

burden of diabetes on the border population (Phase 1).

The purpose of the project is to diminish the impact of diabetes on the border population by conducting activities in two related and chronological phases (prevalence study and intervention program). Phase 1, which will assess the prevalence of diabetes, related behavioral risk factors, and assess the health services for the border population, was completed in October 2002. Phase 2 will be implemented in eleven pilot communities, where persons living with diabetes will be randomized to either intervention group participant (IGP) or delayed intervention control group participant (DICGP). The DICGP will receive usual diabetes self management education by the health care provider in

a community health center setting, and the IGP will be assigned to receive diabetes self management education reinforcement and coaching social support at the community home level, by a Community Health Worker/Promotor de Salud (CHW/PdS). These programs will be culturally and linguistically appropriate and will include the participation of community health workers (promotores) and primary healthcare providers working as a team approach.

Activities for Phase 2 will include implementation of community interventions that will provide weekly site visits to the person living with diabetes and provide follow-up and support for the participant and their family. Two family members, found with the highest risk factor ratio will

also be interviewed by the CHW/PdS. The CHW/PdS will reinforce educational messages on balanced nutrition and physical activity and provide social support and coaching to the person living with diabetes and their family members. An equal number of participants will be in the delayed intervention control group. This group and their high risk family members will complete an initial household survey and a final household survey at the end of 18 months. The CHW/PdS will be trained in diabetes and community mobilization skills. The household survey will be repeated in the fifth year of the project for evaluation purposes. There is no cost to respondents. The estimated annualized burden is 3,960 hours.

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hrs)
Intervention group participants	330	2	1
IGP family members	660	2	1
Delayed intervention control group participants	330	2	1
DICGP family members	660	2	1

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Diane Allen,

Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30 Day-56-04]

Proposed Data Collections Submitted for Public Comment and Recommendations

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 498-1210 or send an e-

mail to omb@cdc.gov. Send written comments to CDC Desk Officer, Human Resources and Housing Branch, New Executive Office Building, Room 10235, Washington, DC 20503 or by fax to (202) 395-6974. Written comments should be received within 30 days of this notice.

Proposed Project

Formative Research on Issues Related to the Use of Mass Media in African American Women—New—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background

Women's health programs, including the National Breast and Cervical Cancer Early Detection Program (NBCCEDP), offer low-cost or free breast cancer screening to uninsured, low-income women. In 1991, CDC established the NBCCEDP to increase breast and cervical cancer screening among uninsured, underserved, low-income women. To date, over 1.5 million

women have received services from NBCCEDP-sponsored programs. Yet NBCCEDP-sponsored programs are estimated to reach only 18% of women 50 years old and older who are eligible for screening services. A research priority for the NBCCEDP is to identify effective strategies to increase enrollment among eligible women who have never received breast or cervical cancer screening. Why women do not participate in this screening is not well understood.

The purpose of this task is to conduct formative research to better understand how low-income African-American women might use TV/radio as sources of health information and identify the particular formats, programs, stations, and hours the targeted women listen. This task will examine how African-American women get information on community issues, services, and events and determine if these can be used as viable means to disseminate information on health services. The only cost to respondent is their time. The estimated annualized burden is 240 hours.

Form name	No. of respondents	No. of responses per respondent	Average burden per response (in hrs)
Call-in Script	120	1	5/60
Eligibility Screener	120	1	5/60
Check-in and Informed Consent	120	1	5/60
Pre-discussion Information Sheet	120	1	15/60

Form name	No. of respondents	No. of responses per respondent	Average burden per response (in hrs)
Group Discussion	120	1	1.5

Dated: June 21, 2004.

Diane Allen,

Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

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[30Day-58-04]

Proposed Data Collections Submitted for Public Comment and Recommendations

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Resources and Housing Branch, New Executive Office Building, Room 10235, Washington, DC 20503 or by fax to (202) 395-6974. Written comments should be received within 30 days of this notice.

Proposed Project

Survey of Veterinarians to Assess Infection Control Practices and Use of Personal Protective Equipment to Reduce Transmission of Zoonotic Diseases—New—National Center for Infectious Diseases (NCID), Centers for Disease Control and Prevention (CDC).

Recent outbreaks of emerging zoonotic diseases in the United States have highlighted the need to better protect the veterinary community from infectious diseases by educating them about personal protective measures. In particular, during the recent 2003 outbreak of monkeypox in the United States associated with prairie dogs and imported rodents, veterinarians or veterinary staff represented over 25% of confirmed and probable human cases. During the height of this outbreak, health officials were tasked with providing information to the medical and veterinary communities to ensure

safety when examining monkeypox-infected patients; a lack of universally accepted infection control and personal protection guidelines within the veterinary community hampered the delivery of effective prevention messages to this vulnerable population.

The proposed survey asks veterinarians about infection control procedures employed in their clinics and the use of personal protective equipment to prevent zoonotic disease transmission.

The proposed study consists of a self-administered, written questionnaire mailed to veterinary clinics in the United States. The American Veterinary Medical Association has volunteered to collaborate on the survey and will provide a list of clinics through their membership mailing list. The study objectives are to describe current knowledge, attitudes, and practices of veterinarians regarding zoonotic disease risks and protection of veterinary clinic staff, and to determine what types of national guidelines on infection control practices in veterinary settings are needed. The estimated annualized burden is 417 hours.

Respondents	Number of respondents	Number of responses/ respondent	Average burden/response (in hours)
Written Questionnaires	2,500	1	10/60

Dated: June 21, 2004.

Diane Allen,

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-57-04]

Proposed Data Collections Submitted for Public Comment and Recommendations

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and

Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 498-1210 or send an e-mail to omb@cdc.gov. Send written comments to CDC Desk Officer, Human Resources and Housing Branch, New Executive Office Building, Room 10235, Washington, DC 20503 or by fax to (202) 395-6974. Written comments should be received within 30 days of this notice.

Proposed Project

NCHS Application for Vital Statistics Training Form, OMB No. 0920-0217—Extension—National Center for Health Statistics (NCHS), Centers for Disease Control and Prevention (CDC). In the United States, legal authority for the registration of vital events, *i.e.*, births, deaths, marriages, divorces, fetal deaths, and induced terminations of pregnancy, resides individually with the States (as

well as cities in the case of New York City and Washington, DC) and Puerto Rico, the Virgin Islands, Guam, American Samoa, and the Commonwealth of the Northern Mariana Islands. These governmental entities are the full legal proprietors of vital records and the information contained therein. As a result of this State authority, the collection of registration-based vital statistics at the national level, referred to as the U.S. National Vital Statistics System (NVSS), depends on a cooperative relationship between the States and the Federal government. This data collection, authorized by 42 U.S.C. 242k, has been conducted by NCHS since it was created in 1960.

NCHS assists in achieving the comparability needed for combining data from all States into national statistics, by conducting a training program for State and local vital