available at http://datawarehouse.hrsa.gov.

Dated: June 19, 2012. Mary K. Wakefield,

Administrator.

[FR Doc. 2012-15819 Filed 6-28-12; 8:45 am]

BILLING CODE 4165-15-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### **National Institutes of Health**

Proposed Collection; Comment Request; National Institute of Nursing Research (NINR) Summer Genetics Institute Alumni Survey

**SUMMARY:** In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, for opportunity for public comment on

proposed data collection projects, the National Institute of Nursing Research (NINR), the National Institutes of Health (NIH) will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB).

Prosposed Collection: Title: NIH/ National Institute of Nursing Research (NINR) Summer Genetics Institute Alumni Survey. Type of Information Collection Request: NEW. Need and Use of Information Collection: The NINR Summer Genetics Institute Alumni Survey will obtain information on the long-term outcomes of this training program for nurse scientists and faculty. Target participants are alumni of this training institute which began in 2000. The survey inquires about career activities, including research, clinical, teaching and educational activities, since completion of the NINR Summer

Genetics Institute. This is a 39-item survey that takes an average of 30 minutes to complete. The findings will provide valuable information on the influence of the Institute in developing genetics research capability among Institute alumni, and development and expansion of clinical practice in genetics among alumni who are nurse clinicians. Frequency of Response: Annual for three (3) years. Affected Public: Individual alumni of the NINR Summer Genetics Institute. Type of Respondents: Nurse scientists, clinicians, and faculty. The annual reporting burden is as follows: Estimated Number of Respondents: 150; Estimated Number of Responses per Respondent: 1; Average Burden Hours per Response: .5; and Estimated Total Annual Burden Hours Requested: 75. There are no Capital Costs, Operating or Maintenance Costs to report.

Type of respondents	Number of respondents	Frequency of response	Average time per response (minutes/hour)	Annual burden hours
Researchers	150	1	0.5	75

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.

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 Dr. Amanda Greene, Science Evaluation Officer, Office of Science Policy and Public Liaison, NINR, Democracy One, 6701 Democracy Blvd., Suite 700, Bethesda, MD 20892, or call non-toll-free number 301–496–9601, or email your request to amanda.greene@nih.gov.

Comments Due Date: Comments regarding this information collection are

best assured of having their full effect if received within 60-days of the date of this publication.

Dated: June 22, 2012.

### Amanda Greene,

NINR Project Clearance Officer, Science Evaluation Officer, NINR, National Institutes of Health.

[FR Doc. 2012–16022 Filed 6–28–12; 8:45 am]

BILLING CODE 4140-01-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### **National Institutes of Health**

Submission for OMB Review; Comment Request: Child Health Disparities Substudy 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 Institute of Child Health and Human Development (NICHD), 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 March 16, 2012, pages 15780-15782 (Volume 77, Number 52) of the Federal Register and allowed 60 days for public comment. No written comments 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: Child Health Disparities Substudy for the National Children's Study (NCS). Type of Information Collection Request: NEW. Need and Use of Information Collection: The Children's Health Act of 2000 (Pub. L. 106–310) states:

(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.

- (c) REQUIREMENT.—The study under subsection (b) shall—
- (1) Incorporate behavioral, emotional, educational, and contextual consequences to enable a complete assessment of the physical, chemical, biological, and psychosocial environmental influences on children's wellbeing;
- (2) Gather data on environmental influences and outcomes on diverse populations of children, which may include the consideration of prenatal exposures; and
- (3) Consider health disparities among children, which may include the consideration of prenatal exposures.

To fulfill the requirements of the Children's Health Act, the Child Health Disparities Substudy will validate measures needed for studying health disparities and selected biomarkers. Utilizing cognitive interview techniques and components of standardized questionnaires, responses will be used to assess and validate measures of health literacy, discrimination, parenting self-efficacy, and health care accessibility. Acceptability and feasibility of saliva collection from a subsample of women and young children will also be evaluated. The incorporation of saliva measurements will increase understanding of biological responses to environmental factors and how these may be correlated with health disparities within this population.

Background: The National Children's Study is a prospective, national longitudinal study of the interaction between environment, genetics on child health and development. The Study defines "environment" broadly, taking a number of natural and man-made environmental, biological, genetic, and psychosocial factors into account. By studying children through their different phases of growth and

development, researchers will be better able to understand the role these factors have on health and disease. Findings from the Study will be made available as the research progresses, making potential benefits known to the public as soon as possible. The National Children's Study is led by a consortium of federal partners: The U.S. Department of Health and Human Services (including the Eunice Kennedy Shriver National Institute of Child Health and Human Development and the National Institute of Environmental Health Sciences of the National Institutes of Health and the Centers for Disease Control and Prevention), and The U.S. Environmental Protection Agency.

To conduct the detailed preparation needed for a study of this size and complexity, the NCS was designed to include a preliminary pilot study known as the Vanguard Study. The purpose of the Vanguard Study is to assess the feasibility, acceptability, and cost of the recruitment strategy, study procedures, and outcome assessments that are to be used in the NCS Main Study. The Vanguard Study begins prior to the NCS Main Study and will run in parallel with the Main Study. At every phase of the NCS, the multiple methodological studies conducted during the Vanguard phase will inform the implementation and analysis plan for the Main Study.

In this information collection request, the NCS requests approval from OMB to perform a multi-center substudy called the Child Health Disparity Substudy. This substudy aims to validate measures needed for studying health disparities and selected biomarkers. Developing optimum measures for studying health disparities is of particular interest to the NCS because studies have shown that health literacy, discrimination,

parenting self-efficacy, health care (access, utilization, and quality) contribute to health disparities. Additionally, aspects of the social environment such as social isolation, lack of control and contingency and social support, violence, discrimination, challenging and changing social relationships, and restricted access to health care are thought to interact with biological processes. Variation in these processes has been associated with negative emotional states, cognitive deficits, problem behavior, and a variety of metabolic and immune-related processes. Alone, or particularly in combination with other commonly collected measures of social forces and family relationships, salivary analytes have the potential to advance our understanding of maternal and child health and development. This project will make its contribution to the NCS Main Study and to the health disparities field as a whole by constructing a validated set of questionnaire measures and biomarker analyses that can be used among pregnant women and mothers of young children for the purpose of investigating disparities.

Frequency of Response: One-time data collection conducted in multiple phases.

Affected Public: Pregnant women, mothers with young children, and their children.

Type of Respondents: Pregnant women, mothers with young children, and their children who are not geographically eligible to enroll in the NCS Vanguard Study.

Annual reporting burden: See Table 1. The annualized cost to respondents is estimated at \$25,000 (based on \$10 per hour). There are no Capital Costs to report. There are no Operating or Maintenance Costs to report.

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN AND COST SUMMARY, CHILD HEALTH DISPARITIES SUBSTUDY

Data colle	ection activity	Type of respondent	Estimated number of respondents	Estimated number of responses per respondent	Average burden per response (in hours)	Estimated total annual burden hours	Estimated total annual respondent cost
Screening for Cog- nitive Interview.	Mothers of children ages 0-5.	Members of NCS target population (not NCS participants).	100	1	5/60	8	\$83
Screening for Pri- mary Data Col- lection.	Women	Members of NCS target population (not NCS participants).	2,000	1	5/60	167	1,667
Screening for Sa- liva Collection.	Women	Members of NCS target population (not NCS participants).	600	1	5/60	50	500
Cognitive Interview	Mothers of children ages 0-5.	Members of NCS target population (not NCS participants).	60	1	75/60	75	750

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN AND COST SUMMARY, CHILD HEALTH DISPARITIES SUBSTUDY—
Continued

Primary Data Col-	Pregnant Women/	Members of NCS tar-	600	2	65/60	1,300	13,000
lection.	Mothers of children ages 0-5.	get population (not NCS participants).					
	Mothers of children ages 0-5.	, , ,	600	1	65/60	650	6,500
Saliva Collection	Pregnant Women/ Mothers of children ages 0–5.	Members of NCS target population (not NCS participants).	200	2	15/60	100	1,000
	Additional mothers of children ages 0–5.	, , ,	200	1	15/60	50	500
	Children ages 0-5		400	1	15/60	100	* 1,000
Total			4,760			2,500	25,000

<sup>\*</sup>The allotted hourly wage rate accounts for the mother's time associated with the data collection activity.

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 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 Management and Budget, Office of Information and Regulatory Affairs, Attn: NIH Desk Officer, by Email 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 Email 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: June 19, 2012.

#### Iamelle E. Banks.

Project Clearance Liaison, Office of Science Policy, Analysis and Communications, National Institute of Child Health and Human Development.

[FR Doc. 2012-16028 Filed 6-28-12; 8:45 am]

BILLING CODE 4140-01-P

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### **National Institutes of Health**

Submission for OMB Review; Comment Request: PHS Applications and Pre-Award Reporting Requirements; Revision

**SUMMARY:** In compliance with the requirement of Section 3507(a)(1)(D) of the Paperwork Reduction Act (PRA) of 1995, the Office of the Director (OD), Office of Extramural Research (OER), 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 March 5, 2012, Volume 77, No. 43, page 13132-13133, and allowed 60 days for public comment. One public comment was received, which asked for clarification about new reporting burdens. It was noted in follow-up that NIH has seen a 21-percent increase in competing applications since the last clearance which has resulted in an increase in the burden hours. We are also transitioning to the Research Performance Progress Report as mandated by OMB.

The purpose of this notice is to allow an additional 30 days for public comment.

Proposed Collection: Title: Public Health Service (PHS) Applications and Pre-award Reporting Requirements. Type of Information Collection Request:

Revision, OMB 0925-0001, Expiration Date 6/30/2012. Form numbers: PHS 398, PHS 416-1, 416-5, and PHS 6031. This collection represents a consolidation of PHS applications and pre-award reporting requirements into a revised data collection under the PRA. Need and Use of Information Collection: This collection includes PHS applications and pre-award reporting requirements: PHS 398 [paper] Public Health Service Grant Application forms and instructions; PHS 398 [electronic] PHS Grant Application component forms and agency-specific instructions used in combination with the SF424 (R&R); PHS Fellowship Supplemental Form and agency-specific instructions used in combination with the SF424 (R&R) forms/instructions for Fellowships [electronic]; PHS 416-1 Ruth L. Kirschstein National Research Service Award (NRSA) Individual Fellowship Application Instructions and Forms used only for a change of sponsoring institution application [paper]; Instructions for a Change of Sponsoring Institution for NRSA Fellowships (F30, F31, F32 and F33) and non-NRSA Fellowships; PHS 416-5 Ruth L. Kirschstein National Research Service Award (NRSA) Individual Fellowship Activation Notice; and PHS 6031 Payback Agreement. The PHS 398 (paper and electronic) is currently approved under 0925-0001; PHS 416-1, 416-5, and PHS 6031 are currently approved under 0925-0002. All forms expire 6/30/2012. Post-award reporting requirements are simultaneously consolidated under 0925-0002, and include the new Research Performance Progress Report (RPPR).

The PHS 398 application is used by applicants to request federal assistance funds for traditional investigator-initiated research projects and to request access to databases and other PHS resources. The PHS 416–1 is used only for a change of sponsoring institution application. PHS Fellowship