Following the presentations, there will be a moderated session during which members of the public can provide oral comments on the DEIS. Commenters will be allowed three minutes to provide comments. Comments will be recorded. Refer to the end of this notice for instructions on how to access the online public meeting.

Virtual Public Meeting Information:
Members of the public may join the
DEIS public meeting by entering the
following information—Meeting ID: 986
1911 2163; Password: CHFBEIS, using
any of the below methods. Note that the
meeting is best viewed through the
Zoom app, and attendees are
encouraged to download the app at the
Zoom website (https://zoom.us) or on
their mobile device, and test their
connection prior to the meeting to
ensure best results.

• From their personal computer by launching the Zoom app (if already installed), and clicking 'Join a Meeting' and entering the above Meeting ID and Password. Attendees should follow the prompts to input their name and email address to access the meeting.

• From their personal computer, by going to the Zoom website at http://zoom.us/join, entering the Meeting ID and Password, and following the prompts to download and install the Zoom app.

• From their mobile device through the Zoom mobile app, by entering the above Meeting ID and Password.

• For attendees who do not have the Zoom app or do not wish to download the app, visit http://zoom.us/join using your computer's browser, enter the Meeting ID, and click the 'Join from your browser' link that is displayed on the landing page. Then, follow the prompts to enter your name and the meeting Password.

Whether joining through the Zoom app or web browser, attendees should follow the prompts to connect their computer audio. Attendees are encouraged to connect through the 'Computer Audio' tab and click 'Join Audio by Computer' under the 'Join Audio' button on the bottom of their screen. Users who do not have a computer microphone and wish to provide public comment during the meeting may connect by following the prompts under the 'Phone Call' tab under the 'Join Audio' button.

For members of the public who do not have access to a personal computer, they may join the meeting audio by dialing the following number: 669–900–9128. When prompted, enter the following information: Meeting ID—986 1911 2163, followed by the pound (#)

key; press pound (#) again when prompted for a participant ID; then enter Password—629071 followed by the pound (#) key. Note, dialing in to the meeting is only necessary if you are not accessing the meeting through your computer or mobile app, or if you would like to provide oral comments during the meeting but do not have a computer microphone. The public meeting will be recorded, and all comments provided will become part of the formal record.

Jared Bradley,

Director, Portfolio Management Division, Pacific Rim Region, Public Buildings Service. [FR Doc. 2020–14710 Filed 7–7–20; 8:45 am]

BILLING CODE 6820-YF-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier CMS-R-153]

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 September 8, 2020.

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:

- 1. Access CMS' website address at website address at https://www.cms.gov/ Regulations-and-Guidance/Legislation/ PaperworkReductionActof1995/PRA-Listing.html.
- 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 N. 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-R-153 Medicaid Drug Use Review (DUR) Program

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: Medicaid Drug Use Review (DUR) Program; Use: States must provide for a review of drug therapy before each prescription is filled or delivered to a Medicaid patient. This review includes screening for potential drug therapy problems due to therapeutic duplication, drug-disease contraindications, drug-drug interactions, incorrect drug dosage or duration of drug treatment, drug-allergy interactions, and clinical abuse/misuse. Pharmacists must make a reasonable effort to obtain, record, and maintain Medicaid patient profiles. These profiles must reflect at least the patient's name, address, telephone number, date of birth/age, gender, history, e.g., allergies, drug reactions, list of medications, and pharmacist's comments relevant to the individual's drug therapy.

The States must conduct RetroDUR which provides for the ongoing periodic examination of claims data and other records in order to identify patterns of fraud, abuse, inappropriate or medically unnecessary care. Patterns or trends of drug therapy problems are identified and reviewed to determine the need for intervention activity with pharmacists and/or physicians. States may conduct interventions via telephone, correspondence, or face-to-face contact.

Annual reports are submitted to CMS for the purposes of monitoring compliance and evaluating the progress of States' DUR programs. The information submitted by States is reviewed and results are compiled by CMS in a format intended to provide information, comparisons, and trends related to States' experiences with DUR. States benefit from the information and may enhance their programs each year based on State reported innovative practices that are compiled by CMS from the DUR annual reports. Form Number: CMS-R-153 (OMB control number: 0938-0659); Frequency: Yearly, quarterly, and occasionally; Affected Public: State, Local, or Tribal Governments; Number of Respondents: 51; Total Annual Responses: 663; Total Annual Hours: 41,004. (For policy questions regarding this collection contact Mike Forman at 410-786-2666.)

Dated: July 2, 2020.

William N. Parham, III,

Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2020–14714 Filed 7–7–20; 8:45 am]

BILLING CODE 4120-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier CMS-10116]

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 ČMS' 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 September 8, 2020.

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:

- 1. Access CMS' website address at website address at https://www.cms.gov/ Regulations-and-Guidance/Legislation/ PaperworkReductionActof1995/PRA-Listing.html
- 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 N. 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-10116 Medicare Program: Conditions for Payment of Power Mobility Devices, including Power Wheelchairs and Power-Operated Vehicles

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: Extension of a currently approved collection; Title of Information Collection: Medicare