

form, as well as in a Portable Document Format (PDF) viewable and downloadable electronic image on OGE's Internet Web site (Uniform Resource Locator address: <http://www.usoge.gov>, under the Ethics Resource Library section) and in future editions of The Ethics CD-ROM. The Office of Government Ethics also will permit departments and agencies to photocopy or have copies printed of the form as well as to develop or utilize, on their own, electronic versions of the form, provided that they precisely duplicate the paper original to the extent possible. As noted above, agencies can also develop their own access forms, provided all the information required by the Ethics Act and OGE regulations is placed on the form, along with the appropriate Privacy Act and paperwork notices with any attendant clearances being obtained therefor.

Public comment is invited on each aspect of the proposed slightly modified OGE Form 201 as set forth in this notice, including specifically views on the need for and practical utility of this proposed modified collection of information, the accuracy of OGE's burden estimate, the enhancement of quality, utility and clarity of the information collected, and the minimization of burden (including the use of information technology).

The Office of Government Ethics, in consultation with OMB, will consider all comments received, which will become a matter of public record.

Approved: June 14, 1999.

Stephen D. Potts,

Director, Office of Government Ethics.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 99N-0791]

Agency Information Collection Activities; Announcement of OMB Approval; Survey of Medical Device Manufacturers for Year 2000 Compliance of Manufacturing Systems

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a collection of information entitled "Survey of Medical Device Manufacturers for Year 2000 Compliance of Manufacturing Systems"

has been approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995.

FOR FURTHER INFORMATION CONTACT: Peggy Schlosburg, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-1223.

SUPPLEMENTARY INFORMATION: In the **Federal Register** of May 10, 1999 (64 FR 25045), the agency announced that the proposed information collection had been submitted to OMB for review and clearance under 44 U.S.C. 3507. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. OMB has now approved the information collection and has assigned OMB control number 0910-0405. The approval expires on November 30, 1999. A copy of the supporting statement for this information collection is available on the Internet at "<http://www.fda.gov/ohrms/dockets>".

Dated: June 11, 1999.

William K. Hubbard,

Acting Deputy Commissioner for Policy.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 99N-1174]

Dietary Supplements; Center for Food Safety and Applied Nutrition Strategy; Public Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing a second public meeting to solicit comments that will assist the Center for Food Safety and Applied Nutrition (CFSAN) to develop an overall strategy for achieving effective regulation of dietary supplements under the Dietary Supplement Health and Education Act (DSHEA). This meeting is intended to give the public an opportunity to comment on the development of the strategy.

DATES: The public meeting will be held on July 20, 1999, from 9 a.m. to 5 p.m. Submit written comments by August 20, 1999.

ADDRESSES: The public meeting will be held at the Oakland Federal Bldg., third

fl. auditorium, north tower, 1301 Clay St., Oakland, CA.

FOR FURTHER INFORMATION CONTACT: Janet B. McDonald, Office of Regulatory Affairs (HFR-PA-145), Food and Drug Administration, 1431 Harbor Bay Pkwy., Alameda, CA 94502-7070, 510-337-6845, FAX 510-337-6708, "e-mail jmcdonal@ora.fda.gov."

SUPPLEMENTARY INFORMATION:

I. Introduction

This public meeting is the second of two meetings to seek stakeholder comments on the development of an overall strategy for achieving effective regulation of dietary supplements under the Federal Food, Drug, and Cosmetic Act, as amended by DSHEA. The first meeting was held on June 8, 1999, in Washington, DC. These two meetings build upon themes that emerged from a broader stakeholder meeting sponsored by CFSAN in June 1998. That meeting addressed the nonfood safety initiative programs that are managed by CFSAN and identified some basic themes including: (1) The need to maintain a credible FDA program, including compliance, enforcement, and consumer outreach activities that will help ensure consumer confidence in FDA regulated products; (2) the need to maintain a solid, science based program staffed with highly qualified scientists; and (3) the recognition that FDA's assistance to consumers and the regulated industry is important.

II. Registration and Requests for Oral Presentations

If you would like to attend the public meeting, you must register with the contact person (address above) by July 9, 1999, by providing your: Name, title, business affiliation, address, telephone, and fax number. To expedite processing, registration information may also be faxed to 510-337-6708. If you need special accommodations due to disability, please inform the contact person when you register.

If you wish to make an oral presentation during the meeting, you must inform the contact person of that desire when you register to attend and submit: (1) A brief written statement of the general nature of the evidence or arguments that you wish to present, (2) the names and addresses of the persons who will give the presentation, and (3) the approximate length of time that you are requesting for your presentation. Depending on the number of people who register to make presentations, we may have to limit the time allotted for each presentation.