determined that 33 unique vendors was a sufficient baseline for estimating the number of respondents. USTRANSCOM further provided that the number of shipments varied from contractor to contractor, ranging from as few as 11 shipments per contractor at the low end, to over 1900 shipments per contractor at the high end. USTRANSCOM also determined that the average number of shipments for FY 2012 (approximately 10,000) by the number of unique vendors (33), was a sufficient baseline, for this estimate, in determining the average number of responses per respondent. Therefore it is estimated that, in accordance FAR 47.208 and the clause at FAR 52.247–68, contractors were required to provide 10 minutes. These revisions require to prepare this notification based on information received from USTRANSCOM, the estimated time require to prepare this notification remains at 10 minutes. These revisions represent an increase from the previously approved information collection.

Respondents: 33.
Responses per Respondent: 303.
Annual Responses: 9,999.
Hours per Response: .167.
Total Burden Hours: 1,670.

Dated: August 8, 2013.
Karlos Morgan.
Acting Director, Federal Acquisition Policy Division, Office of Governmentwide Acquisition Policy, Office of Acquisition Policy, Office of Governmentwide Policy. [FR Doc. 2013–19568 Filed 8–12–13; 8:45 am]

BILLING CODE 6820–EP–P

GENERAL SERVICES ADMINISTRATION
[OMB Control No. 3090–00XX; Docket No. 2013–0001; Sequence 8]

Information Collection; MyUSA
AGENCY: Office of Citizen Services and Innovative Technologies (OCSIT), General Services Administration (GSA).
ACTION: Notice of request for comments regarding a new information collection.

SUMMARY: Under the provisions of the Paperwork Reduction Act (44 U.S.C. chapter 35), the Regulatory Secretariat Division will be submitting to the Office of Management and Budget (OMB) a request to review and approve a new information collection requirement regarding MyUSA.

DATES: Submit comments on or before October 15, 2013.

ADDRESSES: Submit comments identified by Information Collection 3090–00XX; MyUSA by any of the following methods:
• Regulations.gov: http://www.regulations.gov. Submit comments via the Federal eRulemaking portal by searching for “Information Collection 3090–00XX; MyUSA”. Select the link “Submit a Comment” that corresponds with “Information Collection 3090–00XX; MyUSA”. Follow the instructions provided at the “Submit a Comment” screen. Please include your name, company name (if any), and “Information Collection 3090–00XX; MyUSA” on your attached document.
• Fax: 202–501–4067.
• Mail: General Services Administration, Regulatory Secretariat (MVEB), 1800 F Street NW., 2nd Floor, Washington, DC 20405–0001, ATTN: Hada Flowers/IC 3090–00XX; MyUSA.

Instructions: Please submit comments only and cite Information Collection 3090–00XX; MyUSA, in all correspondence related to this collection. All comments received will be posted without change to http://www.regulations.gov, including any personal and/or business confidential information provided.

FOR FURTHER INFORMATION CONTACT: Sarah Crane, Director, Office of Citizen Services and Innovative Technologies, General Services Administration, at telephone number 202–208.5855, or via email to Sarah.Crane@gsa.gov.

SUPPLEMENTARY INFORMATION:
A. Purpose
MyUSA (https://my.usa.gov) provides an account to users that gives them control over their interactions with government agencies and how government uses and accesses their personal information. Users have the option of creating a personal profile that can be reused across government to personalize interactions and streamline common tasks such as filling out forms. Government agencies can build applications that can request permission from the user to access their MyUSA Account and read their personal profile. The information in the system is contributed voluntarily by the user and cannot be accessed by the government without explicit consent of the user; information is not shared between government agencies, except when the user gives explicit consent to share his or her information, and as detailed in the MyUSA System of Records Notice (http://www.gpo.gov/fdsys/pkg/FR-2013-07-05/pdf/2013-16124.pdf).

The information collected is basic profile information, and may include: name, home address, phone number, gender, marital status and basic demographic information such as whether the individual is married, a veteran, a small business owner, a parent or a student.

Use of the system, and contribution of personal information, is completely voluntary.

B. Public Comments
Pursuant to section 3506(c)(2)(A) of the PRA, GSA specifically solicits comments and information to enable it to:
1. Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the Agency, including whether the information will have practical utility;
2. Evaluate the accuracy of the Agency’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;
3. Enhance the quality, utility, and clarity of the information to be collected; and
4. Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses.

C. Annual Reporting Burden
Respondents: 10,000.
Responses per Respondent: 1.
Total annual responses: 10,000.
Hours per Response: .25.
Total Burden Hours: 2,500.
Obtaining Copies of Proposals: General Services Administration,
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2013–D–0880]

Agency Information Collection Activities; Announcement of Office of Management and Budget Approval; Communicating Composite Scores in Direct-to-Consumer Advertising

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing the availability of the draft guidance for industry entitled “Frequently Asked Questions About Medical Foods; Second Edition.” The draft guidance, when finalized, will provide responses to additional questions regarding the definition, labeling, and availability of medical foods and updates to some of the existing responses.

DATES: Although you can comment on any guidance at any time (see 21 CFR 10.115(g)(5)), to ensure that the Agency considers your comment on the draft guidance before it begins work on the final version of the guidance, submit either electronic or written comments on the draft guidance by October 15, 2013.

ADDRESSES: Submit written requests for single copies of this draft guidance to the Office of Nutrition, Labeling, and Dietary Supplements, Center for Food Safety and Applied Nutrition (HFS–850), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740. Send two self-addressed adhesive labels to assist that office in processing your request. See the SUPPLEMENTARY INFORMATION section for electronic access to the draft guidance.

Submit electronic comments on the draft guidance to http://www.regulations.gov. Submit written comments to the Division of Dockets Management (see ADDRESSES). It is only necessary to send one set of comments. Identify comments with the docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday, and will be posted to the docket at http://www.regulations.gov.

III. Comments

Interested persons may submit either electronic comments regarding this draft guidance to http://www.regulations.gov or written comments to the Division of Dockets Management (see ADDRESSES).