quarter to the Payment Management Services, HHS at: http://www.dpm.psc.gov. It is recommended that the applicant also send a copy of the FFR (SF–425) report to the Grants Management Specialist. Failure to submit timely reports may cause a disruption in timely payments to the organization.

Grantees are responsible and accountable for accurate information being reported on all required reports: the Progress Reports and Federal Financial Report.

C. Federal Subaward Reporting System (FSRS)

This award may be subject to the Transparency Act subaward and executive compensation reporting requirements of 2 CFR part 170. The Transparency Act requires the OMB to establish a single searchable database, accessible to the public, with information on financial assistance awards made by Federal agencies. The Transparency Act also includes a requirement for recipients of Federal grants to report information about first-tier subawards and executive compensation under Federal assistance awards.

IHS has implemented a Term of Award into all IHS Standard Terms and Conditions, NoAs and funding announcements regarding the FSRS reporting requirement. This IHS Term of Award is applicable to all IHS grant and cooperative agreements issued on or after October 1, 2010, with a $25,000 subaward obligation dollar threshold met for any specific reporting period. Additionally, all new (discretionary) IHS awards (where the project period is made up of more than one budget period) and where: (1) The project period start date was October 1, 2010 or after and (2) the primary awardee will have a $25,000 subaward obligation dollar threshold during any specific reporting period will be required to address the FSRS reporting. For the full IHS award term implementing this requirement and additional award applicability information, visit the DGM Grants Policy Web site at: https://www.ihs.gov/dgm/index.cfm?module=dsp_dgm_policy_topics.

Telecommunication for the hearing impaired is available at: TTY (301) 443–6394.

VII. Agency Contacts

1. Questions on the programmatic issues may be directed to:
Ms. Nancy Bill, Program Manager,
Injury Prevention Program, IHS, 801 Thompson Ave, TMP Suite 610,
Rockville, MD 20852, Phone: (301) 443–0105, Fax: (301) 443–7538, E-Mail: Nancy.Bill@ihs.gov

2. Questions on grants management and fiscal matters may be directed to: Pallop Chareonvootitam, Senior Grant Management Specialist, 801 Thompson Avenue, TMP Suite 360–78, Rockville, MD 20852, Phone: (301) 443–2195; or the DGM main line (301) 443–5204, Fax: (301) 443–9602, E-Mail: Pallop.Chareonvootitam@ihs.gov

3. Questions on systems matters may be directed to:
Paul Gettys, Grant Systems Coordinator, 801 Thompson Avenue, TMP Suite 360, Rockville, MD 20852, Phone: (301) 443–2114; or the DGM main line (301) 443–5204, Fax: (301) 443–9602, E-Mail: Paul.Gettys@ihs.gov

VIII. Other Information

The Public Health Service strongly encourages all cooperative agreement and contract recipients to provide a smoke-free workplace and promote the non-use of all tobacco products. In addition, Public Law 103–227, the Pro-Children Act of 1994, prohibits smoking in certain facilities (or in some cases, any portion of the facility) in which regular or routine education, library, day care, health care, or early childhood development services are provided to children. This is consistent with the HHS mission to protect and advance the physical and mental health of the American people.

Dated: April 3, 2015,
Robert G. McSwain,
Acting Director, Indian Health Service.

[FR Doc. 2015–08605 Filed 4–13–15; 8:45 am]
BILLING CODE 4165–16–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review: Notice of Meeting

Pursuant to section 10(a) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of a meeting of the Center for Scientific Review Advisory Council.

The meeting will be open to the public, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the Contact Person listed below in advance of the meeting.

Name of Committee: Center for Scientific Review Advisory Council.

Date: May 18, 2015.
Time: 8:30 a.m. to 3:00 p.m.

Agenda: Provide advice to the Director. Center for Scientific Review (CSR), on matters related to planning, execution, conduct, support, review, evaluation, and receipt and referral of grant applications at CSR.

Place: National Institutes of Health, 6701 Rockledge Drive, Room 3091, Bethesda, MD 20822.

Contact Person: Rene Etcheberrrigay, MD. Deputy Director, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 3030, MSC 7776, Bethesda, MD 20892, (301) 435–1111, etcheber@csr.nih.gov.

Any interested person may file written comments with the committee by forwarding the statement to the Contact Person listed on this notice. The statement should include the name, address, telephone number and when applicable, the business or professional affiliation of the interested person.

In the interest of security, NIH has instituted stringent procedures for entrance into NIH buildings. Visitors will be asked to show one form of identification (for example, a government-issued photo ID, driver’s license, or passport) and to state the purpose of their visit.

Information is also available on the Institute’s/Center’s home page: http://public.csr.nih.gov/about csr/CSRAdvisoryCommittee.aspx, where an agenda and any additional information for the meeting will be posted when available.


Dated: April 8, 2015.
Carolyn A. Baum,
Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2015–08459 Filed 4–13–15; 8:45 am]
BILLING CODE 4140–01P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Interagency Coordinating Committee on the Validation of Alternative Methods; Notice of Public Meeting; Request for Public Input

SUMMARY: The Interagency Coordinating Committee on the Validation of Alternative Methods (ICCVAM) will hold a public forum to share information and facilitate direct communication of ideas and suggestions from stakeholders. Interested persons may attend in person or remotely. Time will be set aside for public statements and questions on the topics discussed.
Registration is requested for both public attendance and oral statements, and required for remote access. Information about the meeting and registration is available at http://ntp.niehs.nih.gov/go/iccvmforum-2015.

DATES: Meeting: May 27, 2015, 9:00 a.m. to approximately 12:00 p.m. Eastern Daylight Time (EDT).

Registration for Onsite Meeting: Deadline is May 15, 2015.

Registration for Webcast: Deadline is May 27, 2015

Submission of Oral Public Statements: Deadline is May 15, 2015.

ADDRESSES:
Meeting Location: William H. Natcher Conference Center, National Institutes of Health, Bethesda, MD 20892.

FOR FURTHER INFORMATION CONTACT: Dr. Warren S. Casey, Director, National Toxicology Program Interagency Center for the Evaluation of Alternative Toxicological Methods (NICEATM); email: warren.casey@nih.gov; telephone: (919) 316–4729.

SUPPLEMENTARY INFORMATION:
Background: ICCVAM promotes the development and validation of chemical safety testing methods that protect human health and the environment while replacing, reducing, or refining animal use.

ICCVAM’s goals include promotion of national and international partnerships between governmental and nongovernmental groups, including academia, industry, advocacy groups, and other key stakeholders. To foster these partnerships ICCVAM initiated annual public forums in 2014 to share information and facilitate direct communication of ideas and suggestions from stakeholders (79 FR 25136).

The second of these forums will be held on May 27, 2015, at the National Institutes of Health (NIH) in Bethesda, MD. The meeting will begin with presentations by NICEATM and ICCVAM members on current activities related to the development and validation of alternative test methods and approaches for assessing acute systemic toxicity, endocrine activity, vaccine safety, and skin sensitization potential, as well as updates on ICCVAM processes. Following each presentation, there will be an opportunity for participants to ask questions of the ICCVAM members. Instructions for submitting questions will be provided to remote participants prior to the webcast. The agenda also includes time for participants to make public oral statements to inform ICCVAM on topics relevant to its mission and current activities.

Preliminary Agenda and Other Meeting Information: The preliminary agenda, ICCVAM roster and other background materials, and public statements submitted prior to the meeting will be posted at http://ntp.niehs.nih.gov/go/iccvmforum-2015 to allow remote participation. Public statements will be distributed to NICEATM and ICCVAM members. Interested individuals are encouraged to visit this Web page to stay abreast of the most current meeting information.

Meeting and Registration: This meeting is open to the public with time scheduled for oral public statements and for questions following ICCVAM’s and NICEATM’s presentations. The public may attend the meeting at NIH, where attendance is limited only by the space available, or view remotely by webcast. Those planning to attend the meeting in person are encouraged to register at http://ntp.niehs.nih.gov/go/iccvmforum-2015 by May 15, 2015, to facilitate planning for appropriate meeting space. Those planning to view the webcast must register at http://ntp.niehs.nih.gov/go/iccvmforum-2015 by May 27, 2015. The URL for the webcast will be provided in the email confirming registration.

Visitor and security information for visitors to NIH is available at http://www.nih.gov/about/visitor/index.htm. Individuals with disabilities who need accommodation to participate in this event should contact Dr. Elizabeth Maull at phone: (919) 316–4668 or email: maull@niehs.nih.gov. TTY users should contact the Federal TTY Relay Service at 800–877–8339. Requests should be made at least five business days in advance of the event.

Request for Oral Public Statements: Time will be allotted during the meeting for oral public statements with associated slides relevant to ICCVAM’s mission and current activities. The number and length of presentations may be limited based on available time. Submitters will be identified by their name and affiliation and/or sponsoring organization, if applicable. Persons submitting public statements and/or associated slides should include their name, affiliation (if any), mailing address, telephone, email, and sponsoring organization (if any) with the document.

Persons wishing to present oral statements are encouraged to indicate the topic(s) on which they plan to speak on their registration form. They should also provide a copy of their statement to Dr. Elizabeth Maull at email: maull@niehs.nih.gov by May 15, 2015, to allow time for review by NICEATM and ICCVAM and posting to the meeting page prior to the forum. Written statements may supplement and expand the oral presentation.

Registration for oral public statements will be available onsite, although onsite registration and time allotted for these statements may be limited based on the number of individuals who register to make statements and available time. If registering onsite and reading from written text, please bring 20 copies of the statement for distribution and to supplement the record.

In addition to in-person oral statements at the meeting, public statements may be presented by teleconference line. Directions for accessing the meeting by teleconference line will be provided to registered participants prior to the meeting date.

Responses to this notice are voluntary. No proprietary, classified, confidential, or sensitive information should be included in statements submitted in response to this notice or presented during the meeting. This request for input is for planning purposes only and is not a solicitation for applications or an obligation on the part of the U.S. Government to provide support for any ideas identified in response to the request. Please note that the U.S. Government will not pay for the preparation of any information submitted or for its use of that information.

Background Information on ICCVAM and NICEATM: ICCVAM is an interagency committee composed of representatives from 15 federal regulatory and research agencies that require, use, generate, or disseminate toxicological and safety testing information. ICCVAM conducts technical evaluations of new, revised, and alternative safety testing methods and integrated testing strategies with regulatory applicability and promotes the scientific validation and regulatory acceptance of testing methods that both more accurately assess the safety and hazards of chemicals and products and replace, reduce, or refine (enhance animal well-being and minimize or prevent pain and distress) animal use.

The ICCVAM Authorization Act of 2000 (42 U.S.C. 285j–3) establishes ICCVAM as a permanent interagency committee of the NIEMS and provides the authority for ICCVAM involvement in activities relevant to the development of alternative test methods. ICCVAM acts to ensure that new and revised test methods are validated to meet the needs of Federal agencies, increase the efficiency and effectiveness of federal
agency test method review, and optimize utilization of scientific expertise outside the federal Government. Additional information about ICCVAM can be found at http://ntp.niehs.nih.gov/go/iccvam.

NICEATM administers ICCVAM, provides scientific and operational support for ICCVAM-related activities, and conducts and publishes analyses and evaluations of data from new, revised, and alternative testing approaches. NICEATM and ICCVAM work collaboratively to evaluate new and improved testing approaches applicable to the needs of U.S. federal agencies. NICEATM and ICCVAM welcome the public nomination of new, revised, and alternative testing approaches for validation studies and technical evaluations. Additional information about NICEATM can be found at http://ntp.niehs.nih.gov/go/niceatm.

Dated: April 6, 2015.

John R. Bucher,
Associate Director, National Toxicology Program.

[FR Doc. 2015–08528 Filed 4–13–15; 8:45 am]
BILLING CODE 4140–01–P

DEPARTMENT OF HOMELAND SECURITY

Transportation Security Administration
[Docket No. TSA–2002–11602]

Intent To Request Renewal From OMB of One Current Public Collection of Information: Security Programs for Foreign Air Carriers

AGENCY: Transportation Security Administration, DHS.

ACTION: 60-Day notice.

SUMMARY: The Transportation Security Administration (TSA) invites public comment on one currently approved Information Collection Request (ICR), Office of Management and Budget (OMB) control number 1652–0005, abstracted below that we will submit to OMB for renewal in compliance with the Paperwork Reduction Act (PRA). The ICR describes the nature of the information collection and its expected burden. This information collection is mandatory for foreign air carriers and must be submitted prior to entry into the United States.

DATES: Send your comments by June 15, 2015.

ADDRESSES: Comments may be emailed to TSAPRA@tsa.dhs.gov or delivered to the TSA PRA Officer, Office of Information Technology (OIT), TSA–11, Transportation Security Administration, 601 South 12th Street, Arlington, VA 20598–6011.

FOR FURTHER INFORMATION CONTACT: Christina A. Walsh at the above address, or by telephone (571) 227–2062.

SUPPLEMENTARY INFORMATION:

Comments Invited

In accordance with the Paperwork Reduction Act of 1995 (44 U.S.C. 3501 et seq.), an agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a valid OMB control number. The ICR documentation is available at http://www.reginfo.gov. Therefore, in preparation for OMB review and approval of the following information collection, TSA is soliciting comments to—

(1) Evaluate whether the proposed information requirement is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;

(2) Evaluate the accuracy of the agency’s estimate of the burden;

(3) Enhance the quality, utility, and clarity of the information to be collected; and

(4) Minimize the burden of the collection of information on those who are to respond, including using appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

Information Collection Requirement

OMB Control Number 1652–0005; Security Programs for Foreign Air Carriers, 49 CFR part 1546. TSA uses the information collected to determine compliance with 49 CFR part 1546 and to ensure passenger safety by monitoring foreign air carrier security procedures. Foreign air carriers must carry out security measures to provide for the safety of persons and property traveling on flights provided by the foreign air carrier against acts of criminal violence and air piracy, and the introduction of explosives, incendiaries, or weapons aboard an aircraft. This information collection is mandatory for foreign air carriers and must be submitted prior to entry into the United States. The information TSA collects includes identifying information on foreign air carriers’ flight crews and passengers. Specifically, TSA requires foreign air carriers to submit the following information: (1) A master crew list of all flight and cabin crew members flying to and from the United States; (2) the flight crew list on a flight-by-flight basis; and (3) passenger information on a flight-by-flight basis. Foreign air carriers are required to provide this information via electronic means. On June 19, 2014, TSA removed the previous security program requirement that foreign air carriers submit information regarding the amount of cargo screened because all foreign air carriers are required to screen 100% of cargo.

Additionally, foreign air carriers must maintain these records, as well as training records for crew members and individuals performing security-related functions, and make them available to TSA for inspection upon request. TSA will continue to collect information to determine foreign air carrier compliance with other requirements of 49 CFR part 1546. TSA estimates that there will be approximately 170 respondents to the information collection, with an annual burden estimate of 1,029,010 hours.

Dated: April 6, 2015.

Christina A. Walsh,
TSA Paperwork Reduction Act Officer, Office of Information Technology.

[FR Doc. 2015–08562 Filed 4–13–15; 8:45 am]
BILLING CODE 9110–05–P

DEPARTMENT OF HOMELAND SECURITY

Transportation Security Administration


AGENCY: Transportation Security Administration, DHS.

ACTION: 30-day notice.

SUMMARY: This notice announces that the Transportation Security Administration (TSA) has forwarded the Information Collection Request (ICR), Office of Management and Budget (OMB) control number 1652–0041 abstracted below, to OMB for review and approval of an extension of the currently-approved collection under the Paperwork Reduction Act (PRA). The ICR describes the nature of the information collection and its expected burden. TSA published a Federal Register notice, with a 60-day comment period, describing the collection of information on December 29, 2014, 79 FR 78099. The collection involves the submission of numerical ratings and written comments about the quality of...