Grant funding also provides awardees with the ability to respond rapidly to emerging health issues, including outbreaks of diseases or pathogens. The PHHS Block Grant program is authorized by sections 1901–1907 of the Public Health Service Act.

CDC currently collects information from Block Grant awardees to monitor their objectives and activities (Preventive Health and Health Services Block Grant, OMB No. 0920–0106, exp. 8/31/2016). Each awardee is required to submit an annual application for funding (Work Plan) that describes its objectives and the populations to be addressed, and an Annual Report that describes activities, progress toward objectives, and Success Stories which highlight the improvements Block Grant programs have made and the value of program activities. Information is submitted electronically through the web-based Block Grant Information Management System (BGMIS).

CDC PHHS Block Grant program has benefited from this system by efficiently collecting mandated information in a format that allows data to be easily retrieved in standardized reports. The electronic format verifies completeness of data at data entry prior to submission to CDC, reducing the number of re-submissions that are required to provide concise and complete information.

The Work Plan and Annual Report are designed to help Block Grant awardees attain their goals and to meet reporting requirements specified in the program’s authorizing legislation. Each Work Plan objective is defined in SMART format (Specific, Measurable, Achievable, Realistic and Time-based), and includes a specified start date and end date. Block Grant activities adhere to the Healthy People (HP) framework established by the Department of Health and Human Services (HHS). The current version of the BGMIS associates each awardee-defined activity with a specific HP National Objective, and identifies the location where funds are applied. Although there are no substantive changes to the information collected (Attachment 4A), the Work Plan guidance document for users (Attachment 4B) has been updated to improve their usability and the clarity of instructions provided to BGMIS users. These changes are summarized in Attachments 4C.

There are no changes to the number of Block Grant awardees (respondents), or the estimated burden per response for the Work Plan or the Annual Report. At this time, the BGMIS does not collect data related to performance measures, but a future information collection request may outline additional reporting requirements related to performance measures.

The PHHS Block Grant program must continue to collect data in order to remain in compliance with legislative mandates. The system allows CDC and Grantees to measure performance, identifying the extent to which objectives were met and identifying the most highly successful program interventions. CDC requests OMB approval to continue the Block Grant information collection for three years. CDC will continue to use the BGMIS to monitor awardee progress, identify activities and personnel supported with Block Grant funding, conduct compliance reviews of Block Grant awardees, and promote the use of evidence-based guidelines and interventions. There are no changes to the number of respondents or the estimated annual burden per respondent. The Work Plan and the Annual Report will be submitted annually. The estimated burden per response for the Work Plan is 20 hours and the estimated burden per response for the Annual Report is 15 hours.

Participation in this information collection is required for Block Grant awardees. There are no costs to respondents other than their time. Awardees continue to submit Success Stories with their Annual Progress reports through BGMIS, without changes.

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
<th>Total burden (in hours)</th>
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<td>Work Plan</td>
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<td>1</td>
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<td></td>
<td>Annual Report</td>
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<tr>
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<td></td>
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<td><strong>2,135</strong></td>
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</tbody>
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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

[Document Identifier: CMS–10406 and CMS–10599]

Agency Information Collection Activities: Proposed Collection; Comment Request

**AGENCY:** Centers for Medicare & Medicaid Services, HHS.

**ACTION:** Notice.

**SUMMARY:** The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (the PRA), federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information (including each proposed extension or reinstatement of an existing collection of information) and to allow 60 days for public comment on the proposed action. Interested persons are invited to send comments regarding our burden estimates or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency’s functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality,
utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

DATES: Comments must be received by April 5, 2016.

ADDRESSES: When commenting, please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in any one of the following ways:
1. Electronically. You may send your comments electronically to http://www.regulations.gov. Follow the instructions for “Comment or Submission” or “More Search Options” to find the information collection document(s) that are accepting comments.
2. By regular mail. You may mail written comments to the following address: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number ____, Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of following:
2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov.
3. Call the Reports Clearance Office at (410) 786–4799.

FOR FURTHER INFORMATION CONTACT: Reports Clearance Office at (410) 786–1326.

SUPPLEMENTARY INFORMATION:

Contents
This notice sets out a summary of the use and burden associated with the following information collections. More detailed information can be found in each collection’s supporting statement and associated materials (see ADDRESSES).

CMS–10406 Medicare Probable Fraud Measurement Pilot; CMS–10599 Medicare Prior Authorization of Home Health Services Demonstration

Under the 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.

The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA requires federal agencies to publish a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice.

1. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Medicare Probable Fraud Measurement Pilot; Use: The Centers for Medicare & Medicaid Services (CMS) is seeking Office of Management and Budget (OMB) approval of the collections required for a probable fraud measurement pilot. The probable fraud measurement pilot would establish a baseline estimate of probable fraud in payments for home health care services in the fee-for-service Medicare program. CMS and its agents will collect information from home health agencies, the referring physicians and Medicare beneficiaries selected in a national random sample of home health claims. The pilot will rely on the information collected along with a summary of the service history of the HHA, the referring provider, and the beneficiary to estimate the percentage of total payments that are associated with probable fraud and the percentage of all claims that are associated with probable fraud for Medicare fee-for-service home health. Form Number: CMS–10406 (OMB Control Number 0938–1192); Frequency: Annually; Affected Public: Individual and Private Sector (Business or other for-profits); Number of Respondents: 6,000; Total Annual Responses: 6,000; Total Annual Hours: 7,500. (For policy questions regarding this collection contact Cecelia Franco at (786) 313–0737.)

2. Type of Information Collection Request: New Collection; Title of Information Collection: Medicare Prior Authorization of Home Health Services Demonstration; Use: Section 402(a)(1)(J) of the Social Security Amendments of 1967 (42 U.S.C. 1395b–1(a)(1)(J)) authorizes the Secretary to “develop or demonstrate improved methods for the investigation and prosecution of fraud in the provision of care or services under the health programs established by the Social Security Act (the Act).” In accordance with this authority, we seek to develop and implement a Medicare demonstration project, which we believe will help in developing improved procedures for the identification, investigation, and prosecution of Medicare fraud occurring among HHAs providing services to Medicare beneficiaries.

This demonstration would help assure that payments for home health services are appropriate before the claims are paid, thereby preventing fraud, waste, and abuse. As part of this demonstration, we propose performing prior authorization before processing claims for home health services in: Florida, Texas, Illinois, Michigan, and Massachusetts. We would establish a prior authorization procedure that is similar to the Prior Authorization of Power Mobility Device (PMD) Demonstration, which was implemented by CMS in 2012. This demonstration would also follow and adopt prior authorization processes that currently exist in other health care programs such as TRICARE, certain state Medicaid programs, and in private insurance.

The information required under this collection is requested by Medicare contractors to determine proper payment or if there is a suspicion of fraud. Medicare contractors will request the information from HHA providers submitting claims for payment from the Medicare program in advance to determine appropriate payment. Form Number: CMS–10599 (OMB Control Number: 0938–NEW); Frequency: Occasionally; Affected Public: Private sector (Business or other for-profits and Not-for-profits); Number of Respondents: 908,740; Number of Responses: 1; Total Annual Hours: 454,370. (For questions regarding this collection contact Carla David (410) 786–4799.)


William N. Parham, III, Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.