a key component of some of the economic statistics it is responsible for tracking. In addition, statutes in several states and U.S. territories refer to, or rely upon, the MIRS or the ARM Index for various purposes.4

The OMB control number for this information collection is 2590–0004. The current clearance for the information collection expires on July 31, 2017.

B. Burden Estimate

The Agency received a total of 1,369 monthly MIRS data submissions from 45 unique survey respondents over the period 2014–2016, representing an average of 456.3 monthly submissions per year from all respondents. Based on that figure and the expectation that it may receive slightly fewer data submissions going forward as compared to the last three years, FHFA estimates that it will receive an average of 450 data submissions annually over the next three years.

Most MIRS respondents submit their monthly MIRS data electronically through FHFA’s MIRS web interface. Several, primarily larger, respondents transmit an electronic data file to FHFA, which then uploads the data to the same web interface. A few respondents still elect to complete FHFA Form #075 and submit it by facsimile. FHFA believes that, on average, a respondent will spend 20 minutes transmitting each monthly MIRS data set.

Thus, FHFA estimates that the annualized hour burden on all respondents imposed by this information collection over the next three years will be 150 hours (450 submissions x 0.33 hours).

C. Comments Request

FHFA requests written comments on the following: (1) Whether the collection of information is necessary for the proper performance of FHFA functions, including whether the information has practical utility; (2) the accuracy of FHFA’s estimates of the burdens of the collection of information; (3) ways to enhance the quality, utility, and clarity of the information collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology.

event title link and enter the required information. Include any requests for special accommodations.

**Note:** Participants who do not plan to attend the meeting in person should not register. No registration is required for participants who plan to participate in the meeting via webcast or teleconference.

Because of staff and resource limitations, we cannot accept comments and presentations by facsimile (FAX) transmission.

**Meeting Location, Webcast, and Teleconference**

The meeting will be held in the Auditorium, CMS Central Office, 7500 Security Boulevard, Woodlawn, Maryland 21244–1850. Alternately, the public may either view this meeting via a webcast or listen by teleconference. During the scheduled meeting, webcasting is accessible online at: http://cms.gov/live. Teleconference dial-in information will appear on the final meeting agenda, which will be posted on our Web site when available at: http://www.cms.gov/Regulations-and-Guidance/Guidance/FACA/AdvisoryPanelonAmbulatoryPaymentClassificationGroups.html.

**News Media**

Representatives must contact our Public Affairs Office at (202) 690–6145.

**Advisory Committees’ Information Lines**

The phone number for the CMS Federal Advisory Committee Hotline is (410) 786–3985.

**Web Sites**

For additional information on the Panel, including the Panel charter, and updates to the Panel’s activities, we refer readers to view our Web site at: http://www.cms.gov/Regulations-and-Guidance/Guidance/FACA/AdvisoryPanelonAmbulatoryPaymentClassificationGroups.html.

Information about the Panel and its membership in the Federal Advisory Committee Act database are also located at: http://facadatabase.gov/.

**FOR FURTHER INFORMATION CONTACT:**

**SUPPLEMENTARY INFORMATION:**

**I. Background**

The Secretary of the Department of Health and Human Services (DHHS) is required by section 1833(f)(9)(A) of the Social Security Act (the Act) and is allowed by section 222 of the Public Health Service Act (PHS Act) to consult with an expert outside panel, such as the Advisory Panel on Outpatient Payment (the Panel), regarding the clinical integrity of the Ambulatory Payment Classification (the APC) groups and relative payment weights. The Panel is governed by the provisions of the Federal Advisory Committee Act (Pub. L. 92–463), as amended (5 U.S.C. Appendix 2), to set forth standards for the formation and use of advisory panels. We consider the technical advice provided by the Panel as we prepare the proposed and final rules to update the Hospital Outpatient Prospective Payment System (OPPS) for the following calendar year.

**II. Agenda**

The agenda for the August 21 through August 22, 2017 Panel meeting will provide for discussion and comment on the following topics as designated in the Panel’s Charter:

- Addressing whether procedures within an APC group are similar both clinically and in terms of resource use.
- Evaluating APC group structure.
- Reviewing the packaging of OPPS services and costs, including the methodology and the impact on APC groups and payment.
- Removing procedures from the inpatient-only list for payment under the OPPS.
- Using single and multiple procedure claims data for CMS’ determination of APC group weights.
- Addressing other technical issues concerning APC group structure.
- Recommending the appropriate supervision level (general, direct, or personal) for individual hospital outpatient therapeutic services.

The Agenda will be posted on our Web site at: https://www.cms.gov/Regulations-and-Guidance/Guidance/FACA/AdvisoryPanelonAmbulatoryPaymentClassificationGroups.html approximately 1 week before the meeting.

**III. Presentations**

The subject matter of any presentation and comment matter must be within the scope of the Panel designated in the Charter. Any presentations or comments outside of the scope of this Panel will be returned or requested for amendment. Unrelated topics include, but are not limited to, the conversion factor, charge compression, revisions to the cost report, pass-through payments, correct coding, new technology applications (including supporting information/documentation), provider payment adjustments, supervision of hospital outpatient diagnostic services, and the types of practitioners that are permitted to supervise hospital outpatient services. The Panel may not recommend that services be designated as nonsurgical extended duration therapeutic services.

The Panel may use data collected or developed by entities and organizations other than DHHS and Centers for Medicare & Medicaid Services (CMS) in conducting its review. We recommend organizations submit data for CMS staff and the Panel’s review.

All presentations are limited to 5 minutes, regardless of the number of individuals or organizations represented by a single presentation. Presenters may use their 5 minutes to represent either 1 or more agenda items.

**Section 508 Compliance**

For this meeting, we are aiming to have all presentations and comments available on our Web site. Materials on our Web site must be Section 508 compliant to ensure access to federal employees and members of the public with and without disabilities. We encourage presenters and commenters to refer to guidance on making 508 compliant form. Such guidance is available at: http://www.cms.gov/Research-Statistics-Data-and-Systems/CMS-Information-Technology/Section508/508-Compliant-doc.html. We will review presentations and comments for 508 compliance, and place compliant materials on our Web site. As resources permit, we will also convert non-compliant submissions to 508 compliant forms and offer assistance to submitters who wish to make their submissions 508 compliant. All non-508 compliant presentations and comments will be shared with the public onsite and through the webcast and made available to the public upon request.

Those wishing to access such materials should contact the DFO (the DFO’s address, email, and phone number are provided in this notice).

In order to consider presentations and/or comments, we will need to receive the following:

1. An email copy of the presentation or comments sent to the DFO mailbox, APCPanel@cms.hhs.gov or, if unable to submit by email, a hard copy sent to the DFO at the address noted under the FOR FURTHER INFORMATION CONTACT section of this notice.
2. Form CMS-20017 with complete contact information that includes name, address, phone number, and email addresses for all presenters and commenters and a contact person that can answer any questions, and provide revisions that are requested, for the presentation. Presenters and commenters must clearly explain the actions that they are requesting CMS to take in the appropriate section of the form. A presenter’s or commenter’s relationship with the organization that they represent must also be clearly listed.

• The form is now available through the CMS Forms Web site at: https://www.cms.gov/Medicare/CMS-Forms/CMS-Forms/downloads/cms20017.pdf.

• We encourage presenters to make efforts to ensure that their presentations and comments are 508 compliant.

IV. Oral Comments

In addition to formal oral presentations, which are limited to 5 minutes total per presentation, there will be an opportunity during the meeting for public oral comments, which will be limited to 1 minute for each individual and a total of 3 minutes per organization.

V. Meeting Attendance

The meeting is open to the public; however, attendance is limited to space available. Priority will be given to those who pre-register and attendance may be limited based on the number of registrants and the space available. Persons wishing to attend this meeting, which is located on federal property, must register by following the instructions in the DATES section of this notice under “Meeting Registration Timeframe”. A confirmation email will be sent to the registrants shortly after completing the registration process.

VI. Security, Building, and Parking Guidelines

The following are the security, building, and parking guidelines:

• Persons attending the meeting, including presenters, must be pre-registered and on the attendance list by the prescribed date.

• Individuals who are not pre-registered in advance may not be permitted to enter the building and may be unable to attend the meeting.

• Attendees must present a government-issued photo identification to the Federal Protective Service or Guard Service personnel before entering the building. Without a current, valid photo ID, persons may not be permitted entry to the building.

• Security measures include inspection of vehicles, inside and out, at the entrance to the grounds.

• All persons entering the building must pass through a metal detector.

• All items brought into CMS, including personal items, for example, laptops and cell phones, are subject to physical inspection.

• The public may enter the building 30 to 45 minutes before the meeting convenes each day.

• All visitors must be escorted in areas other than the lower and first-floor levels in the Central Building.

• The main-entrance guards will issue parking permits and instructions upon arrival at the building.

• Foreign nationals visiting any CMS facility require prior approval. If you are a foreign national and wish to attend the meeting onsite, in addition to registering for the meeting, you must also send a separate email to APCPanel@cms.hhs.gov prior to the close of registration to request authorization to attend as a foreign national.

Note: As of March 30, 2015, the “Real ID Act” requires a second form of identification from those whose government issued photo identification or government issued driver’s license was issued by American Samoa, Arizona, Louisiana, Maine, Minnesota, and New York. Attendees with a government issued photo identification or driver’s license issued by the states previously mentioned may need to provide alternative or additional approved proof of identification in order to comply with the “Real ID Act.”

VII. Special Accommodations

Individuals requiring special accommodations must include the request for these services during registration.

VIII. Panel Recommendations and Discussions

The Panel’s recommendations at any Panel meeting generally are not final until they have been reviewed and approved by the Panel on the last day of the meeting, before the final adjournment. These recommendations will be posted to our Web site after the meeting.

IX. Collection of Information Requirements

This document does not impose information collection requirements, that is, reporting, recordkeeping, or third-party disclosure requirements. Consequently, there is no need for review by the Office of Management and Budget under the authority of the Paperwork Reduction Act of 1995 (44 U.S.C. 3501 et seq.).

Dated: May 18, 2017.

Seema Verma,

Administrator, Centers for Medicare & Medicaid Services.

[FR Doc. 2017–10683 Filed 5–24–17; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration


Agency Information Collection Activities; Announcement of Office of Management and Budget Approvals

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is publishing a list of information collections that have been approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995.

FOR FURTHER INFORMATION CONTACT: Ila S. Mizrachi, Office of Operations, Food and Drug Administration, Three White Flint North, 11601 Landsdown Street, North Bethesda, MD 20852, 301–796–7726, PRAS staff@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: The following is a list of FDA information collections recently approved by OMB under section 3507 of the Paperwork Reduction Act of 1995 (44 U.S.C. 3507). The OMB control number and expiration date of OMB approval for each information collection are shown in table 1. Copies of the supporting statements for the information collections are available on the Internet at https://www.reginfo.gov/public/do/PRAMain. An Agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number.