The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Member Conflict: Immune Mechanisms.

Date: March 4, 2016.

Time: 3:00 p.m. to 4:00 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda MD 20892.

(Direct Mail)

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meetings.

Name of Committee: National Cancer Institute Board of Scientific Advisors.

Date: March 30, 2016.

Time: 8:30 a.m. to 12:00 p.m.

Agenda: Director’s Report: Ongoing and New Business; Reports of Program Review Group(s): Budget Presentations; Reports of Special Initiatives; RFA and RFP Concept Reviews; and Scientific Presentations.

Place: National Institutes of Health, 31 Center Drive, Building 31, C-Wing, 6th Floor, Room 10, Bethesda, MD 20892.

Contact Person: Paulette S. Gray, Ph.D., Director, Division of Extramural Activities, National Cancer Institute—Shady Grove, National Institutes of Health, 9609 Medical Center Drive, 7th Floor, Rm. 7W444, Bethesda, MD 20892, 240–276–6340, grayp@mail.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 onto the NIH campus. All visitor vehicles, including taxicabs, hotel, and airport shuttles will be inspected before being allowed on campus. 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://deainfo.nci.nih.gov/advisory/bsa/bsa.htm, where an agenda and any additional information for the meeting will be posted when available.

(Catalogue of Federal Domestic Assistance Program Nos. 93.392, Cancer Construction; 93.393, Cancer Cause and Prevention Research; 93.394, Cancer Detection and Diagnosis Research; 93.395, Cancer Treatment Research; 93.396, Cancer Biology Research; 93.397, Cancer Centers Support; 93.398, Cancer Research Manpower; 93.399, Cancer Control, National Institutes of Health, HHS)


Melanie J. Gray.
Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2016–04103 Filed 2–25–16; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Draft National Toxicology Program Monograph on Immunotoxicity Associated With Exposure to Perfluorooctanoic Acid or Perfluorooctane Sulfonate; Availability of Document; Request for Comments; Notice of Meeting

SUMMARY: The notice announces a meeting to peer review the draft NTP monograph on immunotoxicity associated with exposure to perfluorooctanoic acid (PFOA) or perfluorooctane sulfonate (PFOS). The Office of Health Assessment and Translation, Division of the National Toxicology Program (DNTP), National Institute of Environmental Health Sciences (NIEHS), prepared the draft NTP monograph. The peer review meeting is open to the public. Registration is requested for both public attendance and oral comment and required to access the webcast. Information about the meeting and registration are available at http://ntp.niehs.nih.gov/go/36639.

DATES: Meeting: July 19, 2016, 9:00 a.m. to approximately 2:00 p.m. Eastern Daylight Time (EDT).

Document Availability: The draft NTP monograph should be available by June
immune-related health effects. The NTP evaluation concept for immunotoxicity associated with exposure to PFOA or PFOS was presented and discussed at the NTP Board of Scientific Counselors (BSC) meeting on December 10, 2014 (79 FR 62640). The NTP evaluation concept, related presentation, and BSC meeting minutes are available at http://ntp.niehs.nih.gov/go/9741. The protocol for conducting this systematic review is available at http://ntp.niehs.nih.gov/go/749926.

Meeting and Registration: The meeting is open to the public with time set aside for oral public comment. Please note that this will be both an in-person and web-based meeting. The chair of the peer review panel and NTP staff will be at the meeting location at NIEHS. The peer review panel members will be attending the meeting via web-based video conferencing. Public attendees are welcome to watch the meeting via webcast or attend in person. Attendance at NIEHS is limited only by the space available.

Registration to attend the meeting in-person or to view the webcast is by July 19, 2016, at http://ntp.niehs.nih.gov/go/36639. Registration is required to view the webcast; the URL for the webcast will be provided in the email confirming registration. Visitor and security information for those attending in-person is available at http://www.niehs.nih.gov/about/visiting/index.cfm. Individuals with disabilities who need accommodation to participate in this event should contact Dr. Yun Xie at phone: (919) 541–3436 or email: yun.xie@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.

The preliminary agenda and draft NTP monograph, preliminary agenda, registration, and other meeting materials will be available at http://ntp.niehs.nih.gov/go/36639.

Webcast: The URL for viewing the webcast will be provided to those who request for in-person or to view the webcast by July 12, 2016. Additional information will be posted when available or may be requested in hardcopy, see FOR FURTHER INFORMATION CONTACT. Persons submitting written comments should include their name, affiliation, mailing address, phone, email, and sponsoring organization (if any) with the document. Written comments received in response to this notice will be posted on the NTP Web site according to NTP’s guidelines for public comments (http://ntp.niehs.nih.gov/ntp/about_ntp/guidelines_public_comments_508.pdf), and the submitter will be identified by name, affiliation, and/or sponsoring organization if applicable.

Public comment at this meeting is welcome, with time set aside for the presentation of oral comments on the draft NTP monograph. Guidance for oral public comments is available at http://ntp.niehs.nih.gov/ntp/about_ntp/guidelines_public_comments_508.pdf. In addition to in-person oral comments at NIEHS, public comments can be presented by teleconference line. There will be 50 lines for this call; availability is on a first-come, first-served basis. The lines will be open from 9:00 a.m. until approximately 2:00 p.m. EDT on July 19, 2016, although oral comments will be received only during the formal public comment periods indicated on the preliminary agenda. The access number for the teleconference line will be provided to registrants by email prior to the meeting. Each organization is allowed one time slot. At least 7 minutes will be allotted to each time slot, and if time permits, may be extended to 10 minutes at the discretion of the chair.

Persons wishing to make an oral presentation are asked to register online at http://ntp.niehs.nih.gov/go/36639 by July 12, 2016, and indicate whether they will present comments in-person or via the teleconference line. If possible, oral public commenters should send a copy of their slides and/or statement or talking points at that time. Written comments can supplement and may expand the oral presentation.

Registration and Oral Comments: Registration for oral comments is available at http://ntp.niehs.nih.gov/go/36639. Additional information about NTP consultation and registration is available at http://ntp.niehs.nih.gov/go/36639. Public comments and any other correspondence on the draft NTP monograph should be sent to the FOR FURTHER INFORMATION CONTACT. Persons submitting written comments should include their name, affiliation, mailing address, phone, email, and sponsoring organization (if any) with the document. Written comments received in response to this notice will be posted on the NTP Web site according to NTP’s guidelines for public comments (http://ntp.niehs.nih.gov/ntp/about_ntp/guidelines_public_comments_508.pdf), and the submitter will be identified by name, affiliation, and/or sponsoring organization if applicable.

Background Information on OHAT: OHAT was established to serve as an environmental health resource to the public and regulatory and health agencies (http://www.niehs.nih.gov/ohat/pmc/articles/PMC3094430). This office conducts evaluations to assess the
evidence that environmental chemicals, physical substances, or mixtures (collectively referred to as "substances") cause adverse health effects and provides opinions on whether these substances may be of concern given what is known about current human exposure levels. OHAT also organizes workshops or state-of-the-science evaluations to address issues of importance in environmental health sciences. Information about OHAT is found at http://ntp.niehs.nih.gov/go/ohat.

Background Information on NTP Peer Review Panels: NTP panels are technical, scientific advisory bodies established on an “as needed” basis to provide independent scientific peer review and advise the NTP on agents of public health concern, new/revised toxicological test methods, or other issues. These panels help ensure transparent, unbiased, and scientifically rigorous input to the program for its use in making credible decisions about human hazard, setting research and testing priorities, and providing information to regulatory agencies about alternative methods for toxicity screening. The NTP welcomes nominations of scientific experts for upcoming panels. Scientists interested in serving on an NTP panel should provide current curriculum vitae to the

FOR FURTHER INFORMATION CONTACT. The authority for NTP panels is provided by 42 U.S.C. 217a; section 222 of the Public Health Service (PHS) Act, as amended. The panel is governed by the Federal Advisory Committee Act, as amended (5 U.S.C. Appendix 2), which sets forth standards for the formation and use of advisory committees.


John R. Bucher,
Associate Director, NTP.

[FR Doc. 2016–04102 Filed 2–25–16; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; 30-Day Comment Request: The National Physician Survey of Precision Medicine in Cancer Treatment (NCI)

Summary: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Cancer Institute, the National Institutes of Health, has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information collection listed below. This proposed information collection was previously published in the Federal Register on November 18, 2015 (80 FR 72077), and allowed 60-days for public comment. No public comments were received. The purpose of this notice is to allow an additional 30 days for public comment. The National Cancer Institute (NCI), National Institutes of Health, may not conduct or sponsor, and the respondent is not required to respond to, an information collection that has been extended, revised, or implemented on or after October 1, 1995, unless it displays a currently valid OMB control number.

Direct Comments to OMB: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the: Office of Management and Budget, Office of Regulatory Affairs, OIRA_submission@omb.eop.gov or by fax to 202–395–6974. Attention: NIH Desk Officer.

Comment Due Date: Comments regarding this information collection are best assured of having their full effect if received within 30 days of the date of this publication.

For Further Information Contact: To obtain a copy of the data collection plans and instruments, or request more information on the proposed project, contact: Janet S. de Moor, Ph.D., MPH, Project Officer, Division of Cancer Control and Population Sciences, 9609 Medical Center Drive, 3E438, MSC 9764, Rockville, MD. 20850 or call non-toll-free number 240–276–6806 or Email your request, including your address to: janet.demoor@nih.gov. Formal requests for additional plans and instruments must be requested in writing.


Need and Use of Information Collection: The purpose of this study is to investigate the current practice of precision medicine in cancer treatment among medical oncologists in the U.S. This is a nationally representative survey designed to assess oncologists’ current and potential use of genomic testing, to inform the development of interventions to facilitate optimal use of genomic testing and to improve patient-physician discussions of the risks, possible benefits, and uncertainties surrounding the use of these tests.

Current knowledge of this topic is limited as there are no nationally-representative studies on this topic to date. There are only two non-federal studies that have examined physicians’ knowledge and attitudes regarding somatic genetic and genomic testing. The survey will be administered by mail and web to medical oncology physicians across the U.S. Non-respondents will be invited to complete a follow-back survey to share their reasons for not participating. The study findings will inform NCI of relevant issues and concerns relating to the application of precision medicine to current and future cancer treatment patterns and practice. This information will also inform the development of new funding initiatives to optimize the use of precision medicine in cancer treatment. Additionally, information collected as part of this survey will be used to develop physician educational materials to address barriers to precision medicine in cancer care delivery.

OMB approval is requested for 2 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 350.

TOTAL ANNUALIZED BURDEN HOURS

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