

Federal Register Citation of Previous Announcement: 74 FR 32934 (July 9, 2009)

PREVIOUSLY ANNOUNCED TIME AND DATE OF THE MEETING: July 14, 2009—10 a.m.

CHANGE: Withdrawal of Item 3 in the Closed Session.

CONTACT PERSON FOR MORE INFORMATION: Karen V. Gregory, Secretary, (202) 523–5725.

Karen V. Gregory,
Secretary.

[FR Doc. E9–16961 Filed 7–14–09; 11:15 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; Comment Request; Program Review of the Division of Acquired Immunodeficiency Syndrome Policy Implementation Program

AGENCY: National Institutes of Health (NIH).

ACTION: Notice.

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 Allergy and Infectious Diseases (NIAID), the National Institutes of Health (NIH) will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget OMB for review and approval.

Proposed Collection: Title: Review of the DAIDS. Policy Implementation Program (DPIP).

Type of Information Collection Request: New. *Need and Use of Information Collection:* The program review of the Division of AIDS (DAIDS) Implementation Program (DPIP), is to be conducted over a three-year period, and it will provide feedback to aid in the understanding of the target population's knowledge, attitudes, and perceptions of the DAIDS Policy Implementation Program (DPIP). The target population is classified as Extramural Researchers (ERs), who are recipients of funding from DAIDS to conduct and review research. This target population is comprised of Site Leaders of Clinical Research Sites (CRSs) and Research Networks and Clinical Site Monitors of the CTUs and CRSs. The researchers are located globally, and may be part of

more than one DAIDS funded research study and/or network. The DPIP is built upon four goals of awareness and accessibility, understandability, applicability, and harmonization of the policies and procedures. The review is to determine DPIP's progression to fulfillment of its program goals. The results of the review will provide DAIDS' Policy, Training, and Quality Assurance Branch (PTQAB) with information to guide optimal deployment of clinical research policies and procedures intended to harmonize, standardize and improve DAIDS funded/sponsored research. The program review will help derive an understanding of whether the DPIP program is implemented and functioning as intended to meet its program goals. The *Estimated number* of respondent is 392. *Frequency of Response:* Web-based survey; annually (once a year). Focus Group; one time. *Affected Public:* Extramural Researchers *Type of Respondents:* Adult professionals. There are no Capital Costs to report. There are no Operating or Maintenance Costs to report.

The annual reporting burden is provided below:

Type of respondents	Data collection instrument	Estimated frequency of response	Estimated average time per response	Estimated annual hour burden
Extramural Researchers	Survey	3	1	392
Extramural Researchers	Focus Groups	1	2	261

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: Ms. Dione Washington, Policy, Training, and Quality Assurance Branch, National Institute of Allergy and Infectious Diseases, NIH, 6700B Rockledge Drive, MSC 7620, Bethesda, MD 20892–7620 United States of America; or E-mail your request, including your address to: washingtondi@niaid.nih.gov.

Comments Due Date: Written comments and recommendations for the proposed data collection must be mailed within 60 days of this notice.

Dated: July 9, 2009.

Judith Brooks,

Chief, Policy, Training, and Quality Assurance Branch, NIAID, National Institutes of Health.

[FR Doc. E9–16832 Filed 7–15–09; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; Comment Request; Evaluation of the NIAID HIV Vaccine Research Education Initiative

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 Allergy and Infectious Diseases (NIAID), the National Institutes of Health (NIH) will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB) for review and approval.

PROPOSED COLLECTION: Title: Evaluation of the NIAID HIV Vaccine Research Education Initiative. *Type of Information Collection Request:* New.

Need and Use of Information Collection: Developing a vaccine that protects against HIV infection is one of NIAID's highest priorities. To address the need for volunteers in HIV vaccine clinical trials, and enable NIAID to fulfill its Congressional mandate to prevent infectious diseases like HIV/AIDS, NIAID created the NIAID HIV Vaccine Research Education Initiative (NHVREI). The goal of NHVREI is to increase knowledge about and support for HIV vaccine research among U.S. populations most heavily affected by HIV/AIDS—in particular, African Americans, Hispanics/Latinos, men who have sex with men (MSM), women and youth, recognizing the intersection of these groups.

The NHVREI program objectives include (1) Increasing awareness of the need for an HIV vaccine in communities most affected and infected by HIV/AIDS, (2) Improving the public's knowledge of and attitudes toward HIV

vaccine research, (3) Enhancing the partnership between community and HIV clinical trial researchers, and (4) Creating support for current and future HIV vaccine trials and fostering an environment that supports clinical trial volunteers.

To determine if the strategies used to meet these objectives were successful, it is necessary to measure the effectiveness of initiative elements. Specifically, the evaluation will assess (a) implementation of NHVREI (*i.e.*, process evaluation) and (b) impact (*i.e.*, outcomes evaluation) of NHVREI on awareness of, knowledge about, and support for HIV vaccine research among NHVREI primary audiences (*i.e.*, partner organizations, key influencers) that work with target populations. The ultimate goal of evaluation planning and implementation is to determine what components/strategies of NHVREI are effective and impact desired outcomes,

so that these components/strategies can be continued, enhanced, and/or expanded if needed.

Evaluation will be conducted through several processes including a survey and multiple focus groups. A survey will be conducted with key influencers of the NHVREI target populations to measure their level of awareness, knowledge about, and support for HIV vaccine research.

Focus groups will also be conducted with representatives of organizations receiving grants through the NHVREI Local Partnership Program (LPP) and National Partnership Program (NPP), as well as representatives from a broader group of organizations called the NHVREI Network. The LPP and NPP are organizations that are funded to raise awareness about HIV vaccine research at either the local or national level. The planned NHVREI network will be composed of leadership organizations

and coalitions that are either influencers of or provide information services to the target populations. The purpose of conducting focus groups with LPP, NPP, and NHVREI Network representatives is to obtain data on their experience implementing NHVREI activities. Questions asked during the group discussions will address efforts implementing educational activities and developing materials, community partnerships developed, engagement of key influencers in program activities, and the types of media outreach and capacity building engaged in. *Frequency of Response:* Twice. *Affected Public:* Individuals. *Type of Respondents:* Key influencers of target populations. The annual reporting burden is shown in the table below. There are no Capital Costs to report. There are no Operating or Maintenance Costs to report.

Type of respondents	Form name	Estimated number of respondents	Number of responses per respondent	Average burden hours per response	Estimated total annual burden hours requested
LPP, NPP, and NHVREI Network Key Influencers	Time 1 Focus Groups	78	1	1	78
	Survey	656	1	0.33	216
	Total Time 1	734			294
LPP, NPP, and NHVREI Network Key Influencers	Time 2 Focus Groups	78	1	1	78
	Survey	590	1	0.33	195
	Total Time 2	668			273
	Total Time 1 & Time 2	1,402			567

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 Katharine Kripke, Assistant Director, Vaccine Research Program, Division of AIDS, NIAID, NIH, 6700B Rockledge Dr., Bethesda, MD 20892-7628, or call non-toll-free number 301-402-0846, or E-mail your request, including your address to kripkek@niaid.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: July 9, 2009.

John J. McGowan,

*Deputy Director for Science Management,
NIAID, National Institutes of Health.*

[FR Doc. E9-16834 Filed 7-15-09; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission to OMB, Comment Request; An Outcome Evaluation of the NIH Director's Pioneer Award (NDPA) Program

SUMMARY: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the Office of the Director, 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 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.