

Dated: February 21, 2003.

Thomas Bartenfeld,

Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-03-44]

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. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call the CDC Reports Clearance Officer on (404) 498-1210.

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 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 automated collection techniques or other forms of information technology. Send comments to Anne O'Connor, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: The Role of Housing in HIV/AIDS Prevention—New—National Center for HIV, STD, and TB Prevention (NCHSTP), Centers for Disease Control and Prevention (CDC).

The Centers for Disease Control and Prevention (CDC) and the Department of Housing and Urban Development (HUD) propose to study the effects of housing for homeless or unstably housed persons on the transmission of HIV and the health of persons living with HIV. Results from the study will be used by policy makers to better understand the types of housing and other affiliated services most likely to reduce HIV transmission and disease progression in the homeless population.

The population to be studied will be drawn from persons living with HIV/AIDS who are seeking housing services from three communities with unmet housing needs as evidenced by a waiting list for services, or other evidence of unmet housing need, through the Housing Opportunities for Persons with AIDS (HOPWA) program. The project will be a longitudinal cohort study, following participants for 18 months. Participants will be randomized into two groups. One group will receive vouchers for housing

subsidies plus a 2-session behavioral intervention; the other group will receive referral to housing resources through participating and other agencies plus the 2-session behavioral intervention. No study participants will be denied access to other housing services that are available through participating agencies or other community resources. Since all participants receive the behavioral intervention, the study technically assesses the effects of housing over and above the behavioral intervention.

A cost study will also be conducted to determine the resources needed for this approach and the cost benefits of providing housing for homeless and unstable housed people living with HIV. The purpose of the cost study is to evaluate the effects of housing affordability and the cost-effectiveness (*i.e.* cost-utility ratio) of this strategy relative to other interventions in other public health and other HIV prevention interventions.

The burden for this collection is estimated to be approximately 90 minutes for the survey at baseline and at 6, 12, and 18 months after baseline and 120 minutes for the interview with HUD site service providers. Blood samples for CD4 and viral load counts will also be collected for all participants. These estimates include the time needed to determine if the respondent is eligible to be interviewed, obtain informed consent, and administer the interview.

There are no costs to respondents for participation in the survey other than their time.

Respondents	Number of respondents	Number of responses/respondent	Average burden/response (in hours)	Total burden (in hours)
HOPWA Program Participants	1000	4	90/60	6000
HUD Site Service Provider	15	1	2	30
Total				6030

Dated: February 21, 2003.

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Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-03-46]

Proposed Data Collections Submitted for Public Comment and Recommendations

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Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call the CDC Reports Clearance Officer on (404) 498-1210.

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 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 automated collection techniques or other forms of information technology. Send comments to Anne O'Connor, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: Restriction on Travel of Persons, (0920-0488)—Extension—National Center for Infectious Diseases (NCID), Centers for Disease Control and Prevention (CDC).

In 2000, the Food and Drug Administration (FDA) and CDC consolidated regulations related to controlling the spread of communicable diseases. FDA formerly administered the regulations contained in part 1240 of Title 21, Code of Federal Regulations, which pertained to interstate control of communicable diseases. These regulations may now be found in part 70 of Title 42, Code of Federal Regulations.

Among the regulations in 21 CFR part 1240, FDA transferred to CDC certain sections that relate to restrictions on interstate travel of any person who is in

the communicable period of cholera, plague, smallpox, typhus, or yellow fever, or who, having been exposed to any such disease, is in the incubation period thereof. One of the sections—formerly 21 CFR 1240.50 and now 42 CFR 70.5 (Certain communicable diseases; special requirements)—contains a requirement for reporting certain information to the Federal government. Specifically, this regulation requires any person who is in the communicable period of cholera, plague, smallpox, typhus or yellow fever, or who, having been exposed to any such disease, is in the incubation period thereof, to apply for and receive a permit from the Surgeon General or his authorized representative in order to travel from one State or possession to another.

Control of disease transmission within the States is considered to be the province of state and local health authorities, with federal assistance being sought by those authorities on a cooperative basis, without application of federal regulations. The regulations formerly administered by FDA and assumed by CDC were developed to facilitate Federal action in the event of large outbreaks requiring a coordinated effort involving several states, or in the

event of inadequate local control. While it is not known whether, or to what extent, situations may arise in which these regulations would be invoked, contingency planning for domestic emergency preparedness is now commonplace. Should this occur, CDC will use the reporting and record-keeping requirements contained in the regulations to carry out quarantine responsibilities as required by law.

Because of the uncertainty about whether a situation will ever arise precipitating CDC's enforcement of this rule, the following data collection burden estimate was prepared using the article *Smallpox: An Attack Scenario*, Tara O'Toole; Emerging Infectious Diseases, Vol. 5, No. 4, Jul-Aug 1999. This article describes the aftermath of a hypothetical domestic public health emergency situation involving smallpox virus. Of the potentially 15,000 persons infected with smallpox, the data collection assumes that one-fourth of these would apply for a permit to move from one state to another while in the communicable period of or having been exposed to smallpox. Should the event be different and/or involve a different number of people, the burden will vary accordingly.

Respondent	Number of responses	Number of responses/respondents	Average burden/response (in hrs.)	Total burden (in hrs.)
Applicants (per application for a permit to move from state to state while in the communicable period of or having been exposed to smallpox	3,750	1	15/60	938
Total	938

Dated: February 21, 2003.

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Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Cooperative Research and Development Agreement

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice.

SUMMARY: The Centers for Disease Control and Prevention (CDC) National Immunization Program (NIP) is seeking

a Cooperative Research and Development Agreement (CRADA) partner for collaboration in the development of a Vaccine Management Software (VACMAN) dynamic link library (DLL) component to interface with immunization information systems. The current DLL is compatible to VACMAN version 2.6x and will not be compatible when VACMAN is upgraded to version 3.

Because CRADAs are designed to facilitate the development of scientific and technological knowledge into useful, marketable products, a great deal of freedom is given to Federal agencies in implementing collaborative research. The CDC may accept staff, facilities, equipment, supplies, and money from the other participants in a CRADA; CDC may provide staff, facilities, equipment, and supplies to the project. CDC may not provide funds to the other participants in a CRADA. This opportunity is available until 30 days

after publication of this notice in the **Federal Register**. Respondents may be provided a longer period of time to furnish additional information if CDC finds this necessary.

FOR FURTHER INFORMATION CONTACT:

Technical: Terry Boyd, Data Management Division, National Immunization Program, Centers for Disease Control and Prevention (CDC), 1600 Clifton Rd., NE., Mailstop E-62, Atlanta, GA 30333, telephone (404) 639-8584.

Business: Janet Kelly, Data Management Division, National Immunization Program, Centers for Disease Control and Prevention (CDC), 1600 Clifton Rd., NE., Mailstop E-62, Atlanta, GA 30333, telephone (404) 639-8735.

SUPPLEMENTARY INFORMATION: The VACMAN application was developed by the CDC NIP and is used by CDC Immunization Grant programs to purchase vaccines through a Vaccine