	Remove abdomen lymph nodes	C	
39000	Exploration of chest	C	
39010	Exploration of chest	C	
39200	Removal chest lesion	C	
39220	Removal chest lesion	C	
39499	Chest procedure	C	
39501	Repair diaphragm laceration	C	
39502	Repair paraesophageal hernia	C	
39503	Repair of diaphragm hernia	C	
39520	Repair of diaphragm hernia	C	
39530	Repair of diaphragm hernia	C	
39531	Repair of diaphragm hernia	C	
39540	Repair of diaphragm hernia	C	
39541	Repair of diaphragm hernia	C	
39545	Revision of diaphragm	C	
39560	Resect diaphragm, simple	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
39561	Resect diaphragm, complex	C	
39599	Diaphragm surgery procedure	C	
41130	Partial removal of tongue	C	
41135	Tongue and neck surgery	C	
41140	Removal of tongue	C	
41145	Tongue removal, neck surgery	C	
41150	Tongue, mouth, jaw surgery	C	
41153	Tongue, mouth, neck surgery	C	
41155	Tongue, jaw, & neck surgery	C	
42426	Excise parotid gland/lesion	C	
42845	Extensive surgery of throat	C	
42894	Revision of pharyngeal walls	C	
42953	Repair throat, esophagus	C	
42961	Control throat bleeding	C	
42971	Control nose/throat bleeding	C	
43045	Incision of esophagus	C	
43100	Excision of esophagus lesion	C	
43101	Excision of esophagus lesion	C	
43107	Removal of esophagus	C	
43108	Removal of esophagus	C	
43112	Removal of esophagus	C	
43113	Removal of esophagus	C	
43116	Partial removal of esophagus	C	
43117	Partial removal of esophagus	C	
43118	Partial removal of esophagus	C	
43121	Partial removal of esophagus	C	
43122	Partial removal of esophagus	C	
43123	Partial removal of esophagus	C	
43124	Removal of esophagus	C	
43135	Removal of esophagus pouch	C	
43279	Lap myotomy, heller	C	
43281	Lap paraesophag hern repair	C	NI
43282	Lap paraesoph her rpr w/mesh	C	NI
43300	Repair of esophagus	C	
43305	Repair esophagus and fistula	C	
43310	Repair of esophagus	C	
43312	Repair esophagus and fistula	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
43313	Esophagoplasty congenital	C	
43314	Tracheo-esophagoplasty cong	C	
43320	Fuse esophagus & stomach	C	
43324	Revise esophagus & stomach	C	
43325	Revise esophagus & stomach	C	
43326	Revise esophagus & stomach	C	
43330	Repair of esophagus	C	
43331	Repair of esophagus	C	
43340	Fuse esophagus & intestine	C	
43341	Fuse esophagus & intestine	C	
43350	Surgical opening, esophagus	C	
43351	Surgical opening, esophagus	C	
43352	Surgical opening, esophagus	C	
43360	Gastrointestinal repair	C	
43361	Gastrointestinal repair	C	
43400	Ligate esophagus veins	C	
43401	Esophagus surgery for veins	C	
43405	Ligate/staple esophagus	C	
43410	Repair esophagus wound	C	
43415	Repair esophagus wound	C	
43425	Repair esophagus opening	C	
43460	Pressure treatment esophagus	C	
43496	Free jejunum flap, microvasc	C	
43500	Surgical opening of stomach	C	
43501	Surgical repair of stomach	C	
43502	Surgical repair of stomach	C	
43520	Incision of pyloric muscle	C	
43605	Biopsy of stomach	C	
43610	Excision of stomach lesion	C	
43611	Excision of stomach lesion	C	
43620	Removal of stomach	C	
43621	Removal of stomach	C	
43622	Removal of stomach	C	
43631	Removal of stomach, partial	C	
43632	Removal of stomach, partial	C	
43633	Removal of stomach, partial	C	
43634	Removal of stomach, partial	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
43635	Removal of stomach, partial	C		
43640	Vagotomy & pylorus repair	C		
43641	Vagotomy & pylorus repair	C		
43644	Lap gastric bypass/roux-en-y	C		
43645	Lap gastr bypass incl small i	C		
43770	Lap place gastr adj device	C		
43771	Lap revise gastr adj device	C		
43772	Lap rmyl gastr adj device	C		
43773	Lap replace gastr adj device	C		
43774	Lap rmyl gastr adj all parts	C		
43775	Lap sleeve gastrectomy	C	NI	
43800	Reconstruction of pylorus	C		
43810	Fusion of stomach and bowel	C		
43820	Fusion of stomach and bowel	C		
43825	Fusion of stomach and bowel	C		
43832	Place gastrostomy tube	C		
43840	Repair of stomach lesion	C		
43843	Gastroplasty w/o v-band	C		
43845	Gastroplasty duodenal switch	C		
43846	Gastric bypass for obesity	C		
43847	Gastric bypass incl small i	C		
43848	Revision gastroplasty	C		
43850	Revise stomach-bowel fusion	C		
43855	Revise stomach-bowel fusion	C		
43860	Revise stomach-bowel fusion	C		
43865	Revise stomach-bowel fusion	C		
43880	Repair stomach-bowel fistula	C		
43881	Impl/re-do electrd, antrum	C		
43882	Revise/remove electrd antrum	C		
44005	Freeing of bowel adhesion	C		
44010	Incision of small bowel	C		
44015	Insert needle cath bowel	C		
44020	Explore small intestine	C		
44021	Decompress small bowel	C		
44025	Incision of large bowel	C		
44050	Reduce bowel obstruction	C		
44055	Correct malrotation of bowel	C		
44110	Excise intestine lesion(s)	C		
44111	Excision of bowel lesion(s)	C		
44120	Removal of small intestine	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
44121	Removal of small intestine	C		
44125	Removal of small intestine	C		
44126	Enterectomy w/o taper, cong	C		
44127	Enterectomy w/taper, cong	C		
44128	Enterectomy cong, add-on	C		
44130	Bowel to bowel fusion	C		
44132	Enterectomy, cadaver donor	C		
44133	Enterectomy, live donor	C		
44135	Intestine transplant, cadaver	C		
44136	Intestine transplant, live	C		
44137	Remove intestinal allograft	C		
44139	Mobilization of colon	C		
44140	Partial removal of colon	C		
44141	Partial removal of colon	C		
44143	Partial removal of colon	C		
44144	Partial removal of colon	C		
44145	Partial removal of colon	C		
44146	Partial removal of colon	C		
44147	Partial removal of colon	C		
44150	Removal of colon	C		
44151	Removal of colon/ileostomy	C		
44155	Removal of colon/ileostomy	C		
44156	Removal of colon/ileostomy	C		
44157	Colectomy w/ileoanal anast	C		
44158	Colectomy w/neo-rectum pouch	C		
44160	Removal of colon	C		
44187	Lap, ileo/jejuno-stomy	C		
44188	Lap, colostomy	C		
44202	Lap, enterectomy	C		
44203	Lap resect s/intestine, addl	C		
44204	Laparo partial colectomy	C		
44205	Lap colectomy part w/ileum	C		
44210	Laparo total proctocolectomy	C		
44211	Lap colectomy w/proctectomy	C		
44212	Laparo total proctocolectomy	C		
44227	Lap, close enterostomy	C		
44300	Open bowel to skin	C		
44310	ileostomy/jejunosomy	C		
44314	Revision of ileostomy	C		
44316	Devise bowel pouch	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
44320	Colostomy	C	
44322	Colostomy with biopsies	C	
44345	Revision of colostomy	C	
44346	Revision of colostomy	C	
44602	Suture, small intestine	C	
44603	Suture, small intestine	C	
44604	Suture, large intestine	C	
44605	Repair of bowel lesion	C	
44615	Intestinal stricturoplasty	C	
44620	Repair bowel opening	C	
44625	Repair bowel opening	C	
44626	Repair bowel opening	C	
44640	Repair bowel-skin fistula	C	
44650	Repair bowel fistula	C	
44660	Repair bowel-bladder fistula	C	
44661	Repair bowel-bladder fistula	C	
44680	Surgical revision, intestine	C	
44700	Suspend bowel w/prosthesis	C	
44715	Prepare donor intestine	C	
44720	Prep donor intestine/venous	C	
44721	Prep donor intestine/artery	C	
44800	Excision of bowel pouch	C	
44820	Excision of mesentery lesion	C	
44850	Repair of mesentery	C	
44899	Bowel surgery procedure	C	
44900	Drain abscess, open	C	
44960	Appendectomy	C	
45110	Removal of rectum	C	
45111	Partial removal of rectum	C	
45112	Removal of rectum	C	
45113	Partial proctectomy	C	
45114	Partial removal of rectum	C	
45116	Partial removal of rectum	C	
45119	Remove rectum w/reservoir	C	
45120	Removal of rectum	C	
45121	Removal of rectum and colon	C	
45123	Partial proctectomy	C	
45126	Pelvic exenteration	C	
45130	Excision of rectal prolapse	C	
45135	Excision of rectal prolapse	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
45136	Excise ileoanal reservoir	C	
45395	Lap, removal of rectum	C	
45397	Lap, remove rectum w/pouch	C	
45400	Laparoscopic proc	C	
45402	Lap proctopexy w/sig resect	C	
45540	Correct rectal prolapse	C	
45550	Repair rectum/remove sigmoid	C	
45562	Exploration/repair of rectum	C	
45563	Exploration/repair of rectum	C	
45800	Repair rect/bladder fistula	C	
45805	Repair fistula w/colostomy	C	
45820	Repair rectourethral fistula	C	
45825	Repair fistula w/colostomy	C	
46705	Repair of anal stricture	C	
46710	Repr per/vag pouch snl proc	C	
46712	Repr per/vag pouch dbl proc	C	
46715	Rep perf anoper fistu	C	
46716	Rep perf anoper/vestib fistu	C	
46730	Construction of absent anus	C	
46735	Construction of absent anus	C	
46740	Construction of absent anus	C	
46742	Repair of imperforated anus	C	
46744	Repair of cloacal anomaly	C	
46746	Repair of cloacal anomaly	C	
46748	Repair of cloacal anomaly	C	
46751	Repair of anal sphincter	C	
47010	Open drainage, liver lesion	C	
47015	Inject/aspirate liver cyst	C	
47100	Wedge biopsy of liver	C	
47120	Partial removal of liver	C	
47122	Extensive removal of liver	C	
47125	Partial removal of liver	C	
47130	Partial removal of liver	C	
47133	Removal of donor liver	C	
47135	Transplantation of liver	C	
47136	Transplantation of liver	C	
47140	Partial removal, donor liver	C	
47141	Partial removal, donor liver	C	
47142	Partial removal, donor liver	C	
47143	Prep donor liver, whole	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
47144	Prep donor liver, 3-segment	C	
47145	Prep donor liver, lobe split	C	
47146	Prep donor liver/venous	C	
47147	Prep donor liver/arterial	C	
47300	Surgery for liver lesion	C	
47350	Repair liver wound	C	
47360	Repair liver wound	C	
47361	Repair liver wound	C	
47362	Repair liver wound	C	
47380	Open ablate liver tumor rf	C	
47381	Open ablate liver tumor cryo	C	
47400	Incision of liver duct	C	
47420	Incision of bile duct	C	
47425	Incision of bile duct	C	
47460	Incise bile duct sphincter	C	
47480	Incision of gallbladder	C	
47550	Bile duct endoscopy add-on	C	
47570	Laparo cholecystoenterostomy	C	
47600	Removal of gallbladder	C	
47605	Removal of gallbladder	C	
47610	Removal of gallbladder	C	
47612	Removal of gallbladder	C	
47620	Removal of gallbladder	C	
47700	Exploration of bile ducts	C	
47701	Bile duct revision	C	
47711	Excision of bile duct tumor	C	
47712	Excision of bile duct tumor	C	
47715	Excision of bile duct cyst	C	
47720	Fuse gallbladder & bowel	C	
47721	Fuse upper gj structures	C	
47740	Fuse gallbladder & bowel	C	
47741	Fuse gallbladder & bowel	C	
47760	Fuse bile ducts and bowel	C	
47765	Fuse liver ducts & bowel	C	
47780	Fuse bile ducts and bowel	C	
47785	Fuse bile ducts and bowel	C	
47800	Reconstruction of bile ducts	C	
47801	Placement, bile duct support	C	
47802	Fuse liver duct & intestine	C	
47900	Suture bile duct injury	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
48000	Drainage of abdomen	C	
48001	Placement of drain, pancreas	C	
48020	Removal of pancreatic stone	C	
48100	Biopsy of pancreas, open	C	
48105	Resect/debride pancreas	C	
48120	Removal of pancreas lesion	C	
48140	Partial removal of pancreas	C	
48145	Partial removal of pancreas	C	
48146	Pancreatectomy	C	
48148	Removal of pancreatic duct	C	
48150	Partial removal of pancreas	C	
48152	Pancreatectomy	C	
48153	Pancreatectomy	C	
48154	Pancreatectomy	C	
48155	Removal of pancreas	C	
48400	Injection, intraop add-on	C	
48500	Surgery of pancreatic cyst	C	
48510	Drain pancreatic pseudocyst	C	
48520	Fuse pancreas cyst and bowel	C	
48540	Fuse pancreas cyst and bowel	C	
48545	Pancreatorrhaphy	C	
48547	Duodenal exclusion	C	
48548	Fuse pancreas and bowel	C	
48551	Prep donor pancreas	C	
48552	Prep donor pancreas/venous	C	
48554	Transpl allograft pancreas	C	
48556	Removal, allograft pancreas	C	
49000	Exploration of abdomen	C	
49002	Reopening of abdomen	C	
49010	Exploration behind abdomen	C	
49020	Drain abdominal abscess	C	
49040	Drain, open, abdom abscess	C	
49060	Drain, open, retroper abscess	C	
49062	Drain to peritoneal cavity	C	
49203	Exc abd tum 5 cm or less	C	
49204	Exc abd tum over 5 cm	C	
49205	Exc abd tum over 10 cm	C	
49215	Excise sacral spine tumor	C	
49220	Multiple surgery, abdomen	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
49255	Removal of omentum	C		
49425	Insert abdomen-venous drain	C		
49428	Ligation of shunt	C		
49605	Repair umbilical lesion	C		
49606	Repair umbilical lesion	C		
49610	Repair umbilical lesion	C		
49611	Repair umbilical lesion	C		
49900	Repair of abdominal wall	C		
49904	Omental flap, extra-abdom	C		
49905	Omental flap, intra-abdom	C		
49906	Free omental flap, microvasc	C		
50010	Exploration of kidney	C		
50040	Drainage of kidney	C		
50045	Exploration of kidney	C		
50060	Removal of kidney stone	C		
50065	Incision of kidney	C		
50070	Incision of kidney	C		
50075	Removal of kidney stone	C		
50100	Revise kidney blood vessels	C		
50120	Exploration of kidney	C		
50125	Explore and drain kidney	C		
50130	Removal of kidney stone	C		
50135	Exploration of kidney	C		
50205	Renal biopsy open	C		
50220	Remove kidney, open	C		
50225	Remove kidney open, complex	C		
50230	Remove kidney open, radical	C		
50234	Removal of kidney & ureter	C		
50236	Removal of kidney & ureter	C		
50240	Partial removal of kidney	C		
50250	Cryoblade renal mass open	C		
50280	Removal of kidney lesion	C		
50290	Removal of kidney lesion	C		
50300	Remove cadaver donor kidney	C		
50320	Remove kidney, living donor	C		
50323	Prep cadaver renal allograft	C		
50325	Prep donor renal graft	C		
50327	Prep renal graft/venous	C		
50328	Prep renal graft/arterial	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
50329	Prep renal graft/ureteral	C		
50340	Removal of kidney	C		
50360	Transplantation of kidney	C		
50365	Transplantation of kidney	C		
50370	Remove transplanted kidney	C		
50380	Reimplantation of kidney	C		
50400	Revision of kidney/ureter	C		
50405	Revision of kidney/ureter	C		
50500	Repair of kidney wound	C		
50520	Close kidney-skin fistula	C		
50525	Repair renal-abdomen fistula	C		
50526	Repair renal-abdomen fistula	C		
50540	Revision of horseshoe kidney	C		
50545	Laparo radical nephrectomy	C		
50546	Laparoscopic nephrectomy	C		
50547	Laparo removal donor kidney	C		
50548	Laparo remove w/ureter	C		
50600	Exploration of ureter	C		
50605	Insert ureteral support	C		
50610	Removal of ureter stone	C		
50620	Removal of ureter stone	C		
50630	Removal of ureter stone	C		
50650	Removal of ureter	C		
50660	Removal of ureter	C		
50700	Revision of ureter	C		
50715	Release of ureter	C		
50722	Release of ureter	C		
50725	Release/revise ureter	C		
50728	Revise ureter	C		
50740	Fusion of ureter & kidney	C		
50750	Fusion of ureter & kidney	C		
50760	Fusion of ureters	C		
50770	Splicing of ureters	C		
50780	Reimplant ureter in bladder	C		
50782	Reimplant ureter in bladder	C		
50783	Reimplant ureter in bladder	C		
50785	Reimplant ureter in bladder	C		
50800	Implant ureter in bowel	C		
50810	Fusion of ureter & bowel	C		
50815	Urine shunt to intestine	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
50820	Construct bowel bladder	C		
50825	Construct bowel bladder	C		
50830	Revise urine flow	C		
50840	Replace ureter by bowel	C		
50845	Appendico-vesicostomy	C		
50860	Transplant ureter to skin	C		
50900	Repair of ureter	C		
50920	Closure ureter/skin fistula	C		
50930	Closure ureter/bowel fistula	C		
50940	Release of ureter	C		
51525	Removal of bladder lesion	C		
51530	Removal of bladder lesion	C		
51550	Partial removal of bladder	C		
51555	Partial removal of bladder	C		
51565	Revise bladder & ureter(s)	C		
51570	Removal of bladder	C		
51575	Removal of bladder & nodes	C		
51580	Remove bladder/revise tract	C		
51585	Removal of bladder & nodes	C		
51590	Remove bladder/revise tract	C		
51595	Remove bladder/revise tract	C		
51596	Remove bladder/create pouch	C		
51597	Removal of pelvic structures	C		
51800	Revision of bladder/urethra	C		
51820	Revision of urinary tract	C		
51840	Attach bladder/urethra	C		
51841	Attach bladder/urethra	C		
51865	Repair of bladder wound	C		
51900	Repair bladder/vagina lesion	C		
51920	Close bladder-uterus fistula	C		
51925	Hysterectomy/bladder repair	C		
51940	Correction of bladder defect	C		
51960	Revision of bladder & bowel	C		
51980	Construct bladder opening	C		
53415	Reconstruction of urethra	C		
53448	Remov/repic ur sphinctr comp	C		
54125	Removal of penis	C		
54130	Remove penis & nodes	C		
54135	Remove penis & nodes	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
54390	Repair penis and bladder	C		
54411	Remov/repic penis pros, compl	C		
54417	Remv/repic penis pros, compl	C		
54430	Revision of penis	C		
54650	Orchiopexy (Fowler-Stephens)	C		
55605	Incise sperm duct pouch	C		
55650	Remove sperm duct pouch	C		
55801	Removal of prostate	C		
55810	Extensive prostate surgery	C		
55812	Extensive prostate surgery	C		
55815	Extensive prostate surgery	C		
55821	Removal of prostate	C		
55831	Removal of prostate	C		
55840	Extensive prostate surgery	C		
55842	Extensive prostate surgery	C		
55845	Extensive prostate surgery	C		
55862	Extensive prostate surgery	C		
55865	Extensive prostate surgery	C		
55866	Laparo radical prostatectomy	C		
56630	Extensive vulva surgery	C		
56631	Extensive vulva surgery	C		
56632	Extensive vulva surgery	C		
56633	Extensive vulva surgery	C		
56634	Extensive vulva surgery	C		
56637	Extensive vulva surgery	C		
56640	Extensive vulva surgery	C		
57110	Remove vagina wall, complete	C		
57111	Remove vagina tissue, compl	C		
57112	Vaginectomy w/nodes, compl	C		
57270	Repair of bowel pouch	C		
57280	Suspension of vagina	C		
57296	Revise vag graft, open abd	C		
57305	Repair rectum-vagina fistula	C		
57307	Fistula repair & colostomy	C		
57308	Fistula repair, transperine	C		
57311	Repair urethrovaginal lesion	C		
57531	Removal of cervix, radical	C		
57540	Removal of residual cervix	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
57545	Remove cervix/repair pelvis	C	
58140	Myomectomy abdom method	C	
58146	Myomectomy abdom complex	C	
58150	Total hysterectomy	C	
58152	Total hysterectomy	C	
58180	Partial hysterectomy	C	
58200	Extensive hysterectomy	C	
58210	Extensive hysterectomy	C	
58240	Removal of pelvis contents	C	
58267	Vag hyst w/urinary repair	C	
58275	Hysterectomy/revise vagina	C	
58280	Hysterectomy/revise vagina	C	
58285	Extensive hysterectomy	C	
58293	Vag hyst w/uro repair, compl	C	
58400	Suspension of uterus	C	
58410	Suspension of uterus	C	
58520	Repair of ruptured uterus	C	
58540	Revision of uterus	C	
58548	Lap radical hyst	C	
58605	Division of fallopian tube	C	
58611	Ligate oviduct(s) add-on	C	
58700	Removal of fallopian tube	C	
58720	Removal of ovary/tube(s)	C	
58740	Adhesiolysis tube, ovary	C	
58750	Repair oviduct	C	
58752	Revise ovarian tube(s)	C	
58760	Fimbrioplasty	C	
58822	Drain ovary abscess, percut	C	
58825	Transposition, ovary(s)	C	
58940	Removal of ovary(s)	C	
58943	Removal of ovary(s)	C	
58950	Resect ovarian malignancy	C	
58951	Resect ovarian malignancy	C	
58952	Resect ovarian malignancy	C	
58953	Tah, rad dissect for debulk	C	
58954	Tah rad debulk/lymph remove	C	
58956	Bso, omentectomy w/tah	C	
58957	Resect recurrent gyn mal	C	
58958	Resect recur gyn mal w/lym	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
58960	Exploration of abdomen	C	
59120	Treat ectopic pregnancy	C	
59121	Treat ectopic pregnancy	C	
59130	Treat ectopic pregnancy	C	
59135	Treat ectopic pregnancy	C	
59136	Treat ectopic pregnancy	C	
59140	Treat ectopic pregnancy	C	
59325	Revision of cervix	C	
59350	Repair of uterus	C	
59514	Cesarean delivery only	C	
59525	Remove uterus after cesarean	C	
59620	Attempted vbac delivery only	C	
59830	Treat uterus infection	C	
59850	Abortion	C	
59851	Abortion	C	
59852	Abortion	C	
59855	Abortion	C	
59856	Abortion	C	
59857	Abortion	C	
60254	Extensive thyroid surgery	C	
60270	Removal of thyroid	C	
60505	Explore parathyroid glands	C	
60521	Removal of thymus gland	C	
60522	Removal of thymus gland	C	
60540	Explore adrenal gland	C	
60545	Explore adrenal gland	C	
60600	Remove carotid body lesion	C	
60605	Remove carotid body lesion	C	
60650	Laparoscopy adrenalectomy	C	
61105	Twist drill hole	C	
61107	Drill skull for implantation	C	
61108	Drill skull for drainage	C	
61120	Burr hole for puncture	C	
61140	Pierce skull for biopsy	C	
61150	Pierce skull for drainage	C	
61151	Pierce skull for drainage	C	
61154	Pierce skull & remove clot	C	
61156	Pierce skull for drainage	C	
61210	Pierce skull, implant device	C	
61250	Pierce skull & explore	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
61253	Pierce skull & explore	C	
61304	Open skull for exploration	C	
61305	Open skull for exploration	C	
61312	Open skull for drainage	C	
61313	Open skull for drainage	C	
61314	Open skull for drainage	C	
61315	Open skull for drainage	C	
61316	Implt cran bone flap to abdo	C	
61320	Open skull for drainage	C	
61321	Open skull for drainage	C	
61322	Decompressive craniotomy	C	
61323	Decompressive lobectomy	C	
61332	Explore/biopsy eye socket	C	
61333	Explore orbit/remove lesion	C	
61340	Subtemporal decompression	C	
61343	Incise skull (press relief)	C	
61345	Relieve cranial pressure	C	
61440	Incise skull for surgery	C	
61450	Incise skull for surgery	C	
61458	Incise skull for brain wound	C	
61460	Incise skull for surgery	C	
61470	Incise skull for surgery	C	
61480	Incise skull for surgery	C	
61490	Incise skull for surgery	C	
61500	Removal of skull lesion	C	
61501	Remove infected skull bone	C	
61510	Removal of brain lesion	C	
61512	Remove brain lining lesion	C	
61514	Removal of brain abscess	C	
61516	Removal of brain lesion	C	
61517	Implt brain chemotx add-on	C	
61518	Removal of brain lesion	C	
61519	Remove brain lining lesion	C	
61520	Removal of brain lesion	C	
61521	Removal of brain lesion	C	
61522	Removal of brain abscess	C	
61524	Removal of brain lesion	C	
61526	Removal of brain lesion	C	
61530	Removal of brain lesion	C	
61531	Implant brain electrodes	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
61533	Implant brain electrodes	C	
61534	Removal of brain lesion	C	
61535	Remove brain electrodes	C	
61536	Removal of brain lesion	C	
61537	Removal of brain tissue	C	
61538	Removal of brain tissue	C	
61539	Removal of brain tissue	C	
61540	Removal of brain tissue	C	
61541	Incision of brain tissue	C	
61542	Removal of brain tissue	C	
61543	Removal of brain tissue	C	
61544	Remove & treat brain lesion	C	
61545	Excision of brain tumor	C	
61546	Removal of pituitary gland	C	
61548	Removal of pituitary gland	C	
61550	Release of skull seams	C	
61552	Release of skull seams	C	
61556	Incise skull/sutures	C	
61557	Incise skull/sutures	C	
61558	Excision of skull/sutures	C	
61559	Excision of skull/sutures	C	
61563	Excision of skull tumor	C	
61564	Excision of skull tumor	C	
61566	Removal of brain tissue	C	
61567	Incision of brain tissue	C	
61570	Remove foreign body, brain	C	
61571	Incise skull for brain wound	C	
61575	Skull base/brainstem surgery	C	
61576	Skull base/brainstem surgery	C	
61580	Craniofacial approach, skull	C	
61581	Craniofacial approach, skull	C	
61582	Craniofacial approach, skull	C	
61583	Craniofacial approach, skull	C	
61584	Obitocranial approach/skull	C	
61585	Obitocranial approach/skull	C	
61586	Resect nasopharynx, skull	C	
61590	Infratemporal approach/skull	C	
61591	Infratemporal approach/skull	C	
61592	Orbitocranial approach/skull	C	
61595	Transtemporal approach/skull	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
61596	Transcochlear approach/skull	C	
61597	Transcondylar approach/skull	C	
61598	Transpetrosal approach/skull	C	
61600	Resect/excise cranial lesion	C	
61601	Resect/excise cranial lesion	C	
61605	Resect/excise cranial lesion	C	
61606	Resect/excise cranial lesion	C	
61607	Resect/excise cranial lesion	C	
61608	Resect/excise cranial lesion	C	
61609	Transect artery, sinus	C	
61610	Transect artery, sinus	C	
61611	Transect artery, sinus	C	
61612	Transect artery, sinus	C	
61613	Remove aneurysm, sinus	C	
61615	Resect/excise lesion, skull	C	
61616	Resect/excise lesion, skull	C	
61618	Repair dura	C	
61619	Repair dura	C	
61624	Transcath occlusion, cns	C	
61630	Intracranial angioplasty	C	
61635	Intracran angioplasty w/stent	C	
61680	Intracranial vessel surgery	C	
61682	Intracranial vessel surgery	C	
61684	Intracranial vessel surgery	C	
61686	Intracranial vessel surgery	C	
61690	Intracranial vessel surgery	C	
61692	Intracranial vessel surgery	C	
61697	Brain aneurysm repr, complex	C	
61698	Brain aneurysm repr, complex	C	
61700	Brain aneurysm repr, simple	C	
61702	Inner skull vessel surgery	C	
61703	Clamp neck artery	C	
61705	Revise circulation to head	C	
61708	Revise circulation to head	C	
61710	Revise circulation to head	C	
61711	Fusion of skull arteries	C	
61735	Incise skull/brain surgery	C	
61750	Incise skull/brain biopsy	C	
61751	Brain biopsy w/ct/mr guide	C	
61760	Implant brain electrodes	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
61850	Implant neuroelectrodes	C	
61860	Implant neuroelectrodes	C	
61863	Implant neuroelectrode	C	
61864	Implant neuroelectrde, addl	C	
61867	Implant neuroelectrode	C	
61868	Implant neuroelectrde, addl	C	
61870	Implant neuroelectrodes	C	
61875	Implant neuroelectrodes	C	
62005	Treat skull fracture	C	
62010	Treatment of head injury	C	
62100	Repair brain fluid leakage	C	
62115	Reduction of skull defect	C	
62116	Reduction of skull defect	C	
62117	Reduction of skull defect	C	
62120	Repair skull cavity lesion	C	
62121	Incise skull repair	C	
62140	Repair of skull defect	C	
62141	Repair of skull defect	C	
62142	Remove skull plate/flap	C	
62143	Replace skull plate/flap	C	
62145	Repair of skull & brain	C	
62146	Repair of skull with graft	C	
62147	Repair of skull with graft	C	
62148	Retr bone flap to fix skull	C	
62161	Dissect brain w/scope	C	
62162	Remove colloid cyst w/scope	C	
62163	Neuroendoscopy w/fb removal	C	
62164	Remove brain tumor w/scope	C	
62165	Remove pituit tumor w/scope	C	
62180	Establish brain cavity shunt	C	
62190	Establish brain cavity shunt	C	
62192	Establish brain cavity shunt	C	
62200	Establish brain cavity shunt	C	
62201	Brain cavity shunt w/scope	C	
62220	Establish brain cavity shunt	C	
62223	Establish brain cavity shunt	C	
62256	Remove brain cavity shunt	C	
62258	Replace brain cavity shunt	C	
63043	Laminotomy, addl cervical	C	
63044	Laminotomy, addl lumbar	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
63050	Cervical laminoplasty	C	
63051	C-laminoplasty w/graft/plate	C	
63077	Spine disk surgery, thorax	C	
63078	Spine disk surgery, thorax	C	
63081	Removal of vertebral body	C	
63082	Remove vertebral body add-on	C	
63085	Removal of vertebral body	C	
63086	Remove vertebral body add-on	C	
63087	Removal of vertebral body	C	
63088	Remove vertebral body add-on	C	
63090	Removal of vertebral body	C	
63091	Remove vertebral body add-on	C	
63101	Removal of vertebral body	C	
63102	Remove vertebral body add-on	C	
63103	Remove vertebral body add-on	C	
63170	Incise spinal cord tract(s)	C	
63172	Drainage of spinal cyst	C	
63173	Drainage of spinal cyst	C	
63180	Revise spinal cord ligaments	C	
63182	Revise spinal cord ligaments	C	
63185	Incise spinal column/nerves	C	
63190	Incise spinal column/nerves	C	
63191	Incise spinal column/nerves	C	
63194	Incise spinal column & cord	C	
63195	Incise spinal column & cord	C	
63196	Incise spinal column & cord	C	
63197	Incise spinal column & cord	C	
63198	Incise spinal column & cord	C	
63199	Incise spinal column & cord	C	
63200	Release of spinal cord	C	
63250	Revise spinal cord vessels	C	
63251	Revise spinal cord vessels	C	
63252	Revise spinal cord vessels	C	
63265	Excise intraspinal lesion	C	
63266	Excise intraspinal lesion	C	
63267	Excise intraspinal lesion	C	
63268	Excise intraspinal lesion	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010			
HCPCS Code	Short Descriptor	SI	CI
63270	Excise intraspinal lesion	C	
63271	Excise intraspinal lesion	C	
63272	Excise intraspinal lesion	C	
63273	Excise intraspinal lesion	C	
63275	Biopsy/excise spinal tumor	C	
63276	Biopsy/excise spinal tumor	C	
63277	Biopsy/excise spinal tumor	C	
63278	Biopsy/excise spinal tumor	C	
63280	Biopsy/excise spinal tumor	C	
63281	Biopsy/excise spinal tumor	C	
63282	Biopsy/excise spinal tumor	C	
63283	Biopsy/excise spinal tumor	C	
63285	Biopsy/excise spinal tumor	C	
63286	Biopsy/excise spinal tumor	C	
63287	Biopsy/excise spinal tumor	C	
63290	Biopsy/excise spinal tumor	C	
63295	Repair of laminectomy defect	C	
63300	Removal of vertebral body	C	
63301	Removal of vertebral body	C	
63302	Removal of vertebral body	C	
63303	Removal of vertebral body	C	
63304	Removal of vertebral body	C	
63305	Removal of vertebral body	C	
63306	Removal of vertebral body	C	
63307	Removal of vertebral body	C	
63308	Remove vertebral body add-on	C	
63700	Repair of spinal herniation	C	
63702	Repair of spinal herniation	C	
63704	Repair of spinal herniation	C	
63706	Repair of spinal herniation	C	
63707	Repair spinal fluid leakage	C	
63709	Repair spinal fluid leakage	C	
63710	Graft repair of spine defect	C	
63740	Install spinal shunt	C	
64752	Incision of vagus nerve	C	
64755	Incision of stomach nerves	C	
64760	Incision of vagus nerve	C	
64809	Remove sympathetic nerves	C	
64818	Remove sympathetic nerves	C	
64866	Fusion of facial/other nerve	C	

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
64868	Fusion of facial/other nerve	C		
65273	Repair of eye wound	C		
69155	Extensive ear/neck surgery	C		
69535	Remove part of temporal bone	C		
69554	Remove ear lesion	C		
69950	Incise inner ear nerve	C		
75900	Intravascular cath exchange	C		
75952	Endovasc repair abdom aorta	C		
75953	Abdom aneurysm endovasc rpr	C		
75954	Iliac aneurysm endovasc rpr	C		
75956	Xray, endovasc thor ao repr	C		
75957	Xray, endovasc thor ao repr	C		
75958	Xray, place prox ext thor ao	C		
75959	Xray, place dist ext thor ao	C		
92970	Cardioassist, internal	C		
92971	Cardioassist, external	C		
92975	Dissolve clot, heart vessel	C		
92992	Revision of heart chamber	C		
92993	Revision of heart chamber	C		
99190	Special pump services	C		
99191	Special pump services	C		
99192	Special pump services	C		
99356	Prolonged service, inpatient	C		
99357	Prolonged service, inpatient	C		
99462	Sbsq nb em per day, hosp	C		
99468	Neonate crit care, initial	C		
99469	Neonate crit care, subsq	C		
99471	Ped critical care, initial	C		
99472	Ped critical care, subsq	C		
99475	Ped crit care age 2-5, init	C		
99476	Ped crit care age 2-5, subsq	C		
99477	Init day hosp neonate care	C		
99478	lc, lbw inf < 1500 gm subsq	C		
99479	lc, lbw inf 1500-2500 g subsq	C		
99480	lc, inf pbw 2501-5000 g subsq	C		
0048T	Implant ventricular device	C		
0050T	Removal circulatory assist	C		
0051T	Implant total heart system	C		
0052T	Replace component heart	C		

ADDENDUM E.—HCPCS CODES THAT ARE PAID AS INPATIENT PROCEDURES FOR CY 2010				
HCPCS Code	Short Descriptor	SI	CI	
	syst			
0053T	Replace component heart syst	C		
0075T	Perq stent/chest vert art	C		
0076T	S&i stent/chest vert art	C		
0078T	Endovasc aort repr w/device	C		
0079T	Endovasc visc extnsn repr	C		
0080T	Endovasc aort repr rad s&i	C		
0081T	Endovasc visc extnsn s&i	C		
0092T	Artific disc addl	C		
0095T	Artific diskectomy addl	C		
0098T	Rev artific disc addl	C		
0157T	Open impl gast curve electrd	C		
0158T	Open remy gast curve electrd	C		
0163T	Lumb artif diskectomy addl	C		
0164T	Remove lumb artif disc addl	C		
0165T	Revise lumb artif disc addl	C		
0166T	Tcath vsd close w/o bypass	C		
0167T	Tcath vsd close w bypass	C		
0169T	Place stereo cath brain	C		
0184T	Exc rectal tumor endoscopic	C		
0195T	Arthrod presac interbody	C		
0196T	Arthrod presac interbody eac	C		
0202T	Post vert arthrpst 1 lumbar	C		
0219T	Fuse spine facet jt cerv	C	NI	
0220T	Fuse spine facet jt thor	C	NI	
G0341	Percutaneous islet celltrans	C		
G0342	Laparoscopy islet cell trans	C		
G0343	Laparotomy islet cell transp	C		
G0406	Telhealth inpt consult 15min	C	CH	
G0407	Telhealth inpt consult 25min	C	CH	
G0408	Telhealth inpt consult 35mins	C	CH	
G0412	Open tx iliac spine uni/bil	C		
G0414	Pelvic ring fx treat int fix	C		
G0415	Open tx post pelvic fxcture	C		
G0425	Inpt telehealth consult 30m	C	NI	
G0426	Inpt telehealth consult 50m	C	NI	
G0427	Inpt telehealth con 70/>-m	C	NI	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
010138		0.0066	SUMTER	01590	01590
010143	*	0.0254	CULLMAN	01210	01210
010146		0.0047	CALHOUN	01070	01070
010150		0.0127	BUTLER	01060	01060
010158	*	0.0023	FRANKLIN	01290	01290
010164	*	0.0211	TALLADEGA	01600	01600
012011		0.0047	CALHOUN	01070	01070
013027		0.0134	BALDWIN	01010	01010
013032		0.0061	ETOWAH	01270	01270
014006		0.0061	ETOWAH	01270	01270
030067		0.0298	LAPAZ	03055	03055
040014	*	0.0199	WHITE	04720	04720
040019	*	0.0258	ST. FRANCIS	04610	04610
040039	*	0.0172	GREENE	04270	04270
040047		0.0117	RANDOLPH	04600	04600
040067		0.0007	COLUMBIA	04130	04130
040071	*	0.0149	JEFFERSON	04340	04340
040076	*	0.1000	HOT SPRING	04290	04290
040081		0.0357	PIKE	04540	04540
040149		0.0199	WHITE	04720	04720
042007		0.0149	JEFFERSON	04340	04340
042011		0.0199	WHITE	04720	04720
043034		0.0036	CHICOT	04080	04080
050002	*	0.0010	ALAMEDA	05000	05000
050007		0.0146	SAN MATEO	05510	05510
050009	*	0.0180	NAPA	05380	05380
050013	*	0.0180	NAPA	05380	05380
050014	*	0.0139	AMADOR	05020	05020
050042	*	0.0162	TEHAMA	05620	05620
050043	*	0.0010	ALAMEDA	05000	05000
050069	*	0.0013	ORANGE	05400	05400
050070		0.0146	SAN MATEO	05510	05510
050073	*	0.0171	SOLANO	05580	05580
050075	*	0.0010	ALAMEDA	05000	05000

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
010005	*	0.0296	MARSHALL	01470	01470
010008		0.0174	CRENSHAW	01200	01200
010010	*	0.0296	MARSHALL	01470	01470
010012	*	0.0186	DE KALB	01240	01240
010015		0.0046	CLARKE	01120	01120
010021		0.0052	DALE	01220	01220
010022	*	0.1128	CHEROKEE	01090	01090
010025	*	0.0389	CHAMBERS	01080	01080
010027		0.0026	COFFEE	01150	01150
010029	*	0.0289	LEE	01400	01400
010032		0.0325	RANDOLPH	01550	01550
010035	*	0.0254	CULLMAN	01210	01210
010038		0.0047	CALHOUN	01070	01070
010040		0.0061	ETOWAH	01270	01270
010045		0.0222	FAYETTE	01280	01280
010046		0.0061	ETOWAH	01270	01270
010047		0.0127	BUTLER	01060	01060
010049		0.0026	COFFEE	01150	01150
010052	*	0.0245	TALLAPOOSA	01610	01610
010059	*	0.0071	LAWRENCE	01390	01390
010061	*	0.0542	JACKSON	01350	01350
010065	*	0.0245	TALLAPOOSA	01610	01610
010078		0.0047	CALHOUN	01070	01070
010083	*	0.0134	BALDWIN	01010	01010
010091		0.0046	CLARKE	01120	01120
010100	*	0.0134	BALDWIN	01010	01010
010101	*	0.0211	TALLADEGA	01600	01600
010109		0.0405	PICKENS	01530	01530
010110		0.0215	BULLOCK	01050	01050
010125		0.0476	WINSTON	01660	01660
010128		0.0046	CLARKE	01120	01120
010129		0.0134	BALDWIN	01010	01010

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
050289		0.0146	SAN MATEO	05510	05510
050291		0.0058	SONOMA	05590	05590
050298		0.0011	BERNARDINO	05460	05460
050300	*	0.0011	BERNARDINO	05460	05460
050305	*	0.0010	ALAMEDA	05000	05000
050313		0.0132	SAN JOAQUIN	05490	05490
050320	*	0.0010	ALAMEDA	05000	05000
050325		0.0033	TUOLUMNE	05650	05650
050327	*	0.0011	BERNARDINO	05460	05460
050335	*	0.0033	TUOLUMNE	05650	05650
050336		0.0132	SAN JOAQUIN	05490	05490
050348	*	0.0013	ORANGE	05400	05400
050366		0.0015	CALAVERAS	05040	05040
050367	*	0.0171	SOLANO	05580	05580
050385		0.0058	SONOMA	05590	05590
050426	*	0.0013	ORANGE	05400	05400
050444		0.0233	MERCED	05340	05340
050488	*	0.0010	ALAMEDA	05000	05000
050512	*	0.0010	ALAMEDA	05000	05000
050517	*	0.0011	BERNARDINO	05460	05460
050526	*	0.0013	ORANGE	05400	05400
050528	*	0.0233	MERCED	05340	05340
050541	*	0.0146	SAN MATEO	05510	05510
050543	*	0.0013	ORANGE	05400	05400
050547		0.0058	SONOMA	05590	05590
050548	*	0.0013	ORANGE	05400	05400
050551	*	0.0013	ORANGE	05400	05400
050567	*	0.0013	ORANGE	05400	05400
050570	*	0.0013	ORANGE	05400	05400
050580	*	0.0013	ORANGE	05400	05400
050586	*	0.0011	SAN	05460	05460

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
050084	*	0.0132	SAN JOAQUIN	05490	05490
050089	*	0.0011	BERNARDINO	05460	05460
050090		0.0058	SONOMA	05590	05590
050099	*	0.0011	BERNARDINO	05460	05460
050101	*	0.0171	SOLANO	05580	05580
050113		0.0146	SAN MATEO	05510	05510
050118		0.0132	SAN JOAQUIN	05490	05490
050122		0.0132	SAN JOAQUIN	05490	05490
050129	*	0.0011	BERNARDINO	05460	05460
050133	*	0.0178	YUBA	05680	05680
050136		0.0058	SONOMA	05590	05590
050140	*	0.0011	BERNARDINO	05460	05460
050150	*	0.0342	NEVADA	05390	05390
050167		0.0132	SAN JOAQUIN	05490	05490
050168	*	0.0013	ORANGE	05400	05400
050173	*	0.0013	ORANGE	05400	05400
050174		0.0058	SONOMA	05590	05590
050193	*	0.0013	ORANGE	05400	05400
050195	*	0.0010	ALAMEDA	05000	05000
050197	*	0.0146	SAN MATEO	05510	05510
050211	*	0.0010	ALAMEDA	05000	05000
050224	*	0.0013	ORANGE	05400	05400
050226	*	0.0013	ORANGE	05400	05400
050230	*	0.0013	ORANGE	05400	05400
050245	*	0.0011	BERNARDINO	05460	05460
050264	*	0.0010	ALAMEDA	05000	05000
050272	*	0.0011	BERNARDINO	05460	05460
050279	*	0.0011	BERNARDINO	05460	05460
050283	*	0.0010	ALAMEDA	05000	05000

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
054074		0.0171	SOLANO	05580	
054093		0.0011	SAN BERNARDINO	05460	
054110		0.0010	ALAMEDA	05000	
054111		0.0011	SAN BERNARDINO	05460	
054122		0.0180	NAPA	05380	
054123		0.0132	SAN JOAQUIN	05490	
054135		0.0013	ORANGE	05400	
054141		0.0171	SOLANO	05580	
060001	*	0.0042	WELD	06610	
060003	*	0.0069	BOULDER	06060	
060027	*	0.0069	BOULDER	06060	
060103	*	0.0069	BOULDER	06060	
060116	*	0.0069	BOULDER	06060	
060121	*	0.0042	WELD	06610	
063033		0.0042	WELD	06610	
064007		0.0069	BOULDER	06060	
070006	*	0.0045	FAIRFIELD	07000	
070010	*	0.0045	FAIRFIELD	07000	
070018	*	0.0045	FAIRFIELD	07000	
070028	*	0.0045	FAIRFIELD	07000	
070033	*	0.0045	FAIRFIELD	07000	
070034	*	0.0045	FAIRFIELD	07000	
074000		0.0045	FAIRFIELD	07000	
074012		0.0045	FAIRFIELD	07000	
074014		0.0045	FAIRFIELD	07000	
080001		0.0044	NEW CASTLE	08010	
080003		0.0044	NEW CASTLE	08010	
082000		0.0044	NEW CASTLE	08010	
083300		0.0044	NEW CASTLE	08010	
084001		0.0044	NEW CASTLE	08010	
084002		0.0044	NEW CASTLE	08010	
084003		0.0044	NEW CASTLE	08010	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
			BERNARDINO		
050589	*	0.0013	ORANGE	05400	
050603	*	0.0013	ORANGE	05400	
050609	*	0.0013	ORANGE	05400	
050618	*	0.0011	SAN BERNARDINO	05460	
050667	*	0.0180	NAPA	05380	
050678	*	0.0013	ORANGE	05400	
050680	*	0.0171	SOLANO	05580	
050690		0.0058	SONOMA	05590	
050693	*	0.0013	ORANGE	05400	
050720	*	0.0013	ORANGE	05400	
050744	*	0.0013	ORANGE	05400	
050745	*	0.0013	ORANGE	05400	
050746	*	0.0013	ORANGE	05400	
050747	*	0.0013	ORANGE	05400	
050748		0.0132	SAN JOAQUIN	05490	
050754		0.0146	SAN MATEO	05510	
050758	*	0.0011	SAN BERNARDINO	05460	
052034		0.0010	ALAMEDA	05000	
052035		0.0013	ORANGE	05400	
052037		0.0011	SAN BERNARDINO	05460	
052039		0.0013	ORANGE	05400	
052040		0.0011	SAN BERNARDINO	05460	
052053		0.0013	ORANGE	05400	
053034		0.0013	ORANGE	05400	
053037		0.0011	SAN BERNARDINO	05460	
053301		0.0010	ALAMEDA	05000	
053304		0.0013	ORANGE	05400	
053306		0.0013	ORANGE	05400	
053308		0.0013	ORANGE	05400	

ADDENDUM L.—CY 2010 OPFS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
110100		0.0790	JEFFERSON	11620	11620
110101		0.0067	COOK	11311	11311
110142		0.0185	EVANS	11441	11441
110146	*	0.0364	CAMDEN	11170	11170
110150	*	0.0227	BALDWIN	11030	11030
110187	*	0.0643	LUMPKIN	11701	11701
110189	*	0.0066	FANNIN	11450	11450
110190		0.0241	MACON	11710	11710
110205		0.0507	GILMER	11471	11471
114018		0.0227	BALDWIN	11030	11030
130003	*	0.0235	NEZ PERCE	13340	13340
130024		0.0675	BONNER	13080	13080
130049	*	0.0319	KOOTENAI	13270	13270
130066		0.0319	KOOTENAI	13270	13270
130067	*	0.0725	BINGHAM	13050	13050
132001		0.0319	KOOTENAI	13270	13270
134010		0.0725	BINGHAM	13050	13050
140001		0.0369	FULTON	14370	14370
140026		0.0315	LA SALLE	14580	14580
140043	*	0.0056	WHITESIDE	14988	14988
140058	*	0.0126	MORGAN	14770	14770
140110	*	0.0315	LA SALLE	14580	14580
140116		0.0014	MC HENRY	14640	14640
140160	*	0.0332	STEPHENSON	14970	14970
140161	*	0.0168	LIVINGSTON	14610	14610
140167	*	0.0632	IROQUOIS	14460	14460
140176		0.0014	MC HENRY	14640	14640
140234		0.0315	LA SALLE	14580	14580
150022		0.0158	MONTGOMERY	15530	15530
150030	*	0.0192	HENRY	15320	15320
150072		0.0105	CASS	15080	15080
150076	*	0.0215	MARSHALL	15490	15490
150088	*	0.0111	MADISON	15470	15470
150091	*	0.0050	HUNTINGTON	15340	15340

ADDENDUM L.—CY 2010 OPFS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
090001		0.0033	THE DISTRICT	09000	09000
090003		0.0033	THE DISTRICT	09000	09000
090004	*	0.0033	THE DISTRICT	09000	09000
090005		0.0033	THE DISTRICT	09000	09000
090006		0.0033	THE DISTRICT	09000	09000
090008		0.0033	THE DISTRICT	09000	09000
090011	*	0.0033	THE DISTRICT	09000	09000
092002		0.0033	THE DISTRICT	09000	09000
092003		0.0033	THE DISTRICT	09000	09000
093025		0.0033	THE DISTRICT	09000	09000
093300		0.0033	THE DISTRICT	09000	09000
094001		0.0033	THE DISTRICT	09000	09000
094004		0.0033	THE DISTRICT	09000	09000
100014	*	0.0047	VOLUSIA	10630	10630
100017	*	0.0047	VOLUSIA	10630	10630
100023	*	0.0031	CITRUS	10080	10080
100045	*	0.0047	VOLUSIA	10630	10630
100047	*	0.0028	CHARLOTTE	10070	10070
100068	*	0.0047	VOLUSIA	10630	10630
100072	*	0.0047	VOLUSIA	10630	10630
100077	*	0.0028	CHARLOTTE	10070	10070
100081	*	0.0022	WALTON	10650	10650
100118	*	0.0177	FLAGLER	10170	10170
100139	*	0.0006	LEVY	10370	10370
100232	*	0.0054	PUTNAM	10530	10530
100236	*	0.0028	CHARLOTTE	10070	10070
100249	*	0.0031	CITRUS	10080	10080
100252	*	0.0151	OKEECHOBEE	10460	10460
100290		0.0338	SUMTER	10590	10590
100292	*	0.0022	WALTON	10650	10650
110023	*	0.0416	GORDON	11500	11500
110029	*	0.0052	HALL	11550	11550
110040	*	0.1455	JACKSON	11610	11610
110041	*	0.0623	HABERSHAM	11540	11540

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
190099		0.0189	AVOUELLES	19040	19040
190106	*	0.0102	ALLEN	19010	19010
190116		0.0085	MOREHOUSE	19330	19330
190133		0.0102	ALLEN	19010	19010
190140		0.0035	FRANKLIN	19200	19200
190144	*	0.0387	WEBSTER	19590	19590
190145		0.0090	LA SALLE	19290	19290
190184	*	0.0075	CALDWELL	19100	19100
190190	*	0.0075	CALDWELL	19100	19100
190191	*	0.0187	ST. LANDRY	19480	19480
190246		0.0075	CALDWELL	19100	19100
190257	*	0.0061	LINCOLN	19300	19300
190222		0.0061	LINCOLN	19300	19300
190206		0.0387	WEBSTER	19590	19590
190234		0.0187	ST. LANDRY	19480	19480
190236		0.0243	TANGIPAHOA	19520	19520
190240		0.0243	TANGIPAHOA	19520	19520
190250		0.0261	ACADIA	19000	19000
190306		0.0187	ST. LANDRY	19480	19480
190344		0.0243	TANGIPAHOA	19520	19520
190347		0.0189	VERMILION	19560	19560
190349		0.0189	VERMILION	19560	19560
190355		0.0075	CALDWELL	19100	19100
190358		0.0085	MOREHOUSE	19330	19330
190363		0.0243	TANGIPAHOA	19520	19520
190367		0.0101	JEFFERSON DAVIS	19260	19260
190368		0.0243	TANGIPAHOA	19520	19520
190369		0.0085	MOREHOUSE	19330	19330
190373		0.0187	ST. LANDRY	19480	19480
190379		0.0243	TANGIPAHOA	19520	19520
190381		0.0261	ACADIA	19000	19000
190388		0.0261	ACADIA	19000	19000
190391		0.0085	IBERIA	19220	19220
190407		0.0387	WEBSTER	19590	19590

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	County Code
150102	*	0.0108	STARKE	15740	15740
150113	*	0.0111	MADISON	15470	15470
150133	*	0.0193	KOSCIUSKO	15420	15420
150146	*	0.0087	NOBLE	15560	15560
153040		0.0215	MARSHALL	15490	15490
154014		0.0193	KOSCIUSKO	15420	15420
154035		0.0105	CASS	15080	15080
154047		0.0215	MARSHALL	15490	15490
160013		0.0179	MUSCATINE	16690	16690
160030		0.0013	STORY	16840	16840
160032		0.0235	JASPER	16490	16490
160080	*	0.0066	CLINTON	16220	16220
170137	*	0.0421	DOUGLAS	17220	17220
170150		0.0166	COWLEY	17170	17170
180012	*	0.0080	HARDIN	18460	18460
180017	*	0.0035	BARREN	18040	18040
180049	*	0.0488	MADISON	18750	18750
180064		0.0314	MONTGOMERY	18860	18860
180066	*	0.0439	LOGAN	18700	18700
180070		0.0240	GRAYSON	18420	18420
180079		0.0259	HARRISON	18480	18480
183028		0.0080	HARDIN	18460	18460
184012		0.0080	HARDIN	18460	18460
190003	*	0.0085	IBERIA	19220	19220
190015	*	0.0243	TANGIPAHOA	19520	19520
190017	*	0.0187	ST. LANDRY	19480	19480
190034		0.0189	VERMILION	19560	19560
190044		0.0261	ACADIA	19000	19000
190050		0.0044	BEAUREGARD	19050	19050
190053		0.0101	JEFFERSON DAVIS	19260	19260
190054		0.0085	IBERIA	19220	19220
190078		0.0187	ST. LANDRY	19480	19480
190086	*	0.0061	LINCOLN	19300	19300
190088	*	0.0387	WEBSTER	19590	19590

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
220062	*	0.0072	WORCESTER	22170	
220063	*	0.0271	MIDDLESEX	22090	
220070	*	0.0271	MIDDLESEX	22090	
220080	*	0.0355	ESSEX	22040	
220082	*	0.0271	MIDDLESEX	22090	
220084	*	0.0271	MIDDLESEX	22090	
220090	*	0.0072	WORCESTER	22170	
220095	*	0.0072	WORCESTER	22170	
220098	*	0.0271	MIDDLESEX	22090	
220101	*	0.0271	MIDDLESEX	22090	
220105	*	0.0271	MIDDLESEX	22090	
220163	*	0.0072	WORCESTER	22170	
220171	*	0.0271	MIDDLESEX	22090	
220174	*	0.0355	ESSEX	22040	
220175	*	0.0271	MIDDLESEX	22090	
220176	*	0.0072	WORCESTER	22170	
222000		0.0271	MIDDLESEX	22090	
222003		0.0271	MIDDLESEX	22090	
222024		0.0271	MIDDLESEX	22090	
222026		0.0355	ESSEX	22040	
222044		0.0355	ESSEX	22040	
222047		0.0355	ESSEX	22040	
222048		0.0072	WORCESTER	22170	
223026		0.0271	MIDDLESEX	22090	
223028		0.0355	ESSEX	22040	
223029		0.0072	WORCESTER	22170	
223033		0.0072	WORCESTER	22170	
224007		0.0271	MIDDLESEX	22090	
224026		0.0072	WORCESTER	22170	
224032		0.0072	WORCESTER	22170	
224033		0.0355	ESSEX	22040	
224038		0.0271	MIDDLESEX	22090	
224039		0.0355	ESSEX	22040	
230002	*	0.0043	WAYNE	23810	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
194065		0.0061	LINCOLN	19300	
194075		0.0101	JEFFERSON DAVIS	19260	
194077		0.0061	LINCOLN	19300	
194081		0.0044	BEAUREGARD	19050	
194082		0.0101	JEFFERSON DAVIS	19260	
194083		0.0085	MOREHOUSE	19330	
194085		0.0261	ACADIA	19000	
194087		0.0061	LINCOLN	19300	
194091		0.0243	TANGIPAHOA	19520	
194092		0.0035	FRANKLIN	19200	
200024	*	0.0094	ANDROSCOGGIN	20000	
200032		0.0367	OXFORD	20080	
200034	*	0.0094	ANDROSCOGGIN	20000	
200050	*	0.0227	HANCOCK	20040	
210001		0.0187	WASHINGTON	21210	
210023		0.0079	ANNE ARUNDEL	21010	
210028		0.0383	ST. MARYS	21180	
210043		0.0079	ANNE ARUNDEL	21010	
210061		0.0188	WORCESTER	21230	
212002		0.0187	WASHINGTON	21210	
214001		0.0079	ANNE ARUNDEL	21010	
214003		0.0187	WASHINGTON	21210	
214015		0.0188	WORCESTER	21230	
220001	*	0.0072	WORCESTER	22170	
220002	*	0.0271	MIDDLESEX	22090	
220010	*	0.0355	ESSEX	22040	
220011	*	0.0271	MIDDLESEX	22090	
220019	*	0.0072	WORCESTER	22170	
220025	*	0.0072	WORCESTER	22170	
220029	*	0.0355	ESSEX	22040	
220033	*	0.0355	ESSEX	22040	
220035	*	0.0355	ESSEX	22040	
220049	*	0.0271	MIDDLESEX	22090	
220058	*	0.0072	WORCESTER	22170	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
230176	*	0.0043	WAYNE	23810	
230195	*	0.0021	MACOMB	23490	
230204		0.0021	MACOMB	23490	
230207	*	0.0025	OAKLAND	23620	
230208	*	0.0095	MONTCALM	23580	
230217		0.0047	CALHOUN	23120	
230222	*	0.0035	MIDLAND	23550	
230227		0.0021	MACOMB	23490	
230244	*	0.0043	WAYNE	23810	
230254	*	0.0025	OAKLAND	23620	
230257		0.0021	MACOMB	23490	
230264		0.0021	MACOMB	23490	
230269	*	0.0025	OAKLAND	23620	
230270	*	0.0043	WAYNE	23810	
230273	*	0.0043	WAYNE	23810	
230277	*	0.0025	OAKLAND	23620	
230279	*	0.0210	LIVINGSTON	23460	
230297	*	0.0043	WAYNE	23810	
230301		0.0025	OAKLAND	23620	
232019		0.0043	WAYNE	23810	
232020		0.0052	BAY	23080	
232023		0.0021	MACOMB	23490	
232025		0.0101	BERRIEN	23100	
232027		0.0043	WAYNE	23810	
232028		0.0047	CALHOUN	23120	
232030		0.0025	OAKLAND	23620	
232031		0.0043	WAYNE	23810	
232032		0.0043	WAYNE	23810	
232034		0.0435	ALLEGAN	23020	
232036		0.0223	JACKSON	23370	
232038		0.0043	WAYNE	23810	
233025		0.0047	CALHOUN	23120	
233027		0.0043	WAYNE	23810	
233028		0.0025	OAKLAND	23620	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
230003	*	0.0220	OTTAWA	23690	
230005		0.0473	LENAWEE	23450	
230013	*	0.0025	OAKLAND	23620	
230015		0.0295	ST. JOSEPH	23740	
230019	*	0.0025	OAKLAND	23620	
230020	*	0.0043	WAYNE	23810	
230021	*	0.0101	BERRIEN	23100	
230022	*	0.0212	BRANCH	23110	
230024	*	0.0043	WAYNE	23810	
230029	*	0.0025	OAKLAND	23620	
230035	*	0.0095	MONTCALM	23580	
230037	*	0.0210	HILLSDALE	23290	
230041		0.0052	BAY	23080	
230047		0.0021	MACOMB	23490	
230053	*	0.0043	WAYNE	23810	
230069	*	0.0210	LIVINGSTON	23460	
230071	*	0.0025	OAKLAND	23620	
230072	*	0.0220	OTTAWA	23690	
230075		0.0047	CALHOUN	23120	
230078	*	0.0101	BERRIEN	23100	
230089	*	0.0043	WAYNE	23810	
230092	*	0.0223	JACKSON	23370	
230093		0.0058	MECOSTA	23530	
230096	*	0.0295	ST. JOSEPH	23740	
230099		0.0231	MONROE	23570	
230104	*	0.0043	WAYNE	23810	
230121	*	0.0678	SHIAWASSEE	23770	
230130	*	0.0025	OAKLAND	23620	
230135	*	0.0043	WAYNE	23810	
230142	*	0.0043	WAYNE	23810	
230146	*	0.0043	WAYNE	23810	
230151	*	0.0025	OAKLAND	23620	
230165	*	0.0043	WAYNE	23810	
230174	*	0.0220	OTTAWA	23690	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
300012	*	0.0049	HILLSBOROUGH	30050	
300017	*	0.0075	ROCKINGHAM	30070	
300020	*	0.0049	HILLSBOROUGH	30050	
300023	*	0.0075	ROCKINGHAM	30070	
300029	*	0.0075	ROCKINGHAM	30070	
300034	*	0.0049	HILLSBOROUGH	30050	
303026		0.0075	ROCKINGHAM	30070	
304001		0.0075	ROCKINGHAM	30070	
310002	*	0.0268	ESSEX	31200	
310009	*	0.0268	ESSEX	31200	
310015	*	0.0199	MORRIS	31300	
310017	*	0.0199	MORRIS	31300	
310018	*	0.0268	ESSEX	31200	
310038	*	0.0209	MIDDLESEX	31270	
310039	*	0.0209	MIDDLESEX	31270	
310050	*	0.0199	MORRIS	31300	
310054	*	0.0268	ESSEX	31200	
310070	*	0.0209	MIDDLESEX	31270	
310076	*	0.0268	ESSEX	31200	
310083	*	0.0268	ESSEX	31200	
310093	*	0.0268	ESSEX	31200	
310096	*	0.0268	ESSEX	31200	
310108	*	0.0209	MIDDLESEX	31270	
310119	*	0.0268	ESSEX	31200	
312018		0.0209	MIDDLESEX	31270	
312020		0.0199	MORRIS	31300	
313025		0.0268	ESSEX	31200	
313300		0.0209	MIDDLESEX	31270	
314010		0.0268	ESSEX	31200	
314011		0.0209	MIDDLESEX	31270	
314016		0.0199	MORRIS	31300	
314020		0.0268	ESSEX	31200	
320003	*	0.0480	SAN MIGUEL	32230	
320011		0.0337	RIO ARRIBA	32190	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
233300		0.0043	WAYNE	23810	
234011		0.0025	OAKLAND	23620	
234021		0.0021	MACOMB	23490	
234023		0.0025	OAKLAND	23620	
234025		0.0276	TUSCOLA	23780	
234028		0.0043	WAYNE	23810	
234034		0.0043	WAYNE	23810	
234035		0.0043	WAYNE	23810	
234038		0.0043	WAYNE	23810	
234039		0.0021	MACOMB	23490	
240018		0.0805	GOODHUE	24240	
240044		0.0625	WINONA	24840	
240064	*	0.0134	ITASCA	24300	
240069	*	0.0267	STEELE	24730	
240071	*	0.0385	RICE	24650	
240117		0.0527	MOWER	24490	
240211		0.0812	PINE	24570	
250023	*	0.0541	PEARL RIVER	25540	
250040	*	0.0021	JACKSON	25290	
250117	*	0.0541	PEARL RIVER	25540	
250128		0.0446	PANOLA	25530	
250162		0.0014	HANCOCK	25220	
252011		0.0446	PANOLA	25530	
260059		0.0077	LACLEDE	26520	
260064	*	0.0089	AUDRAIN	26030	
260097		0.0300	JOHNSON	26500	
260116	*	0.0087	ST. FRANCOIS	26930	
260160		0.0144	STODDARD	26985	
260163		0.0087	ST. FRANCOIS	26930	
264005		0.0087	ST. FRANCOIS	26930	
264027		0.0087	CEDAR	26190	
280077	*	0.0080	DODGE	28260	
290002	*	0.0277	LYON	29090	
300011	*	0.0049	HILLSBOROUGH	30050	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
330259	*	0.0123	NASSAU	33400	
330264		0.0642	ORANGE	33540	
330276		0.0036	FULTON	33280	
330277	*	0.0056	STEUBEN	33690	
330331	*	0.0123	NASSAU	33400	
330332	*	0.0123	NASSAU	33400	
330372	*	0.0123	NASSAU	33400	
330386	*	0.0745	SULLIVAN	33710	
334017		0.0642	ORANGE	33540	
334061		0.0642	ORANGE	33540	
340020		0.0156	LEE	34520	
340021	*	0.0162	CLEVELAND	34220	
340024		0.0177	SAMPSON	34810	
340027	*	0.0128	LENOIR	34530	
340037		0.0162	CLEVELAND	34220	
340038		0.0253	BEAUFORT	34060	
340039	*	0.0101	IREDELL	34480	
340068		0.0087	COLUMBUS	34230	
340069	*	0.0015	WAKE	34910	
340070		0.0395	ALAMANCE	34000	
340071	*	0.0226	HARNETT	34420	
340073	*	0.0015	WAKE	34910	
340085	*	0.0250	DAVIDSON	34280	
340096	*	0.0250	DAVIDSON	34280	
340104		0.0162	CLEVELAND	34220	
340114	*	0.0015	WAKE	34910	
340126	*	0.0100	WILSON	34970	
340129	*	0.0101	IREDELL	34480	
340133		0.0260	MARTIN	34580	
340138	*	0.0015	WAKE	34910	
340144	*	0.0101	IREDELL	34480	
340145	*	0.0336	LINCOLN	34540	
340151		0.0052	HALIFAX	34410	
340173	*	0.0015	WAKE	34910	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
320018		0.0024	DONA ANA	32060	
320085		0.0024	DONA ANA	32060	
320088		0.0024	DONA ANA	32060	
322004		0.0024	DONA ANA	32060	
323025		0.0480	SAN MIGUEL	32230	
323032		0.0024	DONA ANA	32060	
324007		0.0024	DONA ANA	32060	
324010		0.0024	DONA ANA	32060	
324012		0.0024	DONA ANA	32060	
330004	*	0.0633	ULSTER	33740	
330008	*	0.0126	WYOMING	33900	
330010		0.0067	MONTGOMERY	33380	
330027	*	0.0123	NASSAU	33400	
330033		0.0223	CHENANGO	33080	
330047		0.0067	MONTGOMERY	33380	
330073	*	0.0151	GENESEE	33290	
330094	*	0.0503	COLUMBIA	33200	
330103		0.0131	CATTARAUGUS	33040	
330106		0.0123	NASSAU	33400	
330126	*	0.0642	ORANGE	33540	
330132		0.0131	CATTARAUGUS	33040	
330135		0.0642	ORANGE	33540	
330144		0.0056	STEUBEN	33690	
330151		0.0056	STEUBEN	33690	
330167	*	0.0123	NASSAU	33400	
330175		0.0260	CORTLAND	33210	
330181	*	0.0123	NASSAU	33400	
330182	*	0.0123	NASSAU	33400	
330191	*	0.0017	WARREN	33750	
330198	*	0.0123	NASSAU	33400	
330205		0.0642	ORANGE	33540	
330224	*	0.0633	ULSTER	33740	
330225	*	0.0123	NASSAU	33400	
330235	*	0.0306	CAYUGA	33050	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
370023		0.0090	STEPHENS	37680	
370065		0.0096	CRAIG	37170	
370072		0.0258	LATIMER	37380	
370083		0.0051	PUSHMATAHA	37630	
370100		0.0100	CHOCTAW	37110	
370149	*	0.0302	POTTAWATOMIE	37620	
370156		0.0121	GARVIN	37240	
370169		0.0163	MCINTOSH	37450	
370172		0.0258	LATIMER	37380	
370214		0.0121	GARVIN	37240	
372017		0.0100	CHOCTAW	37110	
372019		0.0302	POTTAWATOMIE	37620	
373032		0.0100	CHOCTAW	37110	
380022	*	0.0067	LINN	38210	
384011		0.0107	UMATILLA	38290	
390008		0.0060	LAWRENCE	39450	
390016	*	0.0060	LAWRENCE	39450	
390030	*	0.0147	SCHUYLKILL	39650	
390031	*	0.0147	SCHUYLKILL	39650	
390039		0.0037	SOMERSET	39680	
390044	*	0.0191	BERKS	39110	
390052		0.0047	CLEARFIELD	39230	
390056		0.0036	HUNTINGDON	39380	
390065	*	0.0532	ADAMS	39000	
390066	*	0.0372	LEBANON	39460	
390079	*	0.0003	BRADFORD	39130	
390086	*	0.0047	CLEARFIELD	39230	
390096	*	0.0191	BERKS	39110	
390110	*	0.0003	CAMBRIA	39160	
390112		0.0037	SOMERSET	39680	
390113	*	0.0053	CRAWFORD	39260	
390117		0.0002	BEDFORD	39100	
390122		0.0053	CRAWFORD	39260	
390125		0.0022	WAYNE	39760	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
344011		0.0015	WAKE	34910	
344014		0.0015	WAKE	34910	
360002		0.0141	ASHLAND	36020	
360010	*	0.0074	TUSCARAWAS	36800	
360013	*	0.0135	SHELBY	36760	
360025	*	0.0077	ERIE	36220	
360036	*	0.0126	WAYNE	36860	
360040		0.0387	KNOX	36430	
360044		0.0127	DARKE	36190	
360055	*	0.0011	TRUMBULL	36790	
360065	*	0.0075	HURON	36400	
360070		0.0005	STARK	36770	
360071		0.0035	VAN WERT	36820	
360084		0.0005	STARK	36770	
360086	*	0.0186	CLARK	36110	
360096		0.0071	COLUMBIANA	36140	
360107		0.0119	SANDUSKY	36730	
360125	*	0.0133	ASHTABULA	36030	
360131		0.0005	STARK	36770	
360151		0.0005	STARK	36770	
360156		0.0119	SANDUSKY	36730	
360161		0.0011	TRUMBULL	36790	
360175	*	0.0183	CLINTON	36130	
360185	*	0.0071	COLUMBIANA	36140	
360187	*	0.0186	CLARK	36110	
360245	*	0.0133	ASHTABULA	36030	
362016		0.0005	STARK	36770	
362032		0.0005	STARK	36770	
363026		0.0011	TRUMBULL	36790	
364031		0.0005	STARK	36770	
364040		0.0186	CLARK	36110	
364043		0.0035	VAN WERT	36820	
370014	*	0.0361	BRYAN	37060	
370015	*	0.0366	MAYES	37480	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
420069	*	0.0052	CLARENDON	42130	
420070	*	0.0051	SUMTER	42420	
420082		0.0002	AIKEN	42010	
420083	*	0.0027	SPARTANBURG	42410	
420098	*	0.0008	GEORGETOWN	42210	
422004		0.0027	SPARTANBURG	42410	
423028		0.0001	YORK	42450	
423029		0.0108	ANDERSON	42030	
424011		0.0108	ANDERSON	42030	
430008		0.0535	BROOKINGS	43050	
430048		0.0129	LAWRENCE	43400	
430094		0.0129	LAWRENCE	43400	
440007		0.0219	COFFEE	44150	
440008		0.0449	HENDERSON	44380	
440012		0.0009	SULLIVAN	44810	
440016		0.0144	CARROLL	44080	
440017		0.0009	SULLIVAN	44810	
440025	*	0.0009	GREENE	44290	
440031		0.0019	ROANE	44720	
440033		0.0027	CAMPBELL	44060	
440035	*	0.0301	MONTGOMERY	44620	
440047		0.0338	GIBSON	44260	
440050		0.0009	GREENE	44290	
440051		0.0082	MC NAIRY	44540	
440057		0.0021	CLAIBORNE	44120	
440060		0.0338	GIBSON	44260	
440063		0.0033	WASHINGTON	44890	
440070		0.0109	DECATUR	44190	
440081		0.0052	SEVIER	44770	
440084		0.0025	MONROE	44610	
440105		0.0033	WASHINGTON	44890	
440109		0.0070	HARDIN	44350	
440115		0.0338	GIBSON	44260	
440137		0.0738	BEDFORD	44010	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
390130	*	0.0003	CAMBRIA	39160	
390138	*	0.0218	FRANKLIN	39350	
390146		0.0022	WARREN	39740	
390150		0.0031	GREENE	39370	
390151	*	0.0218	FRANKLIN	39350	
390162	*	0.0217	NORTHAMPTON	39590	
390173		0.0037	INDIANA	39390	
390183	*	0.0147	SCHUYLKILL	39650	
390201	*	0.1170	MONROE	39550	
390236		0.0003	BRADFORD	39130	
390313	*	0.0147	SCHUYLKILL	39650	
390316		0.0191	BERKS	39110	
392030		0.0532	ADAMS	39000	
392031		0.0003	CAMBRIA	39160	
392034		0.0217	NORTHAMPTON	39590	
393026		0.0191	BERKS	39110	
393050		0.0217	NORTHAMPTON	39590	
394014		0.0191	BERKS	39110	
394016		0.0022	WARREN	39740	
394020		0.0372	LEBANON	39460	
394052		0.0191	BERKS	39110	
420002		0.0001	YORK	42450	
420007	*	0.0027	SPARTANBURG	42410	
420019		0.0158	CHESTER	42110	
420020	*	0.0008	GEORGETOWN	42210	
420027	*	0.0108	ANDERSON	42030	
420030	*	0.0069	COLLETON	42140	
420036	*	0.0064	LANCASTER	42280	
420039	*	0.0110	UNION	42430	
420043		0.0157	CHEROKEE	42100	
420053		0.0035	NEWBERRY	42350	
420054		0.0002	MARLBORO	42340	
420062	*	0.0125	CHESTERFIELD	42120	
420068	*	0.0027	ORANGEBURG	42370	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
450370	*	0.0235	COLORADO	45312	
450389	*	0.0618	HENDERSON	45640	
450395		0.0441	POLK	45850	
450419	*	0.0024	TARRANT	45910	
450438	*	0.0235	COLORADO	45312	
450451		0.0536	SOMERVELL	45893	
450460		0.0053	TYLER	45942	
450497		0.0375	MONTAGUE	45800	
450539		0.0067	HALE	45582	
450547	*	0.0195	WOOD	45974	
450563	*	0.0024	TARRANT	45910	
450565	*	0.0509	PALO PINTO	45841	
450573		0.0126	JASPER	45690	
450596	*	0.0743	HOOD	45653	
450615		0.0033	CASS	45260	
450639	*	0.0024	TARRANT	45910	
450641		0.0375	MONTAGUE	45800	
450672	*	0.0024	TARRANT	45910	
450675	*	0.0024	TARRANT	45910	
450677	*	0.0024	TARRANT	45910	
450698		0.0127	LAMB	45751	
450747	*	0.0126	ANDERSON	45000	
450755		0.0276	HOCKLEY	45652	
450770	*	0.0182	MILAM	45795	
450779	*	0.0024	TARRANT	45910	
450813		0.0126	ANDERSON	45000	
450838		0.0126	JASPER	45690	
450872	*	0.0024	TARRANT	45910	
450880	*	0.0024	TARRANT	45910	
450884		0.0049	UPSHUR	45943	
450886	*	0.0024	TARRANT	45910	
450888		0.0024	TARRANT	45910	
452018		0.0024	TARRANT	45910	
452019		0.0024	TARRANT	45910	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
440144	*	0.0219	COFFEE	44150	
440148	*	0.0296	DE KALB	44200	
440174	*	0.0312	HAYWOOD	44370	
440176		0.0009	SULLIVAN	44810	
440180		0.0027	CAMPBELL	44060	
440181		0.0365	HARDEMAN	44340	
440182		0.0144	CARROLL	44080	
440184		0.0033	WASHINGTON	44890	
440185	*	0.0230	BRADLEY	44050	
442016		0.0009	SULLIVAN	44810	
443027		0.0009	SULLIVAN	44810	
444006		0.0033	WASHINGTON	44890	
444008		0.0365	HARDEMAN	44340	
450032	*	0.0254	HARRISON	45620	
450039	*	0.0024	TARRANT	45910	
450052	*	0.0276	BOSQUE	45160	
450059		0.0075	COMAL	45320	
450064	*	0.0024	TARRANT	45910	
450087	*	0.0024	TARRANT	45910	
450090		0.0650	COOKE	45340	
450099	*	0.0145	GRAY	45563	
450135	*	0.0024	TARRANT	45910	
450137	*	0.0024	TARRANT	45910	
450144	*	0.0559	ANDREWS	45010	
450163		0.0054	KLEBERG	45743	
450192		0.0271	HILL	45651	
450194		0.0213	CHEROKEE	45281	
450210		0.0151	PANOLA	45842	
450224	*	0.0195	WOOD	45974	
450236		0.0389	HOPKINS	45654	
450270		0.0271	HILL	45651	
450283	*	0.0653	VAN ZANDT	45947	
450347	*	0.0370	WALKER	45949	
450348	*	0.0059	FALLS	45500	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
494029		0.0003	SMYTH	49860	
500003	*	0.0166	SKAGIT	50280	
500007	*	0.0166	SKAGIT	50280	
500019		0.0131	LEWIS	50200	
500039	*	0.0094	KITSAP	50170	
500041	*	0.0020	COWLITZ	50070	
510012		0.0124	MASON	51260	
510018	*	0.0188	JACKSON	51170	
510047	*	0.0269	MARION	51240	
520028	*	0.0286	GREEN	52220	
520035		0.0076	SHEBOYGAN	52580	
520044		0.0076	SHEBOYGAN	52580	
520045		0.0022	WINNEBAGO	52690	
520048		0.0022	WINNEBAGO	52690	
520057		0.0193	SAUK	52550	
520059	*	0.0195	RACINE	52500	
520071	*	0.0161	JEFFERSON	52270	
520076	*	0.0146	DODGE	52130	
520095	*	0.0193	SAUK	52550	
520096	*	0.0195	RACINE	52500	
520102	*	0.0242	WALWORTH	52630	
520116	*	0.0161	JEFFERSON	52270	
520198		0.0022	WINNEBAGO	52690	
522005		0.0195	RACINE	52500	
523302		0.0022	WINNEBAGO	52690	
524002		0.0022	WINNEBAGO	52690	
673026		0.0075	COMAL	45320	
673035		0.0024	TARRANT	45910	

ADDENDUM L.—CY 2010 OPPTS OUT-MIGRATION ADJUSTMENT					
Provider Number	Reclassified for FY 2010	Out-Migration Adjustment	Qualifying County Name	County Code	
452028		0.0024	TARRANT	45910	
452088		0.0024	TARRANT	45910	
452099		0.0024	TARRANT	45910	
452106		0.0075	COMAL	45320	
452110		0.0024	TARRANT	45910	
453040		0.0024	TARRANT	45910	
453041		0.0024	TARRANT	45910	
453042		0.0024	TARRANT	45910	
453089		0.0126	ANDERSON	45000	
453094		0.0024	TARRANT	45910	
453300		0.0024	TARRANT	45910	
453303		0.0024	TARRANT	45910	
454009		0.0213	CHEROKEE	45281	
454012		0.0024	TARRANT	45910	
454051		0.0024	TARRANT	45910	
454052		0.0024	TARRANT	45910	
454061		0.0024	TARRANT	45910	
454072		0.0024	TARRANT	45910	
454086		0.0024	TARRANT	45910	
454101		0.0067	HALE	45582	
460001		0.0001	UTAH	46240	
460013		0.0001	UTAH	46240	
460017		0.0383	BOX ELDER	46010	
460023		0.0001	UTAH	46240	
460039	*	0.0383	BOX ELDER	46010	
460043		0.0001	UTAH	46240	
460052		0.0001	UTAH	46240	
462005		0.0001	UTAH	46240	
490002		0.0003	RUSSELL	49830	
490019	*	0.1088	CULPEPER	49230	
490038		0.0003	SMYTH	49860	
490084		0.0187	ESSEX	49280	
490105		0.0003	SMYTH	49860	
490110		0.0185	MONTGOMERY	49600	

96154	Interv hltm/behav, fam w/pt		Q3	0432	0034
M0064	Visit for drug monitoring	CH	Q3	0607	0034
93619	Electrophysiology evaluation		Q3	0085	8000
93620	Electrophysiology evaluation		Q3	0085	8000
93650	Ablate heart dysrhythm focus		Q3	0085	8000
93651	Ablate heart dysrhythm focus		Q3	0086	8000
93652	Ablate heart dysrhythm focus		Q3	0086	8000
55875	Transper needle place, pros		Q3	0163	8001
77778	Apply interstt radiat compl		Q3	0651	8001
99205	Office/outpatient visit, new		Q3	0608	8002
99215	Office/outpatient visit, est		Q3	0607	8002
G0379	Direct refer hospital observ		Q3	0604	8002
99284	Emergency dept visit		Q3	0615	8003
99285	Emergency dept visit		Q3	0616	8003
99291	Critical care, first hour		Q3	0617	8003
G0384	Lev 5 hosp type B ED visit	CH	Q3	0630	8003
76604	Us exam, chest		Q3	0265	8004
76700	Us exam, abdom, complete		Q3	0266	8004
76705	Echo exam of abdomen		Q3	0266	8004
76770	Us exam abdo back wall, comp		Q3	0266	8004
76775	Us exam abdo back wall, lim		Q3	0266	8004
76776	Us exam k transpl w/doppler		Q3	0266	8004
76831	Echo exam, uterus		Q3	0267	8004
76856	Us exam, pelvic, complete		Q3	0266	8004
76857	Us exam, pelvic, limited		Q3	0265	8004
76870	Us exam, scrotum		Q3	0266	8004
70450	Ct head/brain w/o dye		Q3	0332	8005 or 8006
70480	Ct orbit/ear/fossa w/o dye		Q3	0332	8005 or 8006
70486	Ct maxillofacial w/o dye		Q3	0332	8005 or 8006
70490	Ct soft tissue neck w/o dye		Q3	0332	8005 or 8006
71250	Ct thorax w/o dye		Q3	0332	8005 or 8006
72125	Ct neck spine w/o dye		Q3	0332	8005 or 8006
72128	Ct chest spine w/o dye		Q3	0332	8005 or 8006
72131	Ct lumbar spine w/o dye		Q3	0332	8005 or 8006
72192	Ct pelvis w/o dye		Q3	0332	8005 or 8006
73200	Ct upper extremity w/o dye		Q3	0332	8005 or 8006
73700	Ct lower extremity w/o dye		Q3	0332	8005 or 8006
74150	Ct abdomen w/o dye		Q3	0332	8005 or 8006
74261	Ct colonography, w/o dye	NI	Q3	0332	8005 or 8006
70460	Ct head/brain w/dye		Q3	0283	8006
70470	Ct head/brain w/o & w/dye		Q3	0333	8006
70481	Ct orbit/ear/fossa w/dye		Q3	0283	8006
70482	Ct orbit/ear/fossa w/o&w/dye		Q3	0333	8006
70487	Ct maxillofacial w/dye		Q3	0283	8006
70488	Ct maxillofacial w/o & w/dye		Q3	0333	8006
70491	Ct soft tissue neck w/dye		Q3	0283	8006

ADDENDUM M.—HCPCS CODES FOR ASSIGNMENT TO COMPOSITE APCs FOR CY 2010					
HCPCS Code	Short Descriptor	CI	SI	Single Code APC Assignment	Composite APC Assignment
90801	Psy dx interview		Q3	0323	0034
90802	Intac psy dx interview		Q3	0323	0034
90804	Psydx, office, 20-30 min		Q3	0322	0034
90805	Psydx, off, 20-30 min w/e&m		Q3	0322	0034
90806	Psydx, off, 45-50 min		Q3	0323	0034
90807	Psydx, off, 45-50 min w/e&m		Q3	0323	0034
90808	Psydx, office, 75-80 min		Q3	0323	0034
90809	Psydx, off, 75-80, w/e&m		Q3	0323	0034
90810	Intac psydx, off, 20-30 min		Q3	0322	0034
90811	Intac psydx, 20-30, w/e&m		Q3	0322	0034
90812	Intac psydx, off, 45-50 min		Q3	0323	0034
90813	Intac psydx, 45-50 min w/e&m		Q3	0323	0034
90814	Intac psydx, off, 75-80 min		Q3	0323	0034
90815	Intac psydx, 75-80 w/e&m		Q3	0323	0034
90845	Psychoanalysis		Q3	0323	0034
90846	Family psydx w/o patient		Q3	0324	0034
90847	Family psydx w/patient		Q3	0324	0034
90849	Multiple family group psydx		Q3	0325	0034
90853	Group psychotherapy		Q3	0325	0034
90857	Intac group psydx		Q3	0325	0034
90862	Medication management		Q3	0606	0034
90865	Narcosynthesis		Q3	0323	0034
90880	Hypnotherapy		Q3	0323	0034
90899	Psychiatric service/therapy		Q3	0322	0034
96101	Psycho testing by psych/phys		Q3	0382	0034
96102	Psycho testing by technician		Q3	0382	0034
96103	Psycho testing admin by comp		Q3	0373	0034
96110	Developmental test, lim		Q3	0373	0034
96111	Developmental test, extend	CH	Q3	0373	0034
96116	Neurobehavioral status exam		Q3	0382	0034
96118	Neuropsych tst by psych/phys		Q3	0382	0034
96119	Neuropsych testing by tec	CH	Q3	0382	0034
96120	Neuropsych tst admin w/comp		Q3	0382	0034
96150	Assess hltm/behav, init		Q3	0432	0034
96151	Assess hltm/behav, subseq		Q3	0432	0034
96152	Intervene hltm/behav, indiv		Q3	0432	0034
96153	Intervene hltm/behav, group		Q3	0432	0034

C8907	MRI w/o cont, breast, bi		Q3	0336	8007 or 8008
C8910	MRA w/o cont, chest		Q3	0336	8007 or 8008
C8913	MRA w/o cont, lwr ext		Q3	0336	8007 or 8008
C8919	MRA w/o cont, pelvis		Q3	0336	8007 or 8008
70542	Mri orbit/face/neck w/dye		Q3	0284	8008
70543	Mri orbit/face/neck w/o & w/dye		Q3	0337	8008
70545	Mr angiography head w/dye		Q3	0284	8008
70546	Mr angiography head w/o&w/dye		Q3	0337	8008
70548	Mr angiography neck w/dye		Q3	0284	8008
70549	Mr angiography neck w/o&w/dye		Q3	0337	8008
70552	Mri brain w/dye		Q3	0284	8008
70553	Mri brain w/o & w/dye		Q3	0337	8008
71551	Mri chest w/dye		Q3	0284	8008
71552	Mri chest w/o & w/dye		Q3	0337	8008
72142	Mri neck spine w/dye		Q3	0284	8008
72147	Mri chest spine w/dye		Q3	0284	8008
72149	Mri lumbar spine w/dye		Q3	0284	8008
72156	Mri neck spine w/o & w/dye		Q3	0337	8008
72157	Mri chest spine w/o & w/dye		Q3	0337	8008
72158	Mri lumbar spine w/o & w/dye		Q3	0337	8008
72196	Mri pelvis w/dye		Q3	0284	8008
72197	Mri pelvis w/o & w/dye		Q3	0337	8008
73219	Mri upper extremity w/dye		Q3	0284	8008
73220	Mri upper extremity w/o&w/dye		Q3	0337	8008
73222	Mri joint upr extrem w/dye		Q3	0284	8008
73223	Mri joint upr extr w/o&w/dye		Q3	0337	8008
73719	Mri lower extremity w/dye		Q3	0284	8008
73720	Mri lower extremity w/o&w/dye		Q3	0337	8008
73722	Mri joint of lwr extr w/dye		Q3	0284	8008
73723	Mri joint lwr extr w/o&w/dye		Q3	0337	8008
74182	Mri abdomen w/dye		Q3	0284	8008
74183	Mri abdomen w/o & w/dye		Q3	0337	8008
75561	Cardiac mri for morph w/dye		Q3	0337	8008
75563	Card mri w/stress img & dye		Q3	0337	8008
C8900	MRA w/cont, abd		Q3	0284	8008
C8902	MRA w/o fol w/cont, abd		Q3	0337	8008
C8903	MRI w/cont, breast, uni		Q3	0284	8008
C8905	MRI w/o fol w/cont, brst, un		Q3	0337	8008
C8906	MRI w/cont, breast, bi		Q3	0284	8008
C8908	MRI w/o fol w/cont, breast,		Q3	0337	8008
C8909	MRA w/cont, chest		Q3	0284	8008
C8911	MRA w/o fol w/cont, chest		Q3	0337	8008
C8912	MRA w/cont, lwr ext		Q3	0284	8008
C8914	MRA w/o fol w/cont, lwr ext		Q3	0337	8008
C8918	MRA w/cont, pelvis		Q3	0284	8008
C8920	MRA w/o fol w/cont, pelvis		Q3	0337	8008

70492	Ci soft tissue neck w/o & w/dye		Q3	0333	8006
70496	Ci angiography, head		Q3	0662	8006
70498	Ci angiography, neck		Q3	0662	8006
71260	Ci thorax w/dye		Q3	0283	8006
71270	Ci thorax w/o & w/dye		Q3	0333	8006
71275	Ci angiography, chest		Q3	0662	8006
72126	Ci neck spine w/dye		Q3	0283	8006
72127	Ci neck spine w/o & w/dye		Q3	0333	8006
72129	Ci chest spine w/dye		Q3	0283	8006
72130	Ci chest spine w/o & w/dye		Q3	0333	8006
72132	Ci lumbar spine w/dye		Q3	0283	8006
72133	Ci lumbar spine w/o & w/dye		Q3	0333	8006
72191	Ci angiography pelv w/o&w/dye		Q3	0662	8006
72193	Ci pelvis w/dye		Q3	0283	8006
72194	Ci pelvis w/o & w/dye		Q3	0333	8006
73201	Ci upper extremity w/dye		Q3	0283	8006
73202	Ci upper extremity w/o&w/dye		Q3	0333	8006
73206	Ci angio upr extrm w/o&w/dye		Q3	0662	8006
73701	Ci lower extremity w/dye		Q3	0283	8006
73702	Ci lwr extremity w/o&w/dye		Q3	0333	8006
73706	Ci angio lwr extr w/o&w/dye		Q3	0662	8006
74160	Ci abdomen w/dye		Q3	0283	8006
74170	Ci abdomen w/o & w/dye		Q3	0333	8006
74175	Ci angio abdom w/o & w/dye		Q3	0662	8006
74262	Ci colonography, w/dye	NI	Q3	0283	8006
75635	Ci angio abdominal arteries		Q3	0662	8006
70336	Magnetic image, jaw joint		Q3	0336	8007 or 8008
70540	Mri orbit/face/neck w/o dye		Q3	0336	8007 or 8008
70544	Mr angiography head w/o dye		Q3	0336	8007 or 8008
70547	Mr angiography neck w/o dye		Q3	0336	8007 or 8008
70551	Mri brain w/o dye		Q3	0336	8007 or 8008
70554	Fmri brain by tech		Q3	0336	8007 or 8008
71550	Mri chest w/o dye		Q3	0336	8007 or 8008
72141	Mri chest spine w/o dye		Q3	0336	8007 or 8008
72146	Mri chest spine w/dye		Q3	0336	8007 or 8008
72148	Mri lumbar spine w/o dye		Q3	0336	8007 or 8008
72195	Mri pelvis w/o dye		Q3	0336	8007 or 8008
73218	Mri upper extremity w/o dye		Q3	0336	8007 or 8008
73221	Mri joint upr extrem w/o dye		Q3	0336	8007 or 8008
73716	Mri lower extremity w/o dye		Q3	0336	8007 or 8008
73721	Mri jnt of lwr extre w/o dye		Q3	0336	8007 or 8008
74181	Mri abdomen w/o dye		Q3	0336	8007 or 8008
75557	Cardiac mri for morph		Q3	0336	8007 or 8008
75559	Cardiac mri w/stress img		Q3	0336	8007 or 8008
C8901	MRA w/o cont, abd		Q3	0336	8007 or 8008
C8904	MRI w/o cont, breast, uni		Q3	0336	8007 or 8008



Federal Register

**Friday,
November 20, 2009**

Part III

Federal Reserve System

**12 CFR Part 205
Electronic Fund Transfers; Proposed Rule**

FEDERAL RESERVE SYSTEM**12 CFR Part 205****[Regulation E; Docket No. R-1377]****Electronic Fund Transfers****AGENCY:** Board of Governors of the Federal Reserve System.**ACTION:** Proposed rule; request for public comment.

SUMMARY: The Board is proposing to amend Regulation E, which implements the Electronic Fund Transfer Act, and the official staff commentary to the regulation, which interprets the requirements of Regulation E. The proposal restricts a person's ability to impose dormancy, inactivity, or service fees for certain prepaid products, primarily gift cards. In addition, the proposal generally prohibits the sale or issuance of such products if they have an expiration date of less than five years. The proposed amendments implement statutory requirements set forth in the Credit Card Accountability Responsibility and Disclosure Act of 2009 that are effective on August 22, 2010.

DATES: Comments must be received on or before December 21, 2009.**ADDRESSES:** You may submit comments, identified by Docket No. R-1377, by any of the following methods:

- *Agency Web Site:* <http://www.federalreserve.gov>. Follow the instructions for submitting comments at <http://www.federalreserve.gov/generalinfo/foia/ProposedRegs.cfm>.
- *Federal eRulemaking Portal:* <http://www.regulations.gov>. Follow the instructions for submitting comments.
- *E-mail:* regs.comments@federalreserve.gov. Include the docket number in the subject line of the message.
- *Fax:* (202) 452-3819 or (202) 452-3102.

- *Mail:* Jennifer J. Johnson, Secretary, Board of Governors of the Federal Reserve System, 20th Street and Constitution Avenue, NW., Washington, DC 20551.

All public comments are available from the Board's Web site at <http://www.federalreserve.gov/generalinfo/foia/ProposedRegs.cfm> as submitted, unless modified for technical reasons. Accordingly, your comments will not be edited to remove any identifying or contact information. Public comments may also be viewed electronically or in paper form in Room MP-500 of the Board's Martin Building (20th and C Streets, NW.) between 9 a.m. and 5 p.m. on weekdays.

FOR FURTHER INFORMATION CONTACT: Ky Tran-Trong, Counsel, Vivian Wong, Senior Attorney, or Mandie Aubrey or Dana Miller, Attorneys, Division of Consumer and Community Affairs, Board of Governors of the Federal Reserve System, Washington, DC 20551, at (202) 452-2412 or (202) 452-3667. For users of Telecommunications Device for the Deaf (TDD) only, contact (202) 263-4869.

SUPPLEMENTARY INFORMATION:**I. Statutory Background**

The Electronic Fund Transfer Act (15 U.S.C. 1693 *et seq.*) (EFTA or Act), enacted in 1978, provides a basic framework establishing the rights, liabilities, and responsibilities of participants in electronic fund transfer (EFT) systems. The EFTA is implemented by the Board's Regulation E (12 CFR part 205). Examples of the types of transactions covered by the EFTA and Regulation E include transfers initiated through an automated teller machine (ATM), point-of-sale (POS) terminal, automated clearinghouse (ACH), telephone bill-payment plan, or remote banking service. The Act and regulation provide for the disclosure of terms and conditions of an EFT service; documentation of EFTs by means of terminal receipts and periodic statements; limitations on consumer liability for unauthorized transfers; procedures for error resolution; and certain rights related to preauthorized EFTs. Further, the Act and regulation restrict the unsolicited issuance of ATM cards and other access devices.

The official staff commentary (12 CFR part 205 (Supp. I)) interprets the requirements of Regulation E to facilitate compliance and provides protection from liability under Sections 915 and 916 of the EFTA for financial institutions and other persons subject to the Act who act in conformity with the Board's commentary interpretations. 15 U.S.C. 1693m(d)(1). The commentary is updated periodically to address significant questions that arise.

On May 22, 2009, the Credit Card Accountability Responsibility and Disclosure Act of 2009 (Credit Card Act) was signed into law.¹ Section 401 of the Credit Card Act amends the EFTA and imposes certain restrictions on a person's ability to impose dormancy, inactivity, or service fees with respect to gift certificates, store gift cards, and general-use prepaid cards. In addition, the Credit Card Act generally prohibits the sale or issuance of such products if they are subject to an expiration date

earlier than five years from the date of issuance of a gift certificate or the date on which funds were last loaded to a store gift card or general-use prepaid card.

The Board must prescribe rules implementing EFTA Section 915 within nine months after enactment of the Credit Card Act. The gift card and related provisions become effective 15 months after enactment, or on August 22, 2010. *See* EFTA Section 915(d)(3); Section 403 of the Credit Card Act.

II. Background

A gift card is a type of prepaid card that is designed to be purchased by one consumer and given to another consumer as a present or expression of appreciation or recognition. When provided in the form of a plastic card, a user of a gift card is able to access and spend the value associated with the device by swiping the card at a point-of-sale terminal, much as a person would use a debit card. Among the benefits of a gift card are the ease of purchase for the gift-giver and the recipient's ability to choose the item or items ultimately purchased using the card. According to one survey, over 95 percent of Americans have received or purchased a gift card.²

There are two distinct types of gift cards: closed-loop cards and open-loop cards. Closed-loop gift cards constitute the majority of the gift card market, both in terms of number of cards issued and the dollar value of the amounts loaded onto or spent with gift cards.³ These cards generally are accepted or honored at a single merchant or a group of affiliated merchants (such as a chain of book stores or clothing retailers) as

² *See* Comdata, 2007 Adult Gift Card Study (available at: http://storedvalue.com/assets/pdf/study/2007/study_adult_gift_card_2007.pdf).

³ There are no consensus industry figures about the overall size of the prepaid card market. *See* Rachel Schneider, "The Industry Forecast for Prepaid Cards, 2009," Center for Financial Services Innovation (March 2009) at 4 (available at: http://www.cfsinnovation.com/research-paper-detail.php?article_id=330539). According to the Federal Reserve's 2007 Electronic Payments Study, \$36.6 billion was spent using closed-loop prepaid cards in 2006, compared to \$13.3 billion spent using open-loop prepaid cards. *See* 2007 Federal Reserve Electronic Payments Study 27-42 (March 2008). Industry studies using different methodologies suggest a larger prepaid card market, but nonetheless confirm that the closed-loop cards make up a substantial portion of the market. *See, e.g.,* Tim Sloane, "Sixth Annual Closed Loop Prepaid Market Assessment," Mercator Advisory Group (October 2009) (estimating that of the \$247.7 billion total amount loaded across all prepaid segments in 2008, 75 percent, or \$187.24 billion, were loaded onto closed-loop cards, including closed-loop gift cards); "Loyalty is Closed-Loop Gift Card's 'Second Wind,'" *The Green Sheet*, at 53 (May 29, 2009) (citing an Aite Group estimate of 904 million closed-loop gift cards sold in 2007).

¹ Public Law 111-24, 123 Stat. 1734 (2009).

payment for goods or services. They have limited functionality and generally can only be used to make purchases at the merchant or group of merchants.

Closed-loop gift cards are typically issued by a merchant, or by a card program sponsor or service provider working with a merchant, and not by a financial institution. These cards may be sold in a predenominated or consumer-specified amount at the merchant itself or distributed through other retail outlets, such as at grocery stores or drug stores. Generally, closed-loop gift cards may not be reloaded with additional value after card issuance. With closed-loop gift cards, the issuer typically does not collect any information regarding the identity of the gift card purchaser or the recipient.

For merchant-issuers, gift cards have largely replaced paper-based gift certificates as a more cost-effective and efficient means of facilitating gift-giving by consumers. In addition to reducing costs associated with the issuance of paper certificates, electronic gift cards may also be less vulnerable to fraud or counterfeiting. Merchants benefit from the sale of items purchased with gift cards, as well as from additional spending by gift card recipients beyond the face amount on the card. Merchants may also derive revenue from the imposition of certain fees, such as from monthly maintenance or transaction-based fees or from interest earned from unused card balances.

Open-loop gift cards differ in several respects from closed-loop gift cards. First, open-loop gift cards typically carry a card network brand logo (such as Visa, MasterCard, American Express, or Discover). Thus, they can be used at a wide variety of merchants that accept or honor cards displaying that brand. Second, open-loop gift cards are generally issued by financial institutions. Third, open-loop gift card transactions are processed over the debit or credit card networks. Fourth, open-loop gift cards may carry additional, and in some cases higher, fees than closed-loop gift cards as a result of higher compliance and customer service costs. Fifth, open-loop gift cards are more likely to offer the capability of being reloaded with additional value (reloadable) than are closed-loop gift cards.

A consumer may obtain gift cards in several ways. Gift cards can be purchased at retail locations, by telephone, or on-line, and used either for the purchaser's own purposes or given to another consumer as a gift. In addition, gift cards can be received through a loyalty, award, or promotional program. For example, a merchant may

distribute its own closed-loop gift card to encourage consumers to visit the store or for customer retention purposes, such as through a loyalty or frequent buyer program. Merchants and product manufacturers may also issue gift cards to consumers to provide a rebate for the consumer's purchase of a particular product instead of sending rebate checks. Employers may provide gift cards to their employees as a reward for good job performance.

Concerns have been raised regarding the amount of fees associated with gift cards, the expiration dates of gift cards, and the adequacy of disclosures. Consumers who do not use the value of the card within a short period of time may be surprised to find that the card has expired or that dormancy or service fees have reduced the value of the card. Even where fees or terms are disclosed on or with the card, the disclosures may not be clear and conspicuous.

At the State level, more than 40 States have enacted laws applicable to gift cards in some fashion. Most commonly, State gift card laws may restrict the circumstances under which dormancy, inactivity, or service fees may be charged and/or restrict the circumstances under which the card or funds underlying the card may expire.⁴ Other State laws simply require the disclosure of fees or expiration dates. Many States have applied abandoned property or escheat laws to funds remaining on gift cards, and some States require that consumers have the option of receiving cash back when the underlying balance falls below a certain amount. However, while all State gift card laws address closed-loop gift cards in some form, many State laws do not apply to open-loop bank-issued cards.⁵

III. Summary of Proposal

Restrictions on Dormancy, Inactivity, or Service Fees

Under the proposed rule, no person may impose a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card, or general-use prepaid card, unless three conditions are satisfied. First, such fees may be imposed only if there has been no activity with respect to the certificate or card within the one-year period prior to

the imposition of the fee. Second, only one such fee may be assessed in a given calendar month. Third, disclosures regarding dormancy, inactivity, or service fees must be clearly and conspicuously stated on the certificate or card, and the issuer or vendor must provide these disclosures to the purchaser before the certificate or card is purchased.

Expiration Date Restrictions

The proposed rule would also provide that a gift certificate, store gift card, or general-use prepaid card may not be sold or issued unless the expiration date of the funds underlying the certificate or card is no less than five years after the date of issuance (in the case of a gift certificate) or five years after the date of last load of funds (in the case of a store gift card or general-use prepaid card). In addition, information regarding whether funds underlying a certificate or card may expire must be clearly and conspicuously stated on the certificate or card and disclosed prior to purchase.

Two proposed alternative approaches are set forth to minimize potential confusion for consumers if the expiration date on a certificate or card and the expiration date for the underlying funds differ. The first alternative would prohibit the sale or issuance of a certificate or card that has a printed expiration date that is less than five years from the date of purchase. The second alternative would require policies or procedures to ensure that a consumer has a reasonable opportunity to purchase a certificate or card that has an expiration date that is at least five years from the date of purchase.

The proposed rule would also require a certificate or card to include a disclosure alerting consumers to the difference between the certificate or card expiration date and the funds expiration date, if any, and that the consumer may contact the issuer for a replacement card. This disclosure must be stated with equal prominence and in close proximity to the certificate or card expiration date. In addition, the proposed rule would prohibit the imposition of any fees for replacing an expired certificate or card to ensure that consumers are able to access the underlying funds for the full five-year period.

Additional Disclosure Requirements Regarding Fees

In addition to the statutory restrictions for dormancy, inactivity, or service fees, the proposed rule would require the disclosure of all other fees imposed in connection with a gift

⁴ See, e.g., Consumers Union, State Gift Card Consumer Protection Laws (available at: http://www.consumersunion.org/pub/core_financial_services/003889.html); National Conference of State Legislatures, Gift Cards and Gift Certificates Statutes and Recent Legislation (available at: <http://www.ncsl.org/programs/banking/GiftCardsandCerts.htm>).

⁵ See, e.g., Ark. Code § 4–88–704; Cal. Civil Code § 1749.45; Fla. Stat. § 501.95; and Md. Commercial Code Ann. § 14–1320.

certificate, store gift card, or general-use prepaid card. These disclosures would have to be provided on or with the certificate or card and disclosed prior to purchase. The proposed rule would also require the disclosure on the certificate or card of a toll-free telephone number and, if one is maintained, a Web site that a consumer may use to obtain fee information or replacement certificates or cards.

Exclusions

Consistent with the statute, the proposed rule excludes certain card products from the definitions of gift certificate, store gift card, or general-use prepaid card. For example, cards, codes, or other devices that are issued in connection with a loyalty, award, or promotional program, or that are reloadable and not marketed or labeled as a gift card or gift certificate, would not be subject to the substantive restrictions on imposing dormancy, inactivity, or service fees, or on expiration dates. However, under the proposal, disclosures of all fees, including any dormancy, inactivity, or service fees, and any expiration date that may apply, would be required for certificates or cards issued through a loyalty, award, or promotional program.

IV. Legal Authority

Section 401 of the Credit Card Act creates a new Section 915 of the EFTA and prohibits any person from charging dormancy, inactivity, or service fees with respect to a gift certificate, store gift card, or general-use prepaid card, as those terms are defined in the Act, unless there have been at least 12 months of inactivity with respect to the certificate or card, not more than one fee is charged in any given month, and certain disclosures regarding such fees are provided to the consumer. *See* EFTA Section 915(b); 15 U.S.C. 1693m(b). In addition, Section 401 of the Credit Card Act makes it unlawful for any person to sell or issue a gift certificate, store gift card, or general-use prepaid card that is subject to an expiration date, unless the expiration date is at least five years after the date on which a gift certificate is issued or five years after funds are last loaded on a store gift card or general-use prepaid card, and the terms of expiration are clearly and conspicuously disclosed. *See* EFTA Section 915(c); 15 U.S.C. 1693m(c).

Section 401(d)(1) of the Credit Card Act requires the Board to prescribe rules to carry out the new requirements. This section also gives the Board the authority to prescribe rules addressing the amount of dormancy, inactivity, or service fees that may be imposed, and

the balance below which such fees may be assessed. *See* EFTA Section 915(d)(1); 15 U.S.C. 1693m(d)(1). In addition, Section 401(d)(2) of the Credit Card Act requires the Board to determine the extent to which the individual definitions and provisions of the EFTA and Regulation E should apply to gift certificates, store gift cards, and general-use prepaid cards. *See* EFTA Section 915(d)(2); 15 U.S.C. 1693m(d)(2). Lastly, Section 402 of the Credit Card Act amends EFTA Section 920 to provide that the EFTA does not preempt any State laws that address dormancy, inactivity, or service fees or expiration dates for gift certificates, store gift cards, or general-use prepaid cards if such State laws provide greater consumer protection than the new gift card provisions.

In addition to the statutory mandates set forth in the Credit Card Act, Section 904(a) of the EFTA authorizes the Board to prescribe regulations necessary to carry out the purposes of the title. The express purposes of the EFTA are to establish “the rights, liabilities, and responsibilities of participants in electronic fund transfer systems” and to provide “individual consumer rights.” *See* EFTA Section 902(b); 15 U.S.C. 1693. Section 904(c) of the EFTA further provides that regulations prescribed by the Board may contain any classifications, differentiations, or other provisions, and may provide for such adjustments or exceptions for any class of electronic fund transfers, that the Board deems necessary or proper to effectuate the purposes of the title, to prevent circumvention or evasion, or to facilitate compliance.

V. Section-by-Section Analysis

Section 205.4 General Disclosure Requirements; Jointly Offered Services

Section 205.4 contains the general disclosure requirements under Regulation E, including provisions relating to the form of disclosure. Section 205.4(a)(1) provides that disclosures required by the regulation shall be clear and readily understandable, in writing, and in a form that the consumer may keep. To clarify that this standard is one of general application, the Board is proposing to revise § 205.4(a)(1) to provide that for certain disclosures required by the regulation, different disclosure standards may apply when specified in the rule.

For example, as further discussed below, the disclosures for certain prepaid cards set forth in this proposal are subject to a “clear and conspicuous” standard, consistent with new Section

915 of the EFTA, rather than the “clear and readily understandable” standard that generally applies under Regulation E. *See* proposed § 205.20, discussed below. Similarly, under § 205.11(c), notices provided by financial institutions to satisfy the error investigation requirements may be provided orally or in writing. *See* comment 11(c)–1.

Section 205.12 Relation to Other Laws

Section 920 of the EFTA (as redesignated by the Credit Card Act) provides that the EFTA does not preempt any State laws relating to electronic fund transfers except to the extent that such laws are inconsistent with the EFTA’s provisions. Section 920 further clarifies that a State law is not inconsistent with the EFTA if the State law provides greater protection for the consumer than under the Act. Accordingly, Section 920 effectively creates a Federal floor for the protections set forth in the Act (floor preemption). Section 205.12(b) of Regulation E implements this provision.

The Credit Card Act amended Section 920 of the EFTA to apply the EFTA’s existing preemption provisions to State laws that address “dormancy fees, inactivity charges or fees, service fees, or expiration dates of gift certificates, store gift cards, or general-use prepaid cards.” *See* Section 402 of the Credit Card Act. Thus, State laws that provide greater protection for consumers than Title IV of the Credit Card Act as codified in the EFTA, are not preempted by the EFTA. The Board is proposing to amend § 205.12(b) of Regulation E and comment 12(b)–1 to conform with the amendments to Section 920 of the EFTA made by the Credit Card Act.

Section 205.20 Requirements for Gift Cards and Gift Certificates

20(a) Definitions

New EFTA Section 915(a)(2) generally defines the scope of gift cards and gift certificates that are subject to the Credit Card Act’s restrictions on dormancy, inactivity, or service fees and the terms of expiration. Specifically, Section 915 applies to gift certificates, store gift cards, and general-use prepaid cards as those terms are defined in the statute. In addition, new EFTA Section 915(a)(1) defines a dormancy fee, inactivity charge or fee, and new EFTA Section 915(a)(3) defines a service fee. *See* 15 U.S.C. 1693m(a). Proposed § 205.20(a) defines the following terms: gift certificate; store gift card; general-use prepaid card; loyalty, award, or promotional gift card; dormancy fee, inactivity charge or fee; and service fee.

The proposed definitions of gift certificate, store gift card, and general-use prepaid card generally track the definitions set forth in the statute. However, the Board is proposing certain adjustments to the statutory definitions pursuant to its authority under EFTA Section 904(c) to provide clarity and to harmonize key terms throughout the rule. In general, these adjustments are not intended to make substantive changes to the statutory definitions.

As an initial matter, the Board notes that new EFTA Section 915 does not use consistent terminology to describe the payment devices covered by the statute. For example, the statutory definition of a general-use prepaid card refers to a “card or other payment code or device,” while the statutory definition of a store gift card refers to an “electronic promise, plastic card, or other payment code or device.”

The Board does not believe that distinguishing the types of products covered by the rule by, for instance, the material that is used to produce a payment card would be consistent with the statute’s overall purpose. The adoption of such distinctions would result in some gift card products being excluded from the rule altogether based on the type of material used to make the card. For example, if the definition of store gift card literally required a card to be made out of plastic, then a reloadable gift card that was made with a different material would neither be a store gift card nor fall under any of the other definitions of covered products.⁶

In addition, the exclusions in EFTA Section 915(a)(2)(D) apply to an “electronic promise, plastic card, or payment code or device” that meets certain specified criteria. The Board does not believe that an issuer that, for example, chooses to use non-plastic biodegradable materials to create a more environmentally-friendly product should be precluded from relying on an exclusion solely because its payment device is not made of plastic. Therefore, the proposed rule generally refers to “cards, codes, or other devices” to avoid such arbitrary distinctions and to provide consistency across the definitions.

Proposed comment 20(a)–1 clarifies that the requirements of § 205.20 generally apply to all cards, codes, or other devices that meet the definition of gift certificate, store gift card, or general-use prepaid card, even if they are not issued in card form. That is, the rule would apply even if a physical card or

certificate is not issued. The proposed comment clarifies that products not issued in card form, such as an account number or bar code that enables the consumer to access underlying funds, would be subject to § 205.20 if they otherwise meet the definition of gift certificate, store gift card, or general-use prepaid card. Similarly, § 205.20 would apply to a device with a chip or other embedded mechanism which links the device to stored funds, such as a mobile phone or sticker containing a contactless chip, if the device otherwise meets the definition of gift certificate, store gift card or general-use prepaid card.

In addition, the term “electronic promise” is used in several places in the statute to refer to a type of payment mechanism or device. See EFTA Sections 915(a)(2)(B), (a)(2)(C), and (a)(2)(D). The Board does not believe, however, that there is a meaningful distinction between electronic promises and cards, codes, or other devices that can be used as payment mechanisms. Instead, the Board views an electronic promise as a commitment to pay that is itself manifested or represented by a “card, code, or other device,” rather than as a distinct payment mechanism. Proposed comment 20(a)–2 clarifies that the term “electronic promise” means “a person’s commitment or obligation communicated or stored in electronic form made to a consumer to provide payment for goods or services for transactions initiated by the consumer.”⁷ The proposed comment further provides that the promise is represented by a card, code, or other device that is issued or honored by the person, reflecting the person’s commitment or obligation to pay. Thus, the proposal contemplates that the term “card, code, or other device” when used in the regulation also incorporates the statutory reference to “electronic promises.” For example, if a merchant issues a code that can be given as a gift and redeemed by the recipient in an on-line transaction for goods or services, that code represents an electronic promise by the merchant and would be a card, code, or other device covered by § 205.20. See proposed comment 20(a)–2.

Last, the statutory definitions of “gift certificate” and “store gift card” refer to products that are “issued in a *specified* amount.” In contrast, the statutory definition of a “general-use prepaid

card” refers to products that are “issued in a *requested* amount.” One way to reconcile the use of these different terms in the statute is to interpret “specified” as referring to cards that are issued in a predenominated amount (e.g., a \$50 gift card), and to interpret “requested” as referring to a consumer-requested amount (e.g., where the consumer states the amount to load on a gift card). Such an interpretation would mean that gift certificates and store gift cards issued in a consumer-requested amount and general-use prepaid cards issued in a predenominated amount would be excluded from the rule. The Board does not believe that such a result would be consistent with the statute’s purpose.

The Board believes that consumers should receive the same protections when purchasing gift cards or gift certificates regardless of whether the amount on the card or certificate is determined by the issuer or the consumer. Thus, the Board is interpreting the statutory definitions of gift certificate, store gift card, and general-use prepaid card broadly to cover both predenominated and consumer-designated certificates or cards. Therefore, the proposed rule uses the term “specified” consistently across all three defined product terms to capture all certificates or cards whether they are issued in predenominated amounts or in a consumer-requested, or variable load, amount.

The Board notes that although the EFTA generally applies only to consumer accounts, the gift card provisions of the Credit Card Act do not expressly limit the scope of the new restrictions to cards issued for non-business purposes. The Board solicits comment on whether it is appropriate to limit the scope of the final rule so that it does not apply to cards issued for business purposes. Any such limitation, however, would presumably not exclude cards that are purchased by a business for the purposes of redistribution or resale to consumers for consumers to use. For example, a program manager may purchase gift cards directly from an issuing merchant and sell those cards through the program manager’s retail outlets. Or, a corporation may give gift cards it has purchased directly from the issuing merchant to consumers pursuant to a reward or other incentive program. In such cases, the Board believes that because the end use of the gift card is for consumer purposes, the consumer protections provided by the Credit Card Act should apply, unless the card is otherwise excluded. (See EFTA Section 915(a)(2)(D) and proposed § 205.20(b), discussed below.) Accordingly, given

⁶Products issued in paper form only are excluded under new EFTA Section 915(a)(2)(D)(v) and proposed § 205.20(b)(5).

⁷ See, e.g., UCC 3–106(a)(12) (defining “promise” as a “written undertaking to pay money signed by the person undertaking to pay. An acknowledgment of an obligation by the obligor is not a promise unless the obligor also undertakes to pay the obligation.”)

that issuers would have to adopt controls and potentially monitor the distribution or sale of gift cards to ensure that the end use is for business purposes, comment is also requested regarding the overall utility of, or need for, such a scope provision in the final rule.

20(a)(1) Gift Certificate

Proposed § 205.20(a)(1) defines the term “gift certificate” as a card, code, or other device that is: (a) Issued to a consumer in a specified amount that may not be increased or reloaded in exchange for payment; and (b) redeemable upon presentation at a single merchant or an affiliated group of merchants for goods or services. The proposed definition generally tracks the definition set forth in the statute, but modifies the terms to simplify and clarify the definition. *See* EFTA Section 915(a)(2)(B).

The term “affiliated group of merchants”—as further discussed below under the definition of “store gift card”—includes two or more merchants or other persons that are related by common ownership or common corporate control and share the same name, mark or logo. The term also includes two or more merchants or other persons that agree among each other to honor any card, code, or other device that bears the same name, mark, or logo (other than the mark or logo of a payment network) for the purchase of goods or services solely at such merchants or persons. *See* proposed comment 20(a)(2)–2.

20(a)(2) Store Gift Card

Proposed § 205.20(a)(2) defines the term “store gift card” as a card, code, or other device that is: (a) Issued to a consumer in a specified amount, whether or not that amount may be increased or reloaded by the cardholder, in exchange for payment; and (b) redeemable upon presentation at a single merchant or an affiliated group of merchants for goods and services. The proposed definition generally tracks the definition set forth in the statute, but modifies the terms to simplify and clarify the definition. *See* EFTA Section 915(a)(2)(C). Under the proposed rule, closed-loop cards generally would be considered “store gift cards” or “gift certificates,” unless one of the exclusions in § 205.20(b), discussed below, applies.

A card, code, or other device that meets the requirements in proposed § 205.20(a)(2) qualifies as a “store gift card,” whether or not the cardholder may later add more funds to the card, code, or other device. Thus, because

“store gift card” includes non-reloadable cards, codes, or other devices that are redeemable at single merchants or affiliated groups of merchants, proposed comment 20(a)(2)–1 clarifies and illustrates by way of example that a gift certificate as defined in § 205.20(a)(1) would be a type of store gift card.

Proposed comment 20(a)(2)–2 provides guidance on the term “affiliated group of merchants.” Under new EFTA Section 915(a)(2), both the definition of “gift certificate” and “store gift card” refer to certificates or cards that are redeemable at a single merchant or “an affiliated group of merchants that share the same name, mark, or logo.” The term “affiliate” is not defined within the statute. However, in other contexts, “affiliate” is used to describe a relationship between two or more companies that is defined by some form of common ownership or common corporate control by one of the companies. *See, e.g.,* 12 CFR 222.3(b) (defining “affiliate” under the Board’s Regulation V (Fair Credit Reporting)); 12 CFR 223.2 (defining “affiliate” under the Board’s Regulation W (Transactions Between Member Banks and Their Affiliates)). The Board believes that such a concept should similarly apply to the term “affiliate” when used in the proposed rules. Accordingly, the terms “gift certificate” and “store gift card” generally include cards, codes, or other devices that are redeemable at some or all of the companies that are related by virtue of common ownership or common corporate control and that share the same name, mark, or logo. An “affiliated group of merchants” would also include franchisees because franchisees generally are subject to a common corporate set of policies or practices under the terms of their franchise licenses.

Under some retail card programs, merchants that honor the same certificate or card may not be owned or otherwise controlled by the same parent company. For instance, two unrelated companies may be engaged in complementary businesses and agree to operate a common gift card program in which cardholders may use the same certificate or card at either of the two businesses. To illustrate, a movie theater chain and a restaurant chain may decide to operate a gift card program that enables cardholders to use the same gift card to pay for movie tickets and for a meal preceding or following the movie. While such companies would not be considered “affiliates” in other contexts, the Board believes that it is appropriate to treat such arrangements like gift card programs operated by retailers with the

same parent company or under common corporate control. Accordingly, proposed comment 20(a)(2)–2 provides that the term “affiliated group of merchants” would include two or more merchants or other persons that agree among each other, by contract or otherwise, to redeem cards, codes, or other devices bearing the same name, mark, or logo for purchases of goods or services solely at the establishments of such merchants or persons. (*See also* proposed comment 20(a)(3)–2 regarding mall cards, discussed below.) The proposed comment clarifies, however, that merchants or other persons would not be considered affiliated merely because they agree to accept a card that bears the mark, logo, or brand of a payment network. Thus, for example, a grocery store would not be considered affiliated with a hardware store merely because they both agree to accept Visa or MasterCard-branded cards.

Proposed comment 20(a)(2)–3 addresses mall cards and cross-references proposed comment 20(a)(3)–2, discussed below.

20(a)(3) General-Use Prepaid Card

Proposed § 205.20(a)(3) defines “general-use prepaid card” as a card, code, or other device that is: (a) Issued to a consumer in a specified amount, whether or not that amount may be increased or reloaded by the cardholder, in exchange for payment; and (b) redeemable upon presentation at multiple, unaffiliated merchants or service providers for goods or services, or usable at ATMs. The proposed definition generally tracks the definition set forth in the statute, but modifies the terms to simplify and clarify the definition. *See* EFTA Section 915(a)(2)(A). Under the proposed rule, open-loop cards generally are considered to be “general-use prepaid cards,” unless one of the exclusions in § 205.20(b), discussed below, applies.

Proposed comment 20(a)(3)–1 clarifies that a card, code, or other device is “redeemable upon presentation at multiple, unaffiliated merchants” if, for example, the merchants agree to honor the card, code, or device if it bears the mark, logo, or brand of a payment network, pursuant to the rules of the payment network.

One popular form of gift card is a mall gift card, which is generally intended to be used or redeemed at participating retailers located within the same shopping mall. In some cases, however, the mall card may also be network-branded which permits the card to be used at any retailer that accepts that card brand, including retailers located outside of the mall. Proposed comment

20(a)(3)–2 provides that a mall card could be considered a store gift card or a general-use prepaid card depending on the locations in which the card may be redeemed. That is, if use of the mall card is limited to the retailers at the associated shopping mall, the card is more likely to be considered a store gift card. If the mall card also carries the brand of a payment network and can be used at any retailer accepting that brand, the card would be considered a general-use prepaid card. Regardless, the substantive and disclosure requirements of § 205.20 would apply to mall cards whether they are considered store gift cards or general-use prepaid cards.

20(a)(4) Loyalty, Award, or Promotional Gift Card

New EFTA Section 915(a)(2)(D)(iii) excludes an electronic promise, plastic card, or payment code or device from the definitions of “gift certificate,” “store gift card,” or “general-use prepaid card” if it is a loyalty, award, or promotional gift card, as such term is defined by the Board. Proposed § 205.20(a)(4) generally defines the term “loyalty, award, or promotional gift card” as a card, code, or other device that: (a) Is issued in connection with a loyalty, award, or promotional program; (b) is redeemable upon presentation at one or more merchants for goods or services, or usable at ATMs; and (c) provides certain disclosures about any fees and expiration dates that may apply to the card, code, or other device.

As an initial matter, the Board notes that the proposed definition generally applies to any card, code, or other device issued pursuant to a loyalty, award, or promotional program, regardless of whether the consumer has provided any form of payment or other value to obtain the card. The proposed definition covers, for example, gift cards mailed to a consumer as a rebate on a product that a consumer has purchased in response to a sales promotion, and gift cards given by a merchant to reward frequent customers. The definition also covers cards provided by employers to reward job performance. Proposed comment 20(a)(4)–1 provides examples of loyalty, award, or promotional programs.

Under proposed § 205.20(b)(3), further discussed below, if a card, code, or other device is deemed to be a loyalty, award, or promotional gift card, it would not be subject to the substantive restrictions on imposing dormancy, inactivity or service fees, or the requirement to have expiration dates of at least five years. Accordingly, to mitigate potential consumer surprise

from unexpected fees or expiration dates for these cards, proposed § 205.20(a)(4)(iii) provides that in order to qualify as a “loyalty, award, or promotional gift card,” certain disclosures regarding the fees and expiration dates applying to such cards must also be provided to the consumer. These disclosures are discussed in more detail below under § 205.20(b)(3).

20(a)(5) Dormancy or Inactivity Fee

New section 915(a)(1) of the EFTA defines a “dormancy fee,” or an “inactivity charge or fee” as “a fee, charge, or penalty for non-use or inactivity of a gift certificate, store gift card, or general-use prepaid card.” Proposed § 205.20(a)(5) implements this definition with non-substantive wording modifications to improve readability. Because the Board believes the terms “charge” and “penalty” are synonymous with “fee” as used in this definition, the proposal simplifies the definition by not including the references to “charge” or “penalty” used in the statute.

20(a)(6) Service Fee

New EFTA Section 915(a)(3)(A) defines a “service fee” as “a periodic fee, charge, or penalty for holding or use of a gift certificate, store gift card, or general-use prepaid card.” Proposed § 205.20(a)(6) implements this definition using substantially the same language as the statute. Because the Board believes the terms “charge” and “penalty” are synonymous with “fee” as used in this definition, the proposal simplifies the definition by not including the statutory references to “charge” or “penalty” used in the statute.

In addition, proposed comment 20(a)(6)–1 clarifies that a periodic fee is a fee that may be imposed from time to time for holding or using a gift certificate, store gift card, or general-use prepaid card. Such fees may include a monthly maintenance fee, a transaction fee, a reload fee, or a balance inquiry fee, whether or not the fee is waived for a certain period of time or is only imposed after a certain period of time. Transaction fees include, for example, fees imposed each time a transaction is conducted with the certificate or card and foreign transaction fees.

The Board considered an alternative interpretation of a “periodic fee” as a fee that is imposed at regular intervals, which would include a monthly maintenance fee, but not transaction fees or reload fees that are triggered by consumer activity. The Board notes, however, that the statutory definition of “service fee” refers to the “use” of a gift

certificate, store gift card, or general-use prepaid card. *See* new EFTA Section 915(a)(3)(A) (15 U.S.C. 1693m(a)(3)(A)). Therefore, the Board believes that Congress intended to also capture consumer-initiated fees such as transaction fees and reload fees in the definition of “service fee.” Moreover, the Board is concerned that a narrow interpretation of “service fee” would lead to circumvention by issuers and result in a shift in fee structures from fees imposed at regular intervals to fees that are imposed for a transaction or service associated with the certificate or card. The Board believes that interpreting the term “service fee” broadly, and thus limiting the imposition of such fees, will improve the transparency and predictability of costs to the consumer.

Consistent with new EFTA Section 915(a)(3)(B), proposed comment 20(a)(6)–1 also clarifies that a one-time initial issuance fee is not a service fee. Proposed comment 20(a)(6)–1 also provides examples of other one-time fees that are not service fees, including cash-out fees.

20(b) Exclusions

New EFTA Section 915(a)(2)(D) states that the terms “general-use prepaid card,” “gift certificate,” and “store gift card” do not include an electronic promise, plastic card, or payment code or device that falls into one of six specified categories. *See* 15 U.S.C. 1593m(a)(2)(D). For example, reloadable cards that are not marketed or labeled as a gift card or gift certificate are excluded from the statutory definitions. Similarly, prepaid cards that are not marketed to the general public are excluded from the statutory definitions. Thus, under the statute, an excluded promise, card, code, or device is not subject to the substantive restrictions regarding when a dormancy, inactivity, or service fee may be imposed, or on expiration dates. These excluded products also are not subject to the disclosure requirements in the statute.

Proposed § 205.20(b) implements the statutory exclusions and provides that the terms “gift certificate,” “store gift card,” and “general-use prepaid card” do not include any cards, codes, or other devices that meet any of the six conditions specified in the statute. As noted above, the proposed rule uses the term “card, code, or other device,” instead of the term “electronic promise, plastic card, or payment code or device” for clarity and no substantive difference is intended.

Proposed comment 20(b)–1 provides guidance on the effect of meeting any of the specified exclusions. The comment

states that an excluded card, code, or other device is not subject to any of the substantive restrictions and disclosure requirements regarding the imposition of dormancy, inactivity, or service fees, or expiration dates. The proposed comment also provides that the additional disclosures in proposed § 205.20(f) regarding other fees imposed in connection with a card, code, or other device do not apply to an excluded card, code, or other device.⁸

Proposed comment 20(b)–2 clarifies that a card, code, or other device may qualify for one or more exclusions. For example, a corporation may award its employees with a gift card that is marketed solely to businesses for incentive-related purposes. Under this example, the card, code, or other device may qualify for the exclusion for loyalty, award, or promotional gift cards, or for the exclusion for cards, codes, or other devices not marketed to the general public. Even if a card, code, or other device does not qualify for a particular exclusion, it may still fall outside the rule under a different exclusion. Thus, for example, if the gift card awarded by the corporation is of a type that can also be purchased directly from a merchant, the gift card may fall outside coverage under the rule because it is a loyalty, award, or promotional gift card (provided that certain disclosures are provided with the card as proposed under § 205.20(a)(4)(iii)), even though the card would not qualify as a card that is not marketed to the general public because it can also be obtained through retail channels. See proposed § 205.20(b)(4), discussed below.

The six specific exclusions are discussed below.

20(b)(1) Usable Solely for Telephone Services

Proposed § 205.20(b)(1) implements the exclusion for cards, codes, or other devices that are usable solely for telephone services. See EFTA Section 915(a)(2)(D)(i). Proposed comment 20(b)(1)–1 contains examples of products that fall within this exclusion, such as prepaid cards for long-distance telephone service and prepaid cards for wireless telephone service. The proposed comment further clarifies that this exclusion also includes prepaid products that may be used for other services analogous in function to a telephone, such as prepaid cards for voice over Internet protocol (VoIP) access time.

The Board notes that mobile phones today are capable of a number of

different functions in addition to voice communications, including providing consumers the ability to send text messages and to access the Internet. Accordingly, the Board solicits comment on whether it should exercise its authority under EFTA Section 904 to expand the proposed exclusion to cover other prepaid cards that may be redeemed for similar or related technology services, such as prepaid cards used to obtain mobile broadband or Internet access time. See, e.g., N.J. Rev. Stat. § 56:8–110 (excluding prepaid telecommunications and technology cards from the definitions of “gift card” and “gift certificate”). The Board is concerned that interpreting the exclusion narrowly may have the unintended effect of reducing the availability or variety of prepaid telephone certificates or cards in the market.

20(b)(2) Reloadable and Not Marketed or Labeled as a Gift Card or Gift Certificate

Proposed § 205.20(b)(2) implements the exclusion for cards, codes, or other devices that are reloadable and not marketed or labeled as a gift card or gift certificate. See EFTA Section 915(a)(2)(D)(ii).

Consistent with the statute, the card, code, or other device must be both reloadable *and* not marketed or labeled as a gift card or gift certificate to qualify for the exclusion. Thus, a non-reloadable card is not excluded, even if it is not marketed or labeled as a gift card or gift certificate, unless a different exclusion applies. Similarly, a reloadable card that is marketed as a gift card or gift certificate does not qualify for the exclusion. Proposed comment 20(b)(2)–1 provides that a card, code, or other device is “reloadable” if it has the capability of having more funds added by a consumer after the initial purchase or issuance.

Proposed comment 20(b)(2)–2 clarifies the meaning of the term “marketed or labeled as a gift card or gift certificate.” Under the proposed comment, the term means directly or indirectly offering, advertising, or otherwise suggesting the potential use of a card, code, or other device as a gift for another person. Moreover, whether the exclusion applies does not depend on the type of entity that is making the promotional message. For example, a card may be marketed or labeled as a gift card or gift certificate if anyone (other than the purchaser of the card),⁹

including the issuer, the retailer, the program manager that may distribute the card, or the payment network on which a card is used, promotes the use of the card as a gift card or gift certificate. A certificate or card, including a general-purpose reloadable card, may also be deemed to be marketed or labeled as a gift card or gift certificate even if it is primarily marketed for another purpose. For example, a reloadable network-branded card would be marketed or labeled as a gift card or gift certificate if the issuer principally advertises the card as a less costly alternative to a bank account but promotes the card in a television, radio, newspaper, or Internet advertisement, or on signage as “the perfect gift” during the holiday season.

Proposed comment 20(b)(2)–3 provides positive and negative examples of the term “marketed or labeled as a gift card or gift certificate.” Positive examples of marketing or labeling as a gift card or gift certificate include displaying the word “gift” or “present,” displaying a congratulatory message, and incorporating gift-giving or celebratory imagery or motifs on the card, certificate or accompanying material, such as documentation, packaging and promotional displays. In contrast, a card, code, or other device is not marketed or labeled as a gift card or gift certificate if the issuer, vendor, or other person represents that the card, code, or other device can be used as a substitute for a checking, savings, or deposit account, as a budgetary tool, or to cover emergency expenses. Similarly, a card, code, or other device is not marketed as a gift card or gift certificate if it is promoted as a substitute for travelers’ checks or cash for personal use, or promoted as a means of paying for a consumer’s health-related expenses. See proposed comment 20(b)(2)–3. The Board solicits comment on whether additional guidance on marketing is necessary to provide clarity with respect to the activities that may trigger coverage under the rule and the activities that would not.

As discussed above, a gift card may be sold directly to the consumer by a merchant at the merchant’s store. In this type of arrangement, the merchant is typically the primary party involved in issuing the card and operating the card program. As such, the issuer can be expected to have substantial control over all facets of the card program, including how the card is sold or marketed.

⁹ Thus, a card would not be deemed to be marketed or labeled as a gift card or gift certificate solely because the purchaser gives the card to another consumer as a “gift.”

⁸ See, however, proposed § 205.20(a)(4)(iii) with respect to loyalty, award, or promotional gift cards.

In other cases, a gift card may be sold to consumers through another merchant or retailer, such as a grocery store or a drug store, on display racks that may make retail gift cards available alongside gift cards from other merchants and other types of prepaid cards, including general-purpose reloadable cards and telephone cards. In this type of arrangement, multiple parties are generally involved in the card distribution process. These parties may include: an issuer (whether it is a merchant or a bank); a program manager who works with issuers to administer any or all aspects of a card program, including transaction processing, distribution, and marketing; and a seller or distributor of the card.¹⁰ A seller or distributor of the card can be an issuer, a program manager, or another party, such as a shopping mall or a retailer. In these arrangements, responsibilities for operating the program, including compliance with applicable laws or payment network rules, are generally allocated by contract.

When multiple parties are involved in a card program, the issuer may not play a significant role in the card distribution process and thus may have less control over how the card is displayed or marketed at the locations where the card is sold. An exclusion that depends upon how a card is marketed therefore poses substantial compliance risk for an issuer that cannot fully control how its prepaid cards are marketed to consumers. For example, where a card is sold in a substantial number of retail outlets, the card issuer cannot verify in every instance how the card is displayed or marketed at each retail outlet to ensure that it is not being marketed as a gift card or gift certificate through signage, advertisements, or otherwise.

To address this issue, proposed comment 20(b)(2)–4 provides that the exclusion for a card, code, or other device that is reloadable and not marketed or labeled as a gift card or gift certificate applies if the individual card, code, or other device is not marketed or labeled as a gift card or gift certificate and if entities subject to the rule maintain policies and procedures reasonably designed to avoid such marketing. The proposed comment provides illustrative examples of procedures that would qualify and not qualify for the exclusion for reloadable

cards, codes, or other devices that are not marketed or labeled as gift cards or gift certificates.

Under the first example, an issuer or program manager distributes a general-purpose reloadable card through retailers and enters into a contract with the retailer to establish the terms and conditions under which the card will be sold and marketed at the retailer. The contract includes restrictions prohibiting the general-purpose reloadable card from being sold or otherwise marketed as a gift card or gift certificate, and requirements for policies and procedures to regularly monitor or otherwise verify that the cards are not being sold or marketed as such. The issuer or program manager then sets up one promotional display at the retailer for gift cards and another physically separated display for excluded products under proposed § 205.20(b), including the general-purpose reloadable cards, such that a reasonable consumer would not believe that the excluded cards are gift cards. Under these circumstances, the exclusion in § 205.20(b)(2) applies even if a retail clerk inadvertently stocks or places some of the general-purpose reloadable cards on the gift card display because the issuer or program manager maintains policies and procedures reasonably designed to avoid the marketing of the general-purpose reloadable card as a gift card or gift certificate. See proposed comment 20(b)(2)–4.i.

In the second example, the same facts apply, except that the issuer or program manager has set up a single promotional display at the retailer on which a variety of prepaid cards, including store gift cards, general-purpose reloadable cards, and wireless telephone cards, are sold. A sign stating “Gift Cards” appears prominently at the top of the display. Under proposed comment 20(b)(2)–4.ii, any general-purpose reloadable cards sold under such circumstances would not qualify for the exclusion in proposed § 205.20(b)(2) because the issuer or program manager does not maintain policies and procedures reasonably designed to avoid the marketing of the general-purpose reloadable cards as gift cards or gift certificates.

The Board solicits comment on whether the proposed comment provides sufficient guidance regarding procedures that could enable an issuer, program manager, or other covered entity to comply with the rule with respect to an excluded product under proposed § 205.20(b)(2). In particular, comment is requested on practical issues that may arise in a retail environment, for example, in areas

where there may not be sufficient space for covered and non-covered products to be separately displayed, such as a checkout lane. Commenters are urged to provide specific examples of measures that may be utilized to ensure that a reasonable consumer would not believe that a card that would otherwise be excluded, such as a general-purpose reloadable card, is a gift card or gift certificate.

Some general-purpose reloadable cards that are not intended to be marketed as a gift card, but rather as an alternative to a bank account (or account substitute), such as for the unbanked, may be initially sold as a non-reloadable open-loop card. After the card is purchased, the cardholder may call the issuer to register the card. Once the issuer has obtained the cardholder's personal information, a new personalized, reloadable card may be sent to the cardholder.

The Board understands that under one model, the cardholder may use the temporary non-reloadable card to conduct transactions immediately after card purchase and up until the card is registered by the consumer and replaced with the personalized, reloadable card. Under another model, the temporary non-reloadable card may not be used by the consumer to make purchases until the consumer calls to register the card. Under the second model, the temporary card can be used after registration until the personalized, reloadable card arrives in the mail and is activated by the cardholder.

Under either model, the temporary card would not appear to qualify for the reloadable and not marketed as a gift card or gift certificate exclusion because it is non-reloadable. If the rule were to provide that such products were to fall within the exclusion notwithstanding the issuance of the initial non-reloadable card, then consumers that elect not to register the card (and therefore do not obtain a reloadable card) would not be given the statutory protections under the Credit Card Act. Conversely, if the rule were to provide that such products do not qualify for the exclusion at any point even if the card is ultimately replaced by a reloadable card, then the exclusion in EFTA Section 915(a)(2)(D)(ii) and proposed § 205.20(b)(2) would effectively be eliminated for most, if not all, general-purpose reloadable cards, given existing business models and other regulatory considerations.

Under a third approach, the restrictions on assessing dormancy, inactivity, or service fees, and on expiration dates could be applied solely to the initial non-reloadable card, but

¹⁰ In addition to these parties, a processor may work with the issuer and the program manager to process card transactions, and in some cases provide Web site and telephone customer service. For open-loop card programs, the payment network operates the network and establishes operating rules for card issuers, processors, and merchants or ATMs that accept the card.

not to the reloadable replacement card. While the third approach may provide certain flexibility for some issuers, the Board is concerned that consumers may be confused or surprised when they receive new terms regarding dormancy, inactivity, or service fees and expiration dates for the reloadable card that differ from the terms previously disclosed at the initial purchase. Given these considerations, the Board solicits comment on the appropriate treatment of these products.

20(b)(3) Loyalty, Award, or Promotional Gift Card

Proposed § 205.20(b)(3) implements the exclusion for cards, codes, or other devices for loyalty, award, or promotional gift cards. *See* EFTA Section 915(a)(2)(D)(iii). As discussed above, proposed § 205.20(a)(4) generally defines a “loyalty, award, or promotional gift card” as a card, code, or other device that is issued in connection with a loyalty, award, or promotional program.

In contrast to gift cards purchased at a store, loyalty, award, and promotional gift cards typically are not funded by direct payment from the consumer, but instead are funded by the entity sponsoring the card program, such as a merchant, an employer, or a company. Prepaid cards issued through such programs may serve as cost-effective substitutes for traditional means of distributing funds through a promotion, such as rebate checks, vouchers, or cash awards.

Much like rebate checks, vouchers, and cash awards, gift cards distributed through a loyalty, award, or promotional program are typically redeemable for a limited period of time. Loyalty, award, or promotional gift cards thus generally carry shorter expiration dates compared to gift cards purchased through retail channels.

From a consumer’s perspective, consumers who receive a gift card redeemable at one merchant as part of a loyalty, award, or promotional program may be surprised to find that the fees and expiration date on the card differ substantially from a card that they may have purchased directly from that same merchant. Improved disclosure of these terms for cards subject to the exclusion may help reduce consumer surprise or confusion.

Consistent with the statutory exclusion in EFTA Section 915(a)(2), the proposed rule does not impose substantive restrictions on dormancy, inactivity, or service fees, or on expiration dates, for cards, codes, or other devices issued pursuant to a loyalty, award, or promotional program.

Nonetheless, the Board believes that clear and conspicuous disclosures of the terms that apply to a loyalty, award, or promotional gift card are necessary to help consumers avoid surprise from unexpected dormancy, inactivity, or service fees or from short expiration dates.

Accordingly, the Board is proposing to exercise its authority under new EFTA Section 915(a)(2)(D)(iii) to define loyalty, award or promotional gift cards to require that consumers are given clear and conspicuous disclosures about any fees, including dormancy, inactivity, or service fees, or expiration dates, that may apply when they receive a gift card through a loyalty, award, or promotional program. This requirement would be implemented in proposed § 205.20(a)(4)(ii). Thus, in order to be deemed a “loyalty, award, or promotional gift card,” and therefore qualify for the exclusion in proposed § 205.20(b)(3), the card, code, or other device must set forth disclosures regarding any fees and expiration dates that may apply to the card, code, or device. While disclosures regarding dormancy, inactivity, or service fees, expiration dates, and a toll-free number and Web site for additional information must be on the card, code or other device, disclosures regarding other fees may accompany the card, code, or other device. *See also* proposed §§ 205.20(d)(2), (e)(2), and (f), discussed below. The proposed rule is intended to strike a balance between the competing considerations of enabling companies to manage the costs of providing consumers gift cards in connection with loyalty, award, or promotional programs, and limiting potential consumer confusion or surprise arising from the different terms that may apply to such cards.

20(b)(4) Not Marketed to the General Public

Proposed § 205.20(b)(4) implements the exclusion for cards, codes, or other devices that are not marketed to the general public. *See* EFTA Section 915(a)(2)(D)(iv). Whether a card is “marketed to the general public” depends on the facts and circumstances, but the term generally describes cards, codes, or other devices that are offered, advertised or otherwise promoted to the general public. *See* proposed comment 20(b)(4)-1. A card, code, or other device may be marketed to the general public regardless of the advertising medium, including television, radio, newspaper, the Internet, or signage.

In determining whether the exclusion applies to a particular card, code, or other device, proposed comment

20(b)(4)-1 provides that a number of factors must be considered, including the means or channel through which the card, code, or device may be obtained by a consumer, the subset of consumers that are eligible to obtain the card, code or device, and whether the availability of the card, code, or device is advertised or otherwise promoted in the marketplace. Thus, the Board does not view the method of distribution by itself as dispositive in determining whether a card, code, or other device is marketed to the general public.

Proposed comment 20(b)(4)-2 provides examples illustrating the exclusion. For instance, a merchant may sell its gift cards at a discount to a business, either directly or indirectly through a third party. The business that purchases the cards may give them to employees or loyal consumers as incentives or rewards. In determining whether the gift card is marketed to the general public, the merchant-issuer must consider whether the card is of a type that is advertised or made available to consumers generally or can be easily obtained elsewhere. If the card may also be purchased through retail channels, the exclusion in § 205.20(b)(4) does not apply, even if the consumer obtained the card as an incentive or reward. *See* proposed comment 20(b)(4)-2.i. In these cases, consumers could be confused when they receive gift cards that appear substantially similar to those that they could have purchased directly from a merchant, but contain different terms and conditions, such as a shorter expiration date. Of course, other exclusions under the proposed rule, such as the exclusion for cards issued in connection with a loyalty, award, or promotional program, may apply to such cards. *See* proposed § 205.20(b)(3).

Similarly, the Board has also considered whether cards issued or sold by a business pursuant to a marketing campaign that targets a specific subset of consumers would fall within the exclusion. The Board is concerned that a broad interpretation of the exclusion for cards not marketed to the general public would create a loophole and undermine the protections afforded to consumers under the rule. For example, a national retail chain could decide to market its gift cards only to members of its frequent buyers program. However, if any member of the general public may become a member of the program, the general public would still be able to obtain the cards. Thus, the Board believes such cards would be covered by the rule in those circumstances, unless another exclusion applies. *See* proposed comment 20(b)(4)-2.ii. Similarly, a reloadable card advertised

to teenagers to help them manage their everyday expenses and for emergencies, or marketed to parents to enable them to monitor spending would be a card marketed to the general public. *See* proposed comment 20(b)(4)-2.iii.

In contrast, where the availability of the card itself is not advertised or otherwise promoted, but rather, is merely used as the means through which funds are delivered to a consumer, the Board believes the card is not marketed to the general public. Proposed comment 20(b)(4)-2 includes four examples of cards that may fall within the exclusion depending on the circumstances: (a) A card containing insurance proceeds provided by an insurance company to a customer to settle a claim; (b) a card containing travel expenses or per diem funds provided by a business to an employee; (c) a card containing store credit provided by a retailer to a customer following a merchandise return if the card states that it is issued for store credit; and (d) a card containing tax refunds provided by a tax preparer to a customer. *See* proposed comments 20(b)(4)-2.iv-vii.

Whether a non-reloadable tax refund card is marketed to the general public will depend upon the facts and circumstances. For example, if a tax preparer merely provides the prepaid card as a mechanism for providing a tax refund to a consumer, and does not advertise or otherwise promote the ability to receive a tax refund through a prepaid card, the card would be excluded because it is not marketed to the general public. However, if the tax preparer engages in a marketing campaign that touts the ability of a consumer to receive a prepaid card for faster access to their tax refund proceeds, the tax refund card would not be exempt under this exclusion. *See* proposed comment 20(b)(4)-2.vii.

20(b)(5) Issued in Paper Form Only

Proposed § 205.20(b)(5) sets forth the exclusion for cards, codes, or other devices that are issued in paper form only. *See* EFTA Section 915(a)(2)(D)(v). As explained in proposed comment 20(b)(5)-1, the exclusion applies where the sole means of issuing the card, code, or other device is by paper. Examples of excluded paper gift certificates or cards include paper certificates distributed by restaurants or spas that are redeemable for a specific service or a specified dollar amount, and paper vouchers valid for tickets or events.

To prevent potential circumvention of the rule, the proposed commentary explains that the exclusion does not apply simply because a card, code, or

other device is reproduced or otherwise printed on paper. For example, a bar code or card or certificate number sent electronically to a consumer and redeemable for goods or services is not issued in paper form, even if it may be reproduced or otherwise printed on paper by the consumer.¹¹ Similarly, § 205.20(b)(5) would not apply where an on-line retailer electronically mails a certificate redeemable for goods or services to a consumer, which the consumer could print out on a home printer. In these circumstances, although the consumer might hold a paper facsimile of the card, code, or other device, the exclusion does not apply because the information necessary to redeem the value was initially issued in electronic form.

The proposal does not, however, preclude a paper certificate bearing a bar code or account number that is given to the consumer at the time of purchase from qualifying for the exclusion. For example, a retailer may generate a bar code on a paper certificate at the time of purchase that enables the retailer to scan the certificate and maintain a record of the certificate electronically, rather than enter the information in a ledger. Because the bar code is not issued to the consumer in any form other than on the paper given to the consumer, this certificate would qualify for the exclusion for cards, codes, or other devices issued in paper form.

Comment is requested regarding whether this aspect of the proposal creates an undue risk of circumvention. For example, a paper certificate or card that is encoded with a magnetic stripe might qualify for the exclusion. Other than the material on which the magnetic stripe is printed or produced, however, there is no meaningful distinction between a plastic card with a magnetic stripe and a paper certificate or card with a magnetic stripe encoded on the paper.

20(b)(6) Redeemable Solely for Admission to Events or Venues

Proposed § 205.20(b)(6) excludes cards, codes, or other devices that are redeemable solely for admission to events or venues at a particular location or group of affiliated locations, or to obtain goods or services, in conjunction with such admission, at the event or venue, or at specific locations affiliated with and in geographic proximity to the

event or venue. *See* EFTA Section 915(a)(2)(D)(vi).

Under the proposed rule, the exclusion in § 205.20(b)(6) is generally limited to cards, codes, or other devices that do not state a specific monetary value but instead are redeemable for an admission to an event or venue, such as a ticket to a sporting event or a pass to enter an amusement park. In addition, the exclusion applies to cards, codes, or other devices that entitle consumers to obtain goods or services, in conjunction with admission to an event or venue. *See* EFTA Section 915(a)(2)(D)(vi). For example, the consumer might purchase a certificate or card that entitles the recipient to one ticket to an amusement park plus a dollar amount that can be spent on concessions at the park.

Consistent with the statute, the proposed exclusion in § 205.20(b)(6) would also cover circumstances where the consumer may obtain goods or services at specific locations affiliated with and in geographic proximity to the event or venue in conjunction with admission. For example, a certificate or card may enable a consumer to gain admission to an amusement park and to receive a souvenir of the occasion at a retailer affiliated with the park and located within or nearby the park.

While the exclusion would apply to cards, codes, or other devices that are redeemable for admission to an event or venue, and for goods or services purchased in conjunction with that admission, the exclusion does not cover cards, codes, or other devices issued in a specified monetary value that could be applied toward such admission. For example, a merchant with an affiliated amusement park could issue a \$25 gift card to a consumer that can be redeemed by the recipient to purchase goods at any of the merchant's retail outlets and its on-line store. Under the terms of the prepaid card program, however, the merchant could also allow the card to be provided as a form of payment to purchase tickets at the amusement park.

The Board is concerned that permitting the exclusion to apply in these circumstances would create opportunities for circumvention because an issuer could simply list the purchase of tickets at the amusement park as one of several permitted uses of a gift card to avoid the consumer protections provided by the Credit Card Act. Accordingly, the proposed rule would not apply the exclusion to a card that can be redeemed in a specified amount towards admission to an event or venue. In this regard, the Board notes that the statute refers to cards, codes, or other devices that are redeemable *solely* for

¹¹ An issuer may, however, replace a gift certificate that was initially issued in paper form only with a plastic card or electronic code (for example, to replace a lost paper certificate) without falling outside the exclusion in § 205.20(b)(5).

admission to events or venues at a particular location or group of affiliated locations. See EFTA Section 915(a)(2)(D)(vi).

The proposed exclusion in § 205.20(b)(6) also would not apply to other payment devices that do not have a specified monetary value but are redeemable for a specified product or service, other than admission to an event or venue. For example, an issuer or retailer may sell a certificate or card that is redeemable for a spa treatment or for a hotel stay. In such circumstances, the certificate or card is not applied to obtain admission to the spa or hotel itself, but is used to pay for services at those locations. The exclusion does not apply to such cards because they are not redeemable solely for admission to an event or venue. See EFTA Section 915(a)(2)(D)(vi). Nonetheless, other exclusions in the rule may apply in these circumstances. See, e.g., proposed § 205.20(b)(3).

Proposed comment 20(b)(6)–1 provides examples to illustrate the exclusion in § 205.20(b)(6). In addition to the examples discussed above, the proposed comment also provides an example of cards that are redeemable solely for membership to a buyer's club or warehouse or to a gym. Such cards would fall within the exclusion in § 205.20(b)(6) because memberships are necessary for entry or admission to those locations. The exclusion would not apply if the card has value that could be applied either for a membership or for goods or services at the warehouse or gym. See comment 20(b)(6)–1.v.

20(c) Form of Disclosures

20(c)(1) Clear and Conspicuous

New EFTA Sections 915(b)(3)(A) and (c)(2)(B) (15 U.S.C. 1693m(b)(3)(A) and (c)(2)(B)), as added by Section 401 of the Credit Card Act, require that the disclosures made pursuant to those paragraphs be clear and conspicuous. The Board believes it is also appropriate to apply the clear and conspicuous standard to the disclosures the Board is proposing under § 205.20(f). Thus, pursuant to the Board's authority under new EFTA section 904, proposed § 205.20(c)(1) applies the clear and conspicuous standard to all disclosures required under § 205.20.

Proposed comment 20(c)(1)–1 clarifies the meaning of the term “clear and conspicuous” for the purposes of this section. Specifically, as the proposed comment explains, disclosures are clear and conspicuous for the purposes of this section if they are readily understandable and, in the case of

written and electronic disclosures, the location and type size are readily noticeable to consumers. Disclosures need not, however, be located on the front of the certificate or card to be considered clear and conspicuous. Disclosures are clear and conspicuous for the purposes of this section if they are in a print that contrasts with and is otherwise not obstructed by the background on which they are printed. For example, disclosures on a card or computer screen are not likely to be conspicuous if obscured by a logo printed in the background. Similarly, the proposed comment states that a disclosure on the back of a card that is printed on top of indentations from embossed type on the front of the card is not likely to be conspicuous if it obstructs the readability of the type. The proposed comment clarifies that oral disclosures, to the extent they are permitted, meet the clear and conspicuous standard when they are given at a volume and speed sufficient for a consumer to hear and comprehend them.

Though the proposal requires that the prescribed disclosures be clear and conspicuous, it does not include a specific type size or prominence requirement, except where otherwise noted. As discussed below in proposed § 205.20(e)(3)(iii), certain disclosures regarding funds expiration are required to be made with equal prominence and in close proximity to the certificate or card expiration date on a certificate or card. The Board included this requirement because of its specific concerns related to customer confusion with respect to a certificate or card expiration date that may differ from the expiration date for the underlying funds. However, the Board believes requiring every disclosure on a certificate or card to have an equal prominence or a minimum type size standard is impractical, because the size of certificates or cards will vary. Therefore, a general type size that is appropriate for one card may not fit on a smaller card, due to the limited amount of space. Moreover, such standards would present issues for disclosures even on standard-sized cards, because the amount of space on such cards is limited.

The Board requests comment on whether description of the clear and conspicuous standard in the final rule should include a type size or prominence requirement for all disclosures and, if so, what standard is appropriate. The Board also requests comment on whether there are alternatives to a type size or prominence requirement that could ensure that

disclosures on a card are clear and conspicuous to a consumer.

Proposed § 205.20(c)(1) states that the disclosures required by this section may contain commonly accepted or readily understandable abbreviations or symbols. Proposed comment 20(c)(1)–2 provides illustrative examples, stating that the use of abbreviations and symbols such as “mo.” for month or a “/” to indicate “per” is permissible. The proposed comment notes that it is sufficient under the clear and conspicuous standard to state, for example, that a particular fee is charged “\$2.50/mo. after 12 mos.”

20(c)(2) Format

Proposed § 205.20(c)(2) states that disclosures required by this section generally must be provided to the consumer in written or electronic form. Because the disclosures are not required to be in written form, proposed comment 20(c)(2)–1 clarifies that electronic disclosures made under this section are not subject to compliance with the consumer consent and other applicable provisions of the Electronic Signatures in Global and National Commerce Act (E-Sign Act) (15 U.S.C. 7001 *et seq.*), which only applies when information is required to be provided to a consumer in writing. The comment clarifies that electronic disclosures may not be provided through a hyperlink or in another manner by which the purchaser can bypass the disclosure. An issuer or vendor is not required to confirm that the consumer has read the electronic disclosures.

Proposed comment 20(c)(2)–2 addresses disclosure requirements in circumstances where no physical certificate or card is issued. Under the proposed comment, disclosures would be required to accompany the code, confirmation, or other written or electronic document provided to the consumer.

Proposed § 205.20(c)(2) states that only disclosures provided under § 205.20(c)(3) may be provided orally. Allowing oral disclosures is necessary because, in some circumstances, disclosures cannot be made prior to purchase unless made orally, such as when a certificate or card is purchased by telephone. Even where oral disclosures are permitted, written or electronic disclosures must still be provided on or with the certificate or card. See proposed §§ 205.20(d)(2), (e)(3), and (f).

20(c)(3) Disclosures Prior to Purchase

New EFTA Section 915(b)(3)(B) (15 U.S.C. 1693m(b)(3)(B)), requires that dormancy, inactivity, or service fees be

disclosed before a gift certificate, or store gift card, or general-use prepaid card is purchased. In addition, the Board proposes to use its authority under EFTA Section 904 to require the disclosure of additional fees under § 205.20(f)(1), discussed below, and the terms and conditions of expiration of the funds prior to purchase of the certificate or card. *See* proposed §§ 205.20(e)(3) and (f)(1), discussed below. These requirements are implemented in proposed § 205.20(c)(3).

The Board believes that consumers contemplating the purchase of a certificate or card need information about all fees and the terms and conditions of expiration before purchasing a certificate or card. Even if the purchaser is not the ultimate user of the certificate or card, the Board believes that a purchaser should be aware of any potential costs to the recipient and the amount of time the recipient has to use the funds underlying the certificate or card. Making this type of information available to purchasers may also foster competition.

Proposed comment 20(c)(3)–1 clarifies that the disclosures required under this paragraph must be provided regardless of whether the certificate or card is purchased in person, on-line, by telephone, or by other means.

20(c)(4) Disclosures on the Certificate or Card

Proposed § 205.20(c)(4) addresses the requirements in § 205.20 that certain disclosures be provided on the certificate or card itself. *See* proposed §§ 205.20(d)(2), 205.20(e)(3), and 205.20(f)(2). The paragraph states that a disclosure made in an accompanying terms and conditions document, on packaging, or on a sticker or other label affixed to the certificate or card does not constitute a disclosure on the certificate or card.

The Board believes this interpretation is consistent with new EFTA Section 915(b)(3)(A), which requires that a gift certificate, store gift card, or general-use prepaid card clearly and conspicuously state any dormancy, inactivity, or service fee and the conditions under which they can be imposed. Requiring the fees and conditions to be disclosed on the certificate or card ensures that the consumer and, if applicable, the gift recipient will always have access to the disclosures, because they cannot be separated from the certificate or card. Moreover, a number of State laws already require certain fee and expiration date disclosures on

certificates or cards.¹² Pursuant to its authority under new EFTA Section 915(d)(1)(A), and as discussed below in §§ 205.20(e)(3) and (f)(2), the Board is proposing to extend the requirement that certain disclosures be on the certificate or card itself to certain additional disclosures. Specifically, the proposal states that the certificate or card itself must state the terms and conditions of expiration of the funds; a toll-free telephone number a consumer may call for fee information or replacement certificates or cards; and, if one is maintained, a Web site a consumer may access for fee information or replacement certificates or cards.

The Board recognizes that the proposed requirements regarding disclosures that must appear on a covered certificate or card may present implementation challenges with respect to certain products, particularly those that are small and have little space on which to print required disclosures. The Board seeks comment regarding any approaches or solutions that could avoid potential impediments to innovation while still providing consumers clear and conspicuous disclosures. The Board also seeks comment regarding how issuers currently provide disclosures and how issuers comply with State laws which have similar disclosure requirements to those set forth in the proposed rules.

20(d) Prohibition on Imposition of Fees or Charges

New EFTA Sections 915(b)(1) and (2) generally prohibit the imposition of a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card or general-use prepaid card unless: (a) There has been no activity for the 12-month period ending on the day the charge is imposed; (b) certain disclosure requirements have been met; (c) only one such fee is charged in any given month; and (d) the certificate or card complies with any additional requirements the Board may establish. *See* 15 U.S.C. 1693m(b)(1) and (2). Regarding the disclosure requirements noted above, new EFTA Section 915(b)(3) provides that before a dormancy, inactivity, or service fee may be imposed, a certificate or card must clearly and conspicuously disclose: (a) That a dormancy, inactivity, or service fee may be charged; (b) the amount of the fee; (c) how often such fee or charge

may be assessed; and (d) that such fee or charge may be assessed for inactivity. *See* 15 U.S.C. 1693m(b)(3). Moreover, the issuer or vendor of such certificate or card must inform the purchaser of such charge or fee before such certificate or card is purchased, regardless of whether the certificate or card is purchased in person, over the Internet, or by telephone. *See* 15 U.S.C. 1693m(b)(3)(B).

Proposed § 205.20(d) generally implements new EFTA Sections 915(b)(1), (2), and (3) while proposed § 205.20(c)(3), discussed above, implements new EFTA Section 915(b)(3)(B).¹³ The Board notes that although “dormancy or inactivity fee” is defined separately from “service fee,” for improved readability, proposed § 205.20(d) and associated commentary refer to these fees collectively as “dormancy, inactivity, or service fees.” As discussed above, proposed § 205.20(c)(3) also requires the issuer or vendor to inform the purchaser about certain other terms prior to purchase.

The Board is proposing several comments to clarify the provisions in § 205.20(d). Proposed comment 20(d)–1 illustrates with examples how to determine when a dormancy, inactivity, or service fee may be imposed. Proposed comment 20(d)–2 clarifies the meaning of “activity” for purposes of proposed § 205.20(d)(1). Specifically, any action by the consumer to increase, decrease or otherwise make use of the funds underlying a certificate or card constitutes activity. For example, the purchase and activation of a card or the reloading of funds onto a card constitutes activity for purposes of § 205.20(d)(1). However, activity with respect to a certificate or card would not include the imposition of a fee, the replacement of an expired, lost, or stolen certificate or card, or a balance inquiry. The Board solicits comment on whether there are any other actions taken by a consumer that should be considered “activity” for purposes of proposed § 205.20(d)(1).

Proposed § 205.20(d)(2) and (c)(3) require similar, but not identical, disclosures. Proposed comment 20(d)–3 clarifies the interaction between these provisions. Specifically, the proposed

¹³ The proposed rule does not separately implement the exclusion in new EFTA Section 915(b)(4) from the dormancy, inactivity, or service fee restrictions for gift certificates distributed pursuant to an award, loyalty, or promotional program and with respect to which there is no money or other value exchanged. The Board believes this exclusion is already effectively implemented through the definition of “gift certificate” in proposed § 205.20(a)(1)(iii) and the exclusion in proposed § 205.20(b)(3) for loyalty, award, or promotional gift cards.

¹² *See, e.g.*, Ark. Code § 4–88–703 and Neb. Rev. Stat. §§ 69–1305.03(e) and (f) (requiring expiration date and certain fees to be disclosed on the gift certificate or card), and Or. Rev. Stat. § 646A.278 (requiring expiration date to be disclosed on the gift card).

comment provides that depending on the context, a single disclosure regarding dormancy, inactivity, or service fees imposed that meets the clear and conspicuous requirement may satisfy both the requirement in § 205.20(d)(2) that the disclosures be provided on the certificate or card and the requirement in § 205.20(c)(3) that the disclosures be provided prior to purchase. For example, if the disclosures on a certificate or card, required by § 205.20(d)(2), are visible to the consumer without having to remove packaging or other materials sold with the certificate or card for a purchase made in person, the disclosures also meet the requirements of § 205.20(c)(3). If, however, the disclosure does not meet the requirements of both §§ 205.20(d)(2) and (c)(3), proposed comment 20(d)–3 states that a dormancy, inactivity, or service fee may need to be disclosed multiple times or in multiple locations to satisfy the requirements of §§ 205.20(d)(2) and (c)(3). For example, if the disclosures on a certificate or card, required by § 205.20(d)(2), are obstructed by packaging or other materials sold with the certificate or card for a purchase made in person, they also must be disclosed on the packaging sold with the certificate or card or in other manner visible to the consumer to meet the requirements of § 205.20(c)(3).

Proposed §§ 205.20(d)(2), (e)(3), and (f)(2) require certain disclosures to be made on the certificate or card itself, as applicable. Proposed comment 20(d)–4 clarifies that in addition to disclosures required under § 205.20(d)(2), any applicable disclosures under §§ 205.20(e)(3) and (f)(2) of this section must also be provided on the certificate or card.

Finally, proposed comment 20(d)–5 clarifies the prohibition in § 205.20(d)(3) against charging more than one dormancy, inactivity, or service fee in any given calendar month. Specifically, proposed comment 20(d)–5 provides that if a dormancy, inactivity, or service fee is already imposed in a given calendar month, a second dormancy, inactivity, or service fee may not be imposed that month. If more than one dormancy, inactivity, or service fee is possible on a given day, the person assessing the fee may choose which dormancy, inactivity, or service fee to impose. The proposed comment also clarifies that the restriction in proposed § 205.20(d)(3) applies only to dormancy, inactivity, or service fees. As a result, a fee that is not a dormancy, inactivity, or service fee may be imposed in addition to a dormancy, inactivity, or service fee in a given month. Proposed comment

20(d)–5 would also provide examples with specific dates to illustrate these concepts.

20(e) Prohibition on Sale of Gift Certificates or Cards With Expiration Dates

New EFTA Section 915(c) prohibits the sale of a gift certificate, store gift card, or general-use prepaid card subject to an expiration date unless: (a) The expiration date is not earlier than five years after the date on which a gift certificate was issued, or the date on which card funds were last loaded to a store gift card or general-use prepaid card; and (b) the terms of expiration are clearly and conspicuously stated. *See* 15 U.S.C. 1693m(c). Proposed § 205.20(e) implements new EFTA Section 915(c).

Application of EFTA Section 915(c) to Certificate or Card Expiration and Funds Expiration

New EFTA Section 915(c) does not specify whether the restrictions apply to the expiration of the certificate or card itself or the underlying funds. It is the Board's understanding that for many general-use prepaid cards, and perhaps some gift certificates and store gift cards, the expiration date for the certificate or card differs from the expiration date for the underlying funds. For example, the underlying funds of some network-branded cards, which are required to have card expiration dates under card network rules and systems, never expire.

In order to ensure that consumers receive the full protection established by the statute with respect to the value of the certificate or card, proposed § 205.20(e)(2) would require that funds be available for the later of: (a) Five years from the date the gift certificate was issued, or the date on which funds were last loaded to a store gift card or general-use prepaid card; or (b) until the certificate or card expiration date.

In addition, to prevent consumer confusion, the proposed rule addresses the potential mismatch and resulting disconnect between a stated expiration or valid through date of the certificate or card and the date the funds expire. Specifically, consumers may assume that once the certificate or card expiration date has passed, the underlying funds are no longer valid or available. Presumably, a certificate or card expiration date that matches the funds expiration date would not cause confusion among consumers. However, a certificate or card expiration date that is identical to the funds expiration date may not be feasible. First, at the time a certificate or card expiration date is printed on a certificate or card, it may

be impossible to predict the funds expiration date, which would, under the Board's proposed rule, depend on when a consumer purchases the certificate or card or adds funds to a reloadable card. For example, if the certificate or card expiration date is printed during the certificate or card manufacturing process, this process may occur several months prior to the date the consumer purchases the certificate or card and activates it for use. Second, because the expiration date required under new EFTA Section 915(c) for store gift cards and general-use prepaid cards must be calculated from the date the funds were last loaded, this would mean, in practice, that funds underlying a reloadable card might never expire.

The Board considered prohibiting the use of expiration or valid through dates for gift certificates, store gift cards, and general-use prepaid cards. However, the Board understands that certain network systems may not be able to support products that do not carry expiration or valid through dates because of fraud and security concerns. In addition, card expiration dates may be necessary for other business reasons, such as to ensure that a card can remain usable for its lifespan. Moreover, merchants have become accustomed to looking for, or, in the case of telephone or on-line purchases, requesting, certificate or card expiration dates. Mandating certificates or cards without expiration or valid through dates could create significant confusion among merchants, which in turn, could result in problems for consumers' use of gift certificates, store gift cards, and general-use prepaid cards at such merchants. Therefore, to harmonize, to the extent feasible, the certificate or card expiration date and the funds expiration date, the Board is proposing two alternative approaches for applying new EFTA Section 915(c) to the expiration of a certificate or card in § 205.20(e)(1).

Under Alternative A of proposed § 205.20(e)(1), the Board is proposing that a person may not sell a gift certificate, store gift card, or general-use prepaid card subject to an expiration date unless the certificate or card expiration date is at least five years after the date the certificate or card is sold or issued to a consumer. The Board understands that there are some issuers and retailers of prepaid cards with systems and procedures currently in place to prevent the sale or issuance of a certificate or card unless there is a minimum amount of time left before the certificate or card expiration date; for example, 12 to 18 months from the date of sale or issuance. These issuers and retailers may currently employ

inventory controls or point-of-sale procedures to prevent sales of certificates or cards that do not meet the minimum time. For these issuers and retailers, compliance with Alternative A would likely only involve altering their systems and procedures to accommodate the five-year time period instead of the current minimum time frame. However, the Board is concerned that it may not be operationally feasible for all issuers and retailers of gift certificates, store gift cards, and general-use prepaid cards to institute these types of systems and procedures by the mandatory compliance date of the final rule.

Alternative B of proposed § 205.20(e)(1) would instead require entities involved in issuing, distributing, and selling certificates or cards to adopt policies and procedures to ensure that a consumer will have a reasonable opportunity to purchase a certificate or card with at least five years remaining until the certificate or card expiration date. Proposed comment 20(e)-1 under Alternative B would set forth positive and negative examples of providing consumers a reasonable opportunity to purchase a certificate or card with at least five years remaining until the certificate or card expiration date. For example, a person subject to this rule would comply with Alternative B of proposed § 205.20(e)(1) if a card is printed with an expiration date that is six years from the date the card was produced and on a display rack at a retail store within six months of the date the card was produced. Similarly, a person would comply with Alternative B of proposed § 205.20(e)(1) if a card is printed with an expiration date that is seven years from the date the card was produced and on a display rack at a retail store within one year and six months of the date the card was produced. However, a person would not comply with Alternative B of proposed § 205.20(e)(1) if a card is printed with a card expiration date six years from the date it was produced and is stored in a distribution warehouse for more than one year before being made available for sale.

Unlike Alternative A of § 205.20(e)(1), Alternative B would not require a person to confirm that a certificate or card is in fact sold or issued to a consumer with at least five years before the certificate or card expiration date. As a result, the expiration date reflected on the certificate or card may, in some cases, be less than five years from the date of sale or issuance. While the consumer would still have use of the underlying funds for a minimum of five years from the date of sale or issuance,

as would be required under proposed § 205.20(e)(2), the Board is concerned that a certificate or card reflecting a certificate or card expiration date earlier than the funds expiration date could prompt consumers to dispose of the certificate or card before the funds expiration date.

The Board believes that Alternative A would provide the greatest precision in matching the certificate or card expiration date with the funds expiration date, though Alternative B may be easier to implement than Alternative A. Given that persons subject to the rule may be able to comply with Alternative B more rapidly than Alternative A, the Board also solicits comment on whether it should consider adopting Alternative B for a transitional period and adopt Alternative A as of a subsequent date in order to provide more time to implement Alternative A.

While either Alternative A or Alternative B may adequately address potential consumer confusion regarding expiration dates with respect to non-reloadable cards, such protections may not be sufficient for reloadable cards where the funds expiration date changes each time the card is reloaded. The Board is proposing to address this issue by requiring certain disclosures related to the expiration of the underlying funds. As discussed more fully below in the supplementary information to proposed § 205.20(e)(3), the Board is proposing that the terms and conditions of expiration of the underlying funds be disclosed on the certificate or card, including, where applicable, a statement that the certificate or card expires, but the underlying funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card. *See* proposed § 205.20(e)(3)(iii).

The Board also solicits comment on whether an additional or alternative substantive solution to the proposed notice in § 205.20(e)(3) may be warranted. Specifically, the Board is requesting comment on whether it should also or alternatively require issuers to automatically issue a replacement card to consumers prior to the card expiration date of a reloadable card if the underlying funds will not expire until after the card expiration date. The Board understands that for some reloadable cards, issuers currently collect certain information from the consumer, including name and address, before the consumer may be permitted to reload funds to the card, or in some cases, use the card at all. Thus, these issuers would have the information necessary to send replacement cards

before the card expiration date, much as issuers currently do for credit cards, which would avoid consumer confusion as to whether the underlying funds may still be available. The Board is concerned, however, that not all issuers of reloadable cards may have the systems in place to collect name and address information and that establishing such systems could be prohibitively expensive for these issuers. Furthermore, if a consumer does not notify the gift card issuer of changes in address, the issuer may not have a reliable current address to which it could send a replacement card. The Board seeks comment on operational considerations and the feasibility of implementing this requirement for reloadable cards.

Disclosures Related to Certificate or Card Expiration and Funds Expiration

New EFTA Section 915(c)(2)(B), which the Board proposes to implement in § 205.20(e)(3), requires that the terms and conditions of expiration be clearly and conspicuously stated. *See* 15 U.S.C. 1693m(c)(2)(B).

Under proposed § 205.20(e)(3), three disclosures must be stated on the certificate or card, as applicable. First, proposed § 205.20(e)(3)(i) provides that the disclosures must state the expiration date for the underlying funds or, if the underlying funds do not expire, that fact. In some instances, the exact expiration date of the underlying funds may not be able to be determined. For example, in the case of reloadable cards, the funds expiration date is determined under the statute and the Board's proposed rule by the date the consumer last loaded funds onto the card. As a result, the funds expiration date adjusts each time the consumer reloads the card. For example, if a consumer purchases a reloadable card on January 15, 2010, the funds may expire on or after January 15, 2015. However, if a consumer loads more funds onto the card on July 15, 2012, the funds may not expire until on or after July 15, 2017. To accommodate this circumstance, proposed comment 20(e)-1 under Alternative A (comment 20(e)-2 under Alternative B) clarifies that § 205.20(e) does not require disclosure of the precise date the funds will expire. It would be sufficient to disclose, for example, "Funds expire 5 years from the date funds last loaded to the card."; "Funds can be used 5 years from the date money was last added to the card."; or "Funds do not expire." The Board requests comment on whether these sample disclosures would effectively communicate how long a consumer has

access to funds underlying a certificate or card.

Proposed comment 20(e)-2 under Alternative A (comment 20(e)-3 under Alternative B) clarifies that if the certificate or card and the underlying funds do not expire, that fact need not be disclosed. The Board believes that disclosing the fact that the underlying funds do not expire is not necessary in these situations because there is no risk of consumers confusing the expiration date of the certificate or card with that of the underlying funds.

Second, proposed § 205.20(e)(3)(ii) provides that the disclosures must also include a toll-free telephone number and, if one is maintained, a Web site that a consumer may use to obtain a replacement certificate or card after the certificate or card expires, if the underlying funds may be available. Requiring a toll-free telephone number to be maintained for purposes of obtaining a replacement card is appropriate because, as discussed above, a certificate or card expiration date may be earlier than the funds expiration date.¹⁴ While the proposed rule does not similarly require that a Web site be maintained for such purposes, if one is maintained, that Web site must also be disclosed under § 205.20(e)(3)(ii). By requiring contact information to be on the certificate or card itself, the Board believes that consumers will more easily be able to obtain a replacement certificate or card should the certificate or card expire before the underlying funds.

Proposed comment 20(e)-3 under Alternative A (comment 20(e)-4 under Alternative B) clarifies that if a certificate or card does not expire, or if the underlying funds are not available after the certificate or card expires, the disclosure required by proposed § 205.20(e)(3)(ii) need not be stated on the certificate or card. A toll-free telephone number and a Web site may still be required to be disclosed, however, pursuant to proposed § 205.20(f)(2) if the certificate or card has fees. Proposed comment 20(e)-4 under Alternative A (comment 20(e)-5 under Alternative B) clarifies that the same toll-free telephone number and Web site may be used to comply with the requirements of §§ 205.20(e)(3)(ii) and (f)(2).¹⁵ In addition, the proposed

comment provides that neither a toll-free number nor a Web site must be maintained or disclosed on a certificate or card if no fees are imposed in connection with the certificate or card, and the certificate or card and underlying funds do not expire.

Finally, proposed § 205.20(e)(3)(iii) would require, if applicable, a statement that the certificate or card expires, but the underlying funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card. This requirement is designed to ensure that consumers are alerted to any distinction between the certificate or card expiration date and the funds expiration date so that they do not mistakenly believe the funds are no longer available during the minimum five-year period set forth in the statute.

Proposed § 205.20(e)(3)(iii) also requires the statement to be disclosed with equal prominence and in close proximity to the certificate or card expiration date. While other required disclosures in this section are not subject to similar prominence and proximity requirements, the Board believes that such requirements are appropriate for the disclosures required under proposed § 205.20(e)(3)(iii). Typically, the expiration date for a certificate or card may be printed on the certificate or card in a prominent location and type size, which enables the merchant to easily verify the validity of the card at point-of-sale and the consumer to find this date when making telephone or on-line purchases. Thus, the Board is concerned that the prominence of the expiration date on the certificate or card (without any additional protections) may lead consumers to assume that once the certificate or card itself expires, the underlying funds will be unavailable. The disclosures proposed under § 205.20(e)(3)(iii) regarding expiration are intended not only to inform consumers of their rights, but also to reduce potential consumer confusion that may occur if an expiration date for a certificate or card differs from the funds expiration date. Therefore, the Board believes disclosures regarding the expiration of the funds require more specific format requirements than other disclosures that are required to be on the certificate or card.

As clarified in proposed comment 20(e)-5 under Alternative A (comment 20(e)-6 under Alternative B), close proximity, in the context of a certificate or card, means that the disclosure must appear on the same side as the certificate or card expiration date so that consumers do not automatically assume

funds are not available after the certificate or card expiration date. For example, many card expiration dates are stated on the front of a card. If the disclosure alerting the consumer to the fact that this expiration date does not apply to the underlying funds is printed on the back of the certificate or card, the consumer may not notice the disclosure if he or she does not have reason to look for an additional disclosure. However, if the disclosure is on the front of the card in close proximity to the card expiration date, the consumer may be more likely to notice it and seek additional information regarding how the consumer could continue to use the card after the card expiration date.

Proposed comment 20(e)-5 under Alternative A (comment 20(e)-6 under Alternative B) also clarifies that if the disclosure is the same type size and is located immediately next to or directly above or below the certificate or card expiration date, without any intervening text or graphical displays, the disclosures would be deemed to be equally prominent and in close proximity. The disclosure need not be embossed on the certificate or card to be deemed equally prominent, even if the expiration date is embossed on the certificate or card. The Board believes these format standards would sufficiently ensure that most consumers can determine whether an expiration date for a certificate or card is different from the funds expiration date.

Proposed comment 20(e)-5 under Alternative A (comment 20(e)-6 under Alternative B) provides examples regarding how a disclosure may inform a consumer of the distinction between the certificate or card expiration and the funds expiration. The disclosure may state on the front of the card, for example, "Valid thru 09/2016. Call for new card."; "Active thru 09/2016. Call for replacement card."; or "Call for new card after 09/2016." The Board believes these disclosures, used in conjunction with other disclosures required to be on the card, such as a toll-free number that a consumer could call for a replacement card, would provide sufficient information to inform consumers that they may be able to continue using their funds after the certificate or card itself has expired.

The Board recognizes that the amount of space available for disclosures near the certificate or card expiration date is limited. The Board requests comment regarding the feasibility of disclosing the sample disclosures or similar statements "in close proximity" to the certificate or card expiration date. The Board also requests comment on whether the "equal prominence"

¹⁴ As discussed below under proposed § 205.20(f), the requirement that the telephone number be toll-free recognizes that the end user of a certificate or card may not reside in the area where the certificate or card was initially purchased.

¹⁵ The contact information may also be the same contact information provided for any or all customer service issues or questions relating to the certificate or card.

standard is appropriate in the context of certificates or cards, or if the Board should prescribe a minimum type-size requirement and, if so, what type size is appropriate. Finally, the Board requests comment on other effective methods of notifying consumers that underlying funds may continue to be available after a certificate or card itself expires.

Finally, the Board notes that proposed §§ 205.20(d)(2), (e)(3), and (f)(2) (as discussed below) require certain disclosures to be made on the certificate or card itself, as applicable. Proposed comment 20(e)–6 under Alternative A (comment 20(e)–7 under Alternative B) thus clarifies that in addition to any disclosures required under § 205.20(e)(3), any applicable disclosures under §§ 205.20(d)(2) and (f)(2) of this section must also be provided on the certificate or card.

Other Protections and Clarifications

To ensure that consumers have full use of the funds loaded on a certificate or card for the minimum five-year period set forth in the statute, the Board proposes to use its authority under EFTA Section 904(c) to restrict the imposition of fees to replace an expired certificate or card if the funds loaded on the certificate or card have not expired. See 15 U.S.C. 1693b(c). Proposed § 205.20(e)(4) under both alternatives thus ensures that consumers retain a cost-free means to access funds if a certificate or card expires before the funds have expired. Proposed § 205.20(e)(4) contains an exception, however, for certificates or cards that have been lost or stolen. As a result, a fee to replace a certificate or card before the expiration date of the funds may be imposed for a lost or stolen certificate or card, to the extent otherwise permitted under law. Proposed comment 20(e)–7 under Alternative A (comment 20(e)–8 under Alternative B) clarifies that although a fee is permitted to be charged to replace a lost or stolen certificate or card under proposed § 205.20(e)(4), the rule does not create a substantive requirement that issuers replace a lost or stolen certificate or card.

Proposed comment 20(e)–8 under Alternative A (comment 20(e)–9 under Alternative B), clarifies that a certificate or card is not considered to be issued or loaded with funds until it has been activated for use. The Board understands that gift card issuers often produce gift cards for display on retail shelves and racks or for mailing to consumers. However, for security reasons, these cards cannot be used until the card has been activated by a retail employee or by telephone. The proposed comment clarifies that

although a certificate or card may have been produced, it is not considered to be “issued” or to have had funds “loaded” for purposes of § 205.20(e) until that card has been activated for use.

20(f) Additional Disclosure Requirements for Gift Certificates or Cards

EFTA Section 905(a)(4) (15 U.S.C. 1693c(a)(4)) and § 205.7(b)(5) of Regulation E require the disclosure of any fees imposed by a financial institution for electronic fund transfers or for the right to make such transfers. Pursuant to its authority under new EFTA Section 915(d)(2) (15 U.S.C. 1693m(d)(2)) to determine the extent to which the individual provisions of the EFTA and Regulation E should apply to gift certificates, store gift cards, and general-use prepaid cards, the Board is proposing § 205.20(f) to require additional fee-related disclosures for such certificates and cards.

20(f)(1) Fee Disclosures

The Board believes it is important for consumers to be aware of the fees that may be imposed before they use a certificate or card. As a result, proposed § 205.20(f)(1) would require that, for each type of fee that may be imposed in connection with a gift certificate, store gift card, or general-use prepaid card, certain information concerning fees must be disclosed on or with the certificate or card. Specifically, the type of fee, the amount of the fee (or an explanation of how the fee will be determined), and the conditions under which the fee may be imposed must be disclosed. The provision excludes dormancy, inactivity, and service fees, which must be disclosed under proposed § 205.20(d)(2). Therefore, fees other than dormancy, inactivity, or service fees, such as one-time initial issuance fees and cash-out fees, must be disclosed under proposed § 205.20(f)(1). Furthermore, in light of the other disclosures that must be provided on the certificate or card itself and because the size of a certificate or card may limit the disclosures that may be clearly and conspicuously disclosed on the certificate or card, the proposal permits this additional information to be disclosed either on or with the certificate or card. In addition, similar to the disclosure requirements for dormancy, inactivity, and service fees, the Board proposes to require the disclosure of these fees prior to purchase, as discussed above in the supplementary information to § 205.20(c)(3).

20(f)(2) Telephone Number for Fee Information

The Board also proposes to use its authority under new EFTA Sections 915(c)(2)(B) and 915(d)(1)(A), and EFTA Section 904 to require that a toll-free telephone number and, if one is maintained, a Web site, for information on fees be disclosed clearly and conspicuously on a gift certificate, store gift card, or general-use prepaid card. See 15 U.S.C. 1693m(c)(2)(B); 15 U.S.C. 1693m(d)(1)(A); 15 U.S.C. 1693b. Under proposed § 205.20(f)(2), a toll-free telephone number must be maintained to provide information on fees required to be disclosed under proposed §§ 205.20(d)(2) and (f)(1). The proposed rule does not similarly require that a Web site be maintained for such purposes, but if one is maintained, that Web site must also be disclosed under § 205.20(f)(2).

As discussed above, given the limited space on a certificate or card, the Board anticipates that issuers may opt to disclose some fee information on materials accompanying the certificate or card, as opposed to on the certificate or card itself. If such information accompanies the certificate or card, the disclosure may become separated from the actual certificate or card. By requiring the reference to the toll-free telephone number and, if one is maintained, the Web site on the certificate or card, the proposal seeks to ensure that consumers have an easy and cost-free means of obtaining fee information related to the certificate or card, even if the consumer no longer has the original disclosure.

Furthermore, the Board believes requiring the telephone number to be toll-free is appropriate. Because gift certificates, store gift cards, and general-use prepaid cards may be given by the purchaser to another person, the end user of the certificate or card may not reside in the area where the certificate or card was initially purchased. In addition, the majority of certificates or cards sold in the United States are issued by large retailers or large banks whose customer service centers are not necessarily located in the area where the certificate or card was purchased or will be used. A toll-free telephone number would provide consumers with a means to access fee and replacement certificate or card information without cost no matter where in the United States the user of the certificate or card may utilize the certificate or card.

Moreover, the Board understands that many issuers already maintain toll-free telephone numbers and Web sites for consumers to contact for further

information. Issuers maintaining toll-free telephone numbers or Web sites often provide this information directly on the certificates or cards they issue. As a result, the Board believes the proposed rule would not impose additional burden on many issuers.

The proposal contains several comments to clarify proposed § 205.20(f). Proposed comment 20(f)–1 clarifies that if a certificate or card does not have any fees, the disclosure required by § 205.20(f)(2) need not be disclosed on the certificate or card. A telephone number and a Web site may still be required to be disclosed pursuant to § 205.20(e)(3)(ii) if funds underlying a certificate or card may be available after the certificate or card expires.

Proposed comment 20(f)–2 clarifies that the same toll-free number and Web site may be used to fulfill the requirements of §§ 205.20(e)(3)(ii) and (f)(2).¹⁶ Neither a toll-free number nor a Web site must be maintained or disclosed if no fees are imposed in connection with a certificate or card, and the certificate or card and underlying funds do not expire.

Proposed §§ 205.20(d)(2), (e)(3), and (f)(2) require certain disclosures to be made on the certificate or card itself, as applicable. Proposed comment 20(f)–3 thus clarifies that in addition to any disclosures required to be made pursuant to § 205.20(f)(2), any applicable disclosures under §§ 205.20(d)(2) and (e)(3) of this section must be disclosed on the certificate or card.

Additional Issues

Authority To Adopt Additional EFTA Protections

New EFTA Section 915(d)(2) directs the Board to determine the extent to which the individual definitions and provisions of the EFTA or Regulation E should apply to general-use prepaid cards, gift certificates, and store gift cards. See 15 U.S.C. 1693m(d)(2). As discussed in proposed § 205.20(f), the Board is proposing to exercise this authority to mandate for each type of fee that may be imposed (such as a transaction fee, a balance inquiry fee, or an issuance fee), disclosure of the type of fee, the amount of the fee, and the conditions under which such fee may be imposed. These disclosures must be provided on or with a gift certificate, store gift card, or general-use prepaid card subject to the rule. This

¹⁶ The contact information may also be the same contact information provided for any or all customer service issues or questions relating to the certificate or card.

requirement is consistent with the requirement in EFTA Section 905(a)(4) (15 U.S.C. 1693c(a)(4)) and Regulation E § 205.7(b)(5) to disclose any charges for EFTs or for the right to make transfers.

The Board is not proposing at this time to apply to gift certificates, store gift cards, or general-use prepaid cards, any other requirements that generally apply to accounts under the EFTA and Regulation E, such as periodic statement disclosures or error resolution obligations. See, e.g., EFTA Sections 906(c) and 908; 15 U.S.C. 1693d(c) and 1693f. The Board believes that it is more appropriate to make any such determination in the context of a broader rulemaking that covers prepaid cards generally to avoid any regulatory gaps or inconsistencies. For example, a requirement to impose some form of periodic statement or error resolution obligations for reloadable gift cards could lead to inconsistent treatment if similar requirements were not simultaneously adopted for general-purpose reloadable cards, which in many cases are marketed as substitutes for accounts subject to the EFTA and Regulation E.

At this time, the Board is also not proposing to exercise the authority under new EFTA Section 915(d)(1) to limit the amount of dormancy, inactivity, or service fees, or the balance below which such fees or charges may be assessed. See 15 U.S.C. 1693m(d)(1). The Board understands that dormancy and inactivity fees in connection with retail gift cards have trended downward over time. For example, the most recent survey by one government agency indicates the median inactivity fee has decreased from \$1.73 per month to \$1.38 per month from 2003 to 2007.¹⁷ Given this trend, there does not appear to be a need for the Board to adopt additional restrictions at this time. Moreover, the statute only permits one such fee per month if there has been no activity over the preceding 12-month period. The Board will continue to monitor the development of the gift card

¹⁷ See Montgomery County Office of Consumer Protection, Gift Card Reports, 2003–2007 (available at: <http://www.montgomerycountymd.gov/ocptmpl.asp?url=/content/ocp/consumer/azgiftcardreports.asp>). One major issuer of a network-branded gift cards has recently announced plans to eliminate monthly fees altogether. See Andrew Martin, “American Express to End Monthly Fees on Gift Cards,” *New York Times*, October 1, 2009, at B2. In addition, the Retail Gift Card Association which is comprised of nine of the top retail merchant issuers of retail closed-loop gift cards includes in its Code of Principles, the elimination of dormancy or inactivity fees and of expiration dates. See Retail Gift Card Association, Code of Principles (available at: http://www.thergca.org/uploads/Code_of_Principles_PDF.pdf).

market and could take action to address dormancy, inactivity, or service fees at a later time, if appropriate.

Transition Issues

As discussed above, the Credit Card Act requires the Board to adopt final rules implementing new EFTA Section 915 within nine months of the date of enactment, or no later than February 22, 2010. These final rules must become effective no later than August 22, 2010.

In light of the pending effective date of the final rule, the Board seeks comment on the potential costs that would be incurred if issuers and other program participants were required to remove and replace card stock, including cards that have already been placed into store inventory, to ensure that all products sold on or after August 22, 2010 fully comply with the new requirements.

The Board also solicits comment on whether it should consider rules to grandfather gift certificates, store gift cards, or general-use prepaid cards, as those terms are defined, that are in the marketplace as of the effective date of the rule from some or all of the requirements set forth in this rulemaking. For example, the Board could require all such certificates or cards to comply with the substantive restrictions on imposing dormancy, inactivity, or service fees, and expiration dates, but otherwise permit such certificates or cards to be sold even if they do not contain the required disclosures. To the extent such relief would be provided, however, the Board believes it would be appropriate to do so only for cards that are sold in physical retail channels, but not to cards that are purchased on-line or by telephone, as they may not present the same operational challenges in replacing existing card stock compared to the former. In addition, if the Board were to permit certificates or cards that are available on retail shelves or in distribution warehouses to be sold to consumers after the effective date, comment is requested regarding how issuers or vendors could alert consumers to the changed terms regarding dormancy, inactivity, or service fees and funds expiration dates. The Board also solicits comment on an appropriate transition period after which all certificates or cards must fully comply with the new rules.

V. Initial Regulatory Flexibility Analysis

The Regulatory Flexibility Act (5 U.S.C. 601 *et seq.*) (RFA) generally requires an agency to perform an

assessment of the impact a rule is expected to have on small entities.

However, under section 605(b) of the RFA, the regulatory flexibility analysis otherwise required under section 604 of the RFA is not required if an agency certifies, along with a statement providing the factual basis for such certification, that the rule will not have a significant economic impact on a substantial number of small entities. Based on its analysis and for the reasons stated below, the Board believes that this proposed rule is not likely to have a significant economic impact on a substantial number of small entities. A final regulatory flexibility analysis will be conducted after consideration of comments received during the public comment period.

1. *Statement of the need for, and objectives of, the proposed rule.* The EFTA was enacted to provide a basic framework establishing the rights, liabilities, and responsibilities of participants in electronic fund transfer systems. The primary objective of the EFTA is the provision of individual consumer rights. 15 U.S.C. 1693. The EFTA authorizes the Board to prescribe regulations to carry out the purpose and provisions of the statute. 15 U.S.C. 1693b(a). The Act expressly states that the Board's regulations may contain "such classifications, differentiations, or other provisions, * * * as, in the judgment of the Board, are necessary or proper to effectuate the purposes of [the Act], to prevent circumvention or evasion [of the Act], or to facilitate compliance [with the Act]." 15 U.S.C. 1693b(c).

The Board is proposing revisions to Regulation E to implement Title IV of the Credit Card Act which would generally prohibit any person from imposing a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card, or general-use prepaid card. Title IV also generally provides that a gift certificate, store gift card, or general-use prepaid card may not be sold or issued unless the expiration date is no less than five years from the date a gift certificate is issued or five years from the date funds were last loaded to a store gift card or general-use prepaid card.

In addition, the proposed rule would require the disclosure of all other fees imposed in connection with a gift certificate, store gift card, or general-use prepaid card. The certificate or card must also state a toll-free telephone number and, if one is maintained, a Web site that a consumer may contact to obtain fee information or replacement certificates or cards.

The Board believes that the revisions to Regulation E discussed above are consistent with the Act, as amended by Title IV of the Credit Card Act, and within Congress's broad grant of authority to the Board to adopt provisions that carry out the purposes of the statute.

2. *Small entities affected by the proposed rule.* The number of small entities affected by this proposal is unknown. Under the proposed rule, a person would be prohibited from imposing a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card, or general-use prepaid card, unless three conditions are satisfied. First, a dormancy, inactivity, or service fee may be imposed only if there has been no activity with respect to the certificate or card within the one-year period prior to the imposition of the fee. Second, only one such fee may be assessed in a given calendar month. Third, disclosures regarding dormancy, inactivity, or service fees must be clearly and conspicuously stated on the certificate or card, and the issuer or vendor must provide these disclosures to the purchaser before the certificate or card is purchased.

The proposed rule would also provide that a gift certificate, store gift card, or general-use prepaid card may not be sold or issued unless the expiration date of the funds underlying the certificate or card is no less than five years after the date of issuance (in the case of a gift certificate) or five years after the date of last load of funds (in the case of a store gift card or general-use prepaid card). In addition, information regarding whether funds underlying a certificate or card may expire must be clearly and conspicuously stated on the certificate or card and given prior to purchase.

Two proposed alternative approaches are set forth to minimize potential confusion for consumers if the certificate or card expires before the underlying funds expire. The first alternative would prohibit the sale or issuance of a certificate or card that has a printed expiration date that is less than five years from the date of purchase. The second alternative would require entities subject to the rule to maintain policies or procedures to ensure that a consumer has a reasonable opportunity to purchase a certificate or card with an expiration date that is at least five years from the date of purchase. The proposed rule would also prohibit the imposition of any fees for replacing an expired certificate or card to ensure that consumers are able to access the underlying funds for the full five-year period.

In addition to the statutory fee restrictions described above, the proposed rule would require the disclosure of all other fees imposed in connection with a gift certificate, store gift card, or general-use prepaid card. These disclosures would have to be provided on or with the certificate or card and given prior to purchase. The proposed rule would also require the disclosure on the certificate or card of a toll-free telephone number and, if one is maintained, a Web site that a consumer may contact to obtain fee information or replacement certificates or cards.

Overall, to comply with the proposed rule, all persons involved in issuing, distributing or selling a gift card program may need to review and potentially revise disclosures that appear on or with a certificate or card. In addition, under either alternative approach to the rule addressing potential inconsistencies between card expiration dates and funds expiration dates, issuers, sellers, and distributors of gift certificates, store gift cards, and general-use prepaid cards will have to review and potentially revise their inventory distribution and management policies and controls to minimize the possibility that a consumer may purchase a card with an expiration date of less than five years from the date of purchase.

For gift certificates and store gift cards in particular, the proposed rule would potentially cover all merchants to the extent that they issue or sell gift certificates or store gift cards. According to the U.S. Census Bureau, there were over 3 million businesses that are involved in retail or food services as of September 2009.¹⁸ These businesses are potential issuers of gift certificates or store gift cards.¹⁹

The Small Business Administration (SBA) has defined a small business as one whose average annual receipts do not exceed \$7 million or who have fewer than 500 employees.²⁰ Of the over 3 million retail or food services businesses, the Board expects that well

¹⁸ See U.S. Census Bureau, Press Release, "Advance Monthly Sales for Retail and Food Services—September 2009," (available at: http://www.census.gov/retail/marts/www/marts_current.pdf).

¹⁹ The Board is unaware of any industry data regarding the number of merchants that issue gift certificates, store gift cards, or general-use prepaid cards. Nonetheless, the Board believes the actual number of merchants that issue such certificates or cards is likely to be far fewer than the number of businesses that are involved in retail or food services overall.

²⁰ See SBA, Summary of Size Standards by Industry (available at: <http://www.sba.gov/contractingopportunities/officials/size/summaryofssi/index.html>).

over 90% of these businesses qualify as small businesses under the SBA's standards.²¹ Consequently, a very large number of small entities across all retail trade or food categories could be subject to the proposed rules.

Nonetheless, the proposed requirements would only apply to the extent that a certificate or card program imposes dormancy, inactivity, or service fees or establishes an expiration date with respect to the underlying funds. In this regard, the Board understands that the vast majority of gift certificates and store gift cards issued by merchants or retailers today do not carry such fees or expiration dates.²² Moreover, smaller merchants are more likely to issue gift certificates in paper form only. Such certificates are excluded from coverage by the statute and proposed rule. *See* proposed § 205.20(b)(5). Thus, the Board believes the proposed rule would not impact a significant number of merchants that issue store gift cards or gift certificates. Similarly, the Board believes the proposed rule also would not significantly impact the entities that distribute or sell such cards or certificates on behalf of merchants. Moreover, the Board understands that given their size, such entities are unlikely to be "small businesses" as defined by the SBA.

In addition, the proposed rule would potentially cover issuers of general-use prepaid cards, primarily financial institutions, card program managers that issue or distribute general-use prepaid cards, and distributors or retailers of such cards. General-use prepaid cards may be more likely to carry dormancy, inactivity, or service fees and expiration dates compared to gift certificates and store gift cards. Consequently, entities that issue, distribute or sell general-use prepaid cards would be more likely to be impacted by the proposed rule.

As an initial matter, the Board notes that cards that would otherwise be considered general-use prepaid cards may in many cases be exempt from the statute and proposed rule because they

are reloadable and not marketed or labeled as a gift card or gift certificate. Moreover, as noted above, open-loop cards, which include general-use prepaid cards, make up a relatively small portion of the total prepaid card market in terms of number of cards issued and the dollar value of the amounts loaded. Thus, although the Board is not aware of any data regarding entities that issue or otherwise sell general-use prepaid cards, the Board does not believe that, overall, the rule is likely to have a significant impact on a substantial number of small entities with respect to the issuance or sale of general-use prepaid cards.

3. *Other Federal rules.* The Board has not identified any Federal rules that duplicate, overlap, or conflict with the proposed revisions to Regulation E.

4. *Significant alternatives to the proposed revisions.* The Board solicits comment on any significant alternatives that would reduce regulatory burden associated with this proposed rule on small entities.

VI. Paperwork Reduction Act

In accordance with the Paperwork Reduction Act (PRA) of 1995 (44 U.S.C. 3506; 5 CFR 1320 Appendix A.1), the Board reviewed the rule under the authority delegated to the Board by the Office of Management and Budget (OMB). The collection of information that is subject to the PRA by this proposed rule is found in 12 CFR part 205. The Federal Reserve may not conduct or sponsor, and an organization is not required to respond to, this information collection unless the information collection displays a currently valid OMB control number. The OMB control number is 7100-0200.

This information collection is required to provide benefits for consumers and is mandatory. *See* 15 U.S.C. 1693 *et seq.* Since the Board does not collect any information, no issue of confidentiality arises. The respondents/recordkeepers are for-profit financial institutions, including small businesses. Institutions are required to retain records for 24 months, but this regulation does not specify types of records that must be retained.

Title IV of the Credit Card Act prohibits any person from imposing a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card, or general-use prepaid card, unless three conditions are satisfied. First, such fees may be imposed only if there has been no activity with respect to the certificate or card within the one-year period prior to the imposition of the fee or charge. Second, only one such fee may be assessed in a given month.

Third, disclosures regarding dormancy, inactivity, or service fees must be clearly and conspicuously stated on the certificate or card, and the issuer or vendor must provide these disclosures before the certificate or card is purchased.

The Credit Card Act also provides that a gift certificate, store gift card, or general-use prepaid card may not be sold or issued unless the expiration date is no less than five years after the date of issuance (in the case of a gift certificate) or five years after the date of last load of funds (in the case of a store gift card or general-use prepaid card). In addition, the statute requires that the terms of expiration must be clearly and conspicuously stated on the certificate or card.

Any entities involved in the issuance, distribution, or sale of gift certificates, store gift cards, or general-use prepaid cards (or the issuance or distribution of loyalty, award, or promotional gift cards) potentially are affected by this collection of information because these entities will be required to provide disclosures regarding the fees imposed in connection with these certificates or cards and when the funds underlying a certificate or card expire. Under the proposed rule, gift certificates, store gift cards, and general-use prepaid cards must state certain disclosures about dormancy, inactivity, or service fees; expiration dates; and a telephone number and Web site, if one is maintained, for additional information. Disclosures about other fees must be provided on or with the certificate or card. In addition, disclosures about fees and expiration dates must be provided to the consumer prior to purchase. Loyalty, award, and promotional gift cards also must state disclosures regarding applicable fees and expiration dates.

Entities subject to the rule will have to review and revise disclosures that are currently provided on or with a certificate or card to ensure that they accurately state any fees and expiration dates that may apply.

The total estimated burden increase, as well as the estimates of the burden increase associated with each major section of the proposed rule as set forth below, represents averages for all respondents regulated by the Federal Reserve. The Federal Reserve expects that the amount of time required to implement each of the proposed changes for a given institution may vary based on the size and complexity of the respondent. Furthermore, the burden estimate for this rulemaking includes the burden addressing overdrafts to Regulation E, as announced in a

²¹ *See* Small Business Administration, Office of the Advocacy, Frequently Asked Questions (available at: <http://web.sba.gov/faqs/faqindex.cfm?areaID=24>); Employer Firms, & Employment by Employment Size of Firm by NAICS Codes, 2006 (available at: http://www.sba.gov/advo/research/us06_n6.pdf).

²² *See* Montgomery County Office of Consumer Protection, Gift Cards 2007 (available at: <http://www.montgomerycountymd.gov/ocptmpl.asp?url=/content/ocp/consumer/a-zgiftcardreports.asp>) (reporting that 18 of 22 retail gift cards surveyed do not carry any fees or expiration dates). *See also* Retail Gift Card Association, Code of Principles (available at: http://www.thergca.org/uploads/Code_of_Principles_PDF.pdf) (recommending as a best practice for retail gift card programs that no fees or expiration dates should apply).

separate final rulemaking (Docket No. R-1343).

Proposed § 205.20(b)(2) implements the exclusion for cards, codes, or other devices that are reloadable and not marketed or labeled as a gift card or gift certificate. As noted in proposed comment 205.20(b)(2)-4.i., institutions would qualify for this exclusion so long as policies and procedures reasonably designed to avoid the marketing of a prepaid card not otherwise subject to the rule, such as a general-purpose reloadable card, as a gift card or gift certificate are established and maintained. The Federal Reserve estimates that the 1,205 respondents regulated by the Federal Reserve would take, on average, 40 hours (one-business week) to implement written policies and procedures and provide training associated with proposed § 205.20(b)(2). The Federal Reserve estimates the annual one-time burden for respondents to be 48,200 hours and believes that, on a continuing basis, respondents would take an average of 8 hours annually to maintain their policies and procedures.

The Federal Reserve is proposing two alternative approaches for applying new EFTA Section 915(c) to the expiration of a certificate or card in § 205.20(e)(1). Alternative A proposes that institutions may not sell a gift certificate, store gift card, or general-use prepaid card subject to an expiration date unless the certificate or card expiration date is at least five years after the date the certificate or card is sold or issued to a consumer. Alternative B would require institutions involved in issuing, distributing, and selling certificates or cards to adopt policies and procedures to ensure that a consumer will have a reasonable opportunity to purchase a certificate or card with at least five years remaining until the certificate or card expiration date. With either alternative the Federal Reserve estimates that the 1,205 respondents regulated by the Federal Reserve would take, on average, 40 hours (one-business week) to implement or modify written policies and procedures and provide training associated with proposed § 205.20(e)(1). The Federal Reserve estimates the annual one-time burden for respondents to be 48,200 hours and believes that, on a continuing basis, respondents would take an average of 8 hours annually to maintain their policies and procedures.

Under proposed § 205.20(e)(3), three disclosures must be stated on the certificate or card, as applicable: (1) Disclosures must state the terms of expiration of the underlying funds or, if the underlying funds do not expire, that fact; (2) Disclosures must also include a toll-free telephone number and, if one is

maintained, a Web site that a consumer may use to obtain a replacement certificate or card after the certificate or card expires, if the underlying funds may be available; (3) The terms and conditions of funds expiration required to be disclosed must also include a statement that the certificate or card expires, but the underlying funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card. The Federal Reserve estimates that the 1,205 respondents regulated by the Federal Reserve would take, on average, 80 hours (two-business weeks) to update their systems to revise disclosures and redesign certificates or cards to comply with the proposed disclosure requirements in section 205.20(e)(3). The Federal Reserve estimates the annual one-time burden for respondents to be 96,400 hours and believes that, on a continuing basis, there would be no additional increase in burden.

The Federal Reserve estimates the proposed rule would impose a one-time increase in the annual burden under Regulation E for all respondents regulated by the Federal Reserve by 192,800 hours, from 526,520 to 719,320 hours. In addition, the Federal Reserve estimates that, on a continuing basis, the proposed requirements would increase the annual burden by 19,280 hours from 526,520 to 545,800 hours. The total annual burden would increase by 212,080 hours, from 526,520 to 738,600 hours.

The other Federal financial agencies are responsible for estimating and reporting to OMB the total paperwork burden for the institutions for which they have administrative enforcement authority. They may, but are not required to, use the Federal Reserve's burden estimation methodology. Using the Federal Reserve's method, the current total estimated annual burden for all persons subject to Regulation E, including Federal Reserve-supervised institutions would be approximately 1,403,459 hours. The above estimates represent an average across all respondents and reflect variations between persons based on their size, complexity, and practices. All covered persons, including depository institutions (of which there are approximately 17,200), potentially are affected by this collection of information, and thus are respondents for purposes of the PRA. The proposed rule would impose a one-time increase in the estimated annual burden for such institutions by 2,752,000 hours. On a continuing basis the proposed rule would increase in the estimated annual

burden for such institutions by 275,200 hours. The proposal total annual burden for the respondents regulated by the Federal financial agencies is estimated to be 4,430,659 hours.

Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the Federal Reserve's functions; including whether the information has practical utility; (b) the accuracy of the Federal Reserve's estimate of the burden of the proposed information collection, including the cost of compliance; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of information collection on respondents, including through the use of automated collection techniques or other forms of information technology. Comments on the collection of information should be sent to Michelle Shore, Federal Reserve Board Clearance Officer, Division of Research and Statistics, Mail Stop 95-A, Board of Governors of the Federal Reserve System, Washington, DC 20551, with copies of such comments sent to the Office of Management and Budget, Paperwork Reduction Project (7100-0200), Washington, DC 20503.

Text of Proposed Revisions

Certain conventions have been used to highlight the proposed changes to the text of the regulation and staff commentary. New language is shown inside bold-faced arrows, while language that would be deleted is set off with bold-faced brackets.

List of Subjects in 12 CFR Part 205

Consumer protection, Electronic fund transfers, Federal Reserve System, Reporting and recordkeeping requirements.

For the reasons set forth in the preamble, the Board proposes to amend 12 CFR part 205 and the Official Staff Commentary, as follows:

PART 205—ELECTRONIC FUND TRANSFERS (REGULATION E)

1. The authority citation for part 205 continues to read as follows:

Authority: 15 U.S.C. 1693b.

2. Section 205.4(a)(1) is revised to read as follows:

§ 205.4 General disclosure requirements; jointly offered services.

(a)(1) *Form of disclosures.* Disclosures required under this part shall be clear and readily understandable, in writing, and in a form the consumer may keep▶, except as otherwise provided in this part◀. The disclosures required by

this part may be provided to the consumer in electronic form, subject to compliance with the consumer-consent and other applicable provisions of the Electronic Signatures in Global and National Commerce Act (E-Sign Act) (15 U.S.C. 7001 *et seq.*). A financial institution may use commonly accepted or readily understandable abbreviations in complying with the disclosure requirements of this part.

* * * * *

3. Section 205.12(b)(1) is revised to read as follows:

§ 205.12 Relation to other laws.

* * * * *

(b) * * *

(1) *Inconsistent requirements.* The Board shall determine, upon its own motion or upon the request of a State, financial institution, or other interested party, whether the act and this part preempt State law relating to electronic fund transfers▶, or to dormancy, inactivity, or service fees, or expiration dates, of gift certificates, store gift cards, or general-use prepaid cards◀.

* * * * *

4. Section 205.20 is added as follows:

§ 205.20 Requirements for gift cards and gift certificates.

(a) *Definitions.* For purposes of this section, except as excluded under paragraph (b), the following definitions apply:

(1) *Gift certificate* means a card, code, or other device that is:

- (i) Issued to a consumer in a specified amount that may not be increased or reloaded in exchange for payment; and
- (ii) Redeemable upon presentation at a single merchant or an affiliated group of merchants for goods or services.

(2) *Store gift card* means a card, code, or other device that is:

(i) Issued to a consumer in a specified amount, whether or not that amount may be increased or reloaded by the cardholder, in exchange for payment; and

(ii) Redeemable upon presentation at a single merchant or an affiliated group of merchants for goods or services.

(3) *General-use prepaid card* means a card, code, or other device that is:

(i) Issued to a consumer in a specified amount, whether or not that amount may be increased or reloaded by the cardholder, in exchange for payment; and

(ii) Redeemable upon presentation at multiple, unaffiliated merchants for goods or services, or usable at automated teller machines.

(4) *Loyalty, award, or promotional gift card* means a card, code, or other device that:

(i) Is issued in connection with a loyalty, award, or promotional program;

(ii) Is redeemable upon presentation at one or more merchants for goods or services, or usable at automated teller machines; and

(iii) Sets forth the disclosures specified in paragraphs (d)(2), (e)(2), and (f)(2) of this section and provides the disclosures specified in paragraph (f)(1) of this section on or with the card, code, or other device.

(5) *Dormancy or inactivity fee.* The terms “dormancy fee” and “inactivity fee” mean a fee for non-use of or inactivity on a gift certificate, store gift card, or general-use prepaid card.

(6) *Service fee.* The term “service fee” means a periodic fee for holding or use of a gift certificate, store gift card, or general-use prepaid card.

(b) *Exclusions.* The terms “gift certificate,” “store gift card,” and “general-use prepaid card”, as defined in paragraph (a) of this section, do not include any card, code, or other device that is:

(1) Useable solely for telephone services;

(2) Reloadable and not marketed or labeled as a gift card or gift certificate;

(3) A loyalty, award, or promotional gift card;

(4) Not marketed to the general public;

(5) Issued in paper form only; or

(6) Redeemable solely for admission to events or venues at a particular location or group of affiliated locations, or to obtain goods or services, in conjunction with admission to such events or venues, at the event or venue or at specific locations affiliated with and in geographic proximity to the event or venue.

(c) *Form of disclosures.* (1) *Clear and conspicuous.* Disclosures made under this section must be clear and conspicuous. The disclosures may contain commonly accepted or readily understandable abbreviations or symbols.

(2) *Format.* Disclosures made under this section generally must be provided to the consumer in written or electronic form. Only disclosures provided under paragraph (c)(3) of this section may be given orally.

(3) *Disclosures prior to purchase.*

Before a gift certificate, store gift card, or general-use prepaid card is purchased, the issuer or vendor of such certificate or card must disclose to the consumer the information required by paragraphs (d)(2), (e)(3), and (f)(1) of this section.

(4) *Disclosures on the certificate or card.* Paragraphs (d)(2), (e)(3), and (f)(2) of this section require that certain

information be disclosed on the certificate or card. A disclosure made in an accompanying terms and conditions document, on packaging surrounding a certificate or card, or on a sticker or other label affixed to the certificate or card does not constitute a disclosure on the certificate or card.

(d) *Prohibition on imposition of fees or charges.*

No person may impose a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card, or general-use prepaid card, unless:

(1) There has been no activity with respect to the certificate or card in the one-year period ending on the date on which the fee is imposed;

(2) The following are stated, as applicable, clearly and conspicuously on the gift certificate, store gift card, or general-use prepaid card:

(i) The amount of any dormancy, inactivity, or service fee that may be charged;

(ii) How often such fee may be assessed; and

(iii) That such fee may be assessed for inactivity; and

(3) Not more than one dormancy, inactivity, or service fee is imposed in any given calendar month.

Alternative A—Paragraph (e)

(e) *Prohibition on sale of gift certificates or cards with expiration dates.* No person may sell or issue a gift certificate, store gift card, or general-use prepaid card with an expiration date, unless:

(1) The certificate or card expiration date, if any, is at least five years after the date the certificate or card was sold or issued to a consumer;

(2) The expiration date for the underlying funds is at least the later of:

(i) Five years after the date the gift certificate was issued, or five years after the date on which funds were last loaded to a store gift card or general-use prepaid card; or

(ii) The certificate or card expiration date, if any;

(3) The following disclosures are provided on the certificate or card, as applicable:

(i) The expiration date for the underlying funds or, if the underlying funds do not expire, that fact;

(ii) A toll-free telephone number and, if one is maintained, a Web site that a consumer may use to obtain a replacement certificate or card after the certificate or card expires if the underlying funds may be available; and

(iii) A statement, disclosed with equal prominence and in close proximity to the certificate or card expiration date, that the certificate or card expires, but

the underlying funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card; and

(4) No fee or charge is imposed on the cardholder for replacing the gift certificate, store gift card, or general-use prepaid card prior to the funds expiration date, unless such certificate or card has been lost or stolen.

Alternative B—Paragraph (e)

(e) *Prohibition on sale of gift certificates or cards with expiration dates.* No person may sell or issue a gift certificate, store gift card, or general-use prepaid card with an expiration date, unless:

(1) The person has policies and procedures in place to ensure that a consumer will have a reasonable opportunity to purchase a certificate or card with at least five years remaining until the certificate or card expiration date;

(2) The expiration date for the underlying funds is at least the later of:

(i) Five years after the date the gift certificate was issued, or the date on which funds were last loaded to a store gift card or general-use prepaid card; or

(ii) The certificate or card expiration date, if any;

(3) The following disclosures are provided on the certificate or card, as applicable:

(i) The expiration date for the underlying funds or, if the underlying funds do not expire, that fact;

(ii) A toll-free telephone number and, if one is maintained, a Web site that a consumer may use to obtain a replacement certificate or card after the certificate or card expires if the underlying funds may be available; and

(iii) A statement, disclosed with equal prominence and in close proximity to the certificate or card expiration date, that the certificate or card expires, but the underlying funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card; and

(4) No fee or charge is imposed on the cardholder for replacing the gift certificate, store gift card, or general-use prepaid card prior to the funds expiration date, unless such certificate or card has been lost or stolen.

(f) *Additional disclosure requirements for gift certificates or cards.* Additional disclosures must be provided in connection with a gift certificate, store gift card, or general-use prepaid card, as applicable, as indicated below:

(1) *Fee disclosures.* For each type of fee that may be imposed in connection

with the certificate or card (other than a dormancy, inactivity, or service fee subject to the disclosure requirements under paragraph (d)(2) of this section), the following information must be provided on or with the certificate or card:

(i) The type of fee;

(ii) The amount of the fee (or an explanation of how the fee will be determined); and

(iii) The conditions under which the fee may be imposed.

(2) *Telephone number for fee information.* A toll-free telephone number and, if one is maintained, a Web site that a consumer may use to obtain information about fees described in paragraphs (d)(2) and (f)(1) of this section must be disclosed on the certificate or card.

5. In Supplement I to part 205.

a. Under § 205.12 Relation to other laws, under (b) *Preemption of inconsistent State laws*, paragraph 1. is revised.

b. Section 205.20—Requirements for Gift Cards and Gift Certificates is added.

Supplement I to Part 205—Official Staff Interpretations

* * * * *

Section 205.12—Relation to Other Laws

* * * * *

(b) Preemption of Inconsistent State Laws

1. *Specific determinations.* The regulation prescribes standards for determining whether State laws that govern EFTs, dormancy, inactivity, or service fees, or expiration dates of gift certificates, store gift cards, or general-use prepaid cards are preempted by the act and the regulation. A State law that is inconsistent may be preempted even if the Board has not issued a determination. However, nothing in section 205.12(b) provides a financial institution with immunity for violations of State law if the institution chooses not to make State disclosures and the Board later determines that the State law is not preempted.

* * * * *

Section 205.20—Requirements for Gift Cards and Gift Certificates

20(a) Definitions

1. *Form of card, code, or device.*

Section 205.20 applies to any card, code, or other device that meets one of the definitions in § 205.20(a)(1) through (a)(3) of this section, even if it is not issued in card form. Section 205.20 would apply, for example, to the issuance of an account number or bar

code that can access underlying funds. Similarly, § 205.20 would apply to a device with a chip or other embedded mechanism, linking the device to stored funds, such as a mobile phone or sticker containing a contactless chip, if the device otherwise meets the definition of gift certificate, store gift card or general-use prepaid card.

2. *Electronic promise.* The term “electronic promise” as used in EFTA Sections 915(a)(2)(B), (a)(2)(C), and (a)(2)(D) means a person’s commitment or obligation communicated or stored in electronic form made to a consumer to provide payment for goods or services for transactions initiated by the consumer. The electronic promise is itself represented by a card, code or other device that is issued or honored by the person, reflecting the person’s commitment or obligation to pay. For example, if a merchant issues a code that can be given as a gift and that entitles the recipient to redeem the code in an on-line transaction for goods or services, that code represents an electronic promise by the merchant and would be a card, code, or other device covered by § 205.20.

Paragraph 20(a)(2)—Store Gift Card

1. *Relationship between “gift certificate” and “store gift card”.* The term “store gift card” in § 205.20(a)(2) includes “gift certificates” as defined in § 205.20(a)(1). For example, a numeric or alphanumeric code representing a specified dollar amount or value that is electronically sent to a consumer as a gift which can be redeemed or exchanged by the recipient to obtain goods or services may be both a “gift certificate” and a “store gift card” if the specified amount or value cannot be increased.

2. *Affiliated group of merchants.* The term “affiliated group of merchants” means two or more affiliated merchants or other persons that are related by common ownership or common corporate control (see, e.g., 12 CFR 227.3(b) and 12 CFR 223.2) and that share the same name, mark, or logo. For example, the term would include franchisees that are subject to a common set of corporate policies or practices under the terms of their franchise licenses. The term also applies to two or more merchants or other persons that agree among each other, by contract or otherwise, to redeem cards, codes, or other devices bearing the same name, mark, or logo (other than the mark, logo, or brand of a payment network), for the purchase of goods or services solely at such merchants or persons. For example, assume a movie theatre chain and a restaurant chain jointly agree to

issue cards that share the same “Flix and Food” logo that can be redeemed solely towards the purchase of movie tickets or concessions at any of the participating movie theatres, or towards the purchase of food or beverages at any of the participating restaurants. For purposes of § 205.20, the movie theatres and the restaurants would be considered to be an affiliated group of merchants, and the cards would be considered to be “store gift cards.”

3. *Mall gift cards.* See comment 20(a)(3)–2.

Paragraph 20(a)(3)—General-Use Prepaid Card

1. *Redeemable upon presentation at multiple, unaffiliated merchants.* A card, code, or other device is redeemable upon presentation at multiple, unaffiliated merchants if, for example, such merchants agree to honor the card, code, or device if it bears the mark, logo, or brand of a payment network, pursuant to the rules of the payment network.

2. *Mall gift cards.* Mall gift cards which are generally intended to be used or redeemed for goods or services at participating retailers within a shopping mall may be considered store gift cards or general-use prepaid cards depending on the locations in which the cards may be redeemed. For example, if a mall card may only be redeemed at merchants within the mall itself, the card is more likely to be considered a store gift card. However, certain mall cards also carry the brand of a payment network and can be used at any retailer that accepts that card brand, including retailers located outside of the mall. Such cards would be considered general-use prepaid cards.

Paragraph 20(a)(4)—Loyalty, Award, or Promotional Gift Card

1. *Examples of loyalty, award, or promotional programs.* Section 205.20(a)(4) defines a loyalty, award, or promotional gift card as a card, code, or other device that is issued in connection with a loyalty, award or promotional program. Such cards, codes, or other devices are excluded from the definitions of “gift certificate,” “store gift card,” and “general-use prepaid card” under § 205.20(b)(3), provided that the disclosures specified in paragraphs (d)(2), (e)(2), and (f) of this section are given to the consumer, on or with the card, as specified in § 205.20(a)(4)(iii). Examples of loyalty, award or promotional programs include:

i. Loyalty or consumer retention programs operated or administered by a merchant that provide to consumers cards redeemable for goods or services

or other monetary value as a reward for certain purchases at or visits to the participating merchant;

ii. Rebate programs operated or administered by a merchant or product manufacturer that provide cards redeemable for goods or services or other monetary value to consumers in connection with the consumer’s purchase of a product or service and the consumer’s completion of the rebate submission process.

iii. Sweepstakes or contests that distribute cards redeemable for goods or services or other monetary value to consumers as an invitation to enter into the promotion for a chance to win a prize.

iv. Referral programs that may provide cards redeemable for goods or services or other monetary value to consumers in exchange for referring other potential consumers to a merchant.

v. Incentive programs through which an employer may provide cards redeemable for goods or services or other monetary value to employees, for example, to recognize job performance, such as increased sales.

Paragraph 20(a)(6)—Service Fee

1. *Service fees.* Under § 205.20(a)(6), a service fee includes a periodic fee for holding or use of a gift certificate, store gift card, or general-use prepaid card. A periodic fee includes any fee that may be imposed on a gift certificate, store gift card, or general-use prepaid card from time to time for holding or using the certificate or card, such as a monthly maintenance fee, a transaction fee, a reload fee, or a balance inquiry fee, whether or not the fee is waived for a certain period of time or is only imposed after a certain period of time. A service fee does not include a one-time fee, such as an initial issuance fee or a cash-out fee.

20(b) Exclusions

1. *Application of exclusion.* A card, code, or other device is excluded from the definition of “gift certificate,” “store gift card,” or “general-use prepaid card” if it meets any of the exclusions in § 205.20(b). An excluded card, code, or other device generally is not subject to any of the requirements of this section. (See, however, § 205.20(a)(4)(iii), requiring certain disclosures for loyalty, award, or promotional gift cards).

2. *Eligibility for multiple exclusions.* A card, code, or other device may fall within more than one exclusion. If a card, code, or other device falls within any exclusion, it generally is not covered by § 205.20, even if another exclusion may not apply. Thus, for

example, a corporation may award its employees with a gift card of a type that can also be purchased directly from the merchant. While the card may not qualify for the exclusion for cards, codes, or other devices not marketed to the general public under § 205.20(b)(4) because the card can also be obtained through retail channels, it may nevertheless be exempt from the substantive requirements of § 205.20 because it is a loyalty, award, or promotional gift card. (See, however, § 205.20(a)(4)(iii), requiring certain disclosures for loyalty, award, or promotional gift cards.)

Paragraph 20(b)(1)—Usable Solely for Telephone Services

1. *Examples of excluded products.* The exclusion for products usable solely for telephone services applies to prepaid cards for long-distance telephone service, prepaid cards for wireless telephone service and prepaid cards for other services analogous in function to a telephone, such as prepaid cards for voice over Internet protocol (VoIP) access time.

Paragraph 20(b)(2)—Reloadable and Not Marketed or Labeled as a Gift Card or Gift Certificate

1. *Reloadable.* A card, code, or other device is “reloadable” if it has the capability of having more funds added by a cardholder after the initial purchase or issuance.

2. *Marketed or labeled as a gift card or gift certificate.* The term “marketed or labeled as a gift card or gift certificate” means directly or indirectly offering, advertising or otherwise suggesting the potential use of a card, code or other device, as a gift for another person. Whether the exclusion applies generally does not depend on the type of entity that makes the promotional message. For example, a card may be marketed or labeled as a gift card or gift certificate if anyone (other than the purchaser of the card), including the issuer, the retailer, the program manager that may distribute the card, or the payment network on which a card is used, promotes the use of the card as a gift card or gift certificate. A card or certificate, including a general-purpose reloadable card, is marketed or labeled as a gift card or gift certificate even if it is only occasionally marketed as a gift card or gift certificate. For example, a reloadable network-branded card would be marketed or labeled as a gift card or gift certificate if the issuer principally advertises the card as a less costly alternative to a bank account but promotes the card in a television, radio, newspaper, or Internet advertisement, or

on signage as “the perfect gift” during the holiday season.

3. *Examples of marketed or labeled as a gift card or gift certificate.* Examples of marketed or labeled as a gift card or gift certificate include:

- i. Displaying the word “gift” or “present” on a card, certificate, or accompanying material, including documentation, packaging and promotional displays;
- ii. Representing or suggesting that a certificate or card can be given to another person, for example, as a “token of appreciation” or a “stocking stuffer,” or displaying a congratulatory message on the card, certificate or accompanying material;
- iii. Incorporating gift-giving or celebratory imagery or motifs, such as a bow, ribbon, wrapped present, candle, or congratulatory message, on a card, certificate, accompanying documentation, or promotional material.

The term does not include:

- i. Representing that a card or certificate can be used as a substitute for a checking, savings, or deposit account;
- ii. Representing that a card or certificate can be used to pay for a consumer’s health-related expenses—for example, a card tied to a health savings account;
- iii. Representing that a card or certificate can be used as a substitute for travelers’ checks or cash by the purchaser;
- iv. Representing that a card or certificate can be used as a budgetary tool or to cover emergency expenses.

4. *Reasonable procedures regarding marketing.* The exclusion for a card, code, or other device is reloadable and is not marketed or labeled as a gift card or gift certificate in § 205.20(b)(2) applies if an individual card, code, or other device is not marketed or labeled as a gift card or gift certificate and if entities subject to the rule maintain policies and procedures reasonably designed to avoid such marketing. The following examples illustrate the application of § 205.20(b)(2):

i. An issuer or program manager of prepaid cards agrees to sell general-purpose reloadable cards through a retailer. The contract between the issuer or program manager and the retailer establishes the terms and conditions under which the cards may be sold and marketed at the retailer. The terms and conditions include restrictions prohibiting the general-purpose reloadable cards from being marketed as a gift card or gift certificate, and requirements for policies and procedures to regularly monitor or otherwise verify that the cards are not

being marketed as such. The issuer or program manager sets up one promotional display at the retailer for gift cards and another physically separated display for excluded products under § 205.20(b), including general-purpose reloadable cards and wireless telephone cards, such that a reasonable consumer would not believe that the excluded cards are gift cards. The exclusion in § 205.20(b)(2) applies even if a retail clerk inadvertently stocks or places some of the general-purpose reloadable cards on the gift card display notwithstanding the issuer or program manager’s maintenance of policies and procedures reasonably designed to avoid the marketing of the general-purpose reloadable cards as gift cards or gift certificates.

ii. Same facts as in i., except that the issuer or program manager sets up a single promotional display at the retailer on which a variety of prepaid cards are sold, including store gift cards, general-purpose reloadable cards, and wireless telephone cards. A sign stating “Gift Cards” appears prominently at the top of the display. The issuer or program manager does not qualify for the exclusion in § 205.20(b)(2) with respect to the general-purpose reloadable card because the issuer or program manager does not maintain policies and procedures reasonably designed to avoid the marketing of the general-purpose reloadable cards as gift cards or gift certificates.

Paragraph 20(b)(4)—Not Marketed to the General Public

1. *Marketed to the general public.* A card, code, or other device is marketed to the general public if the potential use of the card, code, or other device is directly or indirectly offered, advertised, or otherwise promoted to the general public. A card, code, or other device may be marketed to the general public regardless of the advertising medium, including television, radio, newspaper, the Internet, or signage. In addition, the method of distribution by itself is not dispositive in determining whether a card, code, or other device is marketed to the general public. Factors that may be considered in determining whether the exclusion applies to a particular card, code, or other device include the means or channel through which the card, code, or device may be obtained by a consumer, the subset of consumers that are eligible to obtain the card, code or device, and whether the availability of the card, code, or device is advertised or otherwise promoted in the marketplace.

2. *Examples illustrating exclusion for cards, codes, or other devices “not*

marketed to the general public.” The following examples illustrate application of the exclusion in § 205.20(b)(4) for cards, codes, or other devices not marketed to the general public.

i. A merchant sells its gift cards at a discount to a business which may give them to employees or loyal consumers as incentives or rewards. In determining whether the gift card falls within the exclusion in § 205.20(b)(4), the merchant must consider whether the card is of a type that is advertised or made available to consumers generally or can be obtained elsewhere. If the card can also be purchased through retail channels, the exclusion in § 205.20(b)(4) does not apply, even if the consumer obtained the card from the business as an incentive or reward. *See, however,* § 205.20(b)(3).

ii. A national retail chain decides to market its gift cards only to members of its frequent buyer program. If any member of the general public may become a member of the program, the card does not fall within the exclusion in § 205.20(b)(4) because the general public has the ability to obtain the cards.

iii. An issuer of prepaid cards advertises a reloadable card to teenagers and their parents promoting the card for use by teenagers for occasional expenses, schoolbooks and emergencies and by parents to monitor spending. Because the card is marketed to and may be sold to any member of the general public, the exclusion in § 205.20(b)(4) does not apply.

iv. An insurance company settles a policyholder’s claim and distributes the insurance proceeds to the consumer by means of a prepaid card. Because the prepaid card is simply the means for providing the insurance proceeds to the consumer and the availability of the card is not advertised to the general public, the exclusion in § 205.20(b)(4) applies.

v. An employer provides a prepaid card to its employees to cover travel expenses and per diem. Because the prepaid card is simply the means for distributing travel expenses and per diem and the availability of the card is not advertised or available to the general public, the exclusion in § 205.20(b)(4) applies.

vi. A merchant provides store credit to a consumer following a merchandise return by issuing a prepaid card that clearly indicates that the card contains funds for store credit. Because the prepaid card is issued for the stated purpose of providing store credit to the consumer and the ability to receive refunds by a prepaid card is not

advertised to the general public, the exclusion in § 205.20(b)(4) applies.

vii. A tax preparation company elects to distribute tax refunds to its clients by issuing non-reloadable prepaid cards, but does not advertise or otherwise promote the ability to receive proceeds in this manner. Because the prepaid card is simply the mechanism for providing the tax refund to the consumer, and the tax preparer does not advertise the ability to obtain tax refunds by a prepaid card, the exclusion in § 205.20(b)(4) applies. However, if the tax preparer promotes the ability to receive tax refund proceeds through a prepaid card as a way to obtain “faster” access to the proceeds, the exclusion in § 205.20(b)(4) does not apply.

Paragraph 20(b)(5)—Issued in Paper Form Only

1. *Exclusion explained.* To qualify for the exclusion in § 205.20(b)(5), the sole means of issuing the card, code, or other device must be in a paper form. Thus, the exclusion generally applies to certificates issued in paper form where solely the paper itself may be used to purchase goods or services. A card, code or other device is not issued solely in paper form simply because it may be reproduced or printed on paper. For example, a bar code or card or certificate number sent electronically to a consumer and redeemable for goods and services is not issued in paper form, even if it may be reproduced or otherwise printed on paper by the consumer. Similarly, an on-line retailer may electronically mail a certificate redeemable for goods or services to a consumer, which the consumer could print out on a home printer. In these circumstances, although the consumer might hold a paper facsimile of the card, code, or other device, the exclusion does not apply because the information necessary to redeem the value was initially issued in electronic form. However, a paper certificate that bears a bar code or account number may fall within the exclusion in § 205.20(b)(5) if the bar code or account number is not issued in any form other than on the paper. In addition, the exclusion in § 205.20(b)(5) would continue to apply in circumstances where an issuer replaces a gift certificate that was initially issued in paper form with a card or electronic code (for example, to replace a lost paper certificate).

Paragraph 20(b)(6)—Redeemable Solely for Admission to Events or Venues

1. *Examples.* The exclusion for payment cards, codes, or other devices that are redeemable solely for admission to events or venues at a particular

location or group of affiliated locations generally applies to cards, codes, or other devices that are not redeemed for a specified monetary value, but rather for admission for entry to an event or venue. The exclusion also covers a card, code, or other device that is usable to purchase of goods or services purchased in addition to entry into the event or the venue, either at the event or venue or at an affiliated location or location in geographic proximity to the event or venue. The following examples illustrate the scope of § 205.20(b)(6):

i. A consumer purchases a prepaid card that entitles the holder to a ticket for entry to an amusement park. The prepaid card does not state a monetary value and may only be used for entry to the park. The card qualifies for the exclusion in § 205.20(b)(6) because it is redeemable solely for admission or entry to an event or venue.

ii. Same facts as in i., except that the gift card also entitles the holder of the gift card to a dollar amount that can be applied towards the purchase of food and beverages or goods or services at the park or at nearby affiliated locations. The card qualifies for the exclusion in § 205.20(b)(6) because it is redeemable for admission or entry and for goods or services in conjunction with that admission.

iii. A consumer purchases a \$25 gift card that the holder of the gift card can use to make purchases at a merchant but alternatively can also apply the value on the card towards the cost of admission to the merchant’s affiliated amusement park. The card is not eligible for the exclusion in § 205.20(b)(6) because it is not redeemable solely for the admission or ticket itself (or for goods and services purchased in conjunction with such admission). The card meets the definition of “store gift card” and is therefore subject to the substantive and disclosure requirements of §§ 205.20(d), (e), and (f), unless a different exclusion applies.

iv. A consumer purchases a gift card that is redeemable for a particular service such as a spa treatment or for a one-night hotel stay. The card is not eligible for the exclusion in § 205.20(b)(6) because it is not redeemable for admission to an event or venue (in this case, the spa or hotel), but instead for a specified service at the spa or hotel. The card meets the definition of “store gift card” and is therefore subject to the substantive and disclosure requirements of §§ 205.20(d), (e), and (f), unless a different exclusion applies.

v. A consumer purchases a gift card that is redeemable solely for a one-year membership to a buyer’s club or warehouse, or to a gym. The card falls

within the exclusion in § 205.20(b)(6) because it is redeemable solely for membership to the club or gym and the membership is necessary for entry or admission to the club or gym. The exclusion would not apply, however, if the card has value that could be applied *either* to membership or for goods or services at the warehouse or gym.

20(c) Form of Disclosures

Paragraph 20(c)(1)—Clear and Conspicuous

1. *Clear and conspicuous standard.* All disclosures required by this section must be clear and conspicuous. Disclosures are clear and conspicuous for purposes of this section if they are readily understandable and, in the case of written and electronic disclosures, the location and type size are readily noticeable to consumers. Disclosures need not be located on the front of the certificate or card to be considered clear and conspicuous. Disclosures are clear and conspicuous for the purposes of this section if they are in a print that contrasts with and is otherwise not obstructed by the background on which they are printed. For example, disclosures on a card or computer screen are not likely to be conspicuous if obscured by a logo printed in the background. Similarly, disclosures on the back of a card that are printed on top of indentations from embossed type on the front of the card are not likely to be conspicuous if it obstructs the readability of the type. To the extent permitted, oral disclosures meet the standard when they are given at a volume and speed sufficient for a consumer to hear and comprehend them.

2. *Abbreviations and symbols.* Disclosures may contain commonly accepted or readily understandable abbreviations or symbols, such as “mo.” for month or a “/” to indicate “per.” Under the clear and conspicuous standard, it is sufficient to state, for example, that a particular fee is charged “\$2.50/mo. after 12 mos.”

Paragraph 20(c)(2)—Format

1. *Electronic disclosures.* Disclosures provided electronically pursuant to this section are not subject to compliance with the consumer-consent and other applicable provisions of the Electronic Signatures in Global and National Commerce Act (E-Sign Act) (15 U.S.C. 7001 *et seq.*). Electronic disclosures may not be provided through a hyperlink or in another manner by which the purchaser can bypass the disclosure. An issuer or vendor is not required to

confirm that the consumer has read the electronic disclosures.

2. *Non-physical certificates and cards.* If no certificate or card is issued, the disclosures must accompany the code, confirmation, or other written or electronic document provided to the consumer.

Paragraph 20(c)(3)—Disclosure Prior to Purchase

1. *Method of purchase.* The disclosures must be provided before a certificate or card is purchased regardless of whether the certificate or card is purchased in person, on-line, by telephone, or by other means.

20(d) Prohibition on Imposition of Fees or Charges

1. *One-year period.* Section 205.20(d) provides, in part, that a person may not impose a dormancy, inactivity, or service fee with respect to a gift certificate, store gift card, or general-use prepaid card until there has been no activity with respect to the certificate or card in the one-year period ending on the date on which the fee is imposed. The following examples illustrate this rule:

i. A certificate or card is purchased on January 15 of year one. If there has been no activity on the certificate or card since the certificate or card was purchased, a dormancy, inactivity, or service fee may not be imposed on the certificate or card until January 15 of year two.

ii. A certificate or card is purchased on February 29 of a leap year. If there has been no activity on the certificate or card since the certificate or card was purchased, a dormancy, inactivity, or service fee may not be imposed on the certificate or card until February 28 of the following year.

iii. Same facts as i., and a fee was imposed on January 15 of year two. Because no more than one dormancy, inactivity, or service fee may be imposed in any given calendar month, the earliest date that another dormancy, inactivity, or service fee may be imposed, assuming there continues to be no activity on the certificate or card, is February 1 of year two. A dormancy, inactivity, or service fee is permitted to be imposed on February 1 of year two because there has been no activity on the certificate or card for the preceding year (February 1 of year one through January 31 of year two), and February is a new calendar month.

iv. Same facts as i., and a fee was imposed on January 15 of year two. On January 31 of year two, the consumer uses the card to make a purchase. Under this circumstance, another dormancy,

or service fee could not be imposed until January 31 of year three at the earliest, assuming there has been no activity on the certificate or card since January 31 of year two.

2. *Activity.* Any action by a consumer to increase, decrease, or otherwise make use of the funds underlying a gift certificate, store gift card, or general-use prepaid card constitutes activity for purposes of § 205.20(d). For example, the purchase and activation of a certificate or card or the reloading of funds onto a store gift card or general-use prepaid card constitutes activity. However, neither the imposition of a fee, the replacement of an expired, lost, or stolen certificate or card, nor a balance inquiry constitutes activity with respect to a gift certificate, store gift card, or general-use prepaid card.

3. *Relationship between §§ 205.20(d)(2) and (c)(3).* Sections 205.20(d)(2) and (c)(3) contain similar, but not identical, disclosure requirements. Section 205.20(d)(2) requires the disclosure of dormancy, inactivity, and service fees on a certificate or card. Section 205.20(c)(3) requires that an issuer or vendor of such certificate or card disclose to a consumer any dormancy, inactivity, and service fees associated with the certificate or card before such certificate or card may be purchased. Depending on the context, a single disclosure that meets the clear and conspicuous requirements of both §§ 205.20(d)(2) and (c)(3) may be used to disclose a dormancy, inactivity, or service fee. For example, if the disclosures on a certificate or card, required by § 205.20(d)(2), are visible to the consumer without having to remove packaging or other materials sold with the certificate or card, for a purchase made in person, the disclosures also meet the requirements of § 205.20(c)(3). Otherwise, a dormancy, inactivity, or service fee may need to be disclosed multiple times or in multiple locations to satisfy the requirements of §§ 205.20(d)(2) and (c)(3). For example, if the disclosures on a certificate or card, required by § 205.20(d)(2), are obstructed by packaging or other materials sold with the certificate or card, for a purchase made in person, they also must be disclosed on the packaging sold with the certificate or card or in other manner visible to the consumer to meet the requirements of § 205.20(c)(3).

4. *Relationship between §§ 205.20(d)(2), (e)(3), and (f)(2).* In addition to any disclosures required under § 205.20(d)(2), any applicable disclosures under §§ 205.20(e)(3) and

(f)(2) of this section must also be provided on the certificate or card.

5. *One fee per month.* Under § 205.20(d)(3), no more than one dormancy, inactivity, or service fee may be imposed in any given calendar month. For example, if a dormancy fee is imposed on January 1, following a year of inactivity, and a consumer makes a balance inquiry on January 15, a balance inquiry fee may not be imposed at that time because a dormancy fee was already imposed earlier that month and a balance inquiry fee is a type of service fee. If, however, the dormancy fee could be imposed on January 1, following a year of inactivity, and the consumer performs a balance inquiry on January 1, the person assessing the fees may choose whether to impose the dormancy fee or the balance inquiry fee on January 1. The restriction in § 205.20(d)(3) does not apply to any fee that is not a dormancy, inactivity, or service fee. For example, assume a service fee is imposed on January 1, following a year of inactivity. If a consumer cashes out the funds on a general-use prepaid card on January 15, a cash-out fee may be imposed at that time because a cash-out fee is not a dormancy, inactivity, or service fee.

20(e) Prohibition on Sale of Gift Certificates or Cards With Expiration Dates

Alternative A

1. *Disclosure of funds expiration—date not required.* Section 205.20(e)(3)(i) does not require disclosure of the precise date the funds will expire. It is sufficient to disclose, for example, “Funds expire 5 years from the date funds last loaded to the card.”; “Funds can be used 5 years from the date money was last added to the card.”; or “Funds do not expire.”

2. *Disclosure not required if no expiration date.* If the certificate or card and underlying funds do not expire, the disclosure required by § 205.20(e)(3)(i) need not be stated on the certificate or card.

3. *Reference to toll-free telephone number and Web site.* If a certificate or card does not expire, or if the underlying funds are not available after the certificate or card expires, the disclosure required by § 205.20(e)(3)(ii) need not be stated on the certificate or card. *See, however,* § 205.20(f)(2).

4. *Relationship to § 226.20(f)(2).* The same toll-free telephone number and Web site may be used to comply with §§ 226.20(e)(3)(ii) and (f)(2). Neither a toll-free number nor a Web site must be maintained or disclosed on a certificate or card if no fees are imposed in

connection with the certificate or card, and the certificate or card and underlying funds do not expire.

5. *Distinguishing between certificate or card expiration and funds expiration.* If applicable, § 205.20(e)(3)(iii) requires a disclosure to be made on the certificate or card that notifies a consumer that the certificate or card expires, but the underlying funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card. The disclosure must be made with equal prominence and in close proximity to the certificate or card expiration date. In the case of a certificate or card, close proximity means that the disclosure must be on the same side as the certificate or card expiration date. If the disclosure is the same type size and is located immediately next to or directly above or below the certificate or card expiration date, without any intervening text or graphical displays, the disclosures would be deemed to be equally prominent and in close proximity. The disclosure need not be embossed on the certificate or card to be deemed equally prominent, even if the expiration date is embossed on the certificate or card. The disclosure may state on the front of the card, for example, “Valid thru 09/2016. Call for new card.”; “Active thru 09/2016. Call for replacement card.”; or “Call for new card after 09/2016.”

6. *Relationship between §§ 205.20(d)(2), (e)(3), and (f)(2).* In addition to disclosures required under § 205.20(e)(3), any applicable disclosures under §§ 205.20(d)(2) and (f)(2) of this section must also be provided on the certificate or card.

7. *Replacement of a lost or stolen certificate or card not required.* Section 205.20 does not require the replacement of a certificate or card that has been lost or stolen.

8. *Date of issuance or loading.* A certificate or card is not issued or loaded with funds until the certificate or card is activated for use.

Alternative B

1. *Reasonable opportunity.* Under § 205.20(e)(1), no person may sell or issue a gift certificate, store gift card, or general-use prepaid card with an expiration date, unless there are policies or procedures in place to ensure that a consumer has a reasonable opportunity to purchase a certificate or card with at least five years remaining until the certificate or card expiration date. The following examples illustrate reasonable and unreasonable opportunities for consumers to purchase a certificate or card with at least five years remaining

until the certificate or card expiration date:

i. A card would comply with § 205.20(e)(1) if it is printed with a card expiration date six years from the date it was produced and is on a display rack of a retail store within six months of the date the card was produced.

ii. A card would comply with § 205.20(e)(1) if it is printed with a card expiration date seven years from the date it was produced and is on a display rack of a retail store within one year and six months of the date the card was produced, the card would comply with § 205.20(e)(1).

iii. A card would not comply with § 205.20(e)(1) if it is printed with a card expiration date six years from the date it was produced and is stored in a distribution warehouse for more than one year after the date the card was produced.

2. *Disclosure of funds expiration—date not required.* Section 205.20(e)(3)(i) does not require disclosure of the precise date the funds will expire. It is sufficient to disclose, for example, “Funds expire 5 years from the date funds last loaded to the card.”; “Funds can be used 5 years from the date money was last added to the card.”; or “Funds do not expire.”

3. *Disclosure not required if no expiration date.* If the certificate or card and underlying funds do not expire, the disclosure required by § 205.20(e)(3)(i) need not be stated on the certificate or card.

4. *Reference to toll-free telephone number and Web site.* If a certificate or card does not expire, or if the underlying funds are not available after the certificate or card expires, the disclosure required by § 205.20(e)(3)(ii) need not be stated on the certificate or card. *See, however,* § 205.20(f)(2).

5. *Relationship to § 226.20(f)(2).* The same toll-free telephone number and Web site may be used to comply with §§ 226.20(e)(3)(ii) and (f)(2). Neither a toll-free number nor a Web site must be maintained or disclosed if no fees are imposed in connection with a certificate or card, and the certificate or card and underlying funds do not expire.

6. *Distinguishing between certificate or card expiration and funds expiration.* If applicable, a disclosure must be made on the certificate or card that notifies a consumer that the certificate or card expires, but the funds either do not expire or expire later than the certificate or card, and that the consumer may contact the issuer for a replacement card. The disclosure must be made with equal prominence and in close proximity to the certificate or card expiration date. In the case of a

certificate or card, close proximity means that the disclosure must be on the same side as the certificate or card expiration date. If the disclosure is the same type size and is located immediately next to or directly above or below the certificate or card expiration date, without any intervening text or graphical displays, the disclosures would be deemed to be equally prominent and in close proximity. The disclosure need not be embossed on the certificate or card to be deemed equally prominent, even if the expiration date is embossed on the certificate or card. The disclosure may state on the front of the card, for example, “Valid thru 09/2016. Call for new card.”; “Active thru 09/2016. Call for replacement card.”; or “Call for new card after 09/2016.”

7. *Relationship between §§ 205.20(d)(2), (e)(3), and (f)(2).* In addition to any disclosures required to be made under § 205.20(e)(3), any applicable disclosures under §§ 205.20(d)(2) and (f)(2) must also be provided on the certificate or card.

8. *Replacement of a lost or stolen certificate or card not required.* Section 205.20 does not require the replacement of a certificate or card that has been lost or stolen.

9. *Date of issuance or loading.* A certificate or card is not issued or loaded with funds until the certificate or card is activated for use.

20(f) Additional Disclosure Requirements for Gift Certificates or Cards

1. *Reference to toll-free telephone number and Web site.* If a certificate or card does not have any fees, the disclosure required by § 205.20(f)(2) need not be stated on the certificate or card. *See, however,* § 205.20(e)(3)(ii).

2. *Relationship to § 226.20(e)(3)(ii).* The same toll-free telephone number and Web site may be used to fulfill §§ 226.20(e)(3)(ii) and (f)(2). Neither a toll-free number nor a Web site must be maintained or disclosed if no fees are imposed in connection with a certificate or card, and the certificate or card and underlying funds do not expire.

3. *Relationship between §§ 205.20(d)(2), (e)(3), and (f)(2).* In addition to any disclosures required to be made pursuant to § 205.20(f)(2), any applicable disclosures under §§ 205.20(d)(2) and (e)(3) must also be provided on the certificate or card.

By order of the Board of Governors of the Federal Reserve System, November 13, 2009.

Jennifer J. Johnson,
Secretary of the Board.

[FR Doc. E9–27717 Filed 11–19–09; 8:45 am]

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S. 475/P.L. 111-97
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S. 509/P.L. 111-98

To authorize a major medical facility project at the Department of Veterans Affairs Medical Center, Walla Walla, Washington, and for other purposes. (Nov. 11, 2009; 123 Stat. 3010)

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