proprietary, competitively sensitive information of a rival SRM supplier supporting Northrop’s missile system business could be transferred to Northrop’s vertically integrated SRM business.

VI. The Consent Agreement

The Consent Agreement remedies the acquisition’s likely anticompetitive effects by requiring, whenever Northrop competes for a missile system prime contract, that Northrop make its SRM products and related services available on a non-discriminatory basis to all other third-party competing prime contractors that wish to purchase them. The non-discrimination prohibitions of the Consent Agreement are comprehensive and apply to any potential discriminatory conduct affecting price, schedule, quality, data, personnel, investment, technology, innovation, design, or risk.

The Consent Agreement requires Northrop to establish firewalls to ensure that Northrop does not transfer or use any proprietary information that it receives from competing missile prime contractors or SRM suppliers in a manner that harms competition. These firewall provisions require that Northrop maintain separate firewalled teams to support offers of SRMs to different third-party missile prime contractors and to maintain these firewalled teams separate from the team supporting Northrop’s missile prime contractor activities. The firewall provisions also prohibit Northrop’s missile business from sharing proprietary information it may receive from third-party SRM suppliers with Northrop’s SRM business.

The Consent Agreement also provides that the DOD’s Under Secretary of Defense for Acquisition and Sustainment shall appoint a compliance officer to oversee Northrop’s compliance with the Order. The compliance officer will have all the necessary investigative powers to perform his or her duties, including the right to interview respondent’s personnel, inspect respondent’s facilities, and require respondents to provide documents, data, and other information. The compliance officer has the authority to retain third-party advisors, at the expense of Northrop, as appropriate to perform his or her duties. Access to these extensive resources will ensure that the compliance officer is fully capable of overseeing the implementation of, and compliance with, the Order.

The purpose of this analysis is to facilitate public comment on the proposed Consent Agreement, and it is not intended to constitute an official interpretation of the proposed Consent Agreement or to modify its terms in any way.

 By direction of the Commission.

Janice Frankle, Acting Secretary.

[FR Doc. 2018–12750 Filed 6–13–18; 8:45 am]

BILLING CODE 6750–01–P

GOVERNMENT ACCOUNTABILITY OFFICE

Request for Nominations for the Physician-Focused Payment Model Technical Advisory Committee (PTAC)

AGENCY: Government Accountability Office (GAO).

ACTION: Request for letters of nomination and resumes.

SUMMARY: The Medicare Access and CHIP Reauthorization Act of 2015 established the Physician-Focused Payment Model Technical Advisory Committee to provide comments and recommendations to the Secretary of Health and Human Services on physician payment models, and gave the Comptroller General responsibility for appointing its members. GAO is now accepting nominations of individuals for this committee.

DATES: Letters of nomination and resumes should be submitted no later than July 20, 2018, to ensure adequate opportunity for review and consideration of nominees prior to appointment. Appointments will be made in October 2018.

ADDRESSES: Submit letters of nomination and resumes by either of the following methods: Email: PTACcommittee@gao.gov. Include PTAC Nominations in the subject line of the message, or Mail: U.S. GAO, Attn: PTAC Nominations, 441 G Street NW, Washington, DC 20548.

FOR FURTHER INFORMATION CONTACT: Greg Giusto at (202) 512–8268 or giustog@gao.gov if you do not receive an acknowledgement within a week of submission or if you need additional information. For general information, contact GAO’s Office of Public Affairs, (202) 512–4800.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Agency for Healthcare Research and Quality, HHS.

ACTION: Notice.

SUMMARY: This notice announces the intention of the Agency for Healthcare Research and Quality (AHRQ) to request that the Office of Management and Budget (OMB) approve the proposed information collection project “Ambulatory Surgery Center Survey on Patient Safety Culture Database.” This proposed information collection was previously published in the Federal Register on March 14th, 2018 and allowed 60 days for public comment. AHRQ received no substantive comments from members of the public. The purpose of this notice is to allow an additional 30 days for public comment.

DATES: Comments on this notice must be received by July 16, 2018.

ADDRESSES: Written comments should be submitted to: AHRQ’s OMB Desk Officer by fax at (202) 395–6974 (attention: AHRQ’s desk officer) or by email at OIRA Submission@omb.eop.gov (attention: AHRQ’s desk officer).

FOR FURTHER INFORMATION CONTACT: Doris Lefkowitz, AHRQ Reports Clearance Officer, (301) 427–1477, or by email at doris.lefkowitz@ahrq.hhs.gov.

SUPPLEMENTARY INFORMATION:

Proposed Project

Ambulatory Surgery Center Survey on Patient Safety Culture Database

In accordance with the Paperwork Reduction Act, 44 U.S.C. 3501–3521, AHRQ invites the public to comment on this proposed information collection. Ambulatory surgery centers (ASCs) are a fast-growing health care setting, demonstrating tremendous growth both in the volume and complexity of procedures being performed. ASCs provide surgical services to patients who are not expected to need an inpatient stay following surgery. The Centers for Medicare and Medicaid Services (CMS) defines ASCs as distinct entities that operate exclusively to provide surgical services to patients who do not require hospitalization and are not expected to need to stay in a surgical facility longer than 24 hours.
How AHRQ’s Mission and Directives Relate to ASCs. As described in its 1999 reauthorizing legislation, Congress directed the Agency for Healthcare Research and Quality (AHRQ) to enhance the quality, appropriateness, and effectiveness of health services, as well as access to such services, by establishing a broad base of scientific research and promoting clinical and health systems practice improvements. The legislation also directed AHRQ to “conduct and support research, evaluations, and training, support demonstration projects, research networks, and multidisciplinary centers, provide technical assistance, and disseminate information on health care and on systems for the delivery of such care, including activities with respect to health statistics, surveys, database development, and epidemiology.” 42 U.S.C. 299a(a)(8).

Shortly after Congress enacted this legislation, the Institute of Medicine (IOM) published “To Err is Human,” a seminal report on medical errors that connects the dots between errors and workplace culture. In it, the IOM called for health care organizations to develop a “culture of safety” such that staffing and system processes are aligned to improve the reliability and safety of patient care. This appeal for safety culture improvements directly relates to AHRQ’s legislative directive and mission (i.e., “to produce evidence to make health care safer, higher quality, more accessible, equitable, and affordable, and to work within the U.S. Department of Health and Human Services and with other partners to make sure that the evidence is understood and used”). Given its legislatively-mandated role, AHRQ is uniquely positioned to support data collection and analyses that will help fuel ASC patient safety culture improvements. The expanding volume and scope of ASC services, the growing attention of federal regulators on patient safety within ASCs, and the resultant implications for public health has prompted AHRQ to present this application to the Office of Management and Budget (OMB). In this request, AHRQ seeks OMB approval to expand its Surveys on Patient Safety Culture (SOPS)TM program by creating an ASC SOPS Database to capture and report on ASC SOPS data voluntarily-submitted by ASCs that have administered the ASC SOPS. The ASC SOPS Database is the newest database for the SOPS program and would be modeled after AHRQ’s existing SOPS Databases for Hospitals, Medical Offices, Nursing Homes, and Community Pharmacies, which have all undergone OMB review and approval.

Background on ASC SOPS. This section provides context for this request to the OMB regarding the need for AHRQ’s requested database. Factors include the continued ASC growth trajectory and increasing public attention on the quality of ASC care—particularly as it relates to patient safety culture.

Rapid ASC Growth. Medicare-certified ASCs have experienced impressive growth in the last 35 years—up from 239 facilities in 1983 to 5,316 in 2010. In recent years, Medicare ASCs have seen continued growth in both their number and scope, as illustrated by the annual average growth rate of 1.1 percent between 2010 to 2014. In 2015, CMS spent $4.1 billion for 3.4 million fee-for-service Medicare beneficiaries to receive care across 5,500 Medicare-certified ASCs. Research suggests that transitioning eligible surgical procedures from inpatient to ASC settings may yield significant and sustained Medicare cost savings.

Federal Attention on ASC Care Quality and Safety Culture. Concern about the quality of ASC care is not new. Following a 2008 Hepatitis C outbreak in Nevada blamed on poor ASC infection control practices, HHS’s Office of the Secretary oversaw a $10 million program for state survey agencies to improve healthcare-associated infection reduction in ASCs. The Centers for Disease Control’s (CDC) National Healthcare Safety Network (NHSN) subsequently expanded its surgical site infection (SSI) surveillance efforts to enable ASC data submission to accommodate state SSI reporting mandates. Through the Affordable Care Act of 2010, Congress also pursued ASC performance improvement by directing the HHS Secretary to implement an ASC-focused Medicare value-based purchasing (VBP) program.

The relationship between patient safety culture and the quality of ASC care has attracted more recent attention from policymakers and regulators. On the national level, the Joint Commission in early 2017 established within its ASC accreditation manual a new chapter on patient safety systems improvement, which includes strategies for “motivating staff to uphold a fair and just safety culture.” CMS, meanwhile, published in November 2017 its Final Rule outlining the ASC Quality Reporting (ASCQR) Program, which ties quality and patient safety performance to reimbursement.

Rationale for the information collection. AHRQ sponsored the development of the ASC SOPS as a new survey in the suite of AHRQ Surveys on Patient Safety Culture. The database will support AHRQ’s goals of promoting improvements in the quality and safety of health care in ASC settings. Like the survey and other toolkit materials, the database results will be made publicly available on AHRQ’s website. Technical assistance is provided by AHRQ through its contractor at no charge to ASCs to facilitate the use of these materials for ASC patient safety and quality improvement. Technical assistance will also be provided to support ASC data submission.
The goal of this project is to create the ASC SOPS Database. This database will:

1. Present results from ASCs that voluntarily submit their data;
2. Present trend data for ASCs that have submitted their data more than once;
3. Provide data to ASCs to facilitate internal assessment and learning in the patient safety improvement process; and
4. Provide supplemental information to help ASCs identify their strengths and areas with potential for improvement in patient safety culture.

This study is being conducted by AHRQ through its contractor, Westat, pursuant to AHRQ's statutory authority to conduct and support research on health care and on systems for the delivery of such care, including activities with respect to the quality, effectiveness, efficiency, appropriateness and value of health care services and with respect to health statistics, surveys, and database development. 42 U.S.C 299a(a)(1) and (8).

Method of Collection

To achieve the goal of this project the following activities and data collections will be implemented:

1. Eligibility and Registration Form—The point-of-contact (POC), often the staff position, tenure, and hours worked per week). The Database Report will include a section on data limitations, emphasizing that the report does not reflect a representative sampling of the U.S. ASC population. Because participating ASCs will choose to voluntarily submit their data into the database and therefore are not a random or national sample of ASCs, estimates based on this self-selected group might be biased. These limitations will be noted in the database report. We will recommend that users review the database results with these caveats in mind.

Each ASC that submits its data will receive a customized survey feedback report that presents their results alongside the aggregated results from other participating ASCs. If an ASC submits data more than once, its survey feedback report will also present trend data.

ASC users of the ASCs SOPS Survey, Database Reports, and Individual ASC Survey Feedback Reports can use these documents to:

- Raise staff awareness about patient safety;
- Diagnose and assess the current status of patient safety culture in their own ASC;
- Identify strengths and areas for patient safety culture improvement;
- Examine trends in patient safety culture change over time; and
- Evaluate the cultural impact of patient safety initiatives and intervention.

Estimated Annual Respondent Burden

Exhibit 1 shows the estimated annualized burden hours for the respondents’ time to participate in the database. Given that this will be the first call for voluntary data submission, participation is initially expected to be modest. An estimated 100 ASC managers (i.e., POCs from ASCs) will complete the database submission steps and forms. Each POC will submit the following:

- Eligibility and registration form (completion is estimated to take about 5 minutes).
- Data use agreement (completion is estimated to take about 3 minutes).
- ASC Site Information Form (completion is estimated to take about 5 minutes).
- Survey data submission will take an average of one hour.

The total burden is estimated to be 121 hours.

Exhibit 2 shows the estimated annualized cost burden based on the respondents’ time to submit their data. The cost burden is estimated to be $5,472.83.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and Prevention
[Docket No. CDC–2018–0057]

Notice of Intent To Prepare an Environmental Impact Statement, Public Scoping Meeting, and Request for Comments: Acquisition of Site for Development of a Replacement Underground Safety Research Program Facility for the Centers for Disease Control and Prevention/National Institute for Occupational Safety and Health (CDC/NIOSH) in Mace, West Virginia

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS)

ACTION: Notice of intent; announcement of public meeting; and request for comments.

SUMMARY: The Centers for Disease Control and Prevention (CDC) within the Department of Health and Human Services (HHS), in cooperation with the General Services Administration (GSA), announces its intent to prepare an Environmental Impact Statement (EIS) to analyze and assess the environmental impacts of the proposed acquisition of a site in Mace, West Virginia, and the development of this site into a replacement of the National Institute for Occupational Safety and Health (NIOSH) Underground Safety Research Program facility (Proposed Action). The current acquisition and development would replace the former Lake Lynn Experimental Mine in Fayette County, Pennsylvania and would support research programs focused on miner health and safety issues. The site being considered for acquisition and development includes 461.35 acres located off of U.S. Route 219 in Randolph and Pocahontas Counties near Mace, West Virginia.

This notice is pursuant to the requirements of the National Environmental Policy Act of 1969 (NEPA) as implemented by the Council on Environmental Quality (CEQ) Regulations (40 CFR parts 1500–1508). CDC, in cooperation with GSA, also intends to initiate consultation, as required by Section 106 of the National Historic Preservation Act (NHPA), to evaluate the potential effects, if any, of the Proposed Action on historic properties. Following the scoping meeting, a Draft EIS will be prepared and circulated for public comment. CDC is the lead federal agency for this Proposed Action.

DATES:
Public Scoping Meeting: A public scoping meeting in open house format will be held on June 26, 2018 in Slatyfork, West Virginia. The meeting will begin at 5:30 p.m. and end no later than 8:30 p.m.

Written comments: Written scoping comments must be submitted by 11:59 p.m. on July 14, 2018.

Accommodations: Persons wishing to participate in the public scoping meeting who need special accommodations should contact Sam Tarr at 770–488–8170 by 5:00 p.m. Eastern Time, June 19, 2018.

ADDRESSES: The public scoping meeting will be held at the Linwood Community Library, 72 Snowshoe Drive, Slatyfork, West Virginia 26291.

Request for Comments
In accordance with the Paperwork Reduction Act, comments on AHRQ’s information collection are requested with regard to any of the following: (a) Whether the proposed collection of information is necessary for the proper performance of AHRQ’s health care research and health care information dissemination functions, including whether the information will have practical utility; (b) the accuracy of AHRQ’s estimate of burden (including hours and costs) of the proposed collection(s) of information; (c) ways to enhance the quality, utility and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information upon the respondents, including the use of automated collection techniques or other forms of information technology.

Comments submitted in response to this notice will be summarized and included in the Agency’s subsequent request for OMB approval of the proposed information collection. All comments will become a matter of public record.

Francis D. Chesley, Jr., Acting Deputy Director.

[FR Doc. 2018–12767 Filed 6–13–18; 8:45 am]

BILLING CODE 4160–90–P