

ESTIMATE OF ANNUALIZED BURDEN TABLE

Form	Number of respondents	Number of responses per respondent	Total responses	Hrs/response	Total burden
ABCs	329	21	6909	10/60	1152
CJD	20	2	40	20/60	13
Cyclospora	55	10	550	15/60	138
Dengue Case Investigation	55	182	10,010	15/60	2,503
Hantavirus Pulmonary Syndrome	40	3	120	20/60	40
Idiopathic CD4+T-lymphocytopenia	10	2	20	10/60	3
Kawasaki Syndrome	55	8	440	15/60	110
Legionellosis Case Report	23	11.7	269	20/60	90
Lyme Disease Report	52	261	20,020	5/60	1,668
Malaria Case Surveillance Report	55	20	1,100	15/60	275
Plague Case Investigation Report	55	0.20	11	20/60	4
Q Fever	55	1	55	10/60	9
Reye's Syndrome Case Surveillance Report	50	1	50	20/60	17
Tick-borne Rickettsial Disease Case Report	55	18	990	10/60	165
Trichinosis Surveillance Case Report	55	0.70	39	20/60	13
Tularemia	55	2.2	121	20/60	40
Typhoid Fever Surveillance Report	55	7	385	20/60	128
Viral Hepatitis Case Record	55	200	11,000	25/60	4,583
Total					10,950

Dated: December 2, 2005.

Joan Karr,

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

[60Day-06-06AK]

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 404-639-4766 or send comments to Seleda Perryman, CDC Assistant 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

Collection of Customer Survey Data Pertaining to the CDC Web site—New—National Center for Health Marketing (NCHM), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Executive Order 12862 directs agencies that provide significant services directly to the public to survey customers to determine the kind and quality of services they need and their level of satisfaction with existing

services. The Centers for Disease Control and Prevention (CDC), National Center for Health Marketing (NCHM), seeks to obtain approval to conduct customer satisfaction surveys and usability tests of the CDC Web site, <http://www.cdc.gov> on an ongoing basis. By collecting customer satisfaction and Web site usability information, CDC will be enabled to serve, and respond to, the ever-changing demands of website users. These users include individuals (patients, educators, students, etc.), interested communities, partners, healthcare providers, and businesses. Survey information will augment current Web content, delivery, and design research which is used to understand the Web user, and more specifically, the CDC user community. Primary objectives are to ensure: (1) CDC's Web site meets its customer needs and (2) the Web site meets the wants, preferences, and needs of its target audiences. Findings will help to: (1) Understand the user community and how to better serve Internet users; (2) discover areas requiring improvement in either content or delivery; (3) determine how to align Web offerings with identified user need(s); and (4) explore methods for offering, presenting and delivering information most effectively. There are no costs to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN TABLE

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total annual burden (in hours)
Web site Users	400,000	1	6/60	40,000

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Joan F. Karr,

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-06-05BF]

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 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) 395-6974. Written comments should be received within 30 days of this notice.

Proposed Project

Human Smoking Behavior—New—National Center for Chronic Disease and Public Health Promotion (NCCDDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

CDC, National Center for Chronic Disease Prevention and Health Promotion (NCCDDPHP), in a joint venture with the National Center for Environmental Health (NCEH), proposes to conduct a 2-year laboratory-based study of human smoking behavior

among established current smokers of the major styles and varieties of cigarettes consumed in the United States. This study will compare how different categories of cigarettes deliver toxic chemicals to smokers in order to further investigate the link between tobacco use and disease.

The major objective of this study is to better understand how human and cigarette variables influence the delivered dose of harmful chemicals in smoke to identify risk factors that result in adverse health effects from smoking. The smoking behavior and biomarkers of 360 smokers will be ascertained. Participants will attend two sessions on consecutive days. Solanesol levels in cigarette filter butts; carbon monoxide boost in breath; carcinogens and nicotine and its metabolites in urine; cotinine in saliva; vent-blocking (as measured by filter stain pattern and visualization of lip and finger placement on the rod using fluorescent markers); smoking topography; and breathing patterns (inhalation and exhalation volume, breath velocity and duration prior to smoking, during smoking and after smoking) will be used to measure dose based on the number of cigarettes smoked, amount of each cigarette smoked, filter vent blocking behavior, smoking behavior and puff characteristics.

Another objective of this study is to define average or “composite” smoking patterns across several of the most popular cigarette categories (ultralight, light, full-flavored menthol and full-flavored non-menthol) from the quantitative and observational data. All current smoking machine methodologies are “one size fits all” approaches to generating cigarette smoke. The composite conditions can be used to establish human behavior-based smoking machine methods for

laboratory studies that require cigarette smoke for chemical or toxicological testing. Currently, laboratory scientists rely on automated smoking machines to generate cigarette smoke for chemical and toxicological testing.

Funding for this study will come from both NCCDDPHP and NCEH. The Centers will share responsibilities, with administrative and technical assistance coming from NCCDDPHP and laboratory support coming from NCEH.

This is a two-year study, and an estimated 500 respondents will be screened by telephone to yield 360 eligible respondents who complete both visits over the two-year study period. The total burden for each respondent who completes screening, visit 1 and visit 2 will be two hours and five minutes. The CATI screening will take five minutes. Visit 1 will take one hour, which includes a short screening item, the informed consent process, biologic sample collection (urine, saliva, and breath carbon monoxide), smoking topography, ventilation hole blocking procedure and breath measurements. Visit 2 will also take approximately one hour, which includes compensation, discussion of quit opportunities if requested, collection of cigarette butts, biologic sample collection (urine, saliva, and breath carbon monoxide), smoking topography, ventilation hole blocking procedure and breath measurements.

The following table summarizes burden on an annualized basis for 500 telephone interviews and 180 eligible respondents (one-half of the total respondents). The 180 eligible respondents estimated to complete visit 2 are the same respondents estimated to complete visit 1.

There are no costs to the respondents other than their time. The total estimated annualized burden hours are 402.

ESTIMATED ANNUALIZED BURDEN TABLE

Respondents	Procedure	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Smokers	CATI Screening	500	1	42
Eligible Smokers	Visit 1, (Day 1)	180	1	180
Eligible Smokers	Visit 2, (Day 2)	180	1	180