C. Legend

Unless disclosure is required by FOIA as amended (5 U.S.C. 552), as determined by the freedom of information officials of DHHS or by a court, data contained in the portions of an application that have been specifically identified by page number, paragraph, etc., by the applicant as containing restricted and/or proprietary information shall not be used or disclosed except for evaluation purposes.

Margaret M. Dotzel,
Acting Associate Commissioner for Policy.
[FR Doc. 00–1539 Filed 1–21–00; 8:45 am]
BILLING CODE 4160–01–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Allergenic Products 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: Allergenic Products Advisory Committee.

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

Date and Time: The meeting will be held on February 10, 2000, 8:30 a.m. to 3:30 p.m.

Location: Holiday Inn, Versailles Ballrooms I and II, 8120 Wisconsin Ave., Bethesda, MD.

Contact Person: William Freas or Pearline K. Muckelvene, Center for Biologics Evaluation and Research (HFM–71), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852, 301–827–0314, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572) in the Washington, DC area, code 12388. Please call the Information Line for up-to-date information on this meeting.

Agenda: The committee will receive an update on the organizational changes of the Laboratory of Immunobiotechnology, its regulatory activities (including reference replacements and lot release statistics) and its research activities. The committee will hear presentations and discuss the following regulatory issues: (1) Potency limits for standardized allergen vaccines, (2) selection of allergen extracts for standardization, and (3) a proposed algorithm for the standardization of new allergens.

Procedure: On February 10, 2000, from 8:30 a.m. to 3:30 p.m., the meeting is open to the public. 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 January 26, 2000. Oral presentations from the public will be scheduled between approximately 11 a.m. and 12 m. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before February 3, 2000, 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).

Linda A. Suydam,
Senior Associate Commissioner.
[FR Doc. 00–1543 Filed 1–21–00; 8:45 am]
BILLING CODE 4160–01–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 98D–0132]

FDA Modernization Act of 1997; Guidance on Medical Device Tracking; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of the revised guidance document entitled “Guidance on Medical Device Tracking.” This guidance document, which replaces the previous guidance issued on February 12, 1999, provides guidelines to manufacturers and distributors concerning their responsibilities for medical device tracking under the Federal Food, Drug, and Cosmetic Act (the act) as amended by the Food and Drug Administration Modernization Act of 1997 (FDAMA).

DATES: Submit written comments at any time.

ADDRESSES: Submit written requests for single copies on a 3.5 diskette of the guidance document entitled “Guidance on Medical Device Tracking” to the Division of Small Manufacturers Assistance (HFZ–220), Center for Devices and Radiological Health, Food and Drug Administration, 1350 Piccadilly Dr., Rockville, MD 20850. Send two self-addressed adhesive labels to assist that office in processing your request, or fax your request to 301–443–8818. Submit written comments on “Guidance on Medical Device Tracking” to the contact person (address below). See the SUPPLEMENTARY INFORMATION section for information on electronic access to the guidance.


SUPPLEMENTARY INFORMATION:

I. Background

Section 211 of FDAMA (Public Law 105–115) amended the tracking provisions of section 519(e) of the act (21 U.S.C. 360i(e)) to authorize FDA, at its discretion, to issue orders that require a manufacturer to track a class II or class III device if: (1) The failure of the device would be reasonably likely to have serious adverse health consequences; (2) the device is intended to be implanted in the body for more than 1 year; or (3) the device is life sustaining or life supporting and used outside a device user facility. The FDA tracking provisions became effective on February 19, 1998.

The revised final guidance replaces the February 1999 guidance and clarifies the devices that must be tracked. Agency experience indicates that industry and other interested parties are confused about the term “replacement heart valves” because there is more than one type. The category of replacement heart valves that must be tracked is limited to mechanical heart valves only and does not include human allograft(tissue) heart valves. The revised guidance document includes this descriptive limitation.

Agency experience also indicates that industry and other interested parties are confused about which infusion pumps are subject to medical device tracking because the types of fluids the pumps are intended to deliver may not be clear from indications for use set out in labeling. The previous guidance stated that infusion pumps, except those designated and labeled for use
exclusively for fluids with low potential risks, such as enteral feeding or anti-infectives, were subject to tracking. The agency has reevaluated the types of infusion pumps subject to tracking and the best way to describe them in the guidance document. The revised guidance explains that tracking is required only for electromechanical infusion pumps that are used outside a user facility. This was the agency’s position in 1993 when tracking was originally implemented (58 FR 43442 at 43449). The phrase “ electromechanical only” will be used to describe the pumps rather than a reference to the classification regulation. FDA believes this will clarify the guidance because the terms used in the classification language for infusion pumps may include types that do not require tracking.

Finally, the agency added abdominal aortic aneurysm stent grafts to the devices that must be tracked. The agency issued tracking orders for these devices on September 28, 1999, which were effective immediately. FDA determined that these devices meet the statutory tracking criteria under section 519(e) of the act because failure of the device would be reasonably likely to have serious adverse health effects. The agency may add or remove devices from the list of tracked devices as a result of its review of premarket applications, recall data, medical device reporting, inspections, petitions, postmarket surveillance, or other information.

II. Significance of Guidance

This guidance document represents the agency’s current thinking on medical device tracking requirements, as amended by FDAMA. It does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the applicable statute, regulations, or both.

The agency has adopted Good Guidance Practices (GGP’s), which set forth the agency’s policies and procedures for the development, issuance, and use of guidance documents (62 FR 8961, February 27, 1997). This guidance document is issued as a Level 2 guidance consistent with GGP’s.

III. Electronic Access

In order to receive “Guidance on Medical Device Tracking” via your fax machine, call the CDRH Facts-On-Demand (FOD) system at 800–899–0381 or 301–827–0111 from a touch-tone telephone. At the first voice prompt press 1 to access DSMA Facts, at second voice prompt press 2, and then enter the document number (169) followed by the pound sign (#). Then follow the remaining voice prompts to complete your request.

Persons interested in obtaining a copy of the guidance may also do so using the Internet. CDRH maintains an entry on the Internet for easy access to information including text, graphics, and files that may be downloaded to a personal computer with access to the Internet. Updated on a regular basis, the CDRH home page includes “Guidance on Medical Device Tracking,” device safety alerts, Federal Register reprints, information on premarket submissions (including lists of approved applications and manufacturers’ addresses), small manufacturers’ assistance, information on video conferencing and electronic submissions, mammography matters, and other device-oriented information. The CDRH home page may be accessed at http://www.fda.gov/cdrh. “Guidance on Medical Device Tracking” will be available at http://www.fda.gov/cdrh/home.html.

IV. Comments

Interested persons may, at any time, submit to the contact person (address above) written comments regarding this guidance. Such comments will be considered when determining whether to amend the current guidance.


Linda S. Kahan,
Deputy Director for Regulations Policy, Center for Devices and Radiological Health.

[FR Doc. 00–1542 Filed 1–21–00; 8:45 am]

BILLING CODE 4160–01–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration
[Docket No. 99D–5297]

Medical Devices; Guidance Document for Premarket Notification Submissions for the Nitric Oxide Delivery Apparatus, Nitric Oxide Analyzer, and Nitrogen Dioxide Analyzer; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a guidance document entitled “Guidance Document for Premarket Notification Submissions for the Nitric Oxide Delivery Apparatus, Nitric Oxide Analyzer, and Nitrogen Dioxide Analyzer.” This guidance will serve as a special control for nitric oxide delivery apparatus; nitric oxide analyzer; and nitrogen dioxide analyzer.

FDA’s Center for Devices and Radiological Health (CDRH) believes that this guidance is necessary to provide reasonable assurance of the safety and effectiveness of these devices. The guidance document includes material specific for the devices, consensus standards for electrical safety, electromagnetic compatibility, software and hardware documentation, and resistance to environmental effects.

DATES: Written comments concerning this guidance document must be received by April 24, 2000.

ADDRESSES: See the SUPPLEMENTARY INFORMATION section for information on electronic access to the guidance document. Submit written requests for single copies of the guidance document entitled “Guidance Document for Premarket Notification Submissions for Nitric Oxide Delivery Apparatus, Nitric Oxide Analyzer, and Nitrogen Dioxide Analyzer” to the Division of Small Manufacturers Assistance (HFF–220), Center for Devices and Radiological Health, Food and Drug Administration, 1350 Piccard Dr., Rockville, MD 20850.

Send two self-addressed adhesive labels to assist that office in processing your request, or fax your request to 301–443–8818. By April 24, 2000, written comments concerning this guidance document must be submitted to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

Comments should be identified with the docket number found in brackets in the heading of this document. After April 24, 2000, comments must be submitted to the contact person identified below.

FOR FURTHER INFORMATION CONTACT:
Michael G. Bazaral, Center for Devices and Radiological Health (HFF–450), Food and Drug Administration, 9200 Corporate Blvd., Rockville, MD 20850, 301–443–8609.

SUPPLEMENTARY INFORMATION:
I. Background

On January 11, 2000, FDA issued an order to Datex–Ohmeda, Inc., under section 513(f)(2) of the Federal Food, Drug, and Cosmetic Act (the act) (21 U.S.C. 360c(f)(2)) classifying the nitric oxide administration apparatus, the nitric oxide gas analyzer, and the nitric dioxide analyzer into class II (special controls). This guidance document is intended to serve as the special control for these devices.

FDA is making this guidance document effective immediately.