

organization or other affiliation, full address and phone, fax, and e-mail information or e-mail this information to *FindYouthInfo@air.org*. Additional identification documents may be required.

Dated: November 4, 2010.

Sherry Glied,

Assistant Secretary for Planning and Evaluation.

Authority: Division F, Pub. L. 111-8; E.O. 13459, 73 FR 8003, February 12, 2008.

[FR Doc. 2010-28396 Filed 11-9-10; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-10-10DE]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC), 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 (33 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 ATSDR 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

Creation of State and Metropolitan Area-based Surveillance Projects for Amyotrophic Lateral Sclerosis (ALS)—New—Agency for Toxic Substances and Disease Registry (ATSDR).

Background and Brief Description

On October 10, 2008, President Bush signed S. 1382: ALS Registry Act which amended the Public Health Service Act to provide for the establishment of an Amyotrophic Lateral Sclerosis (ALS) Registry. The activities described are part of the effort to create the National ALS Registry. The purpose of the registry is to: (1) Better describe the incidence and prevalence of ALS in the United States; (2) examine appropriate factors, such as environmental and occupational, that might be associated with the disease; (3) better outline key demographic factors (such as age, race or ethnicity, gender, and family history of individuals diagnosed with the disease); and (4) better examine the connection between ALS and other motor neuron disorders that can be confused with ALS, misdiagnosed as ALS, and in some cases progress to ALS. The registry will collect personal health information that may provide a basis for further scientific studies of potential risks for developing ALS.

This project purposes to collect information-specific data related to

ALS. The objective of this project is to develop state-based and metropolitan area-based surveillance projects for ALS. The primary goal of the state-based and metropolitan area-based surveillance project is to use these data to evaluate the completeness of the National ALS Registry. The secondary goal of the surveillance project is to obtain reliable and timely information on the incidence and prevalence of ALS and to better describe the demographic characteristics (e.g., age, race, sex, and geographic location) of those with ALS.

Neurologists or their staff will complete an ALS Case Reporting Form on each of their ALS patients. This will be transmitted to the state or metropolitan health department. The contract surveillance staff assigned to the state and metropolitan area health departments will train medical personnel how to complete the ALS Case Reporting Form (Attachment 3) and assist with abstracting records as requested. An ALS Medical Record Verification Form will be collected on a subset of cases reported. Each medical provider reporting source should keep a line listing of individuals diagnosed with or thought to have ALS along with information on whether or not the case was reported and if not, the reason. Surveillance items to be collected include information to make sure that there are no duplicates. There are no costs to the respondents other than their time. The estimated annualized burden hours are 703.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of data collection instrument	Type of respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Training	Medical Personnel/Neurologist	243	1	30/60
ALS Case Reporting Form	Medical Personnel/Neurologist	2,250	1	5/60
ALS Medical Record Verification Form	Neurologist	450	1	20/60
Line Listing (record keeping)	Medical Personnel	243	1	1

Dated: November 4, 2010.

Carol E. Walker,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention/Agency for Toxic Substances and Disease Registry.

[FR Doc. 2010-28337 Filed 11-9-10; 8:45 am]

BILLING CODE P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB review; comment request; NCCAM Office of Communications and Public Liaison Communications Program Planning and Evaluation Research

SUMMARY: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Center for Complementary and Alternative Medicine (NCCAM), the

National Institutes of Health (NIH), has submitted to the Office of Management and Budget (OMB) a request to review and approve the information collection listed below. This proposed information collection was previously published in the **Federal Register** on August 25, 2010 (Vol. 75, No. 164, p. 52349) and allowed 60-days for public comment. There was one public comments received during this time. 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: NCCAM Office of Communications and Public Liaison Communications Program Planning and Evaluation Research. *Type of Information Collection Request:* Extension. *Need and Use of Information Collection:* To carry out NCCAM's legislative mandate to educate and disseminate information about complementary and alternative medicine (CAM) to a wide variety of audiences and organizations, the NCCAM Office of Communications and Public Liaison (OCPL) requests clearance to carry out (1) formative and (2) evaluative research of a variety of print and online materials, outreach activities, and messages to maximize their impact and usefulness.

OCPL wishes to continue to carry out formative research to further understand the knowledge, attitudes, and behaviors of its core constituent groups: members of the general public, researchers, and providers of both conventional and CAM health care. In addition, it seeks to test newly formulated messages and identify barriers and impediments to the effective communication of those messages. With this formative audience

research, OCPL test audience responses to NCCAM's fact sheets, Web content, and other materials and messages.

Clearance is also requested to continue evaluative research on existing materials and messages, as part of OCPL's ongoing effort to develop a comprehensive program of testing and evaluation of all of its communications strategies. This evaluative research will include pilot testing of recently developed messages and information products such as consumer fact sheets and brochures. It will address the need to evaluate the processes by which new materials and messages were developed, the effectiveness of an outreach activity or the extent to which behaviors were changed by the message, and the impact of a message on health knowledge and behaviors.

The tools to collect this information have been selected to minimize burden on NCCAM's audiences, produce or refine messages that have the greatest potential to influence target audience attitudes and behavior in a positive manner, and to use Government resources efficiently. They may include individual in-depth interviews, focus group interviews, intercept interviews, self-administered questionnaires, gatekeeper reviews, and omnibus surveys.

The data will enhance OCPL's understanding of the unique information needs and distinct health-information-seeking behaviors of its core constituencies, and the segments within these constituencies with special information needs (for example, among the general public these segments include cancer patients, the chronically ill, minority and ethnic populations, the elderly, users of dietary supplements, and patients integrating complementary therapies with conventional medical treatments).

Frequency of Response: On occasion. *Affected Public:* Individuals and households; non-profit institutions; Federal Government; State, Local, or Tribal Government. *Type of Respondents:* Adult patients; members of the public; health care professionals; organizational representatives. The annual reporting burden is as follows: *Estimated Number of Respondents:* 2,500; *Estimated Number of Responses per Respondent:* 1; *Average Burden Hours per Response:* 0.58; and *Estimated Total Burden Hours Requested:* 2,109 for the 3-year clearance period (approximately 703 hours annually). The annualized cost to respondents is estimated at \$18,123. There are no Capital Costs, Operating Costs, or Maintenance Costs to report.

TABLE 1—ANNUAL BURDEN HOURS

Type of respondents	Estimated number of respondents	Estimated number of responses per respondent	Average burden hours per response	Estimated total annual burden hours requested
In-depth interviews with general public	30	1	.75	23
Focus groups	20	1	1.5	30
Omnibus surveys	1,900	1	0.25	475
Intercept interviews with public and healthcare professionals	300	1	.25	75
In-depth interviews health professionals	50	1	.50	25
Self-administered questionnaires with health professionals	200	1	.25	50
Total	2,500	678

Request for Comments: Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will have practical utility; (2) The accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) Ways to enhance the quality, utility, and clarity of the information to be collected; and (4) Ways to minimize the burden of the

collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

Direct Comments to OMB: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the Attention: NIH Desk Officer, Office of Management and Budget, at *OIRA_submission@omb.eop.gov* or by fax to 202-395-6974. To request more information on the proposed project or

to obtain a copy of the data collection plans and instruments, contact Christy Thomsen, Director, Office of Communications and Public Liaison, NCCAM, 31 Center Drive, Room 2B11, Bethesda, MD 20892, or fax your request to 301-402-4741, or e-mail *thomsenc@mail.nih.gov*. Ms. Thomsen can be contacted by telephone at 301-451-8876.

Comments Due Date: Comments regarding this information collection are best assured of having their full effect if received within 30 days of the date of this publication.

Dated: November 1, 2010.

Christy Thomsen,
 Director, Office of Communications and
 Public Liaison, National Center for
 Complementary and Alternative Medicine,
 National Institutes of Health.

[FR Doc. 2010-28290 Filed 11-9-10; 8:45 am]

BILLING CODE 4140-01-P

**DEPARTMENT OF HEALTH AND
 HUMAN SERVICES**

**Administration for Children and
 Families**

**Proposed Information Collection
 Activity; Comment Request**

Proposed Projects

Title: Strengthening Communities
 Fund (SCF) Evaluation.

OMB No.: New Collection.

Description: This proposed
 information collection activity is to
 obtain information from participants in
 two Strengthening Communities Fund
 (SCF) programs: The Nonprofit Capacity
 Building Program and the State, Local,
 and Tribal Government Capacity
 Building Program. Both programs are
 designed to contribute to the economic
 recovery as authorized in the American
 Recovery and Reinvestment Act of 2009
 (ARRA). The SCF evaluation is an
 important opportunity to examine
 outcomes achieved by the Strengthening
 Communities Fund and progress toward
 the objective of improving the capacity
 of organizations served by program
 grantees to address broad economic
 recovery issues in their communities.

The evaluation will be designed to
 assess progress and measure increased
 organizational capacity of each
 participating organization. The purpose
 of this request is to receive approval of
 the data collection instruments that will
 be used in this study.

A significant amount of information is
 already being collected through
 program-specific OMB-approved PPR
 forms or is available through secondary
 sources. Proposed surveys and phone
 interviews are very brief to reduce the
 burden on respondents.

Respondents: SCF grantees, and faith-
 based and Community Organizations
 (FBCOs).

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
An on-line survey of SCF grantees	84	1	0.25	21
Telephone interview of SCF grantees	84	1	1.50	126
On-line survey of faith-based and community organizations (FBCOs) that received capacity building services from the SCF grantees	1,000	1	0.50	500

*Estimated Total Annual Burden
 Hours:* 647

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. 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: November 4, 2010.

Robert Sargis,
 Reports Clearance Officer.
 [FR Doc. 2010-28304 Filed 11-9-10; 8:45 am]
BILLING CODE 4184-01-P

**DEPARTMENT OF HEALTH AND
 HUMAN SERVICES**

Food and Drug Administration

[Docket No. FDA-2009-P-0273]

**Determination That Amphetamine
 Sulfate, 5 and 10 Milligram Tablets,
 Was Not Withdrawn From Sale for
 Reasons of Safety or Effectiveness**

AGENCY: Food and Drug Administration,
 HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
 Administration (FDA) has determined
 that Amphetamine sulfate, 5 and 10
 milligram (mg) tablets, was not
 withdrawn from sale for reasons of
 safety or effectiveness. This
 determination will allow FDA to
 approve abbreviated new drug

applications (ANDAs) for Amphetamine
 sulfate, 5 mg and 10 mg tablets, if all
 other legal and regulatory requirements
 are met.

FOR FURTHER INFORMATION CONTACT:
 Patrick Raulerson, Center for Drug
 Evaluation and Research, Food and
 Drug Administration, 10903 New
 Hampshire Ave., Bldg. 51, Rm. 6368,
 Silver Spring, MD 20993-0002, 301-
 796-3522.

SUPPLEMENTARY INFORMATION: In 1984,
 Congress enacted the Drug Price
 Competition and Patent Term
 Restoration Act of 1984 (Pub. L. 98-417)
 (the 1984 amendments), which
 authorized the approval of duplicate
 versions of drug products approved
 under an ANDA procedure. ANDA
 applicants must, with certain
 exceptions, show that the drug for
 which they are seeking approval
 contains the same active ingredient in
 the same strength and dosage form as
 the "listed drug," which is a version of
 the drug that was previously approved.
 ANDA applicants do not have to repeat
 the extensive clinical testing otherwise
 necessary to gain approval of a new
 drug application (NDA). The only
 clinical data required in an ANDA are
 data to show that the drug that is the
 subject of the ANDA is bioequivalent to
 the listed drug.