OMB Circular A-130, the National Security Agency, the Privacy Act, and other Federal agencies; (2) executes the Agency's Risk Management Program, evaluates and assists with the implementation of safeguards to protect major information systems, and IT infrastructure; (3) manages the development, implementation, and evaluation of the HRSA information technology security and privacy training program to meet the requirements as mandated by OMB Circular A-130, the Computer Security Act, and Privacy Act; (4) assesses all new emerging technologies and impact on technology integration on HRSA missions and program objectives; (5) provides leadership for strategic planning that leverages information systems security, program strategies, and advanced technology integration to achieve program objectives through innovative technology use; (6) the HRSA Incident Response Center (HIRC) provides a centralized, responsive resource for computer security incident reporting, management, and situational awareness of the Department's information security posture; (7) provides services include computer security situational awareness reports, computer forensics, cyberrelated advisories, as well as cyber alerts, warnings, and Block/Watch lists are utilized and disseminated; (8) the HIRC coordinates with other Agencies and organizations for computer security and maintains a lab where new products are tested to insure that HRSA is utilizing state of the art, cutting edge technologies to ensure the secure operation of the HRSA infrastructure; and (9) provides leadership for ongoing cyber protection and incident detection response, reporting, and handling in accordance with OMB and departmental guidance.

Division of IT Operational Support Services (RB58)

The Division of IT Operational Support Services (ITOSS) (1) provides leadership, consultation, training, and management services for HRSA's enterprise computing environment; (2) directs and manages the support and acquisition of HRSA network and desktop hardware, servers, wireless communication devices, and software licenses; (3) is responsible for the HRSA Data Center and the operation and maintenance of a complex, highavailability network infrastructure on which mission-critical applications are made available 24 hours per day, 7 days per week; (4) controls infrastructure

configuration management, installations and upgrades, security perimeter protection, and system resource access; (5) coordinates IT activities for Continuity of Operations Planning (COOP) Agency-wide including provisioning and maintaining IT infrastructure and hardware at designated COOP locations to support emergency and COOP requirements; (6) maintains workstation hardware and software configuration management controls; (7) the Chief Technology Officer (CTO), reporting to the ITOSS Division Director is responsible for assessing emerging technologies and the subsequent impact on current infrastructure restraints and program objectives; (8) coordinates and engages with all OIT Divisions and Branches to insure that advanced technology is being utilized to achieve program objectives through innovative technology use; and (9) provides leadership and establishes policy and provides oversight for Agency IT configuration management.

Section RB5–30, Delegations of Authority

All delegations of authority and redelegations of authority made to HRSA officials that were in effect immediately prior to this reorganization, and that are consistent with this reorganization, shall continue in effect pending further re-delegation.

This reorganization is effective upon date of signature.

Dated: September 30, 2011.

Mary K. Wakefield,

Administrator.

[FR Doc. 2011–26007 Filed 10–6–11; 8:45 am]

BILLING CODE 4165-15-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request; A Generic Submission for Theory Development and Validation (NCI)

SUMMARY: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Cancer Institute (NCI), 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. This proposed information collection was previously published in the Federal Register on August 2, 2011

(76 FR 46307) and allowed 60-days for public comment. No public comment were received. The purpose of this notice is to allow an additional 30 days for public comment. The 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.

Proposed Collection: Title: A Generic Submission for Theory Development and Validation (NCI). Type of Information Collection Request: NEW. *Need and Use of Information Collection:* In order to carry out NCI's legislative mandate to conduct and support research with respect to the causes and prevention of cancer, it is beneficial for NCI, through initiatives in the Behavioral Research Program (BRP), to conduct and support behavioral research informed by and informing theory. Formative research in the area of theory development and validation would provide the basis for developing effective cancer prevention and control strategies, allow for a better understanding of theoretical constructs that influence decisions and actions related to cancer, and ultimately contribute to reducing the U.S. cancer burden. Data collections that result from this generic clearance would inform and clarify the use of theory in BRPsupported initiatives and funding announcements. Specifically, this research would allow NCI to conduct research to: (1) Identify psychological, biobehavioral, demographic, and individual difference predictors of cancer prevention and control behaviors and outcomes; (2) Develop and refine integrative theories; (3) Identify and observe theoretical and innovative trends in cancer prevention and control research; and (4) Determine feasibility and usefulness of collaborative and multidisciplinary approaches to cancer prevention and control. Frequency of Response: Will be determined by each project. Affected Public: Individuals or households; Businesses or other for profit; Not-for-profit institutions; Federal Government; State, Local, or Tribal Government. Type of Respondents: Members of the public including, but not limited to health professionals, physicians, and researchers. Table 1 outlines the estimated burden hours and cost required for a three-year approval of this generic submission.

TABLE A.12–1—ESTIMATES OF BURDEN HOURS FOR THREE YEARS [Generic Study]

Type of respondents	Number of respondents	Frequency of response	Average time per response (minutes/hour)	Total burden hours
General Public	2,000	1	15/60 (0.25)	500
Physicians	6,000	1	30/60 (0.5)	3,000
Health Professionals	1,000	1	60/60 (1)	1,000
Researchers	1,000	1	90/60 (1.5)	1,500
Total	11,500			6,000

Request For Comments: Written comments and/or suggestions from the public and affected agencies are invited on 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 minimize 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.

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 Attention: NIH Desk Officer, Office of Management and Budget at OIRA submission@omb.eop.gov or by fax to 202-395-6974. To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Richard P. Moser, Ph.D., Science of Research and Technology Branch, Behavioral Research Program, Division of Cancer Control and Population Sciences, National Cancer Institute/NIH, 6130 Executive Blvd., Rockville, MD 20892, call non-toll-free number 301-496-0273 or e-mail your request, including your address to: moserr@mail.nih.gov.

Comments 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.

Dated: October 3, 2011.

Vivian Horovitch-Kelley,

NCI Project Clearance Liaison, National Institutes of Health.

[FR Doc. 2011–26043 Filed 10–6–11; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of Environmental Health Sciences; Cancellation of Meeting

Notice is hereby given of the cancellation of the Interagency Breast Cancer and Environmental Research Coordinating Committee, October 12, 2011, 1 p.m. to 3 p.m., NIEHS/National Institutes of Health, Building 4401, East Campus, 79 T.W. Alexander Drive, Research Triangle Park, NC 27709 which was published in the **Federal Register** on August 12, 2011, 76 FR 50234.

Dated: September 30, 2011.

Jennifer S. Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. 2011–26000 Filed 10–6–11; 8:45 am] BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Heart, Lung, and Blood Institute; 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: Heart, Lung, and Blood Initial Review Group, Clinical Trials Review Committee.

Date: October 24–25, 2011. Time: 8 a.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: Hyatt Regency Bethesda, One Bethesda Metro Center, 7400 Wisconsin Avenue, Bethesda, MD 20814.

Contact Person: Keary A Cope, Ph.D., Scientific Review Officer, Review Branch/DERA, National Heart, Lung, and Blood Institute, 6701 Rockledge Drive, Room 7190, Bethesda, MD 20892–7924, 301–435–2222, copeka@mail.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.233, National Center for Sleep Disorders Research; 93.837, Heart and Vascular Diseases Research; 93.838, Lung Diseases Research; 93.839, Blood Diseases and Resources Research, National Institutes of Health, HHS)

Dated: October 3, 2011.

Jennifer S. Spaeth,

Director, Office of Federal Advisory Committee Policy.

[FR Doc. 2011-25999 Filed 10-6-11; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Cancer Institute; Notice of Meetings

Pursuant to section 10(a) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is