

(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>. The guidance document entitled "Amended Procedures for Advisory Panel Meetings" will be available at <http://www.fda.gov/cdrh>.

A text-only version of the CDRH Web site is also available from a computer or VT-100 compatible terminal by dialing 800-222-0185 (terminal settings are 8/1/N). Once the modem answers, press Enter several times and then select menu choice 1: FDA BULLETIN BOARD SERVICE. From there follow instructions for logging in, and at the BBS TOPICS PAGE, arrow down to the FDA home page (do not select the first CDRH entry). Then select Medical Devices and Radiological Health. From there select CENTER FOR DEVICES AND RADIOLOGICAL HEALTH for general information, or arrow down for specific topics.

Dated: March 25, 1998.

D.B. Burlington,

Director, Center for Devices and Radiological Health.

[FR Doc. 98-8301 Filed 3-30-98; 8:45 am]

BILLING CODE 4160-01-F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 98D-0173]

PMA/510(k) Expedited Review Guidance for Industry and Center for Devices and Radiological Health Staff; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a draft guidance entitled "PMA/510(k) Expedited Review Guidance for Industry and the Center for Devices and Radiological Health (CDRH) Staff." FDA believes it is in the interest of the public health to review premarket approval applications (PMA's) and premarket notifications (510(k)'s) for certain medical devices in an expedited manner. The expedited review will generally be considered when a device offers a potential for clinically meaningful benefit as compared to the existing alternatives (preventive,

diagnostic, or therapeutic) or when the new medical device promises to provide a revolutionary advance (not incremental advantage) over currently available alternative modalities.

DATES: Submit written comments concerning this guidance by June 29, 1998. After the close of the comment period, written comments may be submitted at any time to one of the contact persons listed in this document.

ADDRESSES: Written comments concerning this guidance that are submitted within the 90-day comment period must be submitted to the Dockets Management Branch (HFA-305), Food and Drug Administration, 12420 Parklawn Dr., rm. 1-23, Rockville, MD 20857. Comments should be identified with the docket number found in brackets in the heading of this document. Submit written requests for single copies of the guidance on a 3.5" diskette to the Division of Small Manufacturers Assistance (DSMA), Center for Devices and Radiological Health (HFZ-220), 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. See the **SUPPLEMENTARY INFORMATION** section for information on electronic access to the guidance.

FOR FURTHER INFORMATION CONTACT:

On expedited review for PMA's: Kathy M. Poneleit, Center for Devices and Radiological Health (HFZ-402), Food and Drug Administration, 9200 Corporate Blvd., Rockville, MD 20850, 301-594-2186.

For expedited review for 510(k)'s: Heather S. Rosecrans, Center for Devices and Radiological Health (HFZ-402), Food and Drug Administration, 9200 Corporate Blvd., Rockville, MD 20850, 301-594-1190.

SUPPLEMENTARY INFORMATION:

I. Background

The criteria and procedures under which expedited review would apply to PMA's and Premarket Notifications (510(k)'s) for medical devices were previously identified in General Program Memorandum #G94-2, "PMA/510(k) Expedited Review." In order to reflect the criteria in section 515(d)(5) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360e(d)(5)), as modified by section 202 of the Food and Drug Administration Modernization Act of 1997 (FDAMA) entitled "Special Review for Certain Devices," the criteria section of the guidance has been

modified. These modifications include rearranging the first three criteria and revising the fourth to track the new statutory language more closely. All other sections of the guidance remain the same. This document rescinds and replaces General Program Memorandum #G94-2, "PMA/510(k) Expedited Review."

These procedures are based upon the Management Action Plan initiative paper entitled "PMA/510(k) Expedited Review Process." This guidance embodies the procedures flowing from that issue paper and implements the principles in that document as the policy of the Office of Device Evaluation (ODE). This Blue Book Memorandum will be used by ODE reviewers in applying procedures for the review of incoming PMA's and 510(k)'s.

FDA believes it is in the interest of the public health to review PMA's and 510(k)'s for certain medical devices in an expedited manner. Expedited review will generally be considered when a device offers a potential for clinically meaningful benefit as compared to the existing alternatives (preventive, diagnostic, or therapeutic) or when the new medical device promises to provide a revolutionary advance (not incremental advantage) over currently available alternative modalities.

Granting of expedited review status means that the marketing application would receive priority review before other pending PMA's and 510(k)'s, i.e., the application will be placed at the beginning of the appropriate review queue. If multiple applications for the same type of medical device offering comparable advantage over existing approved alternatives have been granted expedited review, the applications will be reviewed with priority according to their respective submission due dates. Once one of the applications is approved, those of the same type still pending will generally lose their expedited review status with regard to review resources but will retain their place in the review queue.

II. Significance of Guidance

This guidance document represents the agency's current thinking on the procedures to be followed for expedited review of PMA's and (510(k)'s). 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 requirements of the applicable statute, regulations, or both.

This guidance document entitled "PMA/510(k) Expedited Review Guidance for Industry and CDRH Staff" is a Level 1 guidance under FDA's Good

Guidance Practice Policy. Public comment prior to implementation of the guidance document is not required because the guidance is needed to implement new statutory requirements enacted by FDAMA.

III. Electronic Access

In order to receive the guidance entitled "PMA/510(k) Expedited Review Guidance for Industry and CDRH Staff" via your fax machine, call the CDRH Facts-On-Demand 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 108, 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 World Wide Web (WWW). CDRH maintains an entry on the WWW for easy access to information including text, graphics, and files that may be downloaded to a personal computer with access to the Web. Updated on a regular basis, the CDRH home page includes the "PMA/510(k) Expedited Review Guidance," 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>. The "PMA/510(k) Expedited Review Guidance" will be available at <http://www.fda.gov/cdrh>.

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IV. Comments

Interested persons may, on or before June 29, 1998, submit to the Dockets Management Branch (address above) written comments regarding this guidance document. 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 guidance document and received comments may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday. After June 29, 1998, written comments may be submitted at any time to one of the contact persons in this document.

Dated: March 25, 1998.

D.B. Burlington,

Director, Center for Devices and Radiological Health.

[FR Doc. 98-8300 Filed 3-30-98; 8:45 am]

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DEPARTMENT OF THE INTERIOR

Fish and Wildlife Service

[PRT-676811]

Notice of Regional Director's Permit Amendment

U.S. Fish and Wildlife Service Endangered Species Permit PRT-676811, issued to the Regional Director—Region 2 is amended by two technical amendments: (1) to extend the expiration date from April 15, 1998, through June 15, 1998, and (2) to conduct scientific research and recovery activities to include "take" for species currently listed in Region 2.

SUMMARY: The expiration date of this permit is being extended to allow for appropriate public comment for the renewal of this permit. The permit is currently being processed for renewal to continue to conduct specific activities with endangered species. This notice is provided pursuant to section 10(a) of the Endangered Species Act of 1973, as amended (16 U.S.C. 1531, *et seq.*).

FOR FURTHER INFORMATION CONTACT: U.S. Fish and Wildlife Service, Ecological Services, Division of Endangered Species/Permits, P.O. Box 1306, Albuquerque, New Mexico 87103 at (505) 248-6649.

Dated: March 25, 1998.

Renne Lohofener,

Assistant Regional Director, Ecological Services, Region 2, Albuquerque, New Mexico.

[FR Doc. 98-8342 Filed 3-30-98; 8:45 am]

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DEPARTMENT OF THE INTERIOR

Bureau of Land Management

(AK-910-0777-51)

Iditarod Advisory Council Meeting

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of Iditarod Advisory Council Meeting.

SUMMARY: The Iditarod Advisory Council will conduct an open meeting Wednesday, May 6, 1998, and Thursday, May 7, 1998, from 9 a.m. until 4 p.m. each day. The purpose of the meeting is to discuss the formation of a non-profit organization to assist in the management of the Iditarod National Historic Trail. The meeting will be held at the Seward Museum, 336 Third Avenue, Seward, Alaska.

Public comments pertaining to management of the Iditarod National Historic Trail will be taken from 1—2 p.m. Wednesday, January 6. Written comments may be submitted at the meeting or mailed to the address below prior to the meeting.

ADDRESSES: Inquiries about the meeting should be sent to External Affairs, Bureau of Land Management, 222 W. 7th Avenue, #13, Anchorage, Alaska 99513-7599.

FOR FURTHER INFORMATION CONTACT: Teresa McPherson, (907) 271-5555.

Dated: March 23, 1998.

Nick Douglas,

Anchorage District Manager.

[FR Doc. 98-8357 Filed 3-30-98; 8:45 am]

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DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[AK-910-0777-74]

Alaska Resource Advisory Council Meeting

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of Alaska Resource Advisory Council Meeting.

SUMMARY: The Alaska Resource Advisory Council will conduct an open meeting Thursday, May 7, 1998, from 9 a.m. until 4:30 p.m. and Friday, May 8, 1998, from 8:30 a.m. until 3 p.m. The council will review BLM land management issues and take public comment on those issues. The meeting will be held at the Alaska Resources Library and Information

Service (ARLIS) building located at 3150 "C" Street, Suite 100, Anchorage, Alaska.