DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Agency Forms Undergoing Paperwork Reduction Act Review

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 requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995. To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639-5960 or send an e-mail to omb@cdc.gov. Send written comments to CDC Desk Officer, Office of Management and Budget, Washington, DC or by fax to (202) 395-5806. Written comments should be received within 30 days of this notice.

Proposed Project


Background and Brief Description

Section 306 of the Public Health Service (PHS) Act (42 U.S.C. 242k), as amended, authorizes that the Secretary of Health and Human Services (DHHS), acting through NCHS, shall undertake and support (by grant or contract) research, demonstrations, and evaluations respecting new or improved methods for obtaining current data to support statistical and epidemiological activities for the purpose of improving the effectiveness, efficiency, and quality of health services in the United States.

The Questionnaire Design Research Laboratory (QDRL) conducts questionnaire pre-testing and evaluation activities for CDC surveys (such as the NCHS National Health Interview Survey, OMB No. 0920–0214) and other federally sponsored surveys. NCHS is requesting 3 years approval of OMB for this extension.

The QDRL conducts cognitive interviews, focus groups, mini field-pretests, and experimental research in laboratory and field settings, both for applied questionnaire evaluation and more basic research on response errors in surveys.

The most common questionnaire evaluation method is the cognitive interview. In a cognitive interview, a questionnaire design specialist interviews a volunteer participant. The interviewer administers the draft survey questions as written, but also probes the participant in depth about interpretations of questions, recall processes used to answer them, and adequacy of response categories to express answers, while noting points of confusion and errors in responding. Interviews are generally conducted in small rounds of 20–30 interviews.

Similar methodology has been adopted by other federal agencies, as well as by academic and commercial survey organizations. There are no costs to respondents other than their time. The total estimated annualized burden hours are 625 hours.

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Type of forms</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
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Maryam I. Daneshvar,
Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

[FR Doc. 2010–1166 Filed 1–21–10; 8:45 am]
System (OMB #0920–0612, exp. 1/31/2010)—Revision—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description
Cardiovascular disease (CVD), which includes heart disease, myocardial infarction, and stroke, is the leading cause of death for women in the United States, and is largely preventable. The WISEWOMAN program (Well-Integrated Screening and Evaluation for Women Across the Nation), administered by the Centers for Disease Control and Prevention (CDC), was established to examine ways of improving the delivery of services for women who have limited access to health care and elevated risk factors for CVD. The program focuses on reducing CVD risk factors and provides screening services for select risk factors such as elevated blood cholesterol, hypertension and abnormal blood glucose levels. The program also provides lifestyle interventions and medical referrals. The WISEWOMAN program serves women who are participating in the National Breast and Cervical Cancer Early Detection Program (NBCCEDP), also administered by CDC. CDC requests OMB approval to continue collecting information from WISEWOMAN grantees for three years, with changes. There will be a net decrease in the total annualized burden hours. Although the number of funded grantees will increase from 15 to 21, the burden per respondent will decrease due to changes in the data collection plan and schedule. The collection of cost information will be discontinued and the Progress Report will be collected semi-annually instead of quarterly.

Twice per year, each grantee will electronically transmit a Minimum Data Elements (MDE) dataset that contains information about the women served through the WISEWOMAN program, including their demographics, health status, CVD risk factors, referrals and participation in lifestyle interventions. The progress reports provide a narrative summary of grantee activities, as well as a discussion of each grantee’s progress toward meeting stated programmatic objectives. The information collected from grantees is used to assess the impact of the WISEWOMAN program. The overall program evaluation is designed to demonstrate how WISEWOMAN can obtain more complete health data on vulnerable populations, promote public education about disease incidence and risk-factors, improve the availability of screening and diagnostic services for under-served women, ensure the quality of services provided to under-served women, and develop strategies for improved interventions.

There are no costs to respondents other than their time. The total estimated annualized burden hours are 1,680.

**ESTIMATED ANNUALIZED BURDEN HOURS**

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<th>Type of respondent</th>
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<td>2</td>
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</table>

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BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration
[Docket No. FDA–2009–N–0246]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Establishing and Maintaining a List of U.S. Dairy Product Manufacturers/Processors With Interest in Exporting to Chile

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

DATES: Fax written comments on the collection of information by February 22, 2010.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202–395–7285, or e-mailed to oira_submission@omb.eop.gov. All comments should be identified with the OMB control number 0910–0509. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Denver Presley Jr., Office of Information Management (HFA–710), Food and Drug Administration, 5600 Fisher Lane, Rockville, MD 20857, 301–796–3793.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

In addition, each grantee will submit two written progress reports per year. The progress reports provide a narrative summary of grantee activities, as well as a discussion of each grantee’s progress toward meeting stated programmatic objectives. The information collected from grantees is used to assess the impact of the WISEWOMAN program. The overall program evaluation is designed to demonstrate how WISEWOMAN can obtain more complete health data on vulnerable populations, promote public education about disease incidence and risk-factors, improve the availability of screening and diagnostic services for under-served women, ensure the quality of services provided to under-served women, and develop strategies for improved interventions.

There are no costs to respondents other than their time. The total estimated annualized burden hours are 1,680.

Establishing and Maintaining a List of U.S. Dairy Product Manufacturers/Processors With Interest in Exporting to Chile (OMB Control Number 0910–0509)—Extension

As a direct result of discussions that have been adjunct to the U.S./Chile Free Trade Agreement, Chile has recognized FDA as the competent U.S. food safety authority and has accepted the U.S. regulatory system for dairy inspections. Chile has concluded that it will not require individual inspections of U.S. firms by Chile as a prerequisite for trade, but will accept firms identified by FDA as eligible to export to Chile. Therefore, in the Federal Register of June 22, 2005 (70 FR 36190), FDA announced the availability of a revised guidance document entitled “Establishing and Maintaining a List of U.S. Dairy Product Manufacturers/Processors With Interest in Exporting to Chile.” The guidance can be found at http://www.cfsan.fda.gov/guidance.html. The guidance document explains that FDA has established a list that is provided to the government of Chile and posted on http://