Agenda items are subject to change as priorities dictate.  
Contact Person for More Information: Price Connor, PhD, NIOSH Health Scientist, 1600 Clifton Road, NE., Mailstop E20, Atlanta, Georgia 30333, Telephone (404) 498–2511, Fax (404) 498–2571.  
The Director, Management Analysis and Services Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities for both the Centers for Disease Control and Prevention, and the Agency for Toxic Substances and Disease Registry.  
Dated: March 25, 2011.  
Andre Tyler,  
Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.  

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DEPARTMENT OF HEALTH AND HUMAN SERVICES  
Centers for Disease Control and Prevention  

Board of Scientific Counselors, Office of Public Health Preparedness and Response (BSC, OPHPR)  

In accordance with section 10 (a) (2) of the Federal Advisory Committee Act (Pub. L. 92–463), CDC announces the following meeting of the aforementioned committee:  

Time And Date: 3 p.m.–5 p.m., April 22, 2011.  
Place: CDC, 1600 Clifton Road, NE., Global Communications Center, Building 19, Rooms 245/246, Atlanta, Georgia 30333.  
This meeting is also accessible by Web Conference. Please contact the BSC Coordinator (see Contact Person for More Information) to obtain further instructions on how to participate by phone and online.  
Status: Open to the public limited only by the space available. The meeting room accommodates approximately 40 people. Participation by Web Conference is limited to 50 ports. Visitors to the CDC Roybal campus must be processed in accordance with established federal policies and procedures and should pre-register for the meeting as described in Additional Information for Visitors.  
Purpose: This Board is charged with providing advice and guidance to the Secretary, Department of Health and Human Services, the Assistant Secretary for Health, the Director, Centers for Disease Control and Prevention (CDC), and the Director, Office of Public Health Preparedness and Response (OPHPR), concerning strategies and goals for the programs and research within OPHPR, monitoring the overall strategic direction and focus of the OPHPR Divisions and Offices, and administration and oversight of peer review of OPHPR scientific programs. For additional information about the Board, please visit: http://emergency.cdc.gov/ cdcrepreparedness/science/counselors.asp.  
Matters To Be Discussed: The agenda items for this meeting include: A briefing to the Board on the findings from the external peer review of OPHPR’s Division of Emergency Operations followed by a vote on final recommendations, and updates from liaison representatives to the Board to share key highlights of their organization’s activities that are relevant to the OPHPR mission, as time permits.  
Additional Information for Visitors: All visitors to the CDC Roybal campus are required to present a valid form of picture identification issued by a State, federal or international government. To expedite the security clearance process for visitors to the CDC Roybal campus, all visitors must pre-register by submitting the following information by e-mail or phone (see Contact Person for More Information) no later than 12 noon (EDT) on Friday, April 15, 2011:  
• Full Name,  
• Organizational Affiliation,  
• Complete Mailing Address,  
• Citizenship, and  
• Phone Number or E-mail Address.  
For foreign nationals or non-U.S. citizens, pre-approval is required. Please contact the BSC Coordinator (see Contact Person for More Information) at least 7 days in advance of the posted pre-registration deadline for additional security requirements that must be met.  
Contact Person for More Information: Matthew Jennings, BSC Coordinator, OPHPR, CDC, 1600 Clifton Road, NE., Mailstop D–44, Atlanta, GA 30333, Telephone: (404) 639–7357; Facsimile: (404) 639–7077; E-mail: OPHPR.BSC.Questions@cdc.gov.  
The Director, Management Analysis and Service Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities for both the Centers for Disease Control and Prevention, and the Agency for Toxic Substances and Disease Registry.  
Dated: March 25, 2011.  
Andre Tyler,  
Acting Director, Management Analysis and Service Office, Centers for Disease Control and Prevention.  

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DEPARTMENT OF HEALTH AND HUMAN SERVICES  
Centers for Medicare & Medicaid Services  

[Document Identifier: CMS–10361]  
Agency Information Collection Activities: Proposed Collection; Comment Request  

AGENCY: Centers for Medicare & Medicaid Services, HHS.  

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS) 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.  

1. Type of Information Collection Request: Extension without change of a currently approved collection; Title of Information Collection: Request for Adjustment to the Medical Loss Ratio Standard for a State’s Individual Market; Use: Under section 2718 of the Public Health Service Act (PHS Act), a health insurance issuer (issuer) offering group or individual health insurance coverage must submit a report to the Secretary beginning in June of 2012 for calendar year 2011. The reported data allows for the calculation of an issuer’s medical loss ratio (MLR) by market (individual, small group, and large group) within each State in which the issuer conducts business. The PHS Act establishes an MLR standard for each market segment that issuers must meet. A health insurance issuer who fails to meet the MLR standard for a plan year must rebate to enrollees, on a pro rata basis, the difference between its MLR and the MLR standard.  

Section 2718(b)(1)(A)(ii) allows the Secretary to lower the 80% MLR standard in the individual market in a State if the application of the 80% MLR may destabilize the individual market in such State. An interim final rule (IFR) implementing the MLR was published on December 1, 2010 (75 FR 74865) and was modified by technical corrections on December 30, 2010 (75 FR 82277), which added Part 158 to Title 45 of the Code of Federal Regulations. The IFR is effective January 1, 2011. Under 45 CFR 158.301 (75 FR 74864, 74930), States requesting that HHS lower the MLR standard must submit information that supports their assertion that the individual market in their State may destabilize absent an adjustment to the MLR. Much of the information requested is currently only available at

BILLING CODE 4163–18–P

Federal Register / Vol. 76, No. 63 / Friday, April 1, 2011 / Notices 18221
the State level. HHIS must have such information in order to ascertain whether market destabilization has a high likelihood of occurring. Form Number: CMS–10361 (OMB Control No. 0938–1114); Frequency: Once; Affected Public: State, local or tribal governments; Number of Respondents: 20; Number of Responses: 20; Average Hours per Response: 185; Total Annual Hours: 3,700. (For policy questions regarding this collection, contact Carol Jimenez at (301) 492–4109. For all other issues regarding this collection, call (410) 786–1326.)

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS’ Web site at http://www.cms.hhs.gov/PaperworkReductionActof1995, or e-mail your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office at 410–786–1326.

In commenting on the proposed information collections please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in one of the following ways by May 31, 2011:

1. Electronically. You may submit your comments electronically to http://www.regulations.gov. Follow the instructions for “Comment or Submission” or “More Search Options” to find the information collection document(s) accepting comments.

2. By regular mail. You may mail written comments to the following address: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number, Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

Dated: March 28, 2011.

Martique Jones,
Director, Regulations Development Group, Division B, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2011–7742 Filed 3–31–11; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier CMS–265–11, CMS–381, and CMS–10123 and 10124]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS) 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.

1. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Independent Renal Dialysis Facility Cost Report; Use: The Independent Renal Dialysis Facility Cost Report, is filed annually by providers participating in the Medicare program to identify the specific items of cost and statistics of facility operation that independent renal dialysis facilities are required to report. The forms are revised in accordance with the End-Stage Renal Disease Prospective Payment System Final Rule published August 12, 2010 which implemented statutory requirements of the Medicare Improvements for Patients and Providers Act (MIPPA), enacted July 15, 2008. Additionally, the forms are revised to incorporate data previously reported on the Provider Cost Report Reimbursement Questionnaire, Form CMS–339, Form Number: CMS–265–94 (OMB#: 0938–0236); Frequency: Yearly; Affected Public: Business or other for-profits and Not-for-profit institutions; Number of Respondents: 2,960 Total Annual Responses: 2,960 Total Annual Hours: 3,700 (For policy questions regarding this collection contact Gail Duncan at 410–786–7278. For all other issues call 410–786–1326.)

2. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Identification of Extension Units of Medicare Approved Outpatient Physical Therapy/Outpatient Speech Pathology (OPT/OSP) Providers and Supporting Regulations in 42 CFR 485.701–485.729; Use: The collected information is used in conjunction with 42 CFR 485.701 through 485.729 governing the operation of providers of outpatient physical therapy and speech-language pathology services. The provider uses the form to report to the State survey agency extension locations that it has added since the date of last report. The form is used by the State survey agencies and by the CMS regional offices to identify and monitor extension locations to ensure their compliance with the Federal requirements for the providers of outpatient physical therapy and speech-language pathology services; Form Number: CMS–381 (OMB #: 0938– 0273); Frequency: Annually; Affected Public: Private Sector; Business or other for-profit and not-for-profit institutions; Number of Respondents: 2,960; Total Annual Responses: 2,960; Total Annual Hours: 740. (For policy questions regarding this collection contact Georgia Johnson at 410–786–6859. For all other issues call 410–786–1326.)

3. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Notice of Provider Non-Coverage (CMS–10123) and Detailed Explanation of Non-Coverage (CMS–10124); Use: The Notice of Medicare Provider Non-Coverage (CMS–10123) is used to inform fee-for-service Medicare beneficiaries of the determination that their provider services will end, and of their right to an expedited review of that determination. The Detailed Explanation of Non-Coverage (CMS–10124) is used to provide beneficiaries who request an expedited determination with detailed information of why the services should end. The revised Notice of Provider Non-Coverage and Detailed Explanation of Provider Non-Coverage will no longer require use of the beneficiary’s Medicare number as a patient identifier. Instead, when applicable, providers may use a number that helps to link the notice with a related claim. Form Number: CMS–10123 and 10124 (OMB# 0938–0953); Frequency: Occasionally; Affected Public: Business or other for-profit, not-for-profit institutions, and Individuals or households; Number of Respondents: