

Grant (NRSA) waivers, significant rebudgeting, 2nd and 3rd no cost extensions, and change of scope. These are all prior approvals as required by the NIH Grants Policy, and need to be reviewed and approved by the NHLBI. ePASS will provide a template to ensure that all specific points are addressed and documented in the official grant file. All information is submitted via the internet, tracked in ePASS, and the documentation will automatically be forwarded to the official grant file. The system will ensure that individuals

authorized by the grantee are submitting requests and that the appropriate NIH staff is receiving the requests. The requests will be template driven so that the grantee is including the minimally required information, thus eliminating the usual back and forth to obtain missing information. Forms will have automatic fill-in capability that will reduce typos in grant numbers and PI names, further reducing approval time. Reminders will be sent to NIH staff within ePASS based on roles to ensure timely responses to the grantee. The

system will facilitate email communication with applicants by automatic notifications when applications are received and when NIH has made a determination regarding a request (approval issued or request denied with explanation for denial).

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 470.

ESTIMATES OF HOUR BURDEN

Type of respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total annual hour burden
NHLBI Grantees	940	1	30/60	470
Totals	940	470

Dated: March 17, 2015.

Lynn Susulske,
NHLBI Project Clearance Liaison, National Institutes of Health.
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Subcommittee on Procedures Review, Advisory Board on Radiation and Worker Health (ABRWH or Advisory Board), 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 for the aforementioned subcommittee:

Time and Date: 11:00 a.m.–5:00 p.m., Eastern Standard Time, April 28, 2015.

Place: Audio Conference Call via FTS Conferencing.

Status: Open to the public. The public is welcome to submit written comments in advance of the meeting, to the contact person below. Written comments received in advance of the meeting will be included in the official record of the meeting. The public is also welcome to listen to the meeting by joining the teleconference at the USA toll-free, dial-in number, 1-866-659-0537 and the passcode is 9933701.

Background: The ABRWH 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 compensation program. Key functions of the ABRWH 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, 2015.

Purpose: The ABRWH 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, providing advice to 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 a reasonable likelihood that such radiation doses may have endangered the health of members of this class. The Subcommittee on Procedures Review was established to aid the ABRWH in carrying out its duty to advise the Secretary, HHS, on dose reconstructions. The Subcommittee on Procedures Review is responsible for overseeing, tracking, and participating in the reviews of all procedures used in the dose reconstruction process by the NIOSH Division of Compensation Analysis and Support (DCAS) and its dose reconstruction contractor (Oak Ridge Associated Universities—ORAU).

Matters for Discussion: The agenda for the Subcommittee meeting includes: discussion of procedures in the following ORAU and DCAS technical documents: Procedures for reconstructing dose associated with potential skin contamination, ORAU Team Technical Information Bulletin (OTIB) 0034 (“Internal Dose Coworker Data for X-10”), OTIB 0054 (“Fission and Activation Product Assignment for Internal Dose-Related Gross Beta and Gross Gamma Analyses”), OTIB 0082 (“Dose Reconstruction Method for Chronic Lymphocytic Leukemia”), Update on Review of ORAU Team Report 0053 (“Stratified Co-Worker Sets”); and a continuation of the comment-resolution process for other dose reconstruction procedures under review by the Subcommittee.

The agenda is subject to change as priorities dictate.

Contact Person For More Information: Theodore Katz, Designated Federal Officer, NIOSH, CDC, 1600 Clifton Road NE., Mailstop E-20, Atlanta Georgia 30333, Telephone (513) 533-6800, Toll Free 1(800)CDC-INFO, Email ocas@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.

Elaine L. Baker,

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

Advisory Committee to the Director (ACD), Centers for Disease Control and Prevention (CDC)

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 committee:

Time and Date: 8:30 a.m.–3:00 p.m., EDT, April 23, 2015.

Place: CDC, Building 19, Auditorium B3, 1600 Clifton Road NE., Atlanta, Georgia 30333.

Status: Open to the public, limited only by the space and phone lines available. The meeting room accommodates approximately 50 people. Advance registration for in-person participation is required by April 16, 2015. The public is welcome to participate during the public comment period, which is tentatively scheduled from 2:35 p.m. to 2:40 p.m. This meeting will also be available by teleconference. Please dial (877) 930-8819 and enter code 1579739.

Web links: Windows Media: <http://wm.onlinevideoservice.com/CDC1>.

Flash: <http://www.onlinevideoservice.com/clients/CDC/?mount=CDC3>.

If you are unable to connect using the link, copy and paste the link into your web browser. For technical support please call: (404) 639-3737.

Purpose: The Advisory Committee to the Director, CDC, shall advise the

Secretary, HHS, and the Director, CDC, on policy and broad strategies that will enable CDC to fulfill its mission of protecting health through health promotion, prevention, and preparedness. The committee recommends ways to prioritize CDC's activities, improve results, and address health disparities. It also provides guidance to help CDC work more effectively with its various private and public sector constituents to make health protection a practical reality.

Matters for Discussion: The Advisory Committee to the Director will receive updates from the State, Tribal, Local and Territorial Subcommittee; the Health Disparities Subcommittee, the Global Workgroup, the Internal and External Laboratory Safety Workgroups, and the Public Health—Health Care Collaboration Workgroup, the Ebola response, global health security, recent viral outbreaks, antimicrobial resistance, as well as an update from the CDC Director.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Carmen Villar, MSW, Designated Federal Officer, ACD, CDC, 1600 Clifton Road NE., M/S D-14, Atlanta, Georgia 30333. Telephone (404) 639-7158, Email: GHickman@cdc.gov. The deadline to register for in-person attendance at this meeting is April 16, 2015. To register, please send an email to GHickman@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.

Elaine L. Baker,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention (CDC).

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifiers: CMS-10549]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

ACTION: Notice.

SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS' intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (PRA), federal agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, and to allow a second opportunity for public comment on the notice. Interested persons are invited to send comments regarding the 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.

DATES: Comments on the collection(s) of information must be received by the OMB desk officer by *May 4, 2015*:

ADDRESSES: When commenting on the proposed information collections, please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be received by the OMB desk officer via one of the following transmissions: OMB, Office of Information and Regulatory Affairs, Attention: CMS Desk Officer, Fax Number: (202) 395-5806 *OR*, Email: OIRA_submission@omb.eop.gov.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of following:

1. Access CMS' Web site address at <http://www.cms.hhs.gov/PaperworkReductionActof1995>.
2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov.
3. Call the Reports Clearance Office at (410) 786-1326.

FOR FURTHER INFORMATION CONTACT: Reports Clearance Office at (410) 786-1326.

SUPPLEMENTARY INFORMATION: Under the Paperwork Reduction Act of 1995 (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. The term "collection of