

CPG entitled "Distributor Medical Device Reporting." Two copies of any comments are to be submitted, except that individuals may submit one copy. Comments are to be identified with the docket number found in brackets in the heading of this document. The agency will review all comments, but in issuing a final CPG, need not specifically address every comment. The agency will make changes to the CPG in response to comments, as appropriate. A copy of the draft CPG and received comments may be seen in the office above between 9 a.m. and 4 p.m., Monday through Friday.

III. Electronic Access

Copies of the draft CPG may also be downloaded to a personal computer with access to the World Wide Web (www). The Office of Regulatory Affairs (ORA) and CDRH Home Pages include the draft CPG and may be accessed at "http://www.fda.gov/ora" or "http://www.fda.gov/cdrh" respectively. The draft CPG will be available on the Compliance References or Compliance Information pages for ORA and CDRH respectively.

Dated: August 15, 1997.

Gary Dykstra,

Acting Associate Commissioner for Regulatory Affairs.

[FR Doc. 97-22702 Filed 8-27-97; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

Name of Committee: Food Advisory Committee.

General Function of the Committee: To provide advice and recommendations to the agency on FDA regulatory issues.

Date and Time: The meeting will be held on September 25 and 26, 1997, 8:30 a.m. to 5 p.m.

Location: Holiday Inn—Eisenhower Metro Center, Eisenhower Station Ballroom, 2460 Eisenhower Ave., Alexandria, VA.

Contact Person: Lynn A. Larsen, Center for Food Safety and Applied

Nutrition (HFS-5), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202-205-4727, or Catherine M. DeRoever, Advisory Committee Staff (HFS-22), 202-205-4251, FAX 202-205-4970, or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area), code 10564. Please call the Information Line for up-to-date information on this meeting.

Agenda: The committee will be conducting an informational meeting during which it will be receiving updates on past issues that were referred to the committee and on other activities related to food safety. There will also be briefings by the current working groups formed to discuss the Final Report from the Keystone National Policy Dialogue on Food, Nutrition, and Health, as well as simultaneous working group sessions. Two working groups are expected to have work products for committee discussion.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person by September 17, 1997. Oral presentations from the public will be scheduled between approximately 4 p.m. and 5 p.m. on September 25, 1997. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before September 17, 1997, and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: August 21, 1997.

Michael A. Friedman,

Deputy Commissioner for Operations.

[FR Doc. 97-22854 Filed 8-27-97; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Mammography Workshop

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

The Food and Drug Administration (FDA) is announcing the following public workshop: Mammography

Workshop. The topics to be discussed are: Update on the Mammography Quality Standards Act (MQSA), State regulations on mammography, the medical physicist's responsibilities, FDA's MQSA compliance, the radiographic processor, and preparation for the MQSA inspection.

Date and Time: The public workshop will be held on Tuesday, September 23, 1997, 8:30 a.m. to 5 p.m.; registration, 8 a.m. to 8:30 a.m. Registration will close on September 16, 1997.

Location: The public workshop will be held at the Medical Forum Bldg., 950 22d St. North, Birmingham, AL 35203, 205-458-8800.

Contact: Ralph T. Trout, Food and Drug Administration (HFR-SE19), 60 Eighth St. NE., Atlanta, GA 30309, 404-347-4001, ext. 5248, FAX 404-347-4349.

Registration: Send registration information (including name, title, firm name, address, telephone, and fax number) to the contact person by Tuesday, September 16, 1997. Space is limited, therefore interested parties are encouraged to register early.

If you need special accommodations due to a disability, please contact Ralph T. Trout at least 7 days in advance.

SUPPLEMENTARY INFORMATION: This workshop is being sponsored by FDA's Southeast Region and the radiological health programs of the States of the Southeast Region. These States are Alabama, Florida, Georgia, Louisiana, Mississippi, North Carolina, South Carolina, and Tennessee, and the Commonwealth of Puerto Rico and the Virgin Islands. The purpose of this workshop is to provide mammography facilities with an update on MQSA and technical training in the area of mammography.

Dated: August 22, 1997.

William B. Schultz,

Deputy Commissioner for Policy.

[FR Doc. 97-22980 Filed 8-25-97; 4:44 pm]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Medicated Feed Good Manufacturing Practices (GMP's) Training Workshop

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA), Office of Regulatory Affairs (ORA), Pacific Region is announcing a training workshop to