Part III of the proposed order prohibits Respondent from making misrepresentations in advertising for any morning food or snack food about the existence, contents, validity, results, conclusions, or interpretations of any test, study or research.

Part IV of the proposed order states that the order does not prohibit Respondent from making representations for any product that are specifically permitted in labeling for that product by regulations issues by the FDA under the Nutrition Labeling and Education Act of 1990.

Parts V through VIII of the proposed order require Respondent to keep copies of relevant advertisements and materials substantiating claims made in the advertisements; to provide copies of the order to certain of their personnel; to notify the Commission of changes in corporate structure that might affect compliance obligations under the order; and to file compliance reports with the Commission. Part IX provides that the order will terminate after twenty (20) years, with certain exceptions.

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

By direction of the Commission.

Donald S. Clark

Secretary.

[FR Doc. E9–9484 Filed 4–24–09: 8:45 am] BILLING CODE 6750–01–S

GENERAL SERVICES ADMINISTRATION

[OMB Control No. 3090-0044]

Public Buildings Service; Information Collection; GSA Form 3453, Application/Permit for Use of Space in Public Buildings and Grounds

AGENCY: Public Buildings Service, GSA. **ACTION:** Notice of request for comments regarding a renewal to an existing OMB clearance.

SUMMARY: Under the provisions of the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35), the General Services Administration will be submitting to the Office of Management and Budget (OMB) a request to review and approve a renewal of a currently approved information collection requirement regarding GSA Form 3453, Application/Permit for Use of Space in Public Buildings and Grounds. The clearance currently expires on April 30, 2009.

Public comments are particularly invited on: Whether this collection of information is necessary 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.

DATES: Submit comments on or before: June 26, 2009.

FOR FURTHER INFORMATION CONTACT:

Frank Giblin, Public Buildings Service, at telephone (202) 501–1856, or via e-mail to frank.giblin@gsa.gov.

ADDRESSES: Submit comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to the Regulatory Secretariat (VPR), General Services Administration, Room 4041, 1800 F Street, NW., Washington, DC 20405. Please cite OMB Control No. 3090–0044, GSA Form 3453, Application/Permit for Use of Space in Public Buildings and Grounds, in all correspondence.

SUPPLEMENTARY INFORMATION:

A. Purpose

The general public uses GSA Form 3453, Application/Permit for Use of Space in Public Buildings and Grounds, to request the use of public space in Federal buildings and on Federal grounds for cultural, educational, or recreational activities. A copy, sample, or description of any material or item proposed for distribution or display must also accompany this request.

B. Annual Reporting Burden

Respondents: 8,000.

Responses per Respondent: 1.

Hours per Response: 0.05.

Total Burden Hours: 400.

Obtaining Copies of Proposals: Requesters may obtain a copy of the information collection documents from the General Services Administration, Regulatory Secretariat (VPR), 1800 F Street, NW., Room 4041, Washington, DC 20405, telephone (202) 501–4755. Please cite OMB Control No. 3090–0044, GSA Form 3453, Application/Permit for Use of Space in Public Buildings and Grounds, in all correspondence.

Dated: April 21, 2009.

Philip E. Klokis,

Acting Chief Information Officer.
[FR Doc. E9–9490 Filed 4–24–09; 8:45 am]
BILLING CODE 6820–YT–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. FDA-2009-D-0179]

Draft Guidance for Industry and Food and Drug Administration Staff: Technical Considerations for Pen, Jet, and Related Injectors Intended for Use With Drugs and Biological Products; Availability

AGENCY: Food and Drug Administration,

HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a draft guidance document entitled "Technical Considerations for Pen, Jet, and Related Injectors Intended for Use with Drugs and Biological Products." The draft guidance document provides technical and scientific information for sponsors to consider in developing information to support a marketing application for a pen, jet, or related injector device intended for use with drugs or biological products. The marketing application would typically be a premarket notification submission (510(k)) or a premarket approval (PMA) application for the injector alone. For a combination product that includes the injector, the marketing application would typically be a new drug application (NDA) or a biological licensing application (BLA).

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 this draft guidance before it begins work on the final version of the guidance, submit written or electronic comments on the draft guidance by July 27, 2009.

ADDRESSES: Submit written requests for single copies of the draft guidance to the Office of Combination Products, 15800 Crabbs Branch Way, Rockville, MD 20855. Send one self-addressed adhesive label to assist the office in processing your requests. The draft guidance may also be obtained by mail by calling the Office of Combination Products at 301–427–1934 or by e-mail to combination@fda.gov. See the SUPPLEMENTARY INFORMATION section for

SUPPLEMENTARY INFORMATION section for electronic access to the draft guidance document.

Submit written comments on the draft guidance to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.regulations.gov.