

Government and who have expertise regarding issues of minority health. To qualify for consideration of appointment to the Committee, an individual must possess demonstrated experience and expertise working on issues impacting the health of racial and ethnic minority populations. The Committee charter stipulates that the racial and ethnic minority groups shall be equally represented on the Committee membership. OMH is seeking candidates who can represent the health interest of Hispanics/Latino Americans; Blacks/African Americans; American Indians and Alaska Natives; and/or Asian Americans, Native Hawaiians, and other Pacific Islanders.

Mandatory Professional/Technical Qualifications: Nominees must meet all of the following mandatory qualifications to be eligible for consideration.

(1) Expertise in minority health and racial and ethnic health disparities.

(2) Expertise in developing or contributing to the development of science-based or evidence based health policies and/or programs. This expertise may include experience in the analysis, evaluation, and interpretation of federal/state health or regulatory policy.

(3) Involvement in national, state, regional, tribal, and/or local efforts to improve the health status or outcomes among racial and ethnic minority populations.

(4) Educational achievement, professional certification(s) in health-related fields (e.g., health professions, allied health, behavioral/mental health, public health, health policy, health administration/management, etc.), and professional experience that will support ability to give expert advice on issues related to improving minority health and eliminating racial and ethnic health disparities.

(5) Expertise in population level health data for racial and ethnic minority groups. This expertise may include survey, administrative, and/or clinical data.

Desirable Qualifications:

(1) Knowledge and experience in health care systems, cultural and linguistic competency, social determinants of health, evidence-based research, data collection (e.g., federal, state, tribal, or local data collection), or health promotion and disease prevention.

(2) Nationally recognized via peer-reviewed publications, professional awards, advanced credentials, or involvement in national professional organizations.

Requirements for Nomination

Submission: Nominations should be

typewritten (one nomination per nominator). Nomination package should include: (1) a letter of nomination that clearly states the name and affiliation of the nominee, the basis for the nomination (i.e., specific attributes which qualify the nominee for service in this capacity), and a statement from the nominee indicating a willingness to serve as a member of the Committee; (2) the nominee's contact information, including name, mailing address, telephone number, and email address; (3) the nominee's curriculum vitae, and (4) a summary of the nominee's experience and qualification relative to the mandatory professional and technical criteria listed above. Federal employees should not be nominated for consideration of appointment to this Committee.

Individuals selected for appointment to the Committee shall be invited to serve four-year term. Committee members will receive a stipend for attending Committee meetings and conducting other business in the interest of the Committee, including per diem and reimbursement for travel expenses incurred.

The Department makes every effort to ensure that the membership of HHS federal advisory committees is fairly balanced in terms of points of view represented and the committee's function. Every effort is made to ensure that a broad representation of geographic areas, females, racial and ethnic and minority groups, and the disabled are given consideration for membership on HHS federal advisory committees. Appointment to this Committee shall be made without discrimination because of a person's race, color, religion, sex (including pregnancy), national origin, age, disability, or genetic information. Nominations must state that the nominee is willing to serve as a member of ACMH and appears to have no conflict of interest that would preclude membership. An ethics review is conducted for each selected nominee. Therefore, individuals selected for nomination will be required to provide detailed information concerning such matters as financial holdings, consultancies, and research grants or contracts to permit evaluation of possible sources of conflict of interest.

Dated: March 26, 2013.

Monica A. Baltimore,

Executive Director, Advisory Committee on Minority Health.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-13-12EG]

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 Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639-7570 or send an email 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

Use of Smartphones to Collect Information about Health Behaviors: Feasibility Study—New—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Despite the high level of public knowledge about the adverse effects of smoking, tobacco use remains the leading preventable cause of disease and death in the U.S., resulting in approximately 443,000 deaths annually. During 2005-2010, the overall proportion of U.S. adults who were current smokers declined from 20.9% to 19.3%. Despite this decrease, smoking rates are still well above Healthy People 2020 targets for reducing adult smoking prevalence to 12%, and the decline in prevalence was not uniform across the population. Timely information on tobacco usage is needed for the design, implementation, and evaluation of public health programs.

New mobile communications technologies provide a unique opportunity for innovation in public health surveillance. Text messaging and smartphone Web access are immediate, accessible, and anonymous, a combination of features that could make smartphones ideal for the ongoing research, surveillance, and evaluation of risk behaviors and health conditions, as well as targeted dissemination of information.

CDC proposes to conduct a feasibility study to evaluate the process of conducting Web surveys by smartphone

and text message surveys by feature phone (cell phones that do not have Web access), the outcomes of the surveys, and the value of the surveys. The universe for this study is English-speaking U.S. residents aged 18–65. The sample frame will consist of a national random digit dial sample of telephone numbers from a frame of known cell phone exchanges. Respondents reached on their cell phones will be asked to complete an initial CATI survey consisting of a short series of simple demographic questions, general health questions, and questions about tobacco and alcohol use. At the conclusion of this brief survey, respondents who have smartphones will be asked to participate in the feasibility study, which consists

of a first follow-up survey and, a week later, a second follow-up survey. Those who agree will receive invitations to participate by text message, which will include a link to the survey. A sample of respondents who have feature phones will be asked to participate in a text message pilot, which also consists of a first follow-up survey and a second follow-up survey. Text message respondents will receive a text message inviting them to participate; respondents who opt in will receive text messages with one survey question at a time. Before initiating the feasibility study, CDC will conduct a brief pre-test of information collection forms and procedures.

This study will evaluate: (1) Response bias of a smartphone health survey by

comparing data collected via CATI to data collected via smartphones/text messages, and data collected via smartphones to data collected via text messages; (2) relative cost-effectiveness of data collected via CATI to data collected via smartphones/text messages; (3) coverage bias associated with restricting the sample to smartphone users; and (4) the utility of smartphones for completing frequent, short interviews (e.g., diary studies to track activities or events).

OMB approval is requested for one year. Participation is voluntary. There are no costs to respondents other than their time. The total estimated annualized burden hours are 306.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hr)
Adults Aged 18 to 65, All cell phone users	Pre-test (CATI Screener/CATI Recruitment ...	20	1	8/60
	Screener/CATI Recruitment	1,990	1	1/60
	Initial CATI Survey	1,590	1	7/60
Adults Aged 18 to 65, Smartphone Users	First Web Survey Follow-up for Smartphone Users.	700	1	3/60
	Second Web Survey Follow-up for Smartphone Users.	595	1	3/60
Adults Aged 18 to 65, Non-smartphone Users	First Text Message Survey Follow-up for non-Smartphone Users.	200	1	3/60
	Second Text Message Survey Follow-up for non-Smartphone Users.	170	1	3/60

Ron A. Otten,

Director, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-13-0600]

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 Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call (404) 639-7570 or send an email to omb@cdc.gov. Send written comments to CDC Desk Officer, Office of Management and Budget, Washington,

DC 20503 or by fax to (202) 395-5806. Written comments should be received within 30 days of this notice.

Proposed Project

CDC Model Performance Evaluation Program (MPEP) for Mycobacterium tuberculosis and Nontuberculous Mycobacteria Drug Susceptibility Testing OMB # 0920-0600 (exp. 5/31/2013).—Revision—National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention (NCHHSTP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

As part of the continuing effort to support domestic public health objectives for treatment of tuberculosis (TB), prevention of multi-drug resistance, and surveillance programs, CDC is requesting approval from the Office of Management and Budget to continue data collection from participants in the Model Performance Evaluation Program for Mycobacterium tuberculosis and Non-tuberculous Mycobacterium Drug Susceptibility Testing. This request includes (a)

changing the title of the data collection to “CDC Model Performance Evaluation (MPEP) for Mycobacterium tuberculosis Drug Susceptibility Testing” to reflect that nontuberculous mycobacteria are no longer included in the test package; (b) replacement of Laboratory Enrollment Form with a Participant Biosafety Compliance Letter of Agreement; (c) revision of the Pre-shipment Email; (d) addition of Instructions to Participants Letter; (e) revision of the MPEP M. tuberculosis Results Worksheet; (f) entering survey results online using a modified data collection instrument; (g) modification of Reminder Email; (h) modification of Reminder Telephone Script; and (i) modification of the Aggregate Report Letter.

While the overall number of cases of TB in the U.S. has decreased, rates still remain high among foreign-born persons, prisoners, homeless populations, and individuals infected with HIV in major metropolitan areas. To reach the goal of eliminating TB, the Model Performance Evaluation Program for Mycobacterium tuberculosis and