the impact of each of the programs. The evaluation tools are primarily protocols for conducting interviews with various staff of the three programs and stakeholders associated with the programs. The interview protocols were tested during a pilot study in 2008. There is also a self-administered form for each of the programs to be completed by Executive Directors or his/her designee. The self-administered form was developed as a result of the

pilot study and, therefore, has not been tested for reliability and validity. It is intended that the clearance process will be a mechanism for determining the reliability, validity, and feasibility of using this instrument.

Respondents: Staff of State Councils on Developmental Disabilities, State Protection and Advocacy Systems for Individuals with developmental disabilities, and University Centers for Excellence in Developmental Disabilities, Education, Research, and Service; individuals with developmental disabilities; parents of individuals with developmental disabilities; siblings of individuals with developmental disabilities; guardians; advocates; policymakers; service providers; university faculty; and others (e.g., DDC chairs, members of Protection and Advocacy boards of directors or commissioners; Consumer Advisory Committee members)

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
DD Council: Executive Director Interview	20	1	4	80
DD Council: Interview with Council Chair/Council Members	60	1	0.75	45
ees	160	1	2	320
UCEDD: Telephone Interview with Current and Graduated Students	100	1	0.75	75
UCEDD: Interview with the Consumer Advisory Committee	60	1	0.75	45
UCEDD: Interview with Peer Researchers and Colleagues	100	1	0.75	75
UCEDD: Interview with Recipients of Community Services or Members of Organizations/Agencies that Are Trained To Provide Community Serv-				
ices	100	1	0.75	75
UCEDD: Self-administered Form	20	1	8	160
P&A: Executive Director Interview	20	1	4	80
P&A: Staff Interview	60	1	0.75	45
P&A: Board of Directors (Commissioners)—Chair and Members	60	1	0.75	45
P&A: Group Interview with Policymakers and Collaborators	160	1	2	320
P&A: Interview with Recipient of Community Education	100	1	0.75	75
P&A: Interview with Clients	100	1	0.75	75
P&A: Self-administered Form	20	1	8	160
UCEDD: Interview with Director	20	1	4	80
DD Council: Group Interview with Recipients of Self-Advocacy and Lead-				
ership Education and Training	100	1	0.75	75
DD Council: Group Interview with Recipients of Education and Training to				
Improve Community Capacity	100	1	0.75	75
DD Council: Self-administered Form	20	1	8	160
DD Council Estimate of Total Burden Hours for Activities to Support Ad-				
ministration of Proposed Information Collection Instruments	20	1	33.50	670
P&A Estimate of Total Burden Hours for Activities to Support Administra-				
tion of Proposed Information Collection Instruments	20	1	33.50	670
UCEDD Estimate of Total Burden Hours for Activities to Support Adminis-				
tration of Proposed Information Collection Instruments	20	1	33.50	670

Estimated Total Annual Burden Hours: 4,075.

Additional Information:

Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. All requests should be identified by the title of the information collection. E-mail address: infocollection@acf.hhs.gov.

OMB Comment:

OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect

if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following:

Office of Management and Budget, Paperwork Reduction Project, Fax: 202– 395–7245, Attn: Desk Officer for the Administration for Children and Families.

Dated: June 30, 2009.

Janean Chambers,

Reports Clearance Officer.

[FR Doc. E9–15841 Filed 7–6–09; 8:45 am]

BILLING CODE 4184-01-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 Alcohol Pharmacotherapy and the Treatment and Prevention of HIV/ AIDS. (RFA AA 09 007/008) and Other AIDS Related Research.

Date: August 6, 2009.

Time: 8 a.m. to 11 a.m.

Agenda: To review and evaluate grant applications.

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

Contact Person: Katrina L Foster, PhD, Scientific Review Officer, National Inst on Alcohol Abuse & Alcoholism, National Institutes of Health, 5635 Fishers Lane, Rm. 2019, Rockville, MD 20852. 301–443–4032. katrina@mail.nih.gov.

(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: June 29, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9-15847 Filed 7-6-09; 8:45 am]

BILLING CODE 4140-01-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of General Medical Sciences; 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: National Institute of General Medical Sciences. Special Emphasis Panel Minority Biomedical Research Support. Date: July 19-20, 2009.

Time: 7 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Hyatt Regency Bethesda, One Bethesda Metro Center, Bethesda, MD 20814.

Contact Person: Margaret J. Weidman, PhD, Scientific Review Officer, Office of Scientific Review, National Institute of General Medical Sciences, National Institutes of Health, 45 Center Drive, Room 3AN18B, Bethesda, MD 20892. 301–594–3663.

weidmanma@nigms.nih.gov.

Name of Committee: National Institute of General Medical Sciences. Special Emphasis Panel MBRS Score.

Date: July 20-21, 2009.

Time: 8 a.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Hyatt Regency Bethesda, One Bethesda Metro Center, Bethesda, MD 20814.

Contact Person: Lisa Dunbar, PhD, Scientific Review Officer, Office of Scientific Review, National Institute of General Medical Sciences, National Institutes of Health, 45 Center Drive, Room 3AN12, Bethesda, MD 20892. 301–594–2849. dunbarl@mail.nih.gov.

Name of Committee: National Institute of General Medical Sciences. Special Emphasis Panel New Innovator Awards.

Date: July 21, 2009.

Time: 1 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

*Place: National Institutes of Health, Natcher Building, 45 Center Drive, Bethesda, MD 20892. (Telephone Conference Call).

Contact Person: Richard T. Okita, PhD, Program Director, Pharmacological and Physiological Sciences Branch, National Institute of General Medical Sciences, National Institutes of Health, Natcher Building, Room 2A5-49, Bethesda, MD 20892. 301-594-4469. okitar@nigms.nih.gov. (Catalogue of Federal Domestic Assistance Program Nos. 93.375, Minority Biomedical Research Support; 93.821, Cell Biology and Biophysics Research; 93.859, Pharmacology, Physiology, and Biological Chemistry Research; 93.862, Genetics and Developmental Biology Research; 93.88, Minority Access to Research Careers; 93.96, Special Minority Initiatives, National Institutes of Health, HHS)

Dated: June 29, 2009.

Jennifer Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. E9–15846 Filed 7–6–09; 8:45 am]

BILLING CODE 4140-01-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institutes of Health Guidelines for Human Stem Cell Research

SUMMARY: The National Institutes of Health (NIH) is hereby publishing final "National Institutes of Health

Guidelines for Human Stem Cell Research" (Guidelines).

On March 9, 2009, President Barack H. Obama issued Executive Order 13505: Removing Barriers to Responsible Scientific Research Involving Human Stem Cells. The Executive Order states that the Secretary of Health and Human Services, through the Director of NIH, may support and conduct responsible, scientifically worthy human stem cell research, including human embryonic stem cell (hESC) research, to the extent permitted by law.

These Guidelines implement
Executive Order 13505, as it pertains to
extramural NIH-funded stem cell
research, establish policy and
procedures under which the NIH will
fund such research, and helps ensure
that NIH-funded research in this area is
ethically responsible, scientifically
worthy, and conducted in accordance
with applicable law. Internal NIH
policies and procedures, consistent with
Executive Order 13505 and these
Guidelines, will govern the conduct of
intramural NIH stem cell research.

DATES: *Effective Date:* These Guidelines are effective on July 7, 2009.

Summary of Public Comments on Draft Guidelines: On April 23, 2009 the NIH published draft Guidelines for research involving hESCs in the **Federal Register** for public comment, 74 FR 18578 (April 23, 2009). The comment period ended on May 26, 2009.

The NIH received approximately 49,000 comments from patient advocacy groups, scientists and scientific societies, academic institutions, medical organizations, religious organizations, and private citizens. The NIH also received comments from members of Congress. This Notice presents the final Guidelines together with the NIH response to public comments that addressed provisions of the Guidelines.

Title of the Guidelines, Terminology, and Background

Respondents felt the title of the NIH draft guidelines was misleading, in that it is entitled "National Institutes of Health Guidelines for Human Stem Cell Research," yet addresses only one type of human stem cell. The NIH notes that although the Guidelines pertain primarily to the donation of embryos for the derivation of hESCs, one Section also applies to certain uses of both hESCs and human induced pluripotent stem cells. Also, the Guidelines discuss applicable regulatory standards when research involving human adult stem cells or induced pluripotent stem cells constitutes human subject research.