

review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act. If the proposed project is research involving human subjects, the applicants must comply with Department of Health and Human Services regulations (45 CFR part 46) and, if applicable, Food and Drug Administration regulations (21 CFR parts 50 and 56), regarding the protection of human subjects. The applicants must ensure that the project will be subject to initial and continuing review by the appropriate institutional review boards. Overall, by providing additional scientific information for the risk assessment process, data generated from this research will support other researchers conducting human health assessments involving these substances.

Below are the mechanisms for implementing SSARP. The status of SSARP in addressing priority data needs of the first 60 priority hazardous substances through these mechanisms was described in a **Federal Register** Notice on December 13, 2005 (70 FR 73749).

#### A. TSCA/FIFRA

In developing and implementing SSARP, ATSDR and EPA established procedures to identify priority data needs of common interest to multiple federal programs. Where practicable, these data needs will be addressed through a program of toxicologic testing under TSCA or FIFRA. This part of the research will be conducted according to established TSCA/FIFRA procedures and guidelines.

#### B. Private-Sector Voluntarism

As part of SSARP, on February 7, 1992, ATSDR announced a set of proposed procedures for conducting voluntary research (57 FR 4758). Revisions based on public comments were published on November 16, 1992 (57 FR 54160). ATSDR strongly encourages private-sector organizations to propose research to address priority data needs at any time until ATSDR announces that research has already been initiated for a specific priority data need. Private-sector organizations may volunteer to conduct research to address specific priority data needs identified in this notice by submitting a letter of intent.

The letter of intent should be a brief statement (1–2 pages) that identifies the priority data need(s) to be filled and the methods to be used. TASARC will review these proposals and recommend to ATSDR the voluntary research projects that should be pursued—and how they should be conducted—with the volunteer organizations. ATSDR will

enter into only those voluntary research projects that lead to high-quality, peer-reviewed scientific work. Additional details regarding the process for voluntary research are in the **Federal Register** Notices cited in this section.

#### C. CERCLA

Those priority data needs not addressed by TSCA/FIFRA or initial voluntarism will be considered for funding by ATSDR through its CERCLA budget. Much of this research program is envisioned to be unique to CERCLA—for example, research on substances not regulated by other programs, or research needs specific to public health assessments.

Mechanisms to address these priority data needs may include a second call for voluntarism. Again, scientific peer review of study protocols and results is a requirement for all research conducted under this auspice.

ATSDR encourages private-sector organizations and other governmental programs to use ATSDR's priority data needs to plan their research activities.

Dated: October 21, 2009.

**Ken Rose,**

*Director, Office of Policy, Planning, and Evaluation, National Center for Environmental Health/Agency for Toxic Substances and Disease Registry.*

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**BILLING CODE 4163–70–P**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### National Institutes of Health

#### National Institute on Alcohol Abuse and Alcoholism; Notice of Closed Meeting

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 meeting.

The meeting 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:* National Institute on Alcohol Abuse and Alcoholism Special Emphasis Panel; AA–1 and AA–4 Study Sections Members Conflict.

*Date:* November 10, 2009.

*Time:* 3 p.m. to 5 p.m.

*Agenda:* To review and evaluate grant applications.

*Place:* National Institutes of Health, 5635 Fishers Lane, Bethesda, MD 20892. (Telephone Conference Call.)

*Contact Person:* Lorraine Gunzerath, PhD, MBA, Scientific Review Officer, National Institute on Alcohol Abuse and Alcoholism, Office of Extramural Activities, Extramural Project Review Branch, 5635 Fishers Lane, Room 2121, Bethesda, MD 20892–9304, 301–443–2369, [Igunzer@mail.nih.gov](mailto:Igunzer@mail.nih.gov).

This notice is being published less than 15 days prior to the meeting due to the timing limitations imposed by the securing of meeting attendees.

(Catalogue of Federal Domestic Assistance Program Nos. 93.271, Alcohol Research Career Development Awards for Scientists and Clinicians; 93.272, Alcohol National Research Service Awards for Research Training; 93.273, Alcohol Research Programs; 93.891, Alcohol Research Center Grants; 93.701, ARRA Related Biomedical Research and Research Support Awards, National Institutes of Health, HHS)

Dated: October 19, 2009.

**Jennifer Spaeth,**

*Director, Office of Federal Advisory Committee Policy.*

[FR Doc. E9–25623 Filed 10–26–09; 8:45 am]

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## 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.

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 Conflicts in Language and Cognition.

*Date:* November 12, 2009.

*Time:* 3:30 p.m. to 5:30 p.m.

*Agenda:* To review and evaluate grant applications.

*Place:* National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Telephone Conference Call).

*Contact Person:* Dana Jeffrey Plude, PhD, Scientific Review Officer, Center for