Performance Standard 10—Program Recommendations

A Unit makes statutory or programmatic recommendations, when warranted, to the State government. In meeting this standard, the following performance indicators will be considered:

A. The Unit, when warranted and appropriate, makes statutory recommendations to the State legislature to improve the operation of the Unit, including amendments to the enforcement provisions of the State code.

B. The Unit, when warranted and appropriate, makes other regulatory or administrative recommendations regarding program integrity issues to the State Medicaid agency and to other agencies responsible for Medicaid operations or funding.

C. The Unit monitors actions taken by the State legislature and the State Medicaid or other agencies in response to recommendations.

D. The Unit reports program recommendations to OIG.

Performance Standard 11—Agreement With Medicaid Agency

A Unit periodically reviews its Memorandum of Understanding (MOU) with the single State Medicaid agency to ensure that it reflects current practice, policy, and legal requirements. In meeting this standard, the following performance indicators will be considered:

A. The MOU reflects current policy and practice by both the Unit and the State Medicaid agency.

B. The MOU meets current Federal legal requirements as contained in law or regulation, including 42 CFR § 455.21, “Cooperation with State Medicaid fraud control units,” and 42 CFR 455.23, “Suspension of payments in cases of fraud.”

C. The MOU is consistent with current Federal and State policy, including any policies issued by OIG or the Centers for Medicare & Medicaid Services (CMS).

D. Consistent with Performance Standard 4, the MOU establishes a process to ensure the receipt of an adequate volume and quality of referrals to the Unit from the State Medicaid agency.

E. The MOU incorporates by reference the CMS Performance Standard for Referrals of Suspected Fraud from a Single State Agency to a Medicaid Fraud Control Unit.

Performance Standard 12—Fiscal Control

A Unit exercises proper fiscal control over Unit resources. In meeting this standard, the following performance indicators will be considered:

A. The Unit director, or the director’s designee, approves and signs the Unit’s budget and estimated expenditures.

B. The Unit director, or the director’s designee, approves and signs all fiscal and administrative reports concerning Unit expenditures.

C. The Unit maintains an equipment inventory that is updated on a regular basis to reflect all property under the Unit’s control.

D. The Unit maintains an effective time and attendance system.

E. The Unit applies generally accepted accounting principles in its control of Unit funding.

F. The Unit employs a financial system in which all funds are assigned to individual accounts according to their source and all expenditure items can be traced to the original funding stream and account.

Performance Standard 13—Training

A Unit maintains an annual training plan for all professional disciplines. In meeting this standard, the following performance indicators will be considered:

A. The Unit maintains a training plan for each professional discipline that includes an annual minimum number of training hours and that is at least as stringent as required for professional certification.

B. The Unit ensures that professional staff complies with its training plans and maintains records of the staff’s compliance.

C. Professional certifications are maintained for all staff, including continuing education requirements.

D. The Unit participates in training offered by OIG, CMS, and other MFCUs, as funding permits.

E. Through cross-training or by other means, Unit staff receive training on the role and responsibilities of the State Medicaid agency and other law enforcement partners.

Daniel R. Levinson, Inspector General.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request; New Proposed Collection, Environmental Science Formative Research Methodology Studies for the National Children’s Study

SUMMARY: Under the provisions of 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 reinstatement of approval of the information collection listed below. This proposed information collection was previously published in the Federal Register on April 27, 2011, pages 23603–23605, and allowed 60 days for public comment. Two written comments and two verbal comments were received. The verbal comments expressed support for the broad scope of the study. The written comments were identical and questioned the cost and utility of the study. 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.


(a) PURPOSE.—It is the purpose of this section to authorize the National Institute of Child Health and Human Development* to conduct a national longitudinal study of environmental influences (including physical, chemical, biological, and psychosocial) on children’s health and development.

(b) IN GENERAL.—The Director of the National Institute of Child Health and Human Development* shall establish a consortium of representatives from appropriate Federal agencies (including the Centers for Disease Control and Prevention, the Environmental Protection Agency) to—

(1) plan, develop, and implement a prospective cohort study, from birth to adulthood, to evaluate the effects of both chronic and intermittent exposures on child health and human development; and

(2) investigate basic mechanisms of developmental disorders and environmental factors, both risk and protective, that influence health and developmental processes.

[FR Doc. 2011–25894 Filed 10–5–11; 8:45 am]
BILLING CODE 4152–01–P

62077 Federal Register / Vol. 76, No. 194 / Thursday, October 6, 2011 / Notices
To fulfill the requirements of the Children’s Health Act, the results of formative research will be used to maximize the efficiency (measured by scientific robustness, participant and infrastructure burden, and cost) of environmental sample collection procedures and technology, storage operations, and study logistics) and cost of NCS Vanguard and Main Studies. With this submission, the NCS seeks to obtain OMB’s generic clearance to collect environmental samples from homes and child care settings, and conduct accompanying short surveys related to the physical and chemical environment. The results from these formative research projects will inform the feasibility (scientific robustness), acceptability (burden to participants and study logistics) and cost of NCS Vanguard and Main Study environmental sample and information collection in a manner that minimizes public information collection burden compared to burden anticipated if these projects were incorporated directly into either the NCS Vanguard or Main Study.

Frequency of Response: Annual [As needed on an on-going and concurrent basis]. Affected Public: Members of the public, researchers, practitioners, and other health professionals. Type of Respondents: Women of child-bearing age, fathers, public health and environmental science professional organizations and practitioners, and schools and child care organizations. These include both persons enrolled in the NCS Vanguard Study and their peers who are not participating in the NCS Vanguard Study.

Annual reporting burden: See Table 1. The annualized cost to respondents is estimated at: $780,000 (based on $10 per hour). There are no Operating or Maintenance Costs to report. There are no Operating or Maintenance Costs to report.

### Table 1—Estimated Annual Reporting Burden Summary, Environmental Science

<table>
<thead>
<tr>
<th>Data collection activity</th>
<th>Type of respondent</th>
<th>Estimated number of respondents</th>
<th>Estimated number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Estimated total annual burden hours requested</th>
</tr>
</thead>
<tbody>
<tr>
<td>Home Air</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>1</td>
<td>1</td>
<td>4,000</td>
</tr>
<tr>
<td>Home Water</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>1</td>
<td>1</td>
<td>4,000</td>
</tr>
<tr>
<td>Home Dust</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>1</td>
<td>1</td>
<td>4,000</td>
</tr>
<tr>
<td>School and Child Care Facility Air</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>1</td>
<td>1</td>
<td>4,000</td>
</tr>
<tr>
<td>School and Child Care Facility Water</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>1</td>
<td>1</td>
<td>4,000</td>
</tr>
<tr>
<td>School and Child Care Facility Dust</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>1</td>
<td>1</td>
<td>4,000</td>
</tr>
<tr>
<td>Small, focused survey and instrument design</td>
<td>NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>4,000</td>
<td>2</td>
<td>1</td>
<td>8,000</td>
</tr>
<tr>
<td>and administration.</td>
<td>Health and Social Service Providers.</td>
<td>2,000</td>
<td>1</td>
<td>1</td>
<td>2,000</td>
</tr>
<tr>
<td>Focus groups</td>
<td>Community Stakeholders .... NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>2,000</td>
<td>1</td>
<td>1</td>
<td>2,000</td>
</tr>
<tr>
<td></td>
<td>Health and Social Service Providers.</td>
<td>2,000</td>
<td>1</td>
<td>1</td>
<td>2,000</td>
</tr>
<tr>
<td>Cognitive interviews</td>
<td>Community Stakeholders .... NCS participants .......... Members of NCS target population (not NCS participants).</td>
<td>2,000</td>
<td>1</td>
<td>1</td>
<td>2,000</td>
</tr>
<tr>
<td></td>
<td>Members of NCS target population (not NCS participants).</td>
<td>500</td>
<td>1</td>
<td>2</td>
<td>1,000</td>
</tr>
</tbody>
</table>
TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN SUMMARY, ENVIRONMENTAL SCIENCE—Continued

<table>
<thead>
<tr>
<th>Data collection activity</th>
<th>Type of respondent</th>
<th>Estimated number of respondents</th>
<th>Estimated number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Estimated total annual burden hours requested</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td></td>
<td>---------------------------------</td>
<td>---------------------------------------------</td>
<td>----------------------------------</td>
<td>-----------------------------------------------</td>
</tr>
</tbody>
</table>

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 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 Office of Information and Regulatory Affairs, Attn: NIH Desk Officer, by E-mail to 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 Ms. Jamelle E. Banks, Public Health Analyst, Office of Science Policy, Analysis and Communication, National Institute of Child Health and Human Development, 31 Center Drive, Room 2A18, Bethesda, Maryland, 20892, or call a non-toll free number (301) 496–1877 or E-mail your request, including your address to banksj@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: September 30, 2011.

Jamelle E. Banks,
Public Health Analyst, Office of Science Policy, Analysis and Communications, National Institute of Child Health and Human Development.

[FR Doc. 2011–25868 Filed 10–5–11; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Cancer 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 a meeting of the Board of Scientific Counselors for Basic Sciences National Cancer Institute.

The meeting will be closed to the public as indicated below in accordance with the provisions set forth in section 552b(c)(6), Title 5 U.S.C., as amended for the review, discussion, and evaluation of individual intramural programs and projects conducted by the National Cancer Institute, including consideration of personnel qualifications and performance, and the competence of individual investigators, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Board of Scientific Counselors for Basic Sciences National Cancer Institute.

Date: November 15, 2011.

Time: 9 a.m. to 4 p.m.

Agenda: To review and evaluate personal qualifications and performance, and competence of individual investigators.

Place: National Institutes of Health, National Cancer Institute, 9000 Rockville Pike, Building 31, Conference Room 6, Bethesda, MD 20892.

Contact Person: Florence E. Farber, PhD, Executive Secretary, Office of the Director, National Cancer Institute, National Institutes of Health, 6116 Executive Boulevard, Room 2205, Bethesda, MD 20892, 301–496–7628, ff@nih.gov.

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/bsc/bs/bs.htm, where an agenda and any additional information for the meeting will be posted when available.

(Departmental) National Cancer Institute, including whether the performance of the function of the agency, including whether the information will have practical utility;

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)

Dated: September 30, 2011.

Jennifer S. Spaeth,
Director, Office of Federal Advisory Committee Policy.

[FR Doc. 2011–25897 Filed 10–5–11; 8:45 am]

BILLING CODE 4140–01–P

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, Fellowships: Physiology and Pathobiology of Musculoskeletal, Oral, and Skin Systems.

Date: October 26, 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: Abdelouahab Aitouche, PhD, Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 4222, MSC 7812, Bethesda, MD 20892, 301–435–2365, aitouchea@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel, PAR Panel: