

removal workers, fabric mill workers, and fire fighters. Regulations of the Environmental Protection Agency (EPA) and the Nuclear Regulatory Commission (NRC) also require the use of NIOSH-approved respirators. These regulations also establish methods for respirator manufacturers to submit respirators for testing under the regulation and have them certified as NIOSH-approved if they meet the criteria given in the above regulation. NIOSH, in accordance with 42 CFR Part 84: (1) Issues certificates of approval for respirators which have met specified construction, performance, and protection requirements; (2) establishes procedures and requirements to be met in filing applications for approval; (3) specifies minimum requirements and methods to be employed by NIOSH and by applicants in conducting inspections, examinations, and tests to determine effectiveness of respirators; (4) establishes a schedule of fees to be charged applicants for testing and certification; and (5) establishes approval labeling requirements. Information is collected from those who request services under 42 CFR part 84 in order to properly establish the scope and intent of request. Information

collected from requests for respirator approval functions includes contact information and information about factors likely to affect respirator performance and use. Such information includes, but is not necessarily limited to, respirator design, manufacturing methods and materials, quality assurance plans and procedures, and user instruction and draft labels, as specified in the regulation.

The main instrument for data collection for respirator approval functions is the SAF, Standard Application for the Approval of Respirators, currently Version 7. A replacement instrument, SAF V.8, which collects the same information is available for applicants without the requisite software environment for V.7. Respirator manufacturers are the respondents (estimated to average 75 each year over the years 2011–2013) and upon completion of the SAF their requests for approval are evaluated. Although there is no cost to respondents to submit an application other than their time to participate, respondents requesting respirator approval are required to submit fees for necessary testing as specified in 42 CFR 84.20–22, 84.66, 84.258 and 84.1102. In calendar

year 2010 \$395,564.00 was accepted. Applicants are required to provide test data that shows that the respirator is capable of meeting the specified requirements in 42 CFR part 84. The requirement for submitted test data is likely to be satisfied by standard testing performed by the manufacturer, and no extra burden is expected.

42 CFR part 84 approvals offer corroboration that approved respirators are produced to certain quality standards. Although 42 CFR part 84, subpart E prescribes certain quality standards, it is not expected that requiring approved quality standards will impose an additional cost burden over similarly effective quality standards that are not approved under 42 CFR part 84. Manufacturers with current approvals are subject to site audits by the Institute or its agents. There is no fee associated with audits. Audits may occur periodically or as a result of a reported issue. An average of 61 site audits were conducted annually over the calendar years 2008–2010, and this rate is expected to continue. Audits take an average of 23.5 burden hours from the respondent.

There are no costs to respondents other than their time.

Form	Number of respondents	No. of responses per respondent	Avg. burden per response (in hrs)	Total burden (in hrs)
Standard Application for the Approval of Respirators	75	8	229	137,400
Audit	60	1	24	1,440
Total				138,840

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Carol E. Walker,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–11–0406]

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 e-mail 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

State and Local Area Integrated Telephone Survey (SLAITS), (OMB No. 0920–0406, Expiration 04/30/2011)—Revision—National Center for Health Statistics (NCHS), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Section 306 of the Public Health Service (PHS) Act (42 U.S.C. 242k), as amended, authorizes that the Secretary of Health and Human Services (DHHS), acting through NCHS, shall collect statistics on the extent and nature of illness and disability of the population of the United States. This revision is to notify the public of a request to

continue the SLAITS mechanism for the 2011 to 2014 survey period. A three year clearance is requested.

SLAITS is an integrated and coordinated survey system that has been conducted since 1997, in accordance with the 1995 initiative to increase the integration of surveys within DHHS. It is designed to collect needed health and well-being data at the national, state, and local levels. Using the large sampling frame of the ongoing National Immunization Survey (NIS) and Computer Assisted Telephone Interviewing (CATI), and when necessary independent samples, mail, and Internet modes to support data collection activities, SLAITS has quickly collected and produced household and person-level data to monitor health-related areas. Questionnaire content is drawn from existing surveys within DHHS and other Federal agencies, or developed specifically to meet project sponsor needs.

Examples of topical areas include infant, child, adolescent, parent, and family health, well-being, and knowledge, attitude, and behaviors; children with special health care needs (CSHCN); functioning; life course and social determinants of health; developmental delays and disabilities; acute and chronic conditions; immunizations; access to and use of health care; program participation; adoption; and changes in health insurance coverage and experiences.

Users of SLAITS data include, but are not limited to, Congressional offices, Federal agencies, state and local

governments, schools of public health, colleges and universities, private industry, nonprofit foundations, professional associations, clinicians, researchers, administrators, advocates, and health planners, to evaluate content and/or programs. SLAITS data continue to be heavily used by Federal and state Maternal and Child Health Bureau Directors to evaluate programs and service needs. Several SLAITS modules provided data for multiple Congressionally-mandated reports on healthcare disparities and quality; at least one report to Congress on health insurance coverage among children; and

reports of the National Academy of Sciences. Within DHHS, the Office of the Assistant Secretary for Planning and Evaluation and the Administration for Children and Families used SLAITS to collect data for the first nationally representative survey of adoptive families across adoption types for children with and without special health care needs, and to assess their post-adoption service use and unmet needs.

There is no cost to respondents other than their time to participate. The total estimated annualized burden hours are 194,675.

ESTIMATED ANNUALIZED BURDEN HOURS

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Household screening	1,800,000	1	2/60
Household interview	306,000	1	25/60
Pilot work, pre-testing, and planning activities	12,300	1	35/60

Dated: March 9, 2011.

Carol E. Walker,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30 Day 11-10GP]

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Proposed Project

Clostridium difficile Infection (CDI) Surveillance—New—National Center for Emerging and Zoonotic Infectious Diseases, (NCEZID), Centers for Disease Control and Prevention, (CDC).

Background and Brief Description

Steady increases in the rate and severity of *Clostridium difficile* infection (CDI) indicate a clear need to conduct longitudinal assessments of the impact of CDI in the United States. *C. difficile* is an anaerobic, spore-forming, gram positive bacillus that produces two pathogenic toxins: A and B. CDI ranges in severity from mild diarrhea to fulminant colitis and death. Transmission of *C. difficile* occurs primarily in healthcare facilities, where environmental contamination by *C. difficile* spores and exposure to antimicrobial drugs are common. No longer limited to healthcare environments, community-associated CDI is the focus of increasing attention. Recently, several cases of serious CDI have been reported in what have been considered low-risk populations, including healthy persons living in the community and peri-partum women.

The surveillance population will consist of persons residing in the

catchment area of the participating Emerging Infections Program (EIP) sites. This surveillance poses no more than minimal risk to the study participants as there will be no interventions or modifications to the care study participants receive. EIP surveillance personnel will perform active case finding from laboratory reports of stool specimens testing positive for *C. difficile* toxin and abstract data on cases using a standardized case report form. For a subset of cases (e.g., community-associated *C. difficile* cases) sites will administer a health interview. Remnant stool specimens from cases testing positive for *C. difficile* toxin will be submitted to reference laboratories for culturing, and isolates will be sent to CDC for confirmation and molecular typing. Outcomes of this surveillance project will include the population-based incidence of community- and healthcare-associated CDI, and a description of the molecular characteristics of *C. difficile* strains and the epidemiology of this infection among the population under surveillance.

There is no cost to respondents to participate in this program. The total annualized burden for this data collection is 5,840 hours.

ESTIMATE OF ANNUALIZED BURDEN HOURS

Respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
CDI Surveillance Case Report Form—Complete	10	437	1