#### FOR FURTHER INFORMATION CONTACT:

National Vaccine Program Office, U.S. Department of Health and Human Services, Room 715–H, Hubert H. Humphrey Building, 200 Independence Avenue SW., Washington, DC 20201. *Phone:* (202) 690–5566; *Fax:* (202) 690–4631; *email: nvpo@hhs.gov.* 

**SUPPLEMENTARY INFORMATION: Pursuant** to Section 2101 of the Public Health Service Act (42 U.S.C. 300aa-1), the Secretary of Health and Human Services was mandated to establish the National Vaccine Program to achieve optimal prevention of human infectious diseases through immunization and to achieve optimal prevention against adverse reactions to vaccines. The National Vaccine Advisory Committee was established to provide advice and make recommendations to the Director of the National Vaccine Program on matters related to the Program's responsibilities. The Assistant Secretary for Health serves as Director of the National Vaccine Program.

The topics to be discussed at the NVAC meeting will include seasonal influenza, implementation of the National Vaccine Plan, and vaccine research and development. The meeting agenda will be posted on the NVAC Web site: http://www.hhs.gov/nvpo/ nvac prior to the meeting. Public attendance at the meeting is limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the National Vaccine Program Office at the address/phone listed above at least one week prior to the meeting. Members of the public will have the opportunity to provide comments at the NVAC meeting during the public comment periods on the agenda. Individuals who would like to submit written statements should email or fax their comments to the National Vaccine Program Office at least five business days prior to the meeting.

Dated: May 1, 2012.

### Bruce Gellin,

Director, National Vaccine Program Office, Executive Secretary, National Vaccine Advisory Committee.

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BILLING CODE 4150-44-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

[30Day-12-12IL]

# 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**

NIOSH Generic Clearance for the Collection of Qualitative Feedback on Agency Service Delivery—NEW—Centers for Disease Control and Prevention (CDC), National Institute for Occupational Safety and Health (NIOSH), Health Hazard Evaluation Program.

As part of a Federal Government-wide effort to streamline the process to seek feedback from the public on service delivery, the CDC has submitted a Generic Information Collection Request (Generic ICR): "Generic Clearance for the Collection of Qualitative Feedback on Agency Service Delivery" to OMB for approval under the Paperwork Reduction Act (PRA) (44 U.S.C. 3501 et sea.).

seq.).
To request additional information, please contact Kimberly S. Lane, Reports Clearance Officer, Centers for Disease Control and Prevention, 1600 Clifton Road, MS–D74, Atlanta, GA 30333 or send an email to omb@cdc.gov.

## SUPPLEMENTARY INFORMATION:

Title: NIOSH Generic Clearance for the Collection of Qualitative Feedback on Agency Service Delivery

Abstract: The information collection activity will garner qualitative customer and stakeholder feedback in an efficient, timely manner, in accordance with the Administration's commitment to improving service delivery. By qualitative feedback we mean information that provides useful

insights on perceptions and opinions, but are not statistical surveys that yield quantitative results that can be generalized to the population of study. This feedback will provide insights into customer or stakeholder perceptions, experiences and expectations, provide an early warning of issues with service, or focus attention on areas where communication, training or changes in operations might improve delivery of products or services. These collections will allow for ongoing, collaborative and actionable communications between the Agency and its customers and stakeholders. It will also allow feedback to contribute directly to the improvement of program management.

Feedback collected under this generic clearance will provide useful information, but it will not yield data that can be generalized to the overall population. This type of generic clearance for qualitative information will not be used for quantitative information collections that are designed to yield reliably actionable results, such as monitoring trends over time or documenting program performance. Such data uses require more rigorous designs that address: The target population to which generalizations will be made, the sampling frame, the sample design (including stratification and clustering), the precision requirements or power calculations that justify the proposed sample size, the expected response rate, methods for assessing potential nonresponse bias, the protocols for data collection, and any testing procedures that were or will be undertaken prior to fielding the study. Depending on the degree of influence the results are likely to have, such collections may still be eligible for submission for other generic mechanisms that are designed to yield quantitative results.

The Agency received no comments in response to the 60-day notice published in the **Federal Register** on December 22, 2010 (75 FR 80542).

This is a new collection of information. Respondents will be screened and selected from individuals and households, businesses, organizations, and/or State, Local or Tribal Government. Below we provide CDC's projected annualized estimate for the next three years. There is no cost to respondents other than their time. The estimated annualized burden hours for this data collection activity are 800.

Type of collection	Average number of respondents per activity	Annual frequency per response	Average number of activities	Average hours per response
Online surveys, Telephone Surveys, Focus Groups, In person observation/ testing	200	1	5	48/60

### Kimberly S. Lane,

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

[FR Doc. 2012-11101 Filed 5-7-12; 8:45 am]

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# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

[60Day-12-0607]

### 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 404-639-7570 or send comments to Kimberly Lane, at CDC, 1600 Clifton Road, MS-D74, Atlanta, GA 30333 or send an email 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**

The National Violent Death Reporting System (NVDRS) OMB# 0920–0607 –Extension—National Center for Injury Prevention and Control (NCIPC), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Violence is an important public health problem. In the United States, homicide and suicide are the second and third leading causes of death, respectively, in the 1-34 year old age group. Unfortunately, public health agencies do not know much more about the problem than the numbers and the sex, race, and age of the victims, all information obtainable from the standard death certificate. Death certificates, however, carry no information about key facts necessary for prevention such as the relationship of the victim and suspect and the circumstances of the deaths, thereby making it impossible to discern anything but the gross contours of the problem. Furthermore, death certificates are typically available 20 months after the completion of a single calendar year. Official publications of national violent death rates, e.g. those in Morbidity and Mortality Weekly Report, rarely use data that is less than two years old. Public health interventions aimed at a moving target last seen two years ago may well miss the mark.

Local and Federal criminal justice agencies such as the Federal Bureau of Investigation (FBI) provide slightly more information about homicides, but they do not routinely collect standardized data about suicides, which are in fact much more common than homicides. The FBI's Supplemental Homicide Report (SHRs) does collect basic information about the victim-suspect relationship and circumstances related to the homicide. SHRs, do not link violent deaths that are part of one incident such as homicide-suicides. It also is a voluntary system in which some 10-20 percent of police departments nationwide do not participate. The FBI's National Incident Based Reporting System (NIBRS)

provides slightly more information than SHRs, but it covers less of the country than SHRs. NIBRS also only provides data regarding homicides. Also, the Bureau of Justice Statistics Reports do not use data that is less than two years old

CDC therefore proposes to continue a state-based surveillance system for violent deaths that will provide more detailed and timely information. It taps into the case records held by medical examiners/coroners, police, and crime labs. Data is collected centrally by each state in the system, stripped of identifiers, and then sent to the CDC. Information is collected from these records about the characteristics of the victims and suspects, the circumstances of the deaths, and the weapons involved. States use standardized data elements and software designed by CDC. Ultimately, this information will guide states in designing programs that reduce multiple forms of violence.

Neither victim families nor suspects are contacted to collect this information. It all comes from existing records and is collected by state health department staff or their subcontractors. Health departments incur an average of 2.5 hours per death in identifying the deaths from death certificates, contacting the police and medical examiners to get copies of or to view the relevant records, abstracting all the records, various data processing tasks, various administrative tasks, data utilization, training, communications, etc.

CDC requests an extension to continue data collection with this system in the 18 funded states, and allow 9 new state health departments to be added if funding becomes available. This may bring the total to 27 by the year 2015. Violent deaths include all homicides, suicides, legal interventions, deaths from undetermined causes, and unintentional firearm deaths. The average state will experience approximately 1,000 such deaths each year.

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