GENERAL SERVICES ADMINISTRATION

AGENCY: Department of Defense (DOD), General Services Administration (GSA), and National Aeronautics and Space Administration (NASA).

ACTION: Notice of Request for public comments regarding a new OMB information collection.

SUMMARY: Under the provisions of the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35), the Regulatory Secretariat will be submitting to the Office of Management and Budget (OMB) a request to review and approve a new information collection requirement regarding Lessor’s Annual Cost Statement. A request for public comments was published in the Federal Register at 74 FR 83704, on December 4, 2009. No comments were received.

Public comments are particularly invited on: Whether this collection of information is necessary for the proper performance of functions of the FAR, and whether it will have practical utility; whether our estimate of the public burden of this collection of information is accurate, and based on valid assumptions and methodology; ways to enhance the quality, utility, and clarity of the information to be collected; and ways in which we can minimize the burden of the collection of information on those who are to respond, through the use of appropriate technological collection techniques or other forms of information technology.

DATES: Submit comments on or before January 28, 2011.

ADDRESSES: Submit comments identified by Information Collection 3090–00xx, Lessor’s Annual Cost Statement by any of the following methods:

- Submit comments via the Federal eRulemaking portal by inputting “Information Collection 3090–00xx, Lessor’s Annual Cost Statement” under the heading “Enter Keyword or ID” and selecting “Search”. Select the link “Submit a Comment” that corresponds with “Information Collection 3090–00xx, Lessor’s Annual Cost Statement”. Follow the instructions provided at the “Submit a Comment” screen. Please include your name, company name (if any), and “Information Collection 3090–00xx, Lessor’s Annual Cost Statement” on your attached document.
- Mail: General Services Administration, Regulatory Secretariat (MVCB), 1275 1st Street, NE., Washington, DC 20417. ATTN: Hada Flowers/IC 3090–00xx.

INSTRUCTIONS: Please submit comments only and cite Information Collection 3090–00xx, Lessor’s Annual Cost Statement, 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: Ms. Deborah Lague, Procurement Analyst, Contract Policy Branch, at telephone (202) 694–8149 or via e-mail to deborah.lague@gsa.gov.

SUPPLEMENTAL INFORMATION:

A. Purpose

In accordance with the proposed GSAR 570.802(d), the GSA Form 1217 is used to obtain information about operating expenses for property being offered for lease to house Federal agencies. These expenses are normally included in the rental payments we make to lessors. The form also provides clarity of the information to be obtained.

B. Annual Reporting Burden

Respondents: 5,733.
Responses per Respondent: 1.
Annual Responses: 5,733.
Hours per Response: 1.
Total Burden Hours: 5,733.
Obtaining Copies of Proposals: Requesters may obtain a copy of the information collection documents from the General Services Administration, Regulatory Secretariat (MVCB), 1275 1st Street, NE., Washington, DC 20417, telephone (202) 501–4755. Please cite OMB Control No. 3090–00XX, Lessor’s Annual Cost Statement, in all correspondence.


Millisa Gary,
Acting Director, Acquisition Policy Division.

BILLING CODE 6820–34–P

PRIVATELY OWNED VEHICLE MILEAGE REIMBURSEMENT RATES

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

ACTION: Notice of FTR Bulletin 11–03, Calendar Year (CY) 2011 Privately Owned Vehicle Mileage Reimbursement Rates.

SUMMARY: The General Services Administration’s (GSA) annual privately owned vehicle (POV) mileage reimbursement rate reviews have resulted in new CY 2011 rates for the use of privately owned automobiles (POA), POAs when Government owned automobiles (GQA) are authorized, and motorcycles for official purposes. No change resulted for the use of privately owned airplanes. FTR Bulletin 11–03 establishes these new CY 2011 mileage reimbursement rates, pursuant to the process discussed below. This notice of subject bulletin is the only notification of revisions to the POV rates to agencies other than the changes posted on the GSA website. GSA determines these rates by reviewing the annual standard automobile study conducted by the Internal Revenue Service, as well as conducting motorcycle and aircraft studies, and/or by applying consumer price index data.

DATES: This notice is effective upon the date of publication and applies to travel performed on or after January 1, 2011, through December 31, 2011.

FOR FURTHER INFORMATION CONTACT: For clarification of content, please contact Mr. Cy Greenidge, Office of Governmentwide Policy, Office of Travel, Transportation, and Asset Management, at (202) 219–2349, or by e-mail at travelpolicy@gsa.gov. Please cite Notice of FTR Bulletin 11–03.

SUPPLEMENTAL INFORMATION:

Change in Standard Procedure

GSA posts the POV mileage reimbursement rates, formerly published in 41 CFR Chapter 301, solely...
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Board of Scientific Counselors, National Center for Health Statistics (BSC, NCHS)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC), NCHS announces the following meeting of the aforementioned committee:

**Times and Dates:** 11 a.m.–5:30 p.m., January 27, 2011. 8:30 a.m.–2 p.m., January 28, 2011.

**Place:** NCHS Headquarters, 3311 Toledo Road, Hyattsville, Maryland 20782.

**Status:** This meeting is open to the public; however, visitors must be processed in accordance with established federal policies and procedures. Pre-approval is required for foreign nationals or non-US citizens. Please contact Athella Harris, 301–458–4261, adw1@cdc.gov or Virginia Cain, vcain@cdc.gov at least 10 days in advance for requirements. All visitors are required to present a valid form of picture identification issued by a State, Federal or international government. The meeting room accommodates approximately 100 people.

**Purpose:** This committee provides advice and makes recommendations to the Secretary, HHS; the Director, CDC; and the Director, NCHS, regarding the scientific and technical program goals and objectives, strategies, and priorities of NCHS.

**Matters to Be Discussed:** The agenda will include welcome remarks by the Director, NCHS; an update on the Health Indicators Warehouse; an update on program reviews; and an open session for comments from the public.

Requests to make oral presentations should be submitted in writing to the contact person listed below. All requests must contain the name, address, telephone number, and organizational affiliation of the presenter. Written comments should not exceed five single-spaced typed pages in length and must be received by January 11, 2011.

The agenda is subject to change as priorities dictate.

**Contact Person for More Information:** Virginia S. Cain, PhD, Director of Extramural Research, NCHS/CDC, 3311 E. Toledo Road, Room 7211, Hyattsville, Maryland 20782, telephone (301) 458–4500, fax (301) 458–4020.

The Director, Management Analysis and Services Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention, and the Agency for Toxic Substances and Disease Registry.


**BILLING CODE 4163–18–P**

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2010–N–0640]

Agency Information Collection Activities; Proposed Collection; Comment Request; Data to Support Food and Nutrition Product Communications, as Used by the Food and Drug Administration

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal Agencies are required to publish notice in the Federal Register concerning each proposed collection of information and to allow 60 days for public comment in response to the notice. This notice solicits comments on a generic clearance to collect information to support communications about nutrition and food products regulated by FDA. This data collection will gauge, informally, public opinion on a variety of subjects related to consumer, patient, or health care professional perceptions and use of nutrition and food products and related materials, including but not limited to, food advertising, food and nutrition labeling, emerging risk communications, online sales of food products, and consumer and professional education.

**DATES:** Submit either electronic or written comments on the collection of information by February 28, 2011.

**ADDRESSES:** Submit electronic comments on the collection of information to http://www.regulations.gov. Submit written comments on the collection of information to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT:

**SUPPLEMENTARY INFORMATION:** Under the PRA (44 U.S.C. 3501–3520), Federal Agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. “Collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes Agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal Agencies to provide a 60-day notice in the Federal Register concerning each proposed collection of information before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on these topics: (1) Whether the proposed collection of information is necessary for the proper performance of FDA’s functions, including whether the information will have practical utility; (2) the accuracy of FDA’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.