Time: 8:30 a.m. to 4:00 p.m. Agenda: To review and evaluate cooperative agreement applications.

Place: Embassy Suites at the Chevy Chase Pavilion, 4300 Military Road NW., Washington, DC 20015.

Contact Person: Anne E. Schaffner, Ph.D., Chief, Scientific Review Branch Division of Extramural Research, National Eye Institute, 5635 Fishers Lane, Suite 1300, MSC 9300, Bethesda, MD 20892–9300, (301) 451–2020, aes@nei.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.867, Vision Research, National Institutes of Health, HHS)

Dated: November 3, 2017.

Natasha M. Copeland,

Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2017–24373 Filed 11–8–17; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; 60-Day Comment Request; CareerTrac

AGENCY: National Institutes of Health, HHS.

ACTION: Notice.

SUMMARY: In compliance with the requirement of the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information collection listed below. The purpose of this notice is to allow 60 days for public comment. The Fogarty International Center (FIC), National Institute of Environmental Health Sciences (NIEHS), including the Superfund Research Program (SRP) within NIEHS, National Institute of

General Medical Science (NIGMS), and 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. DATES: Comments regarding this information collection are best assured of having their full effect if received within 60-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: Dr. Rachel Sturke, Evaluation Officer, Division of Science Policy, Planning, and Evaluation, FIC, NIH, 16 Center Drive, Bethesda, MD 20892 or call non-toll-free number (301) 496—1491 or Email your request, including your address to: rachel.sturke@nih.gov. Formal requests for additional plans and instruments must be requested in writing.

SUPPLEMENTARY INFORMATION: Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 requires: Written comments and/or suggestions from the public and affected agencies are invited to address one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will have practical utility; (2) The accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) Ways to enhance the quality, utility, and clarity of the information to be

collected; and (4) Ways to minimizes the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

Proposed Collection Title: CareerTrac, 0925–0568, Expiration Date: 06/30/2019—REVISION, Fogarty International Center (FIC), National Institute of Environmental Health Sciences (NIEHS), National Institute of General Medical Science (NIGMS), National Cancer Institute (NCI), National Institutes of Health (NIH).

Need and Use of Information Collection: This data collection system is being used to track, evaluate and report short and long-term outputs, outcomes and impacts of trainees involved in health research training programs-specifically tracking this for at least ten years following training by having Principal Investigators enter data after trainees have completed the program. The data collection system provides a streamlined, web-based application permitting principal investigators to record career achievement progress by trainee on a voluntary basis. FIC, NIEHS, NCI and NIGMS management will use this data to monitor, evaluate and adjust grants to ensure desired outcomes are achieved, comply with OMB Part requirements, respond to congressional inquiries, and as a guide to inform future strategic and management decisions regarding the grant program.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 16,154.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Number of respondents	Number of responses per respondent	Average time per response (in hours)	Total annual burden hour
FIC Grantee	80	90	30/60	3,600
NIEHS Grantee	60	45	30/60	1,350
NCI/CRCHD Grantee	264	22	30/60	2,904
NIGMS Grantee	80	150	30/60	6,000
Superfund Grantee	20	105	30/60	1,050
Trainees	5000	1	15/60	1,250
Total	5,504	34,808		16,154

Dated: October 15, 2017.

Celia Wolfman,

Project Clearance Liaison, FIC, NIH. [FR Doc. 2017–24362 Filed 11–8–17; 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; Generic Clearance To Support the Safe To Sleep® Campaign (Eunice Kennedy Shriver National Institute of Child Health and Human Development)

AGENCY: National Institutes of Health,

HHS.

ACTION: Notice.

SUMMARY: In compliance with the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information collection listed below.

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

ADDRESSES: 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: Desk Officer for NIH.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact: Lorena Kaplan, M.P.H., CHES, Office of Communications, Eunice Kennedy Shriver National Institute of Child Health and Human Development, National Institutes of Health, 31 Center Drive, Room 2A32, Bethesda, Maryland 20892, or call non-toll free number (301) 496-6670 or Email your request, including your address to lorena.kaplan@nih.gov. Formal requests for additional plans and instruments must be requested in writing. **SUPPLEMENTARY INFORMATION: This**

proposed information collection was

previously published in the Federal

Register on Monday, August 28, 2017,

page 40776–40777 (82 FR 40776–40777) and allowed 60 days for public comment. NICHD received one comment in response to the 60-Day **Federal Register** Notice. The purpose of this notice is to allow an additional 30 days for public comment.

The Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD), 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.

In compliance with section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information collection listed below.

Proposed Collection: Generic Clearance to Support the Safe to Sleep® Campaign 0925–0701, REINSTATEMENT WITH CHANGE at the Eunice Kennedy Shriver National Institute for Child Health and Human Development (NICHD), National Institutes of Health (NIH).

Need and Use of Information Collection: This is a request to reinstate with change a generic clearance that would be used for submissions specific to the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD) Safe to Sleep® (STS) public education campaign. Submissions for the STS campaign will be used to assess the understanding and reach of STS campaign materials and messages, and to monitor and improve campaign activities such as training workshops and overall implementation. The purpose of this information collection is to monitor and modify campaign activities, to plan future campaign activities, to develop messages and materials, and to develop distribution and outreach strategies that are effective at communicating their message to bring about the intended response, awareness, and/or behavioral change for the target audiences. This generic clearance will enable the NICHD to: (1) More efficiently assess the implementation of campaign activities; (2) better understand the target audiences' knowledge, attitudes, and beliefs toward STS messages and materials; (3) better understand how the campaign activities have influenced the target audiences' behaviors and

practices; and (4) monitor and improve activities such as trainings, materials, and messages. Having a way to gather feedback on the STS campaign activities is critical to assessing the reach and effect of campaign efforts. Data collected for the campaign can inform where future STS campaign resources can produce the most meaningful results.

Data collected for the STS campaign generic clearance will be used by a number of audiences, including STS campaign staff, NICHD leadership, STS campaign collaborators, Federal SUID/ SIDS Workgroup members, SUID/SIDS stakeholders, clinical and maternal and child health professionals. These audiences may use the information collections to: (1) Develop new campaign messages, materials, and/or training curricula; (2) monitor and improve campaign activities; (3) make decisions about campaign activities; (4) inform current campaign activities; and (5) inform and/or change practices and behaviors of program participants.

Examples of the types of information collections that could be included under this generic clearance include: Focus groups and in-depth interviews with parents/caregivers and/or health professionals to get feedback on distribution and outreach activities, and/or campaign messages; and Surveys with parents/caregivers and/or health professionals to: (1) Assess the usefulness of the new STS campaign materials, including print and on-line materials and a video, (2) track outreach experiences of program participants, (3) assess training participants' changes in knowledge related to safe infant sleep behavior and implementation of outreach methods taught, and (4) assess program participants' resource needs.

The sub-studies for this generic clearance will be small scale, designed to obtain results frequently and quickly to guide campaign development and implementation, inform campaign direction, and be used internally for campaign management purposes.

NICHD's current scope and capacity for STS generic sub-studies is non-existent and this request would fill this gap.

Changes have been made to the annualized burden hours to reflect the anticipated data collections during the next 3 years.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 12,920.