

DC or by fax to (202) 395-5806. Written comments should be received within 30 days of this notice.

Proposed Project

The National Electronic Disease Surveillance System (NEDSS) (OMB Number 0920-0728 exp. 2/28/2011)—Extension—National Center for Public Health Informatics (NCPHI), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

CDC is responsible for the dissemination of nationally notifiable diseases information and for monitoring and reporting the impact of epidemic influenza on mortality, Public Health Services Act (42 U.S.C. 241). Since April 1984, CDC National Center for Public Health Informatics Epidemiology Program Office (EPO) began working with the Council of State and Territorial Epidemiologists (CSTE) to demonstrate the efficiency and effectiveness of computer transmission of surveillance data between CDC and the State health departments.

By 1989, all 50 States were using this computerized disease surveillance system, which was then renamed the National Electronic Telecommunications System for Surveillance (NETSS) to reflect its

national scope (OMB numbers 0920-0447 and 0920-0007).

Beginning in 1999, CDC, Epidemiology Program Office (EPO) worked with CSTE, State and local public health system staff, and other CDC disease prevention and control program staff to identify information categories and information technology standards to support integrated disease surveillance. That effort is now focused on development and completion of the National Electronic Disease Surveillance System (NEDSS), coordinated by CDC's National Center for Public Health Informatics, Division of Integrated Surveillance Systems and Services (DISSS).

States will continue to use portions of NETSS to transmit data to CDC. One of the reasons for providing NETSS to NEDSS data mapping is to identify what data elements in NETSS correspond to data elements in NEDSS. Those elements mapped from NETSS to NEDSS were collected in OMB number 0920-0007.

NEDSS will electronically integrate and link together a wide variety of surveillance activities and will facilitate more accurate and timely reporting of disease information to CDC and State and local health departments. Consistent with recommendations supported by our State and local

surveillance partners and described in the 1995 report, *Integrating Public Health Information and Surveillance Systems*, NEDSS includes data standards, an Internet based communications infrastructure built on industry standards, and policy-level agreements on data access, sharing, burden reduction, and protection of confidentiality.

To support NEDSS, CDC has developed an information system, the NEDSS Base System (NBS), which uses NEDSS technical and information standards. The NBS is currently deployed to 16 States, including AL, AR, ID, MD, ME, MT, NE, NM, NV, RI, SC, TN, TX, VA, VT, and WY.

CDC is requesting a three-year OMB clearance extension of collecting the NEDSS data. The table below outlines the annualized burden which consists of two components. The first component is "weekly reporting" (52 weeks annually). The second component is an end of year report titled "annual reporting". The two components collectively represent the estimated annualized hours for the submitting jurisdictions.

There are no costs to respondents other than their time. The total estimated annual burden hours for the Weekly Morbidity Reports and the Annual Summary Report is 9,384.

ESTIMATED ANNUALIZED BURDEN HOURS

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Weekly Reporting			
States	50	52	3
Territories	5	52	1.5
Cities	2	52	3
Annual Reporting			
States	50	1	16
Territories	5	1	10
Cities	2	1	16

Dated: October 7, 2010.

Carol Walker,

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

[FR Doc. 2010-25916 Filed 10-14-10; 8:45 am]

BILLING CODE 4163-18-P

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 on (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: 2011 National Survey on Drug Use and Health (OMB No. 0930-0110)—Revision

The National Survey on Drug Use and Health (NSDUH), formerly the National Household Survey on Drug Abuse (NHSDA) is a survey of the civilian, non-institutionalized population of the United States 12 years old and older. The survey is used to determine the prevalence of use of tobacco products, alcohol, illicit substances, and illicit use of prescription drugs. The survey is also used to collect information on mental health problems and the utilization of substance abuse and mental health services. The results are used by SAMHSA, ONDCP, Federal government agencies, and other organizations and researchers to establish policy, direct program activities, and better allocate resources.

The 2011 NSDUH will continue conducting a follow-up clinical interview with a subsample of approximately 1,500 respondents. The design of this study is based on the recommendations from a panel of expert consultants convened by the Center for Mental Health Services (CMHS), SAMHSA, to discuss mental health surveillance data collection strategies.

The goal is to create a statistically sound measure that may be used to estimate the prevalence of Serious Mental Illness (SMI) among adults (age 18+).

For the 2011 NSDUH, no questionnaire changes are proposed.

As with all NSDUH/NHSDA surveys conducted since 1999, the sample size of the survey for 2011 will be sufficient to permit prevalence estimates for each of the fifty states and the District of Columbia.

Because the NSDUH collects data on substance use, mental health and the utilization of substance abuse and mental health services, it is an appropriate and convenient vehicle to measure the impact of the Deepwater Horizon oil spill on residents of that region. Therefore, SAMHSA is planning to expand the NSDUH by oversampling the geographic region impacted by the oil spill. The current NSDUH sample design will be implemented and an oversampling method that results in an additional 2,000 completed interviews in the gulf coast region will be employed. The additional interviews will be concentrated in the coastal counties of Alabama, Florida, Louisiana, and Mississippi. All survey instruments and protocols will be identical for this additional sample. The total number of respondents for the 2011 NSDUH will be 69,500, or 2,000 cases more than the planned sample size for 2010.

Though there will be some increase in the sample for all four states involved in the Deepwater Horizon event (Alabama, Florida, Louisiana, and Mississippi), specific counties in the gulf coast region were chosen for focused oversampling. These counties were chosen based on the following criteria:

- Claims activity to BP for economic and related health needs;

- County involvement with Department of Education and Administration for Children and Families programming; and
- State assessment of impacted counties based on consultation with SAMHSA during the preparation of aid applications.

COUNTIES DESIGNATED AS THE MOST AFFECTED AREAS

State name	County/parish name
Alabama	Baldwin
Alabama	Clarke
Alabama	Escambia
Alabama	Mobile
Alabama	Monroe
Alabama	Washington
Florida	Bay
Florida	Escambia
Florida	Franklin
Florida	Gulf
Florida	Okaloosa
Florida	Santa Rosa
Florida	Wakulla
Florida	Walton
Louisiana	Iberia
Louisiana	Jefferson
Louisiana	Lafayette
Louisiana	Lafourche
Louisiana	Orleans
Louisiana	Plaquemines
Louisiana	St. Bernard
Louisiana	St. Martin
Louisiana	St. Mary
Louisiana	St. Tammany
Louisiana	Terrebonne
Louisiana	Vermilion
Mississippi	George
Mississippi	Hancock
Mississippi	Harrison
Mississippi	Jackson
Mississippi	Pearl River
Mississippi	Stone

The total annual burden estimate is shown below:

Instrument	Number of respondents	Responses per respondent	Hours per response	Total burden hours	Hourly wage rate	Annualized hourly costs
Household Screening	196,720	1	0.083	16,328	\$14.64	\$239,042
Interview	69,500	1	1.000	69,500	14.64	1,017,480
Clinical Follow-up Certification	90	1	1.000	90	14.64	1,318
Clinical Follow-up Interview	1,500	1	1.000	1,500	14.64	21,960
Screening Verification	5,560	1	0.067	373	14.64	5,461
Interview Verification	10,425	1	0.067	698	14.64	10,219
Total	196,810			88,489		1,295,480

Send comments to Summer King, SAMHSA Reports Clearance Officer, Room 8–1099, One Choke Cherry Road, Rockville, MD 20857 AND e-mail a copy to summer.king@samhsa.hhs.gov. Written comments should be received within 30 days of this notice.

Dated: October 8, 2010.

Elaine Parry,

Director, Office of Management, Technology and Operations.

[FR Doc. 2010–26077 Filed 10–14–10; 8:45 am]

BILLING CODE 4162–20–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Child Care and Development Block Grant Reporting Requirements—ACF–700.

OMB No.: 0980–0241.

Description: The Child Care and Development Fund (CCDF) report requests annual Tribal aggregate information on services provided through the CCDF, which is required by the CCDF Final Rule (45 FR parts 98 and

99). Tribal Lead Agencies (TLAs) are required to submit annual aggregate data appropriate to Tribal programs on children and families receiving CCDF-funded child care services. The CCDF statute and regulations also require TLAs to submit a supplemental narrative as part of the ACF–700 report. This narrative describes child care activities and actions in the TLA’s service area. Information from the ACF–700 and supplemental narrative report will be included in the Secretary’s Report to Congress, as appropriate, and will be shared with all TLAs to inform them of CCDF-funded activities in other Tribal programs.

Respondents: Tribal Governments.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
ACF–700 Report	260	1	38	9,880

Estimated Total Annual Burden Hours: 9,880

Additional Information: Copies of the proposed collection may be obtained 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. All requests should be identified by the title of the information collection. E-mail address: infocollection@acf.hhs.gov.

OMB Comment: OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, Fax: 202–395–7285, E-mail: OIRA_SUBMISSION@OMB.EOP.GOV, Attn: Desk Officer for the Administration for Children and Families.

Dated: October 12, 2010.

Robert Sargis,

Reports Clearance Officer.

[FR Doc. 2010–26052 Filed 10–14–10; 8:45 am]

BILLING CODE 4184–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request; Multi-Ethnic Study of Atherosclerosis (MESA) Event Surveillance

SUMMARY: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Heart, Lung, and Blood Institute (NHLBI), the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval the information collection listed below. This proposed information collection was previously published in the **Federal Register** on August 4, 2010, pages 46945–6, and allowed 60-days for public comment. Only one comment was received. The purpose of this notice is to allow an additional 30 days for public comment. The National Institutes of Health may not conduct or sponsor, and the respondent is not required to respond to, an information collection that has been extended, revised, or implemented on or after October 1, 1995, unless it displays a currently valid OMB control number.

Proposed Collection: Title: Multi-Ethnic Study of Atherosclerosis (MESA) Event Surveillance. *Type of Information Request:* Renewal (OMB No. 0925–

0493). *Need and Use of Information Collection:* The study, MESA, is identifying and quantifying factors associated with the presence and progression of subclinical cardiovascular disease (CVD)—that is, atherosclerosis and other forms of CVD that have not produced signs and symptoms. The findings provide important information on subclinical CVD in individuals of different ethnic backgrounds and provide information for studies on new interventions to prevent CVD. The aspects of the study that concern direct participant evaluation received a clinical exemption from OMB clearance (CE–99–11–08) in April 2000. OMB clearance is being sought for the contact of physicians and participant proxies to obtain information about clinical CVD events that participants experience during the follow-up period. *Frequency of response:* Once per CVD event. *Affected public:* Individuals. *Types of Respondents:* Physicians and selected proxies of individuals recruited for MESA. The annual reporting burden is as follows: *Estimated Number of Respondents:* 74; *Estimated Number of Responses per Respondent:* 1.0; *Average Burden Hours Per Response:* 0.20; and *Estimated Total Annual Burden Hours Requested:* 14.7. The annualized cost to respondents is estimated at: \$500. There are no capital, operating, or maintenance costs to report.