

rehabilitation, stroke specialist care, and primary care providers.

When Congress directed the CDC to establish the Paul Coverdell National Acute Stroke Program (PCNASP) in 2001, CDC intended to monitor trends in stroke and stroke care, with the ultimate mission of improving the quality of care for stroke patients in the United States. Since 2015, CDC has funded and provided technical assistance to nine state health departments to develop comprehensive stroke systems of care. A comprehensive system of care improves quality of care by creating seamless transitions for individuals experiencing stroke. In such a system, pre-hospital providers, in-hospital providers, and early post-hospital providers coordinate patient hand-offs and ensure continuity of care. CDC contracted with RTI International to conduct an assessment of the state health departments awarded grants in 2015 to assess their implementation in their state-based contexts and progress toward short- and intermediate-term outcomes.

CDC and RTI International propose to collect information from all nine funded PCNASP grantees to gain insight into the effectiveness of implementation of their quality improvement strategies, development (and use) of a data integrated management system, and partner collaboration in building comprehensive state-wide stroke systems of care. The information collection will focus on describing PCNASP specific contributions to effective state-based stroke systems of care and the costs associated with this work. Two components of the information collection include: (1) Program implementation cost data collection from program partners using a cost and resource utilization tool; and (2) telephone interviews with key program stakeholders, such as the PCNASP principal investigator, program manager, quality improvement specialist, data analyst/program evaluator, and partner support staff. Cost data collection will focus on a stratified sample of partners' cumulative spending to support PCNASP activities,

spending by reporting period, and spending associated with specific PCNASP strategies related to building comprehensive state-wide stroke systems of care. Interview questions will target how each grantee implemented its strategies, challenges encountered and how they were overcome, factors that facilitated implementation, lessons learned along the way, and observed outcomes and improvements. The information to be collected does not currently exist for large scale, statewide programs that employ multiple combinations of strategies led by state public health departments to build comprehensive stroke systems of care. The insights to be gained from this data collection will be critical to improving immediate efforts and achieving the goals of spreading and replicating state-level strategies that are proven programmatically and are cost-effective in contributing to a higher quality of care for stroke patients.

The total estimated annual burden hours are 328. There are no costs to the respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Partner Program Manager	Cost Resource and Utilization Tool	137	1	2
Principal Investigator	Telephonic Interviews	3	1	1
Grantee Program Manager	Telephonic Interviews	3	1	1
Quality Improvement Specialist	Telephonic Interviews	3	1	1
Data Analyst/Program Evaluator	Telephonic Interviews	3	1	1
Partner Support Staff	Telephonic Interviews	6	1	1

Leroy A. Richardson,
 Chief, Information Collection Review Office,
 Office of Scientific Integrity, Office of the
 Associate Director for Science, Office of the
 Director, Centers for Disease Control and
 Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-18-18XG; Docket No. CDC-2018-0034]

Proposed Data Collection Submitted for Public Comment and Recommendations

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice with comment period.

SUMMARY: The Centers for Disease Control and Prevention (CDC), as part of its continuing effort to reduce public burden and maximize the utility of government information, invites the general public and other Federal agencies to take this opportunity to comment on proposed and/or continuing information collections, as required by the Paperwork Reduction Act of 1995. This notice invites comment on a proposed information collection project titled "Evaluation of the third decade of the National Occupational Research Agenda (NORA) Council Effectiveness". This is a survey to collect information from NORA council members and leaders about council activities and satisfaction with council functioning.

DATES: Written comments must be received on or before June 18, 2018.

ADDRESSES: You may submit comments, identified by Docket No. CDC-2018-0034 by any of the following methods:

- *Federal eRulemaking Portal: Regulations.gov.* Follow the instructions for submitting comments.

- *Mail:* Leroy A. Richardson, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS-D74, Atlanta, Georgia 30329.

Instructions: All submissions received must include the agency name and Docket Number. CDC will post, without change, all relevant comments to *Regulations.gov*.

Please note: Submit all Federal comments through the Federal eRulemaking portal (regulations.gov) or by U.S. mail to the address listed above.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the information collection plan and instruments, contact Leroy A. Richardson, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton

Road, NE, MS-D74, Atlanta, Georgia 30329; phone: 404-639-7570; Email: omb@cdc.gov.

SUPPLEMENTARY INFORMATION: Under the Paperwork Reduction Act of 1995 (PRA) (44 U.S.C. 3501-3520), Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. In addition, the PRA also requires Federal agencies to provide a 60-day notice in the **Federal Register** concerning each proposed collection of information, including each new proposed collection, each proposed extension of existing collection of information, and each reinstatement of previously approved information collection before submitting the collection to OMB for approval. To comply with this requirement, we are publishing this notice of a proposed data collection as described below.

The OMB is particularly interested in comments that will help:

1. Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;
2. Evaluate 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. Enhance the quality, utility, and clarity of the information to be collected; and
4. Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology,

e.g., permitting electronic submissions of responses.

5. Assess information collection costs.

Proposed Project

Evaluation of the third decade of the National Occupational Research Agenda (NORA) Council Effectiveness—New—National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The National Institute for Occupational Safety and Health (NIOSH) is responsible for conducting research and making recommendations to prevent worker injury and illness, as authorized in Section 20(a)(1) of the Occupational Safety and Health Act (29 U.S.C. 669). In 1995-1996, NIOSH saw an opportunity to enhance its ability to accomplish its mission through partnerships that involved a broad national stakeholder base in occupational safety and health. With stakeholder input, NIOSH developed and launched a partnership program titled the National Occupational Research Agenda (NORA) in 1996. Participation in NORA includes stakeholders from universities, large and small businesses, professional societies, government agencies, and worker organizations. NORA runs in ten year cycles, with the first decade running 1996-2006, the second 2006-2016, and the third 2016-2026.

The structure of NORA has evolved over time, and now, in the third decade, it is organized into ten industry sectors based on major areas of the U.S. economy, and seven health and safety cross-sectors organized according to the major health and safety issues affecting the U.S. working population. The work of the sectors and cross-sectors is managed through a partnership

structure of councils. Each of the 17 councils develops and maintains an agenda for the decade for its sector. The sector agendas become part of the national agenda for improvements in occupational safety and health through research and partnerships. Representing all stakeholders, the councils use an open process to set research objectives, share information, encourage partnerships, and promote improved workplace practices.

NIOSH seeks to request a 12-month Office of Management and Budget (OMB) approval to administer a survey to NORA council members and leaders. As the steward of NORA, it is NIOSH's responsibility to ensure that councils, which are central to the work of NORA, are operating well. Without this data collection, NIOSH's internal review of NORA would lack critical stakeholder input from its many non-Federal partners.

The target population is all current and former members and leaders of each of the 17 NORA councils in the third decade of NORA. The web-based survey requests information on council activities, the effectiveness of the council and its processes, and suggestions for improving the effectiveness and impact of NORA councils in the future. Without this data collection, NIOSH's internal management review of NORA would lack critical stakeholder input from its many non-Federal partners.

NIOSH has developed a 17-item survey and will send to approximately 425 non-Federal NORA Sector council members or leaders. NIOSH estimates that it will take 12 minutes to complete the survey.

There are no costs to respondents other than their time. The total estimated time burden is 85 hours.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
Non-federal NORA Council members or leaders.	Council Survey	425	1	12/60	85
Total	85

Leroy A. Richardson,
*Chief, Information Collection Review Office,
 Office of Scientific Integrity, Office of the
 Associate Director for Science, Office of the
 Director, Centers for Disease Control and
 Prevention.*

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**DEPARTMENT OF HEALTH AND
 HUMAN SERVICES**

**Centers for Disease Control and
 Prevention**

[30Day-18-0278]

**Agency Forms Undergoing Paperwork
 Reduction Act Review**

In accordance with the Paperwork Reduction Act of 1995, the Centers for Disease Control and Prevention (CDC) has submitted the information collection request titled National Hospital Ambulatory Medical Care Survey (NHAMCS) to the Office of Management and Budget (OMB) for review and approval. CDC previously published a “Proposed Data Collection Submitted for Public Comment and Recommendations” notice on November 27, 2017 to obtain comments from the public and affected agencies. CDC received one comment related to the previous notice. This notice serves to allow an additional 30 days for public and affected agency comments.

CDC will accept all comments for this proposed information collection project. The Office of Management and Budget is particularly interested in comments that:

(a) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;

(b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;

(c) Enhance the quality, utility, and clarity of the information to be collected;

(d) Minimize the burden of the collection of information on those who are to respond, including, through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses; and

(e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639-7570 or send an email to *omb@cdc.gov*. Direct written comments and/or suggestions regarding the items contained in this notice to the Attention: CDC Desk Officer, Office of Management and Budget, 725 17th Street NW, Washington, DC 20503 or by fax to (202) 395-5806. Provide written comments within 30 days of notice publication.

Proposed Project

National Hospital Ambulatory Medical Care Survey (NHAMCS) (OMB Control Number 0920-0278, Expiration 02/28/2018)—Reinstatement with change—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. The National Hospital Ambulatory Medical Care Survey (NHAMCS) has conducted annually since 1992. NCHS is seeking OMB approval to reinstate this survey for an additional three years, following a brief discontinuation on February 28, 2018.

The target universe of the NHAMCS is in-person visits made to emergency departments (EDs) of non-Federal, short-stay hospitals (hospitals with an average length of stay of less than 30 days) that have at least six beds for inpatient use, and with a specialty of general and

medical, maternity, children’s general, or long term acute care.

NHAMCS was initiated to complement the National Ambulatory Medical Care Survey (NAMCS, OMB Control Number 0920-0234, Expiration 03/31/2019), which provides similar data concerning patient visits to physicians’ offices. NAMCS and NHAMCS are the principal sources of data on ambulatory care provided in the United States.

NHAMCS provides a range of baseline data on the characteristics of the users and providers of hospital ambulatory medical care. Data collected include patients’ demographic characteristics, reason(s) for visit, providers’ diagnoses, diagnostic services, medications, and disposition. These data, together with trend data, may be used to monitor the effects of change in the health care system, for the planning of health services, improving medical education, determining health care work force needs, and assessing the health status of the population.

Starting 2018, CDC will implement just the ED component of NHAMCS. However, once reinstated the 2017 survey will run concurrently with the 2018 survey until the final months of pending 2017 data collection have been completed. This is typical with any data collection cycle: It begins in the last month of the preceding year and ends around the middle of the following year. For the 2017 data collection, CDC will collect information on all three settings (ED, OPD, and ASL). For this three-year request, CDC does not expect substantive changes or supplements for the survey.

Users of NHAMCS 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 are no costs to the respondents other than their time. The total estimated annualized burden hours are 1,251.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Hospital Chief Executive Officer	Hospital Induction 2017 Data Collection	20	1	75/60
Hospital Chief Executive Officer	Hospital Induction 2018+ Data Collection	340	1	45/60
Ancillary Service Executive	Ambulatory Unit Induction (ED, OPD and ASL).	840	1	15/60
Ancillary Service Executive	Ambulatory Unit Induction (ED only)	578	1	15/60