

building permits, sediment and erosion control permits, grading permits, and stormwater management permits.

Background

In 1981, GSA completed an EIS that analyzed the impacts from the construction of new laboratory space at the MRC and the consolidation of four facilities in the Washington, DC, metro area and other sites in St. Louis, MO, and Cincinnati, OH. In 1990, Congress enacted the Food and Drug Revitalization Act that gave GSA the authority to grant contracts to consolidate FDA facilities. To accommodate future growth and further consolidate FDA operations, GSA is preparing an EIS to assess the impacts of development on the MRC and an increase in the employee population of up to approximately 1,800 employees, over a period of 20 years.

The purpose of the proposed action is to provide a Master Plan for the MRC to guide future site development. The proposed action is needed to accommodate projected growth at the MRC and provide the necessary office and laboratory space for FDA to conduct complex and comprehensive research and reviews.

Schedule for Decision-Making

A Draft EIS is expected to be released for public review in June 2021. The GSA will hold a public hearing on the impacts of the proposed action in July 2021, and will seek preliminary approval of the MRC Master Plan from the National Capital Planning Commission (NCPC) at NCPC's September 2021 hearing. A Final EIS will be prepared that will take into consideration all comments received on the Draft EIS, and a Record of Decision is anticipated in spring 2022. Pending completion of NEPA compliance and review by NCPC, GSA anticipates adopting the MRC Master Plan in spring 2022.

Scoping Process

In accordance with NEPA, a scoping process will be conducted to aid in determining the alternatives to be considered and the scope of issues to be addressed, as well as for identifying the significant issues related to the proposed MRC Master Plan. Scoping will be accomplished through a virtual public scoping meeting; direct mail correspondence to potentially interested persons, agencies, and organizations; and meetings with agencies having an interest in the Master Plan. It is important that Federal, regional, State, and local agencies, and interested

individuals take this opportunity to identify environmental concerns that should be addressed during the preparation of the EIS.

Public Scoping Meeting

Due to the ongoing COVID-19 pandemic and state/local requirements for social distancing, a pre-recorded presentation will be available at www.gsa.gov/ncrnepa in lieu of a traditional in-person public scoping meeting. A project phone line [410-777-9537] has also been set up to listen to the presentation and to leave comments on the proposed Master Plan. The pre-recorded presentation and phone line will be available from January 4, 2021, through February 11, 2021. The GSA is publishing notices in the Washington Post and Prince George's Post announcing the meeting.

Written Comments

Agencies and the public are encouraged to provide comments on identification of potential alternatives, information, and analyses relevant to the proposed action. Comments may be provided in writing via mail or email. Verbal comments may also be provided via the project phone line. Written comments regarding the environmental analysis for the proposed MRC Master Plan must be postmarked by February 11, 2021, and sent to the following: Mr. Marshall Popkin, NEPA Compliance Specialist, Office of Planning and Design Quality, Public Buildings Service, U.S. General Services Administration, 1800 F Street NW, Room 4400, Washington, DC 20405.

Email: marshall.popkin@gsa.gov using the subject line: FDA MRC Master Plan EIS Comment.

Kristi Tunstall Williams,

Deputy Director, Office of Planning and Design Quality, Public Buildings Service, National Capital Region, U.S. General Services Administration.

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GENERAL SERVICES ADMINISTRATION

[OMB Control No. 3090-0163; Docket No. 2020-0001; Sequence No. 11]

Submission for OMB Review; General Services Acquisition Regulation; Information Specific to a Contract or Contracting Action (Not Required by Regulation)

AGENCY: Office of Acquisition Policy, General Services Administration (GSA).

ACTION: Notice of request for public comments regarding an extension to an existing OMB information collection.

SUMMARY: Under the provisions of the Paperwork Reduction Act, the Regulatory Secretariat Division will be submitting to the Office of Management and Budget (OMB) a request to review and approve an extension of a previously approved information collection requirement regarding information specific to a contract or contracting action that is not required by regulation.

DATES: Submit comments on or before: January 21, 2021.

ADDRESSES: Written comments and recommendations for this 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: Mr. Clarence Harrison, Procurement Analyst, GSA Acquisition Policy Division, at telephone 202-227-7051 or email GSARPolicy@gsa.gov.

SUPPLEMENTARY INFORMATION:

A. Purpose

GSA has various mission responsibilities related to the acquisition and provision of supplies, transportation, information technology, telecommunications, real property management, and disposal of real and personal property. These mission responsibilities generate requirements that are realized through the solicitation and award of public contracts.

Most GSA procurement-related information collections are required by the Federal Acquisition Regulation (FAR) or General Services Administration Acquisition Regulation (GSAR); each clause requiring such a collection must be individually approved by OMB. However, some solicitations require contractors to submit information specific to that contracting action, such as information needed to evaluate offers (e.g. specific instructions for technical and price proposals, references for past performance) or data used to administer resulting contracts (e.g. project management plans).

This information collection is currently associated with GSA's information collection requirements contained in solicitations issued in accordance with the Uniform Contract Format under FAR Part 14, Sealed Bidding (see GSAR 514.201-1); FAR

Part 15, Contracting by Negotiation (see GSAR 552.215–73); and solicitations under FAR Part 12, Acquisition of Commercial Items (see GSAR 512.301). This includes information collection requirements found in GSA Federal Supply Schedule (FSS) solicitations.

B. Annual Reporting Burden

Respondents: 2,597,377.

Responses per Respondent: 1.

Total Responses: 2,597,377.

Hours Per Response: .40.

Total Burden Hours: 1,038,950.

C. Public Comments

A notice was published in the **Federal Register** at 85 FR 62731 on October 5, 2020. Two comments were received. No changes were made to the information collection requirements or supporting statement as a result of the public comments, because they were not applicable to the policy.

Obtaining Copies of Proposals:

Requesters may obtain a copy of the information collection documents from the GSA Regulatory Secretariat Division, by calling 202–501–4755 or emailing GSARegSec@gsa.gov. Please cite OMB Control No. 3090–0163, Information Specific to a Contract or Contracting Action (Not Required by Regulation), in all correspondence.

Jeffrey Koses,

Senior Procurement Executive, Office of Acquisition Policy, Office of Government-wide Policy.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Solicitation for Nominations for Members of the U.S. Preventive Services Task Force (USPSTF)

AGENCY: Agency for Healthcare Research and Quality (AHRQ), HHS.

ACTION: Solicits nominations for new members of the USPSTF.

SUMMARY: The Agency for Healthcare Research and Quality (AHRQ) invites nominations of individuals qualified to serve as members of the U.S. Preventive Services Task Force (USPSTF).

DATES: Nominations must be received electronically by March 15th of a given year to be considered for appointment to begin in January of the following year.

ADDRESSES: Submit your responses electronically via: <https://uspstfnominations.ahrq.gov/register>.

FOR FURTHER INFORMATION CONTACT: Lydia Hill at coordinator@uspstf.net; 301–427–1587.

SUPPLEMENTARY INFORMATION:

Arrangement for Public Inspection

Nominations and applications are kept on file at the Center for Evidence and Practice Improvement, AHRQ, and are available for review during business hours. AHRQ does not reply to individual nominations, but considers all nominations in selecting members. Information regarded as private and personal, such as a nominee's social security number, home and email addresses, home telephone and fax numbers, or names of family members will not be disclosed to the public in accord with the Freedom of Information Act. 5 U.S.C. 552(b)(6); 45 CFR 5.31(f).

Nomination Submissions

Nominations must be submitted electronically, and should include:

1. The applicant's current curriculum vitae and contact information, including mailing address, and email address; and
2. A letter explaining how this individual meets the qualification requirements and how he or she would contribute to the USPSTF. The letter should also attest to the nominee's willingness to serve as a member of the USPSTF.

AHRQ will later ask people under serious consideration for USPSTF membership to provide detailed information that will permit evaluation of possible significant conflicts of interest. Such information will concern matters such as financial holdings, consultancies, non-financial scientific interests, and research grants or contracts.

To obtain a diversity of perspectives, AHRQ particularly encourages nominations of women, members of underrepresented populations, and persons with disabilities. Interested individuals can nominate themselves. Organizations and individuals may nominate one or more people qualified for membership on the USPSTF at any time. Individuals nominated prior to March 15, 2020, who continue to have interest in serving on the USPSTF should be re-nominated.

Qualification Requirements

To qualify for the USPSTF and support its mission, an applicant or nominee should, at a minimum, demonstrate knowledge, expertise and national leadership in the following areas:

1. The critical evaluation of research published in peer-reviewed literature and in the methods of evidence review;

2. Clinical prevention, health promotion and primary health care; and
3. Implementation of evidence-based recommendations in clinical practice including at the clinician-patient level, practice level, and health-system level.

Additionally, the Task Force benefits from members with expertise in the following areas:

- Public Health.
- Health Equity and The Reduction Of Health Disparities.
- Application of Science to Health Policy.
- Dissemination and Implementation.
- Behavioral Medicine/Clinical Health Psychology.
- Communication of Scientific Findings to Multiple Audiences Including Health Care Professionals, Policy Makers and the General Public.

Candidates with experience and skills in any of these areas should highlight them in their nomination materials.

Applicants must have no substantial conflicts of interest, whether financial, professional, or intellectual, that would impair the scientific integrity of the work of the USPSTF and must be willing to complete regular conflict of interest disclosures.

Applicants must have the ability to work collaboratively with a team of diverse professionals who support the mission of the USPSTF. Applicants must have adequate time to contribute substantively to the work products of the USPSTF.

Nominee Selection

Nominated individuals will be selected for the USPSTF on the basis of how well they meet the required qualifications and the current expertise needs of the USPSTF. It is anticipated that new members will be invited to serve on the USPSTF beginning in January, 2022. All nominated individuals will be considered; however, strongest consideration will be given to individuals with demonstrated training and expertise in the areas of Family Medicine, Internal Medicine, Pediatrics, Obstetrics and Gynecology, and Advanced Practice Nursing. AHRQ will retain and may consider for future vacancies nominations received this year and not selected during this cycle.

Some USPSTF members without primary health care clinical experience may be selected based on their expertise in methodological issues such as meta-analysis, analytic modeling, or clinical epidemiology. For individuals with clinical expertise in primary health care, additional qualifications in methodology would enhance their candidacy.