

In the **Federal Register** of September 9, 2009 (74 FR 46430), FDA published a 60-day notice requesting public comment on the proposed collection of information. No comments were received.

Dated: December 23, 2009.

David Horowitz,

Assistant Commissioner for Policy.

[FR Doc. E9-31199 Filed 1-4-10; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; Comment Request; Process Evaluation of the NIH's Roadmap Interdisciplinary Research Work Group Initiatives

SUMMARY: In compliance with the requirements of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, for opportunity for public comment on proposed data collection projects, the Office of the Director, 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: The National Institute of Dental and Craniofacial Research of the National Institutes of Health requests a three-year clearance for the "Process Evaluation of the NIH Roadmap Interdisciplinary Research Work Group Initiatives," a new collection. This study will be used to determine whether the NIH's Interdisciplinary Research Work Group initiatives have been, and are being, conducted as planned, whether the expected outputs are being produced, and how the activities and processes associated with the initiatives can be improved. Information collected during the evaluation will be used to assess whether and how these initiatives differed from existing initiatives to determine whether these unique initiatives or mechanisms are necessary, to make decisions about whether to continue and/or to modify the programs, and to make decisions about structural or procedural changes within NIH that may be necessary to support cross-cutting interdisciplinary programs. The frequency of response is once for most respondents, and twice for a limited group. The affected public includes a limited number of individuals; *Type of respondents:* principal investigators, other grant investigators, and Initiative trainees. The annual reporting burden is as follows: *Estimated number of*

respondents: 450; *Estimated number of responses per respondent:* PIs, 2; Other Investigators, 1; Trainees, 1; *Average burden hours per response:* 30 minutes; and *Estimated total annual burden hours requested:* 250 hours. The total annualized cost to respondents (calculated as the number of respondents * frequency of response * average time per response * approximate hourly wage rate) is estimated to be \$4,565.

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 Sue Hamann, Ph.D., Science Evaluation Officer, Office of Science Policy Officer and Analysis, NIDCRD, NIH. You may reach Dr. Hamann by telephone on 301-594-4849 (this is not a toll-free number), or you may e-mail your request to Dr. Hamann at Sue.Hamann@nih.hhs.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: December 22, 2009.

Sue Hamann,

Science Evaluation Officer, OSPA, NIDCR, National Institutes of Health.

[FR Doc. E9-31234 Filed 1-4-10; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60 Day-10-0004]

Proposed Data Collections Submitted for Public Comment and Recommendations

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 Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. Alternatively, to obtain a copy of the data collection plans and instrument, call 404-639-5960 and send comments to Maryam I. Daneshvar, CDC Acting Reports Clearance Officer, 1600 Clifton Road NE., MS-D74, Atlanta, Georgia 30333; comments may also be sent by e-mail to omb@cdc.gov.

Comments are invited on (a) whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have a practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of information technology. Written comments should be received within 60 days of this notice.

Proposed Project

National Disease Surveillance Program II. Disease Summaries (0920-0004 Exp. 5/31/2010)—Revision—National Center for Emerging and Zoonotic Infectious Diseases (NCEZID) (proposed), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Surveillance of the incidence and distribution of disease has been an important function of the U.S. Public Health Service (PHS) since 1878. Through the years, PHS/CDC has formulated practical methods of disease control through field investigations. The CDC National Disease Surveillance Program is based on the premise that diseases cannot be diagnosed, prevented, or controlled until existing knowledge is expanded and new ideas developed and implemented. Over the years, the mandate of CDC has broadened to include preventive health