Coordinator for Health Information Technology;
(2) Shall have complied with all the requirements under this section;
(3) In the case of a private entity, shall be incorporated in and maintain a primary place of business in the United States, and in the case of an individual, whether participating singly or in a group, shall be a citizen or permanent resident of the United States; and
(4) May not be a Federal entity or Federal employee acting within the scope of their employment.

An individual or entity shall not be deemed ineligible because the individual or entity used Federal facilities or consulted with Federal employees during a competition if the facilities and employees are made available to all individuals and entities participating in the competition on an equitable basis.

Registered participants shall be required to agree to assume any and all risks and waive claims against the Federal Government and its related entities, except in the case of willful misconduct, for any injury, death, damage, or loss of property, revenue, or profits, whether direct, indirect, or consequential, arising from or related to competition, whether the injury, death, damage, or loss arises through negligence or otherwise.

Participants shall be required to obtain liability insurance or demonstrate financial responsibility, in amounts determined by the head of the Office of the National Coordinator for Health Information Technology, for claims by—

(1) A third party for death, bodily injury, or property damage, or loss resulting from an activity carried out in connection with participation in a competition, with the Federal Government named as an additional insured under the registered participant’s insurance policy and registered participants agreeing to indemnify the Federal Government against third party claims for damages arising from or related to competition activities; and
(2) The Federal Government for damage or loss to Government property resulting from such an activity.

Participants must be teams of at least two people.

All participants are required to provide written consent to the rules upon or before submitting an entry.

DATES:

Phase I
- Submission Period Begins: 12:01am, EDT, July 15th, 2011.
- Submission Period for Initial Entries Ends: 11:59 pm, EDT, August 26th, 2011.

Phase II
- Final Submission Period Begins: 12:01am, E.D.T., October 3rd, 2011.
- Final Submission Period Ends: 11:59pm, E.S.T., November 18th, 2011.

Registration Process for Participants
To register for this challenge participants should:
- Access the ONC Investing in Innovation (i2) Challenge Web site at:
  - http://www.health2challenge.org/
category/onc/.
  - A registration link for the challenge can be found on the landing page under the challenge description.

Amount of the Prize
Phase I
- Four semi-finalists will receive prizes of $10,000 each.

Phase II
- Two finalist teams will receive awards of $20,000 each.

Awards may be subject to Federal income taxes and HHS will comply with IRS withholding and reporting requirements, where applicable.

Basis Upon Which Winner Will Be Selected
The judging panel will make selections based upon the following criteria:
1. Impact on cancer prevention and control.
2. Use of cancer-related data.
3. Integration with existing health records.
4. Innovation.
5. Usability.

Additional Information
Ownership of intellectual property is determined by the following:
- Each entrant retains title and full ownership in and to their submission. Entrants expressly reserve all intellectual property rights not expressly granted under the challenge agreement.
- By participating in the challenge, each entrant hereby irrevocably grants to Sponsor and Administrator a limited, non-exclusive, royalty free, worldwide, license and right to reproduce, publicly perform, publicly display, and use the Submission to the extent necessary to administer the challenge, and to publicly perform and publicly display the Submission, including, without limitation, for advertising and promotional purposes relating to the challenge.

Dated: July 18, 2011

Farzad Mostashari,
National Coordinator for Health Information Technology.

[FR Doc. 2011–18728 Filed 7–25–11; 8:45 am]
BILLING CODE 4150–45–

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[[60Day–11–0572]

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 404–639–5960 and send comments to, Daniel Holcomb, CDC Reports Clearance Officer, 1600 Clifton Road, MS–D74, Atlanta, GA 30333 or send an 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 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. Written comments should be received within 60 days of this notice.

Proposed Project

Health Message Testing System, (0920–0572, exp. 11/30/2011)—
Revision—Office of the Associate Director for Communication, Centers for Disease Control and Prevention (CDC).
Background and Brief Description

Before CDC disseminates a health message to the public, the message always undergoes scientific review. However, even though the message is based on sound scientific content, there is no guarantee that the public will understand a health message or that the message will move people to take recommended action. Communication theorists and researchers agree that for health messages to be as clear and influential as possible, target audience members or representatives must be involved in developing the messages and provisional versions of the messages must be tested with members of the target audience.

However, increasingly there are circumstances when CDC must move swiftly to protect life, prevent disease, or calm public anxiety. Health message testing is even more important in these instances, because of the critical nature of the information need.

CDC receives a mandate from Congress with a tight deadline for communicating with the public about a specific topic. For example, Congress gave CDC 120 days to develop and test messages for a public information campaign about Helicobacter pylori, a bacterium that can cause stomach ulcers and increase cancer risk if an infected individual is not treated with antibiotics.

In the interest of timely health message dissemination, many programs forgo the important step of testing messages on dimensions such as clarity, salience, appeal, and persuasiveness (i.e., the ability to influence behavioral intention). Skipping this step avoids the delay involved in the standard OMB review process, but at a high potential cost. Untested messages can waste communication resources and opportunities because the messages can be perceived as unclear or irrelevant. Untested messages can also have unintended consequences, such as jeopardizing the credibility of Federal health officials.

The Health Message Testing System (HMTS), a generic information collection, will enable programs across CDC to collect the information they require in a timely manner to:

- Ensure quality and prevent waste in the dissemination of health information by CDC to the public.
- Refine message concepts and to test draft materials for clarity, salience, appeal, and persuasiveness to target audiences.
- Guide the action of health communication officials who are responding to health emergencies, Congressionally-mandated campaigns with short timeframes, media-generated public concern, time-limited communication opportunities, trends, and the need to refresh materials or dissemination strategies in an ongoing campaign.

Each testing instrument will be based on specific health issues or topics. Although it is not possible to develop one instrument for use in all instances, the same kinds of questions are asked in most message testing. This package includes generic questions and formats that can be used to develop health message testing data collection instruments. These include a list of screening questions, comprised of demographic and introductory questions, along with other questions that can be used to create a mix of relevant questions for each proposed message testing data collection method. However, programs may request to use additional questions if needed.

Message testing questions will focus on issues such as comprehension, impressions, personal relevance, content and wording, efficacy of response, channels, and spokesperson/sponsor. Such information will enable message developers to enhance the effectiveness of messages for intended audiences.

Data collection methods proposed for HMTS include intercept interviews, telephone interviews, focus groups, online surveys, and cognitive interviews. In almost all instances, data will be collected by outside organizations under contract with CDC.

There is no cost to the respondents other than their time.

### TABLE A12A—ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<tr>
<th>Data collection methods</th>
<th>Number of respondents per method</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Central Location Intercept Interviews, Telephone Interviews, Individual In-depth Interview (Cognitive Interviews), Focus Group Screenings, Focus Groups, Online Surveys ..................................................</td>
<td>18,525</td>
<td>1</td>
<td>8/60</td>
<td>2,470</td>
</tr>
<tr>
<td>Total ...............................................................................................................</td>
<td>18,525</td>
<td></td>
<td></td>
<td>2,470</td>
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</tbody>
</table>

Dated: July 19, 2011.

Daniel Holcomb,
Reports Clearance Officer, Centers for Disease Control and Prevention.

[FR Doc. 2011–18807 Filed 7–25–11; 8:45 am]
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–11–11FK]

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) 639–5960 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