

## ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Grant Application Data Summary (GADS) .....	500	1	0.50	250

*Estimated Total Annual Burden Hours: 250*

In compliance with the requirements of Section 506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. E-mail address: [infocollection@acf.hhs.gov](mailto:infocollection@acf.hhs.gov). All requests should be identified by the title of the information collection.

The Department specifically requests comments 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) 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. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Dated: May 12, 2009.

**Janean Chambers,**

*Reports Clearance Officer.*

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Centers for Disease Control and Prevention

[30Day-09-0234]

#### Agency Forms Undergoing Paperwork 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 requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995. To request a copy of these requirements, call the CDC Reports Clearance Officer at 404-639-5960 or send comments to CDC/ATSDR Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D74, Atlanta, GA 30333 or send an e-mail to [omb@cdc.gov](mailto:omb@cdc.gov). Written comments should be received within 30 days of this notice.

#### Proposed Project

National Ambulatory Medical Care Survey (NAMCS) (OMB No. 0920-0234)—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 "utilization of health care" in the United States. NAMCS was conducted annually from 1973 to 1981, again in 1985, and resumed as an annual survey in 1989. The purpose of NAMCS is to meet the needs and demands for statistical information about the provision of ambulatory medical care services in the United States. NCHS is seeking OMB approval to extend this survey for three years.

Ambulatory services are rendered in a wide variety of settings, including physician offices and hospital outpatient and emergency departments. The NAMCS target universe consists of all office visits made by ambulatory patients to non-Federal office-based physicians (excluding those in the specialties of anesthesiology, radiology, and pathology) who are engaged in direct patient care.

In 2006, physicians and mid-level providers (*i.e.*, nurse practitioners, physician assistants, and nurse midwives) practicing in community health centers (CHCs) were added to the NAMCS sample, and these data will continue to be collected. To complement NAMCS data, NCHS initiated the National Hospital Ambulatory Medical Care Survey (NHAMCS, OMB No. 0920-0278) in

1992 to provide data concerning patient visits to hospital outpatient and emergency departments. NAMCS and NHAMCS are the principal sources of data on ambulatory care provided in the United States.

NAMCS provides a range of baseline data on the characteristics of the users and providers of ambulatory medical care. Data collected include the patients' demographic characteristics, reason(s) for visit, provider diagnoses, diagnostic services, medications, and visit disposition. In addition, information on cervical cancer screening practices in physician offices will continue to be collected through the Cervical Cancer Screening Supplement (CCSS), which was added in 2006. It will allow CDC's National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP) to evaluate cervical cancer screening methods and the use of Human Papillomavirus DNA tests.

A supplemental mail survey on the adoption and use of electronic medical records (EMRs) in physician offices was added to NAMCS in 2008, and will continue. These data were requested by the Office of the National Coordinator for Health Information Technology (ONC), Department of Health and Human Services, to measure progress toward goals for EMR adoption. The mail survey will collect information on characteristics of physician practices and the capabilities of EMRs used in those practices.

In 2009, in addition to conducting the on-going survey, NAMCS will include an additional sample of 70 physicians to pretest additional questionnaire items on laboratory values. These new items were requested by the Division of Heart Disease and Stroke Prevention within NCCDPHP to better understand the extent to which ambulatory health care providers identify and control abnormal values before and after cardiovascular disease. Users of NAMCS 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, and health planners.

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

estimated annualized burden hours are 5,932.

ESTIMATED ANNUALIZED BURDEN TABLE

Type of form	Type of respondent	Form name	Number of respondents	Number of responses per respondent	Hours per response
Core NAMCS Forms .....	Office-based physicians/CHC providers.	Physician Induction Interview (NAMCS-1).	3,657	1	28/60
	Community Health Center Directors.	Community Health Center Induction Interview (NAMCS-201).	104	1	20/60
	Office-based physicians/CHC providers/staff.	Patient Record form (NAMCS-30).	738	30	9/60
	Office/CHC staff .....	Pulling, re-filing Patient Record form (NAMCS-30).	650	30	1/60
	Office-based physicians/CHC providers/staff.	Cervical Cancer Screening Supplement (NAMCS-CCS).	464	1	15/60
Lab Values Pre-test Forms ...	Office-based physicians .....	EMR/EHR Mail Survey .....	1,143	1	20/60
	Office-based physicians .....	Physician Induction Interview (NAMCS-1).	23	1	28/60
	Office-based physicians/staff	Patient Record form (NAMCS-30).	8	30	9/60

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Substance Abuse and Mental Health Services Administration**

**Agency Information Collection Activities: Proposed Collection; Comment Request**

In compliance with Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 concerning opportunity for public comment on proposed collections of information, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the information collection plans, call the SAMHSA Reports Clearance Officer at (240) 276-1243.

Comments are invited on: (a) Whether the proposed collections of information are 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.

**Proposed Project: Recovery Services for Adolescents and Families—New**

The Substance Abuse and Mental Health Services Administration's (SAMHSA) Center for Substance Abuse Treatment will conduct a data collection on the helpfulness of recovery support services for young people and their families after leaving substance abuse treatment. Specifically, the Recovery Services for Adolescents and Families (RSAF) project is trying to understand whether or not young people and their families find the following recovery support services helpful: (1) Telephone/text message support; (2) a recovery-oriented social networking site; and (3) a family program. Approximately 200 adolescent respondents will be asked to complete 4 data collection forms (some repeated) during 5 interviews (baseline and 4 follow-ups) over a 12 month period after enrollment or discharge from treatment. Approximately 200 collateral respondents (*i.e.*, a parent/guardian/concerned other) will be asked to complete 7 data collection forms (some repeated) during 5 interviews (baseline and 4 follow-ups) over a 12 month period after their adolescent's enrollment or discharge from treatment. Approximately 15 to 20 project staff respondents, including Project Coordinators, Telephone Support Volunteers, a Social Network Site Moderator, Family Program Clinicians, and a Support Services Supervisor, will be asked to complete between 2 and 5

data collection forms at varying intervals during the delivery of recovery support services. Across all respondents, a total of 26 data collection forms will be used. Depending on the time interval and task, information collections will take anywhere from about 5 minutes to 2 hours to complete. A description of each data collection form follows:

*Follow-Up Locator Form—Participant (FLF-P; Adolescent Respondent).* The FLF-P contains over 50 items that are a combination of yes/no, multiple choice, and open-ended formats. Data are gathered about an adolescent's contact information, personal contacts, criminal justice contacts, school/job contacts, hang-out information, internet contacts, and identifying information in order to locate and interview that adolescent over multiple follow-up intervals.

*Global Appraisal of Individual Needs—Initial (GAIN-I 5.6.0 Full; Adolescent Respondent).* The GAIN is an evidence-based assessment used with both adolescents and adults and in outpatient, intensive outpatient, partial hospitalization, methadone, short-term residential, long-term residential, therapeutic community, and correctional programs. There are over 1,000 questions in this initial version that are in multiple formats, including multiple choice, yes/no, and open-ended. Eight content areas are covered: Background, Substance Use, Physical Health, Risk Behaviors and Disease Prevention, Mental and Emotional Health, Environment and Living Situation, Legal, and Vocational. Each section contains questions on the recency of problems, breadth of