the meeting that will cover briefings and BSC deliberation on the following topics: Interval updates from the OPHPR Director and OPHPR Divisions and Offices; updates from the Biological Agent Containment working group; discussion of Industry, Private Sector, and Public Health Interactions Supporting Emergency Preparedness and Response; and Preparedness Updates from Liaison Representatives.

Day two of the meeting will cover briefings and BSC deliberation on the following topics: OPHPR Office of Policy, Planning and Evaluation activities; CDC’s Data Preparedness activities; Public Health System Perspectives on Hurricanes Response; and Excellence in Response Operations Initiative. Agenda items are subject to change as priorities dictate.

The Director, Management Analysis and Services Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Elaine Baker,
Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2018–06546 Filed 3–30–18; 8:45 am]
BILLING CODE 4163–19–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS–10191]

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 the necessity and utility of the proposed information collection for the proper performance of the agency’s functions, the accuracy of the estimated burden, ways to enhance the quality, utility, and clarity of the information to be collected, and the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

DATES: Comments must be received by June 1, 2018.

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) 786–1326.

FOR FURTHER INFORMATION CONTACT: William Parham at (410) 786–4669.

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–10191 Medicare Parts C and D Program Audit and Timeliness Monitoring Data Requests

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.

Information Collection

1. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Medicare Parts C and D Program Audit and Timeliness Monitoring Data Requests; Use: Medicare Part D plan sponsors and Medicare Advantage organizations (collectively referred to as sponsoring organizations) are required to comply with all Medicare Parts C and D program requirements. In 2010, the explosive growth of these sponsoring organizations precipitated the need for CMS to develop an annual audit strategy to ensure that we evaluate sponsoring organizations compliance with the program requirements. In addition to describing how sponsoring organizations are selected for audit and which program areas will be audited, CMS’ annual audit strategy reflected a move to a more targeted, data-driven, and risk-based audit approach. Since 2010, CMS has continued to focus on assisting the industry with improving their operations to ensure beneficiaries receive appropriate access to care. CMS has developed audit protocols that focus on high-risk areas that have the greatest potential for beneficiary harm. CMS’ program audit protocols are posted to the CMS website each year for use by sponsoring organizations to prepare for their audit. Currently CMS utilizes the following 5 protocols to audit sponsoring organizations’ performance: Compliance Program Effectiveness (CPE), Formulary Administration (FA); Coverage Determinations, Appeals, and Grievances (CDAG); Organization Determinations, Appeals, and Grievances (ODAG), Special Needs Program Model of Care (SNP–MOC) (only administered on organizations who operate SNPs). Beginning in audit
year 2019, the SNP–MOC program area has been more accurately renamed Special Needs Program Care Coordination Quality Improvement Performance Evaluation (SNP–CCQIPE). In addition, the Medication Therapy Management (MTM) pilot protocol has been suspended until further notice. For that reason, it is no longer posted to the CMS website.

Beginning in audit year 2019, the data collected via program-specific record layouts, and collected via impact analyses on an as-needed basis, will be consolidated into each program area data request document. The pre-audit issue summary was updated for technical terminology changes. Three of the questionnaires and the power point template that previously have been distributed as part of our CPE audits will remain. However, the CPE self-assessment questionnaire and the CDAG and ODAG questionnaires have been removed. We have added new questionnaires for FA and SNP–CCQIPE. A revised template for collecting root cause analyses from organizations on an as-needed basis during the program audit has been included in this package.

We have also included a new independent validation audit work plan template that will be collected from sponsors that are required to undergo an independent validation audit. The validation audit is part of our robust audit process where CMS requires sponsoring organizations that have been audited and found to have deficiencies to undergo a validation audit to ensure correction. The validation audit utilizes the same audit protocols, but only tests the elements where deficiencies were found, as opposed to re-administering the entire audit. This validation audit work plan template will be populated by the sponsoring organization’s independent auditing firm to describe how it plans to test for correction of the deficiencies identified during the program audit.

To assist in improving the audit process, we have also included an audit feedback questionnaire that is representative of the survey link we send to sponsoring organizations at the end of each program audit. Completion of this questionnaire is optional for sponsoring organizations to provide feedback on the audit process.

The proposed changes to each data collection instrument, along with the new FA and SNP–CCQIPE questionnaires, root cause template, validation audit work plan template and audit feedback questionnaire are included in the posted PRA package.

Finally, separate from the audit process and in order to address sponsoring organizations’ concerns regarding undue harm in Star Ratings during audit years. The number of sponsoring organizations that are required to submit universes annually for their coverage/organization determinations and appeals increased. In 2016, CMS expanded this annual collection to all MA and Part D sponsoring organizations. The universes are submitted in the same format as required for audits under the Part D CIDAG protocol and the Part C ODAG protocol. The universes are then analyzed for timelines on an annual basis, across all sponsoring organizations, to allow a more comprehensive review of the accuracy of Part C and D appeals data to calculate Star Ratings. Form Number: CMS–10191 (OMB control number: 0938–1000); Frequency: Yearly; Affected Public: Private Sector (business or other for-profit and not-for-profit institutions); Number of Respondents: 166; Total Annual Responses: 211; Total Annual Hours: 51,548. (For policy questions regarding this collection contact Brenda Hudson at 443–743–9299.)


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

[FR Doc. 2018–06645 Filed 3–30–18; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Community Living

Agency Information Collection Activities; Submission for OMB Review; Public Comment Request; Older American Act Title III and Title VII (Chapters 3 and 4) Annual State Program Reporting (Annual Performance Data Collection); This is a Revision to the Existing State Program Report (OMB Approval 0985–0008)

AGENCY: Administration for Community Living (ACL), HHS.

ACTION: Notice.

SUMMARY: The Administration for Community Living is announcing that the proposed collection of information listed above has been submitted to the Office of Management and Budget (OMB) for review and clearance as required under section 506(c)(2)(A) of the Paperwork Reduction Act of 1995. This 30-Day notice collects comments on the information collection requirements related to annual performance data from State grantees under the Older Americans Act related to Title III and Title VII (Chapters 3 and 4) of that act. Title III includes, for example, home delivered and congregate meal services, transportation and caregiver service; and Title VII includes Elder Abuse Prevention and Legal Assistance Development (ICR Rev).

DATES: Submit written comments on the collection of information by May 2, 2018.

ADDRESSES: Submit written comments on the collection of information by:

(a) Email to: OIRA_submission@omb.eop.gov, Attn: OMB Desk Officer for ACL;
(b) Fax to 202.395.5806, Attn: OMB Desk Officer for ACL; or
(c) By mail to the Office of Information and Regulatory Affairs, OMB, New Executive Office Bldg., 725 17th St. NW, Rm. 10235, Washington, DC 20503. Attn: OMB Desk Officer for ACL.

FOR FURTHER INFORMATION CONTACT: ACL’s Office of Performance and Evaluation at SPRredesign.comments@acl.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, ACL has submitted the following proposed collection of information to OMB for review and clearance. This collection is a revision of the 2016 approved version of the State Program Report and incorporates significant reduction in data collected. This data collection is essential to provide performance measures as required by Congress and the GPRA Modernization Act of 2010 (GPRAMA). Significant revisions to the SPR were last implemented in 2005. This proposed collection is a revision of the currently approved version (effective 2016–2019). The factors that influenced the proposed revision of the SPR, include: (1) The need to modernize the data structure to allow for more efficient reporting and the ability to use current technology for reporting and analysis; (2) the interest in aligning data elements within and across data collections; (3) the need to consider alternative data elements that reflect the current Aging Network and long-term care services and supports; and (4) the need to reduce reporting burden while enhancing data quality. The proposed SPR revision reduces the number of data elements reported by 70% and the amount of time for completion by 30% as compared to the current 2016–2019 SPR. This is a reduction of 874 hours from the previous version.

[FR Doc. 2018–06645 Filed 3–30–18; 8:45 am]