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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-12-09BK]

Agency for Toxic Substances and Disease Registry; Agency Forms Undergoing Paperwork Reduction Act Review

The Agency for Toxic Substances and Disease Registry (ATSDR) 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/ATSDR 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 20503 or by fax to (202) 395-5806. Written comments should be received within 30 days of this notice.

Proposed Project

Registration of Individuals Displaced by the Hurricanes Katrina and Rita (Pilot Project)—New—Agency for Toxic Substances and Disease Registry (ATSDR), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

On August 29, 2005, Hurricane Katrina made landfall on the coast of the Gulf of Mexico near New Orleans, Louisiana, and became one of the most

deadly and destructive storms in U.S. history. Also occurring in 2005, Hurricane Rita was the fourth-most intense Atlantic hurricane ever recorded and the most intense tropical cyclone ever observed in the Gulf of Mexico. Following the initial phase of the response, the Federal Emergency Management Agency (FEMA) assumed the primary role for housing displaced persons over the intermediate term. To support those needing temporary housing, FEMA provided over 130,000 travel trailers, park homes, and mobile homes for persons displaced by the above mentioned storms. However, some persons living in trailers complained of an odor or of eye or respiratory tract irritation.

FEMA entered into an Interagency Agreement with the Centers for Disease Control and Prevention (CDC)/ATSDR on August 16, 2007 to conduct a comprehensive public health assessment, based on objective and credible research, of air quality conditions present in FEMA housing units to guide FEMA policy makers and inform the public as to the actual conditions in the field and any actions required to better promote a safe and healthful environment for the disaster victims FEMA housed in the units. FEMA's agreement with the CDC includes an initial formaldehyde exposure assessment as well as a subsequent long-term study of the health effects among resident children. Formaldehyde testing conducted and evaluated by the CDC pursuant to the initial exposure assessment has identified the need to evaluate the feasibility of establishing a national registry to identify and monitor the health of disaster victims who occupied FEMA-provided temporary housing units. The establishment of such a registry would complement the long-term health effects study set forth in the FEMA-CDC Interagency Agreement.

The proposed pilot registry will have two goals: Primary Goal: Test the feasibility and cost of contacting and

enrolling members in a registry by collecting and verifying phone interview data. Secondary Goal: Test the difference in prevalence rates of health conditions compared to national surveys (*i.e.*, NHANES and NHIS).

The data collected in the pilot registry and the evaluation of the pilot registry will be used to determine the feasibility and estimate the costs of developing and populating a more complete registry of people affected by Hurricanes Katrina and Rita. In addition, comparisons of prevalence rates of health outcomes obtained through the pilot registry with estimates from national surveys will help determine the utility of conducting a full registry. For example, if all or most health outcomes do not appear to be in excess, the value of a full registry may be questionable.

A pre-registration datasets will be created before enrollment. This dataset will be populated with contact information of the occupants of temporary housing units provided by FEMA. FEMA provided the datasets for this pilot registry.

A computer-assisted telephone interview (CATI) system based on a paper questionnaire will be used during all interviews to collect data for this project. The first part will consist of screening questions to determine eligibility for enrollment. The second part will contain contact information of the registrant and other household members, demographics, and health status questions, focusing on respiratory outcomes and mental health.

The two minute screening questionnaire will be administered to a total of 8,000 respondents. Annualized over a two year period, 4,000 will be screened. The 25 minute main questionnaire will be administered to a total of 5,000 respondents. Annualized over a two year period, 2,500 occupants will complete the main questionnaire.

There are no costs to the respondents other than their time. The total estimated annual burden hours are 1,176.

ESTIMATED ANNUALIZED BURDEN HOURS

Respondents	Form	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Temporary and Non-Temporary housing unit occupants.	Screening	4,000	1	2/60
Main questionnaire	2,500	1	25/60	

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

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Agency Forms Undergoing Paperwork Reduction Act Review

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Proposed Project

HIV Prevention among Latino MSM: Evaluation of a locally developed intervention—New—National Center for HIV/AIDS, Viral Hepatitis, STD, TB Prevention (NCHHSTP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Latinos are the largest and fastest growing ethnic minority group in the U.S. and have the second highest rate of HIV/AIDS diagnoses of all racial/ethnic groups in the country. From the

beginning of the epidemic through 2007, Latinos accounted for 17% of all AIDS cases reported to the CDC. Among Latino males, male-to-male sexual contact is the single most important source of HIV infection, accounting for 46% of HIV infections in U.S.-born Latino men from 2001 to 2005, and for more than one-half of HIV infections among South American, Cuban, and Mexican-born Latino men in the U.S. (CDC, 2007a; 2007b). In 2006, male-to-male sex accounted for 72% of new HIV infections among Latino males. Relative to other men who have sex with men (MSM), the rate of HIV infection among Latino MSM is twice the rate recorded among whites (43.1 vs. 19.6 per 100,000).

Despite the high levels of infection risk that affect Latino MSM, no efficacious interventions to prevent infection by HIV and other sexually transmitted diseases (STDs) are available for this vulnerable population. CDC's Prevention Research Synthesis group, whose role is to identify HIV prevention interventions that have met rigorous criteria for demonstrating evidence of efficacy, has not identified any behavioral interventions for Latino MSM that meet current efficacy criteria, and no such interventions are listed in CDC's 2011 update of its Compendium of Evidence-Based HIV Behavioral Interventions (<http://www.cdc.gov/hiv/topics/research/prs/compendium-evidence-based-interventions.htm>). There is an urgent need for efficacious, culturally congruent HIV/STD prevention interventions for Latino MSM.

The purpose of this project is to test the efficacy of an HIV prevention intervention for reducing sexual risk among Latino men who have sex with men in North Carolina. The HOLA en

Grupos intervention is a Spanish-language, small-group, 4-session intervention that is designed to increase consistent and correct condom use and HIV testing among Latino MSM and to affect other behavioral and psychosocial factors that can increase their vulnerability of HIV/STD infection. This study will use a randomized controlled trial design to assess the efficacy of the HOLA en Grupos intervention compared to a general health comparison intervention.

CDC is requesting approval for a 3-year clearance for data collection. The data collection system involves screening of potential study participants for eligibility, collection of participants' contact information, and measures of intervention and comparison participants' socio-demographic characteristics, health seeking actions, HIV/STD and substance use-related risk behaviors, and psychosocial factors at baseline before intervention delivery and 6 months after intervention delivery. An estimated 350 men will be screened for eligibility in order to enroll the 300 men required for the study. The baseline and the 6-month follow-up assessments will be similar. However, the 6-month assessment will ask study participants fewer questions because there is no need to ask all questions during both assessments. Collection of eligibility information from potential participants will require about 10 minutes; collection of baseline assessment information will require about 1 hour and 45 minutes; and collection of the 6-month follow-up assessment information will require about 1 hour.

The total estimated annual burden hours are 883. There is no cost to participants other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Form name	Number of respondents	Number responses per respondent	Average burden per respondent (in hours)
Prospective Study Participant	Participant Screening Form	350	1	10/60
Enrolled Study Participant	Baseline Assessment	300	1	1.75
Enrolled Study Participant	6-month follow-up assessment	300	1	1

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