# Estimated Costs to the Federal Government

The total cost to the government for this activity is estimated to be \$1,000,000.00.

# **Request for Comments**

In accordance with the above-cited legislation, comments on AHRQ's information collection are requested with regard to any of the following: (a) Whether the proposed collection of information is necessary for the proper performance of health care research and information dissemination functions of AHRQ, including whether the information will have practical utility; (b) the accuracy of AHRQ's estimate of burden (including hours and costs) of the proposed collection(s) 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 upon the respondents, including the use of automated collection techniques or other forms of information technology.

Comments submitted in response to this notice will be summarized and included in the request for OMB approval of the proposed information collection. All comments will become a matter of public record.

#### References

<sup>1</sup>Brennan TA, Leape LL, Laird NM, et al. Incidence of adverse events and negligence in hospitalized patients: Results of the Harvard Medical Practice Study. N Engl J Med 1991; 324:370–376.

<sup>2</sup> McDonald CJ, Weiner M, Hui SL. Deaths due to medical errors are exaggerated in the Institute of Medicine Report. JAMA 2000; 284:93–95.

<sup>3</sup>Leape LL. Institute of Medicine medical error figures are not exaggerated. JAMA. 2000; 28:95–97.

<sup>4</sup> Hayward RA, Hofer TP. Estimating hospital deaths due to medical errors: preventability is in the eye of the reviewer. JAMA. 2001; 286:415–420.

<sup>5</sup> Institute of Medicine. To Err is Human: Building a Safer Health System. Washington, DC: National Academy Press, 2000.

<sup>6</sup> Institute of Medicine. Crossing the Quality Chasm: A New System for the 21st Century. Washington, DC: National Academy Press, 2001.

<sup>7</sup> Institute of Medicine. Patient Safety: Achieving a New Standard for Care. Washington, DC: National Academy Press, 2004.

<sup>8</sup> http://www.blsmeetings.net/

*PatientSafetyandHIT/* (Accessed August 11, 2005).

<sup>9</sup> Green LA, Fryer GE, Yawn BP, Lanier D, Dovey SM: The ecology of medical care revisited. N Engl J Med 2001; 344:2021–2025.

<sup>10</sup> Uribe CL, Schweikhart SB, Pathak DS, Dow M, Marsh GB. Perceived barriers to medical-error reporting: an exploratory investigation. J Healthcare Management. 2002;47(4):263–79.

Dated: December 1, 2006. **Carolyn M. Clancy,**  *Director.* [FR Doc. 06–9643 Filed 12–11–06; 8:45 am] **BILLING CODE 4160–90–M** 

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

# [30Day-07-0008]

# 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 or by fax to (202) 395–6974. Written comments should be received within 30 days of this notice.

## **Proposed Project**

Emergency Epidemic Investigations (0920–0008)—Revision—Office of Workforce and Career Development (OWCD), Centers for Disease Control and Prevention (CDC).

## Background and Brief Description

One of the objectives of CDC's epidemic services is to provide for the prevention and control of epidemics and protect the population from public health crises such as man made or natural biological disasters and chemical emergencies. This objective is carried out, in part, by training investigators, maintaining laboratory capabilities for identifying potential problems, collecting and analyzing data, and recommending appropriate actions to protect the public's health. When state, local, or foreign health authorities request help in controlling an epidemic or solving other health problems, CDC dispatches skilled epidemiologists from the Epidemiologist Intelligence Service (EIS) to investigate and resolve the problem.

The purpose of the Emergency Epidemic Investigation surveillance is to collect data on the conditions surrounding and preceding the onset of a problem. The data must be collected in a timely fashion so that information

can be used to develop prevention and control techniques, to interrupt disease transmission and to help identify the cause of an outbreak. Since the events necessitating the collections of information are of an emergency nature, most data collection is done by direct interview or written questionnaire and are one-time efforts related to a specific outbreak or circumstance. If during the emergency investigation, the need for further study is recognized, a project is designed and separate OMB clearance is required. Interviews are conducted to be as unobtrusive as possible and only the minimal information necessary is collected. The Emergency Epidemic Investigations is the principal source of data on outbreaks of infectious and noninfectious diseases, injuries, nutrition, environmental health and occupational problems.

Each investigation does contribute to the general knowledge about a particular type of problem or emergency, so that data collections are designed to take into account similar situations in the past. Some questionnaires are standardized, such as investigations of outbreaks aboard aircraft or cruise vessels.

The Emergency Epidemic Investigations provides a range of data on the characteristics of outbreaks and those affected by them. Data collected include demographic characteristics, exposure to the causative agent(s), transmission patterns and severity of the outbreak on the affected population. These data, together with trend data, may be used to monitor the effects of change in the health care system, planning of health services, improving the availability of medical services and assessing the health status of the population.

<sup>1</sup> Users of the Emergency Epidemic Investigations data include, but are not limited to EIS Officers in investigating the patterns of disease or injury, investigating the level of risky behaviors, identifying the causative agent and identifying the transmission of the condition and the impact of interventions.

Epi Trip Reports are delivered to the state health agency official requesting assistance shortly after completion of the Emergency Epidemic Investigation. The official can comment on both the timeliness and the practical utility of the recommendations from the investigation. CDC is requesting that a new form be added to the current clearance. Upon completion of the Emergency Epidemic Investigation, requesting officials at the state or local health department will be asked to complete a brief questionnaire to assess the promptness of the investigation and the usefulness of the recommendations.

The total burden hours are 3,775. This slight increase over the last request for clearance is due to additional data that will be collected from the requesting state or local officials described above. *Estimated Annualized Burden Table:* 

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
General Public	15,000	1	15/60
State and Local Officials	100		15/60

Dated: December 6, 2006.

#### Joan F. Karr,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention. [FR Doc. E6–21117 Filed 12–11–06; 8:45 am] BILLING CODE 4163–18–P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

#### [30Day-07-0603]

# 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) 639–4766 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) 395–6974. Written comments should be received within 30 days of this notice.

## **Proposed Project**

Information Network (REACH IN)– Extension-National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

# Background and Brief Description

Racial and Ethnic Approaches to Community Health 2010 (REACH 2010) currently funds forty local coalitions to establish community based programs and culturally appropriate interventions to eliminate racial and ethnic health disparities. Two previously funded grantees also retain access to the system. Communities served by REACH 2010 include: African American, American Indian, Hispanic American, Asian American, and Pacific Islander. These communities select among infant mortality, deficits in breast and cervical cancer screening and management, cardiovascular diseases, diabetes, HIV/ AIDS, and deficits in childhood and adult immunizations to focus their interventions. Guided by logic models, each community articulates goals, objectives, and related activities; tracks whether goals and objectives are met, ongoing, or revised; and evaluates all program activities. This information is then entered into the REACH Information Network (REACH IN). REACH IN is a customized internetbased support system that allows REACH 2010 grantees to perform remote data entry and retrieval of data.

This support system is designed to create on-demand graphs and reports of grantees' activities and accomplishments, monitor progress toward the achievement of goals and objectives, and share and synthesize information across grantees' activities. Both quantitative and qualitative analyses can be performed. These analyses relate primarily to three stages of the REACH 2010 logic model: capacity building, targeted actions (interventions), and community and systems change and change among change agents. Users are supported with technical assistance and training, covering the usage of the system from a content/project goals perspective, and technical operations.

The annualized estimated burden is based on 42 respondents, including 40 currently funded grantees and two that were funded previously who retain access to the system. It is estimated that they each use the system four times a year to enter data, each data entry taking about 30 minutes.

There are no costs to the respondents other than their time. The total estimated annualized burden hours are 84.

Estimated Annualized Burden Table:

Type of responses or kinds of respondents	Nunber of respondents	Number of responses per respondent	Average burden per response (in hours)
REACH 2010 grantees	42	4	30/60

Dated: December 6, 2006.

## Deborah Holtzman,

Reports Clearance Officer, Centers for Disease Control and Prevention. [FR Doc. E6–21118 Filed 12–11–06: 8:45 am]

BILLING CODE 4163–18–P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

[60Day-07-07AC]

## 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 Seleda Perryman, CDC Assistant Reports Clearance