The goal of this ICR is to collect information from awardees funded under the Prescription Drug Overdose Prevention for States (CDC–RFA–CE15–1501) cooperative agreement, for program monitoring and improvement among funded state health departments.

Information to be collected will provide crucial data for program performance monitoring and budget tracking, and provide CDC with the capacity to respond in a timely manner to requests for information about the program from the Department of Health and Human Services (HHS), the White House, Congress, and other sources. Awardees will report progress and activity information to CDC on an annual schedule using an Excel-based fillable electronic templates, prepopulated to the extent possible by CDC

staff, to be submitted via Grant Solutions. Each awardee will submit an Annual reporting Progress Report Tool. The estimated burden per response is 4 hours for each Annual reporting Progress Report Tool. In addition, each awardee will submit an Annual reporting Evaluation Plan Tool. The estimated burden per response is 3 hours for each Annual reporting Evaluation Plan Tool.

In Year 1, each awardee will have additional burden related to initial collection of the reporting tools. Initial Collection Annual Progress Report Tool is estimated to be 20 hours per response, Initial population of the tools is a one-time activity which is annualized over the 3 years of the information collection request. After completing the initial population of the tools, pertinent

information only needs to be updated for each annual report. The same instruments will be used for all information collection and reporting.

CDC will use the information collected to monitor each awardee's progress and to identify facilitators and challenges to program implementation and achievement of outcomes.

Monitoring allows CDC to determine whether an awardee is meeting performance and budget goals and to make adjustments in the type and level of technical assistance provided to them, as needed, to support attainment of their performance measures.

OMB approval is requested for three years. Participation in the information collection is required as a condition of funding. There are no costs to 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)	Total burden (in hours)
State and Territorial Health Department Program Awardees.	Initial Collection Annual Progress Report Tool.	16	1	20	320
	Annual reporting—Progress Report Tool.	16	1	4	64
	Annual reporting Evaluation Plan Tool.	16	1	4	64
Total					448

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-15-15UJ]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) has submitted the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995. The notice for the proposed information collection is published to obtain comments from the public and affected agencies.

Written comments and suggestions from the public and affected agencies

concerning the proposed collection of information are encouraged. Your comments should address any of the following: (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*. Written comments and/or suggestions regarding

the items contained in this notice should be directed to the Attention: CDC 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

Examining How Local Health Departments Can Leverage Age-Friendly Cities Initiatives to Build Resilience in Elderly Populations—New—Office of Public Health Preparedness and Response, Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Despite considerable progress in efforts to define and build community resilience (CR), critical gaps remain in addressing the needs of older adults (age 60+), which is expected to rise to 25% by 2050. Age Friendly Initiatives (AFIs), including Senior Villages (SV) represent a promising strategy for U.S. communities and cities to support older adults aging in place, and could potentially build CR. However, few AFIs have wholly incorporated the critical element of emergency preparedness and

resilience. Even when these domains have been included, there is no evaluation of whether these efforts have resulted in improved resilience outcomes among seniors (e.g., greater self-sufficiency). This study will quantify the contribution that AFIs and SVs have made to improving resilience outcomes for older adults and provide guidance to local health departments (LHDs) for improving their engagement with AFIs/SVs.

The Office of Public Health Preparedness and Response proposes to conduct a new information collection, Examining How Local Health Departments Can Leverage Age-Friendly Cities Initiatives to Build Resilience in Elderly Populations. Information collection activities will target four groups. Respondents will include AFI Staff, Village Directors, LHD Representatives, and adults aged 65+ within the AFI and SV communities.

The study will outline where current AFIs and CR efforts align; conduct interviews in AFIs and SVs across the U.S. to understand relationships with LHDs; clarify the process through which policymakers can incorporate CR into AFIs; survey test sites in a quasi-experimental design of AFIs currently

underway; and develop a toolkit to help LHDs identify the need for AFIs, evaluate and monitor AFIs ability to improve resilience, develop effective and efficient partnerships with AFIs to expand AFI–LHD efforts across the U.S to build community resilience.

OMB approval is requested for two years. Participation in the survey is voluntary. There are no costs to respondents other than their time. The total estimated annual burden hours are 302. A summary of annualized burden hours is below.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hrs)
Age Friendly Initiative Staff	Interview Guide for Age Friendly Initiative Staff.	16	1	30/60
Senior Village Director	Interview Guide for Senior Village Director	15	1	30/60
Local Health Department Representative	Interview Guide for Local Health Department Representative.	8	1	30/60
Older Adult—Screened Out	Senior Village Survey	716	1	2/60
Older Adult—Participant	Senior Village Survey	775	1	20/60

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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BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[CMS-6059-N3]

Medicare, Medicaid, and Children's Health Insurance Programs:
Announcement of the Extended Temporary Moratoria on Enrollment of Ambulance Suppliers and Home Health Agencies in Designated Geographic Locations

AGENCY: Centers for Medicare & Medicaid Services (CMS), HHS. **ACTION:** Extension of temporary moratoria.

SUMMARY: This document announces the extension of temporary moratoria on the enrollment of new ambulance suppliers and home health agencies, subunits, and branch locations in specific locations within designated metropolitan areas in Florida, Illinois, Michigan, Texas, Pennsylvania, and New Jersey to

prevent and combat fraud, waste, and abuse.

DATES: Effective Date: July 29, 2015. **FOR FURTHER INFORMATION CONTACT:** Belinda Gravel, (410) 786–8934.

News media representatives must contact CMS' Public Affairs Office at (202) 690–6145 or email them at *press@cms.hhs.gov*.

SUPPLEMENTARY INFORMATION:

I. Background

A. CMS' Imposition of Temporary Enrollment Moratoria

Section 6401(a) of the Affordable Care Act added a new section 1866(j)(7) to the Social Security Act (the Act) to provide the Secretary with authority to impose a temporary moratorium on the enrollment of new Medicare, Medicaid, or CHIP providers and suppliers, including categories of providers and suppliers, if the Secretary determines a moratorium is necessary to prevent or combat fraud, waste, or abuse under these programs. For a more detailed explanation of these authorities, please see the July 31, 2013 notice (78 FR 46339) or February 4, 2014 extension and establishment of a temporary moratoria document (hereinafter referred to as the February 4, 2014 moratoria document or notice) (79 FR 6475).

Based on this authority and our regulations at § 424.570, we initially

imposed moratoria to prevent enrollment of new home health agencies, subunits, and branch locations 1 (hereafter referred to as HHAs) in Miami-Dade County, Florida and Cook County, Illinois, as well as surrounding counties, and part B ambulance suppliers in Harris County, Texas and surrounding counties, in a notice issued on July 31, 2013 (78 FR 46339). We then exercised this authority again in a notice published on February 4, 2014 (79 FR 6475) when we extended the existing moratoria for an additional 6 months and expanded it to include enrollment of HHAs in Broward County, Florida; Dallas County, Texas; Harris County, Texas; and Wayne County, Michigan and surrounding counties, and enrollment of ground ambulance suppliers in Philadelphia, Pennsylvania and surrounding counties. Then, we further extended the previously mentioned moratoria in moratoria documents issued on August 1, 2014 (79 FR 44702) and February 2, 2015 (80 FR 5551).

¹As noted in the preamble to the final rule implementing the moratorium authority (February 2, 2011, CMS–6028–FC (76 FR 5870), home health agency subunits and branch locations are subject to the moratoria to the same extent as any other newly enrolling home health agency.