Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Key functions of the Advisory Board include providing advice on the development of probability of causation guidelines that have been promulgated by the Department of Health and Human Services (HHS) as a final rule; advice on methods of dose reconstruction, which have also been promulgated by HHS as a final rule; advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program; and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC).

In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to CDC. NIOSH implements this responsibility for CDC. The charter was issued on August 3, 2001, renewed at appropriate intervals, and will expire on August 3, 2013.

Purpose: The Advisory Board is charged with (a) providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the scientific validity and quality of dose reconstruction efforts performed for this program; and (c) upon request by the Secretary, HHS, advise the Secretary on whether there is a class of employees at any Department of Energy facility who were exposed to radiation but for whom it is not feasible to estimate their radiation dose, and on whether there is reasonable likelihood that such radiation doses may have endangered the health of members of this class. The Subcommittee for Dose Reconstruction Reviews was established to aid the Advisory Board in carrying out its duty to advise the Secretary, HHS, on dose reconstruction.

Matters To Be Discussed: The agenda for the Subcommittee meeting includes: discussion of dose reconstruction cases under review (sets 7–9 and claims with Category A findings from sets 10–13); dose reconstruction quality management and assurance activities, including overview of contractor quality management and internal dose reconstruction blind reviews; dose reconstruction issues from NIOSH 10-year review, including review of resource impact of possible changes to efficiency process and plans for claimant favorability analysis.

The agenda is subject to change as priorities dictate.

In the event an individual cannot attend, written comments may be submitted. Any written comments received will be provided at the meeting and should be submitted to the contact person below well in advance of the meeting.

Contact Person for More Information: Theodore Katz, Executive Secretary, NIOSH, CDC, 1600 Clifton Road, Mailstop E–20, Atlanta, Georgia 30333, Telephone (513) 535–6800, Toll Free 1 (800) CDC–INFO, Email oca@cdc.gov.

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.


Cathy Ramadei,
Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2012–11388 Filed 5–10–12; 8:45 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Council for the Elimination of Tuberculosis Meeting (ACET)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC) announces the following meeting for the aforementioned council:

Time and Date: 11:00 a.m.–3:30 p.m., June 5, 2012.

Place: This meeting is accessible by Web conference, Toll-free +1 (800) 369–1742, Toll +1 (517) 308–9167; Participant Code: ACET 2012. Participants can join the event directly at: https://www.mymeetings.com/nc/join/.

Audience passcode: ACET 2012.

Participants may join the meeting as follows:

Conference number: PW5343563.

Status: Open to the public limited only by Web conference. Participation by Web conference is limited by the number of 500 ports available.

Purpose: This council advises and makes recommendations to the Secretary of Health and Human Services, the Assistant Secretary for Health, and the Director, CDC, regarding the elimination of tuberculosis. Specifically, the Council makes recommendations regarding policies, strategies, objectives, and priorities; addresses the development and application of new technologies; and reviews the extent to which progress has been made toward eliminating tuberculosis.

Matters To Be Discussed: Agenda items include the following topics: (1) Developing a 5–year Strategic Plan for ACET; (2) Modeling TB Epidemiology; (3) Tuberculosis Outbreaks in Special Populations; (4) The Restructuring of United States Tuberculosis Program (TRUST)–How to Manage Budget Cuts; and (5) other tuberculosis-related issues.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Margie Scott-Coelh, Centers for Disease Control and Prevention, 1600 Clifton Road, NE, M/S E–07, Atlanta, Georgia 30333, telephone (404) 639–8317; Email: zkr@cdc.gov.

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.


Cathy Ramadei,
Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2012–11405 Filed 5–10–12; 8:45 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Safety and Occupational Health Study Section (SOHSS), National Institute for Occupational Safety and Health (NIOSH)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC) announces the following meeting of the aforementioned council:

Times and Dates: 8:00 a.m.–5:00 p.m., June 21, 2012 (Closed); 8:00 a.m.–5:00 p.m., June 22, 2012 (Closed).


Status: The meeting will be closed to the public in accordance with provisions set forth in Section 552(b)(4) and (6), Title 5 U.S.C., and the Determination of the Director, Management Analysis and Services Office, CDC, pursuant to Public Law 92–463.

Purpose: The Safety and Occupational Health Study Section will review, discuss, and evaluate grant application(s) received in response to the Institute’s standard grants review and funding cycles pertaining to research issues in occupational safety and health, and allied areas.

It is the intent of NIOSH to support broad-based research endeavors in keeping with the Institute’s program goals. This will lead to improved understanding and appreciation for the magnitude of the aggregate health burden associated with occupational injuries and illnesses, as well as to support more focused research projects, which will lead to improvements in the delivery of occupational safety and health services, and the prevention of work-related injury and illness. It is anticipated that research funded will promote these program goals.

Matters To Be Discussed: The meeting will convene to address matters related to the conduct of Study Section business and for the study section to consider safety and
occupational health-related grant applications.

These portions of the meeting will be closed to the public in accordance with provisions set forth in Section 552b(c)(4) and (6), Title 5 U.S.C., and the Determination of the Director, Management Analysis and Services Office, Centers for Disease Control and Prevention, pursuant to Section 10(d) Public Law 92–463.

Agenda items are subject to change as priorities dictate.

**Contact Person for More Information:** Price Connor, Ph.D., NIOSH Health Scientist, 2400 Executive Parkway, Mailstop E–20, Atlanta, Georgia 30345, 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.


**Cathy Ramadei,**
Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2012–11401 Filed 5–10–12; 8:45 am]

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Medicare & Medicaid Services**

[Document Identifier: CMS–643 and CMS–10425]

**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:** Hospice Survey and Deficiencies Report Form and Supporting Regulations. **Use:** CMS uses the information collected as the basis for certification decisions for hospices that wish to obtain or retain participation in the Medicare and Medicaid programs. The information is used by CMS regional offices, which have the delegated authority to certify Medicare facilities for participation, and by State Medicaid agencies, which have comparable authority under Medicaid. The information on the Hospice Survey and Deficiencies Report Form is coded for entry into the OSCAR system. The data is analyzed by the CMS regional offices and by the CMS central office components for program evaluation and monitoring purposes. The information is also available to the public upon request. **Form Number:** CMS–643 (OCN 0938–0379). **Frequency:** Yearly. **Affected Public:** State, Local, or Tribal Governments. **Number of Respondents:** 3,644. **Total Annual Responses:** 1,217. **Total Annual Hours:** 1,217. *(For policy questions regarding this collection contact Kim Roche at 410–786–3524. For all other issues call 410–786–1326.)*

2. **Type of Information Collection Request:** New collection. **Title of Information Collection:** Evaluation of Patient Satisfaction and Experience of Care for Medicare Beneficiaries with End-Stage Renal Disease (ESRD): Impact of the ESRD Prospective Payment System (PPS) and ESRD Quality Incentive Program (QIP). **Use:** The Medicare Prescription Drug Improvement, and Modernization Act of 2003 (MMA) required the Secretary of HHS to submit to Congress a report detailing the elements and features for the design and implementation of a bundled ESRD PPS, specifying that such a system should include the bundling of separately billed drugs, clinical laboratory tests, and other items “to maximum extent feasible”. The Medicare Improvements for Patients and Providers Act of 2008 (MIPPA) directed the Secretary of HHS to implement a payment system under which a single payment is made to a provider of services or a renal dialysis facility for renal dialysis services in lieu of any other payment. The ESRD PPS combines composite rate dialysis services with separately billable services under a single payment adjusted to reflect patient differences in resource needs or case-mix. The MIPPA also stipulated the development of quality incentives for the ESRD program. CMS has established the End-Stage Renal Disease Quality Incentive Program (ESRD QIP) to address this provision of the legislation.

In order to assess the impact of the Final Rule on ESRD beneficiary experiences, satisfaction, and health outcomes, CMS is requesting OMB approval to conduct data collection to obtain input on the effect of the Final Rule on our ESRD beneficiaries. The purposes of this data collection effort are to assess beneficiary satisfaction and experience of care in terms of access to services, quality of care, outcomes, and cost. This will be measured through telephone surveys with ESRD beneficiary and through interviews with key stakeholders in the renal health care community. The information obtained from both the beneficiary respondents and key stakeholders will be used both to provide an initial reporting of the ESRD PPS/QIP’s effects on beneficiary satisfaction and experience of care and to inform the Centers for Medicare & Medicaid Services (CMS) of the impact of the ESRD PPS/QIP on patient satisfaction and experience of care, including unintended consequences, for consideration of future modification of the programs.

**Form Number:** CMS–10425 (OCN: 0938–New). **Frequency:** Yearly. **Affected Public:** Individuals. **Number of Respondents:** 2,340. **Number of Responses:** 2,340. **Total Annual Hours:** 1,287. *(For policy questions regarding this collection contact Steve Blackwell at 410–786–6852. For all other issues 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 address at [http://www.cms.hhs.gov/PaperworkReductionActof1995](http://www.cms.hhs.gov/PaperworkReductionActof1995), or Email your request, including your address, phone number, OMB number, and CMS document identifier, to [Paperwork@cms.hhs.gov](mailto:Paperwork@cms.hhs.gov), or call the Reports Clearance Office on (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 July 10, 2012:

1. **Electronically.** You may submit your comments electronically to [http://www.regulations.gov](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