its terms. The contractor will not have a right to a hearing or judicial review regarding CMS’s renewal or non-renewal decision.

§ 455.238 Conflict of Interest.
(a) Offerors for Medicaid integrity audit program contracts, and Medicaid integrity audit program contractors, are subject to the following requirements:
(1) The conflict of interest standards and requirements of the Federal Acquisition Regulation organizational conflict of interest guidance, found under 48 CFR subpart 9.5.
(2) The standards and requirements that are contained in each individual contract awarded to perform activities described under section 1936 of the Act.
(b) Post-award conflicts of interest: CMS considers that a post-award conflict of interest has developed if, during the term of the contract, one of the following occurs:
(1) The contractor or any of its employees, agents, or subcontractors received, solicited, or arranged to receive any fee, compensation, gift (defined at 5 CFR 2635.203(b)), payment of expenses, offer of employment, or any other thing of value from any entity that is reviewed, audited, investigated, or contacted during the normal course of performing activities under the Medicaid integrity audit program contract.
(2) CMS determines that the contractor’s activities are creating a conflict of interest.
(c) If CMS determines that a conflict of interest exists during the term of the contract, among other actions, CMS may:
(1) Not renew the contract for an additional term.
(2) Modify the contract.
(3) Terminate the contract.

§ 435.240 Conflict of Interest Resolution.
(a) Review Board: CMS may establish a Conflicts of Interest Review Board to assist in resolving organizational conflicts of interest.
(b) Resolution: Resolution of an organizational conflict of interest is a determination by the contracting officer that:
(1) The conflict is mitigated.
(2) The conflict precludes award of a contract to the offeror.
(3) The conflict requires that CMS modify an existing contract.
(4) The conflict requires that CMS terminate an existing contract.
(5) It is in the best interest of the government to contract with the offeror or contractor even though the conflict of interest exists and a request for waiver is approved in accordance with 48 CFR 9.503.

In the August 22, 2003 Federal Register (68 FR 50756), we published a notice announcing a new SOR titled Automated Survey Processing Environment (ASPEN) Complaint/Incidents Tracking System (ACTS), HHS/CMS, System No. 09–70–0565.

In the May 23, 2006 Federal Register (71 FR 29643) we published a notice that modified the ACTS SOR. This notice included all modifications and the full text of this system of records. ACTS is a Windows-based program whose primary purpose is to track and process complaints and incidents reported against health care facilities regulated by CMS and State agencies. These facilities include Clinical Laboratory Improvement Amendment (CLIA)-certified laboratories, skilled nursing facilities, hospitals, home health agencies (HHAs), end stage renal disease (ESRD) facilities, hospices, rural health clinics (RHGs), comprehensive outpatient rehabilitation facilities (CORFs), outpatients physical therapy services, community mental health centers (CMHCS), ambulatory surgical centers (ASCs), suppliers of portable x-ray services, and intermediate care facilities for persons with mental retardation (ICF/MRs). ACTS contains identifiable information on individuals, who are complainants, residents, patients, clients, contacts or witnesses. It also may include alleged perpetrators, survey team members, laboratory directors, laboratory owners, and employees and directors of the health care facilities noted previously. ACTS is designed to manage all operations associated with complaint and incident tracking and processing, from initial intake and investigation through the final disposition.

B. The Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HTIS), HHS/CMS, System No. 09–70–0544.

In the July 6, 2005 Federal Register (70 FR 38944), we published a notice announcing a new SOR titled Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HTIS), HHS/CMS, System No. 09–70–0544.

In general, HTIS consists of an electronic repository of information, documents, and supplementary paper document files resulting from investigations of alleged violations of the transactions and code sets, security, and unique identifier provisions of HIPAA. HTIS’ purpose is to support investigations of complainants, determinations as to whether there were violations as charged in the original complaint, referral of violations to law enforcement entities as necessary, and maintenance and retrieval of records that contain the results of the complaint investigations. The system of records

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Office of the Secretary
45 CFR Part 5b
[CMS–0029–F]
RIN 0938–A069
Exemption of Certain Systems of Records Under the Privacy Act
AGENCY: Office of the Secretary, HHS.
ACTION: Final rule.
SUMMARY: This final rule exempts four systems of records (SORs) from subsections (c)(3), (d)(1) through (d)(4), (e)(4)(G) and (H), and (l) of the Privacy Act pursuant to 5 U.S.C. 552a(k)(2): The Automated Survey Processing Environment (ASPE) Complaint/Incidents Tracking System (ACTS), HHS/CMS, System No. 09–70–0565; the Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HTIS), HHS/CMS, System No. 09–70–0544; the Organ Procurement Organizations System (OPOS), HHS/CMS, System No. 09–70–0575; and the Fraud Investigation Database (FID), HHS/CMS, System No. 09–70–0527.
DATES: Effective Date: These regulations are effective on October 27, 2008.
FOR FURTHER INFORMATION CONTACT: Walter Stone, (410) 786–3577.
SUPPLEMENTARY INFORMATION:
I. Background
The four systems of records (SORs) that are the subject of this final rule and the May 25, 2007 proposed rule are as follows:
A. The Automated Survey Processing Environment Complaints/Incidents Tracking System (ACTS), HHS/CMS, System No. 09–70–0565
In the August 22, 2003 Federal Register (68 FR 50756), we published a notice announcing a new SOR titled Automated Survey Processing Environment (ASPEN) Complaint/Incidents Tracking System (ACTS), HHS/CMS, System No. 09–70–0565.

In the May 23, 2006 Federal Register (71 FR 29643) we published a notice that modified the ACTS SOR. This notice included all modifications and the full text of this system of records. ACTS is a Windows-based program whose primary purpose is to track and process complaints and incidents reported against health care facilities regulated by CMS and State agencies. These facilities include Clinical Laboratory Improvement Amendment (CLIA)-certified laboratories, skilled nursing facilities, hospitals, home health agencies (HHAs), end stage renal disease (ESRD) facilities, hospices, rural health clinics (RHGs), comprehensive outpatient rehabilitation facilities (CORFs), outpatient physical therapy services, community mental health centers (CMHCS), ambulatory surgical centers (ASCs), suppliers of portable x-ray services, and intermediate care facilities for persons with mental retardation (ICF/MRs). ACTS contains identifiable information on individuals, who are complainants, residents, patients, clients, contacts or witnesses. It also may include alleged perpetrators, survey team members, laboratory directors, laboratory owners, and employees and directors of the health care facilities noted previously. ACTS is designed to manage all operations associated with complaint and incident tracking and processing, from initial intake and investigation through the final disposition.

B. The Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HTIS), HHS/CMS, System No. 09–70–0544.

In the July 6, 2005 Federal Register (70 FR 38944), we published a notice announcing a new SOR titled Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HTIS), HHS/CMS, System No. 09–70–0544.
covers individuals who have submitted complaints alleging violations of the provisions of HIPAA. Investigative files maintained in HITS are received either as electronic documents or as paper records that are compiled for law enforcement purposes.

C. The Organ Procurement Organizations System (OPOS), HHS/CMS, System No. 09–70–0575

In the May 22, 2006 Federal Register (71 FR 29336), we published a notice announcing a new SOR titled Organ Procurement Organizations System (OPOS), HHS/CMS, System No. 09–70–0575. OPOS is a Windows based program whose purpose is to track and process complaints and incidents reported against Organ Procurement Organizations. Section 701 of the Organ Procurement Organization System Certification Act of 2000 (Pub. L. 106–505) gave the Department the authority to collect and maintain individually identifiable information pertaining to allegations filed by a complainant, beneficiary, or provider of services against Organ Procurement Organizations. This information includes information gathered during all aspects of an investigation, including initial complaints, findings, results, disposition, and relevant correspondence.

D. The Fraud Investigation Database (FID), HHS/CMS, System No. 09–70–0527

In the October 28, 2002 Federal Register (70 FR 65795), we published a notice that modified, among other things, the name of a SOR entitled “CMS Utilization Review Investigatory Files, System No. 09–70–0527” to “CMS Fraud Investigation Database (FID).” The notice included the full text of the FID system of records. The FID system of records contains the name, work address, work phone number, social security number, Unique Provider Identification Number (UPIN), and other identifying demographics of individuals alleged to have violated provisions of the Social Security Act (the Act) related to Medicare, Medicaid, HMO/Managed Care, and the Children’s Health Insurance Program. The FID system of records also contains the contact information and other identifying demographics of individuals alleged to have violated other criminal or civil statutes connected with the Act and the Act’s programs. Here, individuals are persons alleged to have abused the Act’s programs. (For example, an individual could be a person alleged to have rendered unnecessary services to Medicare beneficiaries or Medicaid recipients, over-used services, or engaged in improper billing.) They are persons whose activities have provided a substantial basis for criminal or civil prosecution, or who are identified as defendants in criminal prosecution cases.

II. Provisions of the Proposed Rule

In the May 25, 2007 Federal Register (72 FR 29289) we published a proposed rule that would exempt the ACTS, HITS, OPOS, and FID systems of records from subsection (c)(3), (d)(1) through (d)(4), (e)(4)(G) and (H), and (f) of the Privacy Act pursuant to 5 U.S.C. 552a(k)(2). These exemptions would apply only to the extent that information in a record is subject to exemption pursuant to 5 U.S.C. 552a(k)(2). We proposed that the ACTS, HITS, OPOS, and FID systems of records would be exempted from the following subsections for the reasons set forth below:

- Subsection (c)(3). Release of an accounting of disclosures to an individual who is the subject of an investigation could reveal the nature and scope of the investigation and could result in the altering or destruction of evidence, improper influencing of witnesses, and other evasive actions that could impede or compromise the investigation.
- Subsection (d)(1). Release of investigative records to an individual who is the subject of an investigation could interfere with pending or prospective law enforcement proceedings, constitute an unwarranted invasion of the personal privacy of third parties, reveal the identity of confidential sources, or reveal sensitive investigative techniques and procedures.
- Subsections (d)(2) through (d)(4). Amendment or correction of investigative records could interfere with pending or prospective law enforcement proceedings, or could impose an impossible administrative and investigative burden by requiring us to continuously retrograde our investigations in an attempt to resolve questions of accuracy, relevance, timeliness, and completeness.
- Subsection (e)(4)(G) and (H). Notifying an individual who is the subject of an investigation or a witness that a system of records contains information about him or her could reveal the nature and scope of the investigation and could result in the altering or destruction of evidence, improper influencing of witnesses, and other evasive actions that could impede or compromise the investigation.
- Subsection (f). Establishing procedures for notification, inspection or amendment of records, or appeals of denials of access to records would interfere with pending or prospective law enforcement proceedings, constitute an unwarranted invasion of the personal privacy of third parties, reveal the identity of confidential sources, or reveal sensitive investigative techniques. Furthermore, these actions could impose an impossible administrative and investigative burden by requiring us to continuously retrograde our investigations in an attempt to resolve questions of accuracy, relevance, timeliness, and completeness.

Accordingly, we proposed to amend 45 CFR 5b.11(b)(2)(iii) of the Privacy Act regulations by adding the following:

- A new paragraph (H) that exempts investigative materials compiled for law enforcement purposes from ACTS.
- A new paragraph (I) that exempts investigative materials compiled for law enforcement purposes from HITS.
- A new paragraph (J) that exempts investigative materials compiled for law enforcement purposes from OPOS.
- A new paragraph (K) that exempts investigative materials compiled for law enforcement purposes from FID.

III. Analysis of and Responses to Public Comments

We solicited and received two timely public comments on the May 25, 2007 proposed rule. The following is a summary of the comments and our responses.

Comment: One commenter believed that 45 CFR 5b.11(d) seems to allow the Department of Health and Human Services to disclose identities of sources who furnished information under an express promise of confidentiality.

Response: We do not disclose information that would reveal the identities of sources who furnish information under an express promise of confidentiality because the promise of confidentiality made to a witness is an agreement with that individual, and such disclosure would be both a violation of that agreement and counterproductive to law enforcement efforts, as it would discourage individuals from coming forward to supply information about alleged misconduct. 45 CFR 5b.11(b) gives the responsible Department official discretion to grant notification of access to a record in a system of records which is exempt under 45 CFR 5b.11(b), unless disclosure to the general public is otherwise prohibited by law. The department does not intend to exercise its discretion to disclose identifying...
information about sources who furnish information under an express promise of confidentiality.

Comment: Commenters requested that the exemptions be narrowed or clarified by defining the terms “investigative materials” and “law enforcement purposes,” including differentiating among kinds of records within each system that constitute “investigatory materials,” as well as describing agency uses that are not consistent with “law enforcement purposes.” A commenter suggested that CMS implement regulatory definitions, criteria, guidelines or other means to effectuate a confidentiality promise to an informant and to recognize whether or not one has been effectuated for purposes of compliance with subsection (k)(2) of the Privacy Act.

Response: We believe that with respect to clarifying what constitutes a confidentiality promise, we continue to rely upon the following language in subsection (k)(2) of the Privacy Act (5 U.S.C. 552a), which permits exemptions from certain subsections of the Privacy Act:

Investigatory material compiled for law enforcement purposes, other than material within the scope of subsection (j)(2) of this section [the Privacy Act]: Provided, however, That if any individual is denied any right, privilege, or benefit that he would otherwise be entitled by Federal law, or for which he would otherwise be eligible as a result of the maintenance of such material, such material shall be provided to such individual, except to the extent that the disclosure of such material would reveal the identity of a source who furnished information to the Government under an express promise that the identity of the source would be held in confidence, or, prior to the effective date of this section, [September 27, 1975] under an implied promise that the identity of the source would be held in confidence;

The (k)(2) exemption covers:

1. Material compiled for criminal investigative law enforcement purposes by an entity that does not have as its principal function the enforcement of criminal law and (2) investigative material compiled for law enforcement purposes that does not fall into the scope of the exemption under 5 U.S.C. 552(j)(2). The material must be investigative and compiled for some “law enforcement” purpose, such as a civil investigation, or a criminal investigation by an agency that does not perform as its principal function the enforcement of criminal law.

Further, since the information in the SORs at issue was collected on or after September 27, 1975, we believe that, with investigative material that would reveal the identity of a confidential source, only express promises to a source that his or her identity would not be revealed will be implicated here. An example of an express promise could occur when a source expressly requests that his or her identity not be revealed as a condition of furnishing the information, and CMS agrees to that condition and documents that promise in writing.

The fourth SORs at issue were established after September 27, 1975, the effective date of the Privacy Act, as follows:

1. The CMS Fraud Investigation Database (FID) was published under its previous name, “ICFJA Utilization Review Investigatory Files,” on December 29, 1988 (53 FR 52792) and republished under its current name on October 28, 2002 (67 FR 65795).
2. The Automated Survey Processing Environment (ASPE), Complaints/Incidents Tracking System (ACTS) was first established on August 22, 2003 (68 FR 50795).
3. The Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HTIS) was first established on July 6, 2005 (70 FR 38944).
4. The Organ Procurement Organizations System (OPOS) was first established on May 22, 2006 (71 FR 29336).

Further information about this exemption can be found in the Office of Management and Budget’s Privacy Act Guidelines, (see the July 9, 1975 Federal Register (40 FR 28972 through 28973)).

IV. Provisions of the Final Rule

After review of the public comments, we are finalizing the provisions of the proposed rule with minor technical changes. We are revising the paragraphs in § 5b.11(b)(2)(ii) so that the SORs are listed in chronological order by the date established.

V. Collection of Information Requirements

This final rule does not impose information collection and recordkeeping requirements. Consequently, it need not be reviewed by the Office of Management and Budget under the authority of the Paperwork Reduction Act of 1995 (44 U.S.C. 35).

VI. Regulatory Impact Statement

We have examined the impact of this rule 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 Act), the Unfunded Mandates Reform Act of 1995 (Pub. L. 104–4), and Executive Order 13132.

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 regulating actions with economically significant effects ($100 million or more in any one year or other substantial adverse economic effects) known as “major rules”. This rule does not meet the “major rule” criteria therefore we are not preparing an RIA.

The RFA requires agencies to analyze options for regulatory relief of small businesses. For purposes of the RFA, small entities include small businesses, nonprofit organizations, and small governmental jurisdictions. Most hospitals and most other providers and suppliers are small entities, either by nonprofit status or by having revenues of $6 million to $29 million in any one year. Individuals and States are not included in the definition of a small entity. We are not preparing an analysis for the RFA because we have determined that this rule will not have a significant economic impact on a substantial number of small entities.

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 604 of the RFA. For purposes of section 1102(b) of the Act, we define a small rural hospital as a hospital that is located outside of a Metropolitan Statistical Area and has fewer than 100 beds. We are not preparing an analysis for section 1102(b) of the Act because we have determined that this rule will not have a significant impact on the operations of a substantial number of small rural hospitals. This analysis must conform to the provisions of section 604 of the RFA. For purposes of section 1102(b) of the Act, we define a small rural hospital as a hospital that is located outside of a Metropolitan Statistical Area and has fewer than 100 beds. We are not preparing an analysis for section 1102(b) of the Act because we have determined that this rule will not have a significant economic impact on a substantial number of small rural hospitals. This analysis must conform to the provisions of section 604 of the RFA. For purposes of section 1102(b) of the Act, we define a small rural hospital as a hospital that is located outside of a Metropolitan Statistical Area and has fewer than 100 beds. We are not preparing an analysis for section 1102(b) of the Act because we have determined that this rule will not have a significant economic impact on a substantial number of small rural hospitals.
rule) that imposes substantial direct requirement costs on State and local governments, preempts State law, or otherwise has Federalism implications. Since this regulation does not impose any costs on State or local governments, the requirements of Executive Order 13132 are not applicable.

In accordance with the provisions of Executive Order 12866, this regulation was reviewed by the Office of Management and Budget.

List of Subjects for 45 CFR Part 5b Privacy.

For the reasons set forth in the preamble, the Department of Health and Human Services amends 45 CFR part 5b as set forth below:

PART 5b—PRIVACY ACT REGULATIONS

1. The authority citation for part 5b continues to read as follows:


2. Section 5b.11 is revised by adding paragraphs (b)(2)(ii)(H), (I), (J), and (K) to read as follows:

§ 5b.11 Exempt Systems

* * * * *

(b) * * * *

(2) * * * *

(ii) * * * *

(H) Investigative materials compiled for law enforcement purposes from the CMS Fraud Investigation Database (FID), HHS/CMS.

(I) Investigative materials compiled for law enforcement purposes from the Automated Survey Processing Environment (ASPEN) Complaints/Incidents Tracking System (ACTS), HHS/CMS.

(J) Investigative materials compiled for law enforcement purposes from the Health Insurance Portability and Accountability Act (HIPAA) Information Tracking System (HITS), HHS/CMS.

(K) Investigative materials compiled for law enforcement purposes from the Organ Procurement Organizations System (OPOS), HHS/CMS.

* * * * *


Kerry Weems,
Acting Administrator, Centers for Medicare & Medicaid Services.

Approved: June 13, 2008.

Michael O. Leavitt,
Secretary.

Editorial Note: This document was received at the Office of the Federal Register on September 16, 2008.

[FR Doc. E8–21909 Filed 9–25–08; 8:45 am]

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FEDERAL COMMUNICATIONS COMMISSION

47 CFR Part 101

[WT Docket No. 00–19; RM–9418; FCC 02–218]

Streamline Processing of Microwave Applications in the Wireless Telecommunications Services

AGENCY: Federal Communications Commission.

ACTION: Correcting amendment.

SUMMARY: In this document, the Federal Communications Commission corrects an inadvertent error that occurred when the Commission adopted final rules pertaining to Streamline Processing of Microwave Applications in the Wireless Telecommunications Services and Telecommunications Industry AssociationPetition for Rulemaking. These rules were published in the Federal Register on Friday, January 31, 2003 (68 FR 4953). Specifically, the error occurred in a table to the rules concerning directional antennas and compliance with antenna standards. As a result of this correction, the table will be amended as intended by the Commission.

DATES: Effective September 26, 2008.

FOR FURTHER INFORMATION CONTACT: John Schauble at 202–418–0797.

SUPPLEMENTARY INFORMATION: This is a correction to a summary of the Commission’s Report and Order in WT Docket No. 00–19, FCC 02–218, adopted on July 18, 2002 and released on July 31, 2002. The Report and Order streamlined, clarified and updated part 101 rules. These actions were to provide increased flexibility to licensees, ensure greater and more efficient use of spectrum bands regulated under part 101, and ensure that our Rules are consistent with international agreements.

Need for Correction

As published, the final rules contain an error in § 101.115 Directional Antennas, Antenna Standards Table. The Commission did not request that the listed entry to the Antenna Standards Table for the frequency of 10,550 to 10,680 MHz be omitted. This correction removes the listing that should have been omitted. The entry for 10,550 to 10,680 MHz, which was adopted in that proceeding, will remain.

List of Subjects in 47 CFR Part 101

Communications equipment, Marine safety, Radio, Reporting and recordkeeping requirements.

Accordingly, 47 CFR part 101 is amended by making the following correcting amendments:

PART 101—FIXED MICROWAVE SERVICES

1. The authority citation for part 101 continues to read as follows:

Authority: 47 U.S.C. 154, 303 unless otherwise noted.

§ 101.115 [Amended]

2. Section 101.115 is amended by removing the entry “10,550 to 10,680 MHz” for both Category A and B in the table following paragraph (b)(2).

Federal Communications Commission.

Marlene H. Dortch,
Secretary.

[FR Doc. E8–22721 Filed 9–25–08; 8:45 am]

BILLING CODE 6712–01–P