**Federal Register** at 82 FR 43022, on September 13, 2017. One comment was received.

**DATES:** Submit comments on or before January 25, 2018.

ADDRESSES: Submit comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to: Office of Information and Regulatory Affairs of OMB, Attention: Desk Officer for GSA, Room 10236, NEOB, Washington, DC 20503. Additionally submit a copy to GSA by any of the following methods:

- Regulations.gov: http://
  www.regulations.gov. Submit comments
  via the Federal eRulemaking portal by
  searching the OMB control number
  9000–0013. Select the link that
  corresponds with "Information
  Collection 9000–0013, Certified Cost or
  Pricing Data and Data Other Than
  Certified Cost or Pricing Data." Follow
  the instructions provided on the screen.
  Please include your name, company
  name (if any), and "Information
  Collection 9000–0013, Certified Cost or
  Pricing Data and Data Other Than
  Certified Cost or Pricing Data", on your
  attached document.
- Mail: General Services
  Administration, Regulatory Secretariat
  Division (MVCB), 1800 F Street NW,
  Washington, DC 20405. ATTN: Ms.
  Mandell/IC 9000–0013, Certified Cost or
  Pricing Data and Data Other Than
  Certified Cost or Pricing Data.

Instructions: Please submit comments only and cite Information Collection 9000–0013, Certified Cost or Pricing Data and Data Other Than Certified Cost or Pricing Data, in all correspondence related to this collection. All comments received will be posted without change to <a href="http://www.regulations.gov">http://www.regulations.gov</a>, including any personal and/or business confidential information provided.

FOR FURTHER INFORMATION CONTACT: Mr. Michael O. Jackson, Procurement Analyst, Federal Acquisition Policy Division, GSA 202–208–4949, or michaelo.jackson@gsa.gov.

### SUPPLEMENTARY INFORMATION:

### A. Purpose

The Truth in Negotiations Act requires the Government to obtain certified cost or pricing data under certain circumstances. Contractors may request an exemption from this requirement under certain conditions and provide other information instead.

#### **B. Public Comment**

A 60 day notice was published in the **Federal Register** at 82 FR 43022, on September 13, 2017. One comment was

received; however, it was not substantive, and did not change the estimate of the burden.

### C. Annual Reporting Burden

Fiscal year 2016 data was obtained from the Federal Procurement Data System to estimate burdens for the provisions and clauses addressed in this information collection notice. This update does not include the requirements at FAR 42.7, Indirect Cost Rates, as this requirement is covered under OMB Control Number 9000-0069. The data for 52.215-20 is for new contract awards in FY 2016. The data for modifications and orders executed in FY 2016 applies to new contract awards as well as to prior multiple year contracts that continue to be active. The following is a summary of the FY 2016

1. Subcontractor C&P Data-Mods (FAR 52.214–28)

Respondents: 8. Responses per Respondent: 1. Total Responses: 8. Hours per Response: 160. Total Burden Hours: 1,280.

2. Subcontractor C&P Data (FAR 52.215–12)

Respondents: 3,832. Responses per Respondent: 1. Total Responses: 3,832. Hours per Response: 160. Total Burden Hours: 613,120.

3. Subcontractor C&P Data-Mods (FAR 52.214–13)

Respondents: 1,292. Responses per Respondent: 1. Total Responses: 1,292. Hours per Response: 160. Total Burden Hours: 206,720.

4. Requirement for C&P Data and Data Other Than C&P Data (FAR 52.215–20)

Respondents: 25,853. Responses per Respondent: 1.69. Total Responses: 117,225. Hours per Response: 143. Total Burden Hours: 6,506,140.

5. Requirement for C&P Data and Data Other Than C&P Data-Mods (FAR 52.215–21)

Respondents: 8,440. Responses per Respondent: 3. Total Responses: 27,623. Hours per Response: 106. Total Burden Hours: 2,432,560.

### 6. Total

Respondents: 39,425. Responses per Respondent: 3.80. Total Responses: 149,980. Hours per Response: 65. Total Burden Hours: 9,759,820.
Obtaining copies of proposals:
Requesters may obtain a copy of the information collection documents from the General Services Administration,
Regulatory Secretariat Division (MVCB),
1800 F Street NW, Washington, DC
20405, telephone 202–501–4755. Please cite OMB Control No. 9000–0013,
Certified Cost or Pricing Data and Data
Other Than Certified Cost or Pricing
Data, in all correspondence.

Dated: December 19, 2017.

#### Lorin S. Curit,

Director, Federal Acquisition Policy Division, Office of Government-wide Acquisition Policy, Office of Acquisition Policy, Office of Government-wide Policy.

[FR Doc. 2017-27672 Filed 12-22-17; 8:45 am]

BILLING CODE 6820-EP-P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Agency for Healthcare Research and Quality

#### **Notice of Meeting**

**AGENCY:** Agency for Healthcare Research and Quality, HHS.

**ACTION:** Notice.

**SUMMARY:** Each Special Emphasis Panel (SEP) meeting will commence in open session before closing to the public for the duration of the meeting "AHRQ RFA-HS17-011, National Research Service Award (NRSA) Institutional Research Training Grant (T32)."

DATES: January 11–12, 2018 (Open on January 11 from 8:00 a.m. to 8:30 a.m. and closed for the remainder of the meeting).

**ADDRESSES:** Hilton Rockville, 1750 Rockville Pike, Rockville, MD 20857.

#### FOR FURTHER INFORMATION CONTACT:

Anyone wishing to obtain a roster of members, agenda or minutes of the nonconfidential portions of this meeting should contact: Mrs. Bonnie Campbell, Committee Management Officer, Office of Extramural Research, Education and Priority Populations, AHRQ, 5600 Fishers Lane, Rockville, Maryland 20857.

Agenda items for this meeting are subject to change as priorities dictate.

### SUPPLEMENTARY INFORMATION: In

accordance with section 10 (a)(2) of the Federal Advisory Committee Act (5 U.S.C. App. 2), announcement is made of an Agency for Healthcare Research and Quality (AHRQ) Special Emphasis Panel (SEP) meeting on AHRQ National Research Service Award (NRSA) Institutional Research Training Grant (T32) " AHRQ RFA-HS17-011, National Research Service Award (NRSA) Institutional Research Training Grant (T32)."

A SEP is a group of experts in fields related to health care research who are invited by AHRQ, and agree to be available on an as needed basis, to conduct scientific reviews of applications for AHRQ support. Individual members of the Panel do not attend regularly scheduled meetings and do not serve for fixed terms or a long period of time. Rather, they are asked to participate in particular review meetings which require their type of expertise.

Each SEP meeting will commence in open session before closing to the public for the duration of the meeting. The SEP meeting referenced above will be closed to the public in accordance with the provisions set forth in 5 U.S.C. App. 2, section 10(d), 5 U.S.C. 552b(c)(4), and 5 U.S.C. 552b(c)(6). Grant applications for the AHRQ RFA-HS17-011, "National Research Service Award (NRSA) Institutional Research Training Grant (T32)," are to be reviewed and discussed at this meeting. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Dated: December 19, 2017.

#### Gopal Khanna,

Director.

[FR Doc. 2017–27664 Filed 12–22–17; 8:45 am]

BILLING CODE 4160-90-P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Agency for Healthcare Research and Quality

## Notice of an Upcoming Challenge Competition

**AGENCY:** Agency for Healthcare Research and Quality (AHRQ), Department of Health and Human Services.

**ACTION:** Notice.

**SUMMARY:** This notice announces the intention of the Agency for Healthcare Research and Quality (AHRQ) to conduct a Challenge Competition in Fall 2018 to develop user-friendly technical tools to collect and integrate patient-reported outcome data in electronic health records or other health information technology products.

#### FOR FURTHER INFORMATION CONTACT:

Janey Hsiao, Health Scientist Administrator, Center for Evidence and Practice Improvement, Agency for Healthcare Research and Quality, 5600 Fishers Lane, Mail Stop 06E73A, Rockville, Maryland, 20857, Email: Janey.hsiao@ahrq.hhs.gov, Phone: (301) 427–1335.

#### SUPPLEMENTARY INFORMATION:

#### **Background**

The patient's perspective is central to healthcare decisions affecting prevention, diagnosis, treatment, and long-term care. Patient-reported outcomes (PROs) critically inform patient-centered outcomes research (PCOR) and can inform clinical management of individuals, shared decision making, patient selfmanagement support, care planning, goal setting and goal attainment. PROs offer a complementary perspective to that of clinician assessments, and may provide greater insights into health status, function, symptom burden, adherence, health behaviors, and quality of life. However, standardized tools that collect PRO data in a way that is meaningful and useful to both patients and clinicians in primary care and ambulatory settings are not widely available.

The limited inclusion of PRO data in electronic health records (EHRs) and other health information technology (IT) solutions reduces the understanding and use of the patient's perspective in research and clinical care. Further, while some EHRs are currently able to capture some structured PRO data, including many of the NIH-funded Patient Reported Outcomes Measurement Information System® (PROMIS®) instruments, this information is not commonly collected in routine care. Thus, these data are often not available for both clinical care and research. Moreover, standards do not exist for collecting and integrating PRO data into health IT systems, thereby limiting the ability to easily share these data across health systems for research or other purposes including quality improvement.

## **Proposed Project**

To fill these gaps, AHRQ intends to support the development of user-friendly, PRO-collection tools that utilize health IT standards, including application programming interfaces (APIs) to collect physical function data in ambulatory care settings (including primary care). Data element and data capture standards would allow for PRO assessments to be conducted and easily shared regardless of what EHR or health

IT solution is being used. It would also allow for consistency in interpretation, and clarify the meaning of results for patient-provider communication and shared decision-making.

The development of user-friendly, PRO-collection tools will be conducted though a multi-phase Challenge Competition in Fall 2018. The statutory authority for this challenge competition is Section 105 of the America COMPETES Reauthorization Act of 2010. Only the winners from each phase can move on to the next phase so the participant pool becomes more limited throughout the competition. Developers will be asked to create tools based on implementation specifications provided by AHRQ. The tools should enable patients to share their physical function data with clinicians and researchers. AHRQ will convene a panel to judge the Challenge Competition. The judges of the Challenge Competition will evaluate the resulting submissions for adhering to the implementation specifications set forth in the Challenge Competition.

AHRQ will manage the Challenge Competition including developing the concept, designing prizes, drafting the **Federal Register** Notice, setting up the Challenge website, answering questions from developers, and giving prizes to winners. The Challenge Competition will be conducted by AHRQ in furtherance of the Secretary's authority to develop interoperable data networks that can link data from multiple sources, including electronic health records. 42 U.S.C. 299b–37(f).

Dated: December 19, 2017.

## Gopal Khanna,

Director.

[FR Doc. 2017–27663 Filed 12–22–17; 8:45 am]

BILLING CODE 4160-90-P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## Centers for Disease Control and Prevention

[30Day-18-0822]

# Agency Forms Undergoing Paperwork Reduction Act Review

In accordance with the Paperwork Reduction Act of 1995, the Centers for Disease Control and Prevention (CDC) has submitted the information collection request titled *The National Intimate Partner and Sexual Violence Survey (NISVS)* to the Office of Management and Budget (OMB) for review and approval. CDC previously published a "Proposed Data Collection Submitted for Public Comment and