INCLUSION AND EXCLUSION CRITERIA BY POPULATION, INTERVENTION, COMPARATOR, OUTCOME, TIMING, SETTING/STUDY DESIGN (PICOTS)—Continued

PICOTS elements	Inclusion criteria	Exclusion criteria		
Timing Settings Study design		 Less than 4 weeks. Hospital and acute care. In vitro studies, nonoriginal data (e.g., narrative reviews, scoping reviews, editorials, letters, or erratum), retrospective cohort studies, case series, qualitative studies, cost-benefit analysis, cross-sectional (i.e., nonlongitudinal) studies, survey. 		
Publications	Relevant systematic reviews, or meta-analyses (used for identifying additional studies). Studies published in English only Studies published in peer-reviewed journals. Studies published at and after the year 2000.	Non-English language studies.		

Abbreviations: CVD = cardiovascular disease; GI = gastrointestinal; HDL = high-density lipoprotein; KQ = Key Question; LDL = low-density lipoprotein PICOTS = populations, interventions, comparators, outcomes, timing, and settings; RCT = randomized controlled trial; TC = total cholesterol; U.S. = United States.

Dated: January 8, 2024.

Marquita Cullom,

Associate Director.

[FR Doc. 2024-00505 Filed 1-11-24; 8:45 am]

BILLING CODE 4160-90-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Submission to OMB for Review and Approval; Public Comment Request; CAREWare Customer Satisfaction and Usage Survey, OMB No. 0906–xxxx–New

AGENCY: Health Resources and Services Administration (HRSA), Department of Health and Human Services.

ACTION: Notice.

SUMMARY: In compliance with of the Paperwork Reduction Act of 1995, HRSA has submitted an Information Collection Request (ICR) to the Office of Management and Budget (OMB) for review and approval. Comments submitted during the first public review of this ICR will be provided to OMB. OMB will accept further comments from the public during the review and approval period. OMB may act on HRSA's ICR only after the 30-day comment period for this notice has closed.

DATES: Comments on this ICR should be received no later than February 12, 2024.

ADDRESSES: Written comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular

information collection by selecting "Currently under Review—Open for Public Comments" or by using the search function.

FOR FURTHER INFORMATION CONTACT: To request a copy of the clearance requests submitted to OMB for review, email Joella Roland, the HRSA Information Collection Clearance Officer at paperwork@hrsa.gov or call (301) 443—3983.

SUPPLEMENTARY INFORMATION:

Information Collection Request Title: CAREWare Customer Satisfaction and Usage Survey, OMB No. 0906–xxxx– New.

Abstract: HRSA developed CAREWare, a software application first released in 2000, to help meet the data collection and reporting needs of Ryan White HIV/AIDS Program (RWHAP) grant recipients. The secure software is a free, electronic health and social support services information system for RWHAP grant recipients and their subrecipients to assist in the data requirement submissions that inform the development of the Ryan White HIV/AIDS Program Service Report, the AIDS Drug Assistance Program Data Report, the Ending the HIV Epidemic Initiative Triannual Report, and the voluntary Clinical Quality Measures Performance Measures module. Over time, the software has evolved into a comprehensive health information system and is now the source of more than half of all the RWHAP client-level data received from recipients and subrecipients of RWHAP grant funding. CAREWare software manages HIV clinical and support service data from more than 360,000 client records in 48 states; Washington, DC; Puerto Rico; and the U.S. Virgin Islands.

The CAREWare software application contains customizable modules for tracking demographic information,

services, medications, laboratory test results, immunization history, diagnoses (updated with International Classification of Diseases, Tenth Revision codes), referrals to outside agencies, and an appointment scheduler. There is a custom report generator and a performance measures module that supports quality of care initiatives at the provider level. The software also has several ways to import data from third-party sources, including commercial labs and other electronic health records (using both Health Level Seven and simple Comma Separated Value-formatted files), HIV surveillance systems, and for RWHAP Part B AIDS Drug Assistance Programs, pharmacy benefit programs. The software and user support materials can be accessed here: https://hab.hrsa.gov/program-grantsmanagement/careware. Finally, CAREWare supports users through an experienced helpdesk with ongoing software maintenance issues and enhancements to the user interface.

HRSA is proposing a customer satisfaction survey to gather feedback from CAREWare users regarding their experiences and satisfaction with the software platform and to obtain suggestions for improvement.

A 60-day **Federal Register** Notice (FRN) was published in the **Federal Register** on October 20, 2023 (Volume 88, No. 202, pages 72493–94). There was one out-of-scope public comment received in response to the FRN.

Need and Proposed Use of the Information: HRSA aims to understand CAREWare users' needs and concerns by collecting information on current software features and inquiring about opportunities to improve the user experience and product features. The survey will address the software's functionality and how well it meets the data collection, reporting, and quality

management needs of the CAREWare user. The feedback will enable HRSA to assess, benchmark, and improve customer satisfaction with RWHAP grant recipients.

Likely Respondents: RWHAP recipients and providers who use CAREWare to produce data files for the Ryan White HIV/AIDS Program Service Report, the AIDS Drug Assistance Program Data Report, the Ending the HIV Epidemic Initiative Triannual Report, and the voluntary Clinical Quality Measures Performance Measures module.

Burden Statement: Burden in this context means the time expended by persons to generate, maintain, retain, disclose, or provide the information requested. This includes the time needed to review instructions; to develop, acquire, install, and utilize technology and systems for the purpose of collecting, validating, and verifying information, processing and maintaining information, and disclosing and providing information; to train personnel and to be able to respond to a collection of information; to search data sources; to complete and review

the collection of information; and to transmit or otherwise disclose the information. The total annual burden hours estimated for this ICR are summarized in the table below.

A pilot with seven RWHAP grant recipients concluded on November 20, 2023, one month after the 60-day FRN publication date of October 20, 2023. The pilot resulted in a lesser burden estimate than initially reported in the 60-day FRN.

TOTAL ESTIMATED ANNUALIZED BURDEN—HOURS

Form name	Number of respondents	Number of responses per respondent	Total responses	Average burden per response (in hours)	Total burden hours
CAREWare User Survey	1,160	1	1,160	0.88	1,021
Total	1,160	1	1,160	0.88	1,021

HRSA specifically requests comments on (1) the necessity and utility of the proposed information collection for the proper performance of the agency's functions, (2) the accuracy of the estimated burden, (3) ways to enhance the quality, utility, and clarity of the information to be collected, and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

Maria G. Button,

Director, Executive Secretariat.
[FR Doc. 2024–00534 Filed 1–11–24; 8:45 am]
BILLING CODE 4165–15–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of Allergy and Infectious Diseases; Notice of Closed Meeting

Pursuant to section 1009 of the Federal Advisory Committee Act, as amended, notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The contract proposals and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the contract proposals, the disclosure of which

would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute of Allergy and Infectious Diseases Special Emphasis Panel; Small Business Innovation Research (SBIR) Phase II Program Contract Solicitation (PHS 2022–1) Topic 109— Development of Monoclonal Antibodymediated Interventions to Combat Malaria (N01).

Date: February 6, 2024.

Time: 1:00 p.m. to 4:00 p.m.

Agenda: To review and evaluate contract proposals.

Place: National Institute of Allergy and Infectious Diseases, National Institutes of Health, 5601 Fishers Lane, Room 3F36, Rockville, MD 20892 (Virtual Meeting).

Contact Person: Noton K. Dutta, Ph.D., Scientific Review Officer, Scientific Review Program, Division of Extramural Activities, National Institute of Allergy and Infectious Diseases, National Institutes of Health, 5601 Fishers Lane, Room 3F36, Rockville, MD 20852, 240–669–2857, noton.dutta@nih.gov. (Catalogue of Federal Domestic Assistance Program Nos. 93.855, Allergy, Immunology, and Transplantation Research; 93.856, Microbiology and Infectious Diseases Research, National Institutes of Health, HHS)

Dated: January 9, 2024.

Lauren A. Fleck,

Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2024-00556 Filed 1-11-24; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of Allergy and Infectious Diseases; Notice of Closed Meeting

Pursuant to section 1009 of the Federal Advisory Committee Act, as amended, notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The contract proposals and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the contract proposals, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute of Allergy and Infectious Diseases Special Emphasis Panel; HHS–NIH–CDC–SBIR PHS 2024–1 Phase I: Alternatives to Benzathine Penicillin for Treatment of Syphilis (Topic 134).

Date: February 12, 2024.

Time: 11:00 a.m. to 3:00 p.m.

Agenda: To review and evaluate contract proposals.

Place: National Institute of Allergy and Infectious Diseases, National Institutes of Health, 5601 Fishers Lane, Room 3F52A Rockville, MD 20892 (Virtual Meeting).

Contact Person: Shilpakala Ketha, Ph.D., Scientific Review Officer, Scientific Review Program, Division of Extramural Activities, National Institute of Allergy and Infectious