PERSON TO CONTACT FOR INFORMATION:

Judith Ingram, Press Officer, Telephone: (202) 694–1220.

Shelley E. Garr,

Deputy Secretary.

[FR Doc. 2015-12064 Filed 5-14-15; 11:15 am]

BILLING CODE 6715-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Mine Safety and Health Research Advisory Committee, National Institute for Occupational Safety and Health (MSHRAC, 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 committee:

Time and Date: 9:30 a.m.–2:30 p.m. EDT, June 8, 2015.

Place: Teleconference and Webinar. Status: Open to public, limited only by the space available on the webinar system, which accommodates a maximum of 100 people. If you wish to attend by phone or webinar, please contact Marie Chovanec by email at MChovanec@cdc.gov or by phone at 412–386–5302 at least 3 days in advance.

Purpose: This committee is charged with providing advice to the Secretary, Department of Health and Human Services; the Director, CDC; and the Director, NIOSH, on priorities in mine safety and health research, including grants and contracts for such research, 30 U.S.C. 812(b)(2), Section 102(b)(2).

Matters for Discussion: The meeting will focus on mining safety and health research projects and outcomes, including refuge alternatives, rock dust, silica exposures, metal mine ground control, and mining survey. The meeting will also include updates from the National Personal Protective Technology Laboratory and the Division of Respiratory Disease Studies; and committee discussion on the program portfolio.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Jeffrey H. Welsh, Executive Secretary, MSHRAC, NIOSH, CDC, 626 Cochrans Mill Road, telephone (412) 386–4040, fax (412) 386–6614.

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.

[FR Doc. 2015-11889 Filed 5-15-15; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Disease, Disability, and Injury Prevention and Control Special Emphasis Panel (SEP): Initial Review

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 a meeting for the initial review of applications in response to Funding Opportunity Announcement Number, (FOA) DP15–008, Health Promotion and Disease Prevention Research Centers: Special Interest Project Competitive Supplements (SIPS).

Time and Date: 11:00 a.m.—6:00 p.m., EDT, June 11, 2015 (Closed).

Place: Teleconference.

Status: 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, CDC, pursuant to Public Law 92– 463.

Matters For Discussion: The meeting will include the initial review, discussion, and evaluation of applications received in response to "Health Promotion and Disease Prevention Research Centers: Special Interest Project Competitive Supplements (SIPS)", FOA DP15–008.

Contact Person For More Information: Brenda Colley Gilbert, Ph.D., M.S.P.H., Director, Extramural Research Program Operations and Services, CDC, 4770 Buford Highway NE., Mailstop F–80, Atlanta, Georgia 30341, Telephone: (770) 488–6295, BJC4@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.

[FR Doc. 2015-11888 Filed 5-15-15; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Community Living

Notice of Intent To Award a Single Source Non-Competing Continuation Cooperative Agreement for Two National Activities Grant Projects Under Section 6 of the Assistive Technology Act of 1998, as Amended (AT Act)

AGENCY: Administration for Community Living, Department of Health and Human Services.

ACTION: Notice.

SUMMARY: The Administration for Community Living (ACL) is transitioning the Rehabilitation Engineering and Assistive Technology Society of North America (RESNA) Catalyst Project Assistive Technology Technical Assistance Center (AT TA Center) and the University of New Hampshire Institute on Disability Assistive Technology Public Internet Site (National AT Web site) to ACL as a result of the Workforce Opportunity Improvement Act (Pub. L. 113–128) signed by President Obama in July 2014.

The RÉSNA Catalyst Project is a national training and technical assistance center for assistive technology programs that provides comprehensive information and state-specific, regional and national resources to entities funded under Sections 4 and 5 of the *AT Act* to improve the implementation and effectiveness of their programs, and to provide appropriate technical assistance and training to entities not funded under the *AT Act* to improve awareness of and access to assistive technology.

The University of New Hampshire Institute on Disability supports the renovation, updating, and maintenance of an accessible National AT Web site that provides the public comprehensive, up-to-date information on accommodating individuals with disabilities and resources related to assistive technology, including but not limited to programs under the *AT Act*.

The RESNA Catalyst Project and New Hampshire National AT Web site previously operated through a cooperative agreement with the U.S. Department of Education, Rehabilitation Services Administration. The Department of Health and Human Services is currently transitioning programs under the *AT Act* to ACL.

DATES: Estimated Project Period— September 30, 2015 through September 30, 2016

SUPPLEMENTARY INFORMATION:

Program Name: Assistive Technology National Activities.

Award Amount: \$640,000 to Rehabilitation Engineering and Assistive Technology Society of North America; \$100,000 to University of New Hampshire Institute on Disability.

Project Period: 9/30/2015 to 9/30/2016

Award Type: Cooperative Agreement.

Statutory Authority: This program is authorized under Section 6 of the *Assistive Technology Act of 1998*, as amended (29 U.S.C. 3005)

Catalog of Federal Domestic Assistance (CFDA) Number: 93.464 Discretionary Projects

Program Description

The purpose of the National Activities cooperative agreements with RESNA and the University of New Hampshire is to continue existing activities designed to support and improve the administration of the *AT Act*. The grantees will continue to use both traditional and innovative approaches that will assist individuals and entities through information dissemination and provide state-specific, regional and national training and technical assistance concerning assistive technology.

Justification: ACL is currently working on transitioning the Assistive Technology National Activities program to ACL. To ensure uninterrupted continuation of the grant goals and objectives, ACL plans to issue a one year non-competing award to both RESNA and the University of New Hampshire.

FOR FURTHER INFORMATION CONTACT: For further information or comments regarding this action, contact Lori Gerhard, U.S. Department of Health and Human Services, Administration for Community Living, Center for Consumer Access and Self-Determination, Office of Integrated Programs, One Massachusetts Avenue NW., Washington, DC 20001; telephone (202) 357–3443; fax (202) 357–3469; email Lori.Gerhard@acl.hhs.gov.

Dated: May 13, 2015.

Kathy Greenlee,

Administrator and Assistant Secretary for Aging.

[FR Doc. 2015–11961 Filed 5–15–15; 8:45 am] BILLING CODE 4154–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2015-N-0012]

Cooperative Agreement to International Council for Harmonization of Technical Requirements for Pharmaceuticals for Human Use

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
Administration (FDA) is announcing the availability of grant funds for the support of the International Council for Harmonization of Technical
Requirements for Pharmaceuticals for Human Use (ICH). The goal of the ICH is to bring together leading global drug regulatory agencies and pharmaceutical manufacturer associations to achieve greater harmonization of technical standards to ensure that safe, effective, and high-quality medicines are developed and registered in the most resource-efficient manner.

DATES: The application due date is September 30, 2015. The expiration date is October 1, 2015.

ADDRESSES: Submit electronic applications to: *http://www.grants.gov*. For more information, see section III of the **SUPPLEMENTARY INFORMATION** section of this notice.

FOR FURTHER INFORMATION CONTACT:

Tracy Porter, Office of Strategic Programs, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, Rm. 1173, Silver Spring, MD 20993, 301–796–7789, Tracy.Porter@fda.hhs.gov; or Lisa Ko, Office of Acquisitions and Grants Services, Food and Drug Administration, 5630 Fishers Lane, Rm. 2037, Rockville, MD 20857, 240–402–7592, Lisa.Ko@fda.hhs.gov.

For more information on this funding opportunity announcement (FOA) and to obtain detailed requirements, please refer to the full FOA located at http://www.grants.gov. Search by Funding Opportunity Number: RFA-FD-15-014. SUPPLEMENTARY INFORMATION:

I. Funding Opportunity Description RFA-FD-15-014

93.103

A. Background

1. Authority

FDA activities to increase the harmonization of regulatory requirements to ensure that safe, effective, and high quality medicines are developed and registered in the most resource-efficient manner are authorized by 21 U.S.C. 383(c) and 393(b).

2. Program Background

The ICH is a globally unique venue with the capability of bringing together the regulatory authorities and pharmaceutical industry to discuss scientific and technical aspects of drug registration. ICH is a programmatic global priority for FDA to achieve its identified strategic priority of globalization. Working through its Center for Drug Evaluation and Research (CDER) and Center for Biologics Evaluation and Research, FDA has played a leading role in ICH since its inception in 1990. ICH, founded to harmonize drug regulatory standards between three regions, the United States, the European Union, and Japan, has gradually evolved to respond to the increasingly global face of drug development, so that the benefits of international harmonization for better global health can be realized worldwide. ICH's mission is to achieve greater harmonization to ensure that safe, effective, and high-quality medicines are developed and registered in the most resource-efficient manner.

Over the past 2 years, FDA played a leadership role in transforming ICH to meet the challenges of 21st century standards development while firmly positioning ICH future work to continue the focus on technical standards harmonization informed by relevant expertise from regulatory agencies and regulated industry. This effort has included: (1) Establishing ICH as a formal legal entity in the form of a nonprofit association under Swiss law; (2) expanding the opportunities for formal participation of other drug regulatory authorities beyond the three founding regions via the ICH Assembly; and (3) ensuring adequate and predictable funding for the ICH harmonization work (which is also critical to FDA's mission).

FDA remains an ICH founding member and completely committed to ICH success as a science-based standards development venue to ensure harmonization globally for safe, effective, and high-quality medicines. As exemplified in the past 25 years, FDA leadership and participation is an