

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. Received comments may be seen in the office above between 9 a.m. and 4 p.m., Monday through Friday. FDA will also place on public display any amendments to, or comments on, the petitioner's environmental assessment without further announcement in the **Federal Register**. If, based on its review, the agency finds that an environmental impact statement is not required and this petition results in a regulation, the notice of availability of the agency's finding of no significant impact and the evidence supporting that finding will be published with the regulation in the **Federal Register** in accordance with 21 CFR 25.40(c).

Dated: June 9, 1995.

Alan M. Rulis,

Acting Director, Office of Premarket Approval, Center for Food Safety and Applied Nutrition.

[FR Doc. 95-14946 Filed 6-19-95; 8:45 am]

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[Docket No. 95D-0114]

Medical Devices; Premarket Notification (510(k)) Procedures/Good Manufacturing Practices; Compliance Program; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of revisions to the standard compliance program for good manufacturing practices (GMP's) (Compliance Program 7382.830). These revisions are intended to refine and refocus FDA's compliance program linking GMP requirements with class I and II premarket notification (510(k)) submissions and other relevant applications. The revisions are being made as part of FDA's reinventing Government initiative and have been incorporated into "Compliance Program 7382.830, Inspection of Medical Device Manufacturers," which supersedes the "Medical Device Reference List" procedures.

ADDRESSES: Submit written requests for single copies of the revisions to the Division of Small Manufacturers Assistance (DSMA) (HFZ-220), Center for Devices and Radiological Health, Food and Drug Administration, 1350 Piccard Dr., Rockville, MD 20850, 301-443-6597 or 1-800-638-2041. Requests

should be identified with the docket number found in brackets in the heading of this document. Send two self-addressed adhesive labels to assist the office in processing your requests. The revisions are available for public examination in the Dockets Management Branch (HFA-305), Food and Drug Administration, rm. 1-23, 12420 Parklawn Dr., Rockville, MD 20857, between 9 a.m. and 4 p.m., Monday through Friday. Copies of a facsimile of the revision are available from CDRH Facts on Demand by requesting the following document numbers and their respective parts: 2702 (compliance program), 3702 (attachment A), 4702 (attachment A-1), 5702 (attachment B through F), 6702 (attachment G), (1-800-899-0281). Copies of the revisions may also be obtained from the Electronic Docket administered by DSMA and available to any one with a video terminal or personal computer (1-800-252-1366).

FOR FURTHER INFORMATION CONTACT: Marje A. Hoban, Center for Devices and Radiological Health (HFZ-306), Food and Drug Administration, 2094 Gaither Rd., MD 20850, 301-594-4695.

SUPPLEMENTARY INFORMATION: In a letter dated April 7, 1995, the Director of the Center for Devices and Radiological Health advised registered medical device companies of changes that FDA was making to its compliance program linking class I and II 510(k) submissions with GMP requirements. These procedural changes became effective May 1, 1995, and have been made part of the standard compliance program for GMP's (Compliance Program 7382.830). FDA is now making the revisions available in conjunction with the April 7, 1995, letter. The general framework of the restructured program includes: (1) Criteria for linking GMP's with marketing clearance for class I or II (510(k)) devices; (2) procedures for notifying firms that clearance of their class I or II (510(k)) submission may be deferred due to serious, related GMP violations; (3) actions FDA will take to reply promptly to a firm's response to an FDA Form 483 and/or GMP Warning Letter; and (4) timeframes for agency action. The changes noted above also apply to PMA supplements that are not subject to the PMA preapproval inspection program, and to export certificates for legally marketed devices.

These changes are being made as part of FDA's reinventing Government initiative. This compliance program supersedes the "Medical Device Reference List" announced in the **Federal Register** of October 26, 1993 (58 FR 57614).

Dated: June 12, 1995.

William B. Schultz,

Deputy Commissioner for Policy.

[FR Doc. 95-14947 Filed 6-19-95; 8:45 am]

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Public Health Service

Centers for Disease Control and Prevention; Statement of Organization, Functions, and Delegations of Authority

Part H, Chapter HC (Centers for Disease Control and Prevention) of the Statement of Organization, Functions, and Delegations of Authority of the Department of Health and Human Services (45 FR 67772-76, dated October 14, 1980, and corrected at 45 FR 69296, October 20, 1980, as amended most recently at 60 FR 17792-17795, dated April 7, 1995) is amended to reflect the merger of the Division of Training and Manpower Development and the Division of Standards Development and Technology Transfer, and the establishment of the Education and Information Division, National Institute for Occupational Safety and Health.

Section HC-B, Organization and Functions, is hereby amended as follows:

After the functional statement for the *Health Effects Laboratory Division (HCC3)*, insert the following:

Education and Information Division (HCC4). (1) Develops from existing scientific and technical information documents containing (a) criteria for recommended occupational safety and health standards, and (b) technical and scientific information relevant to a variety of occupational safety and health issues; (2) develops recommended health and safety standards under the Occupational Safety and Health Act of 1970 and the Federal Mine Safety and Health Act of 1977; (3) prepares and coordinates with the Office of the Director comments and testimony on regulations proposed by the Department of Labor and other departments or agencies that pertain to occupational diseases or injuries; (4) assists the Institute Director in establishing and operating a priority system for research, surveillance, document development, and recommended standards; (5) prepares and at least annually revises the legislatively mandated toxic substance list; (6) establishes and maintains a library and a clearinghouse for receiving, storing, retrieving, and disseminating technical information on occupational safety and health; (7) provides risk evaluations for NIOSH

policy recommendations; (8) develops and tests occupational safety and health training materials, technologies, strategies, and courses; (9) determines occupational safety and health workforce needs on a nationwide basis and develops strategies to meet those needs; (10) develops methods, through research, to evaluate and monitor the effectiveness of training, including program features, faculty, training methods, and outcome measures; (11) conducts the NIOSH summer intern training program for minority students; (12) serves as the NIOSH printing office; (13) provides graphic design and audio-visual standards and support for the Institute; (14) serves as the NIOSH Docket Office; (15) evaluates in coordination with other divisions the economic and societal burden of occupationally induced diseases and injuries; (16) establishes and maintains the NIOSH archives; and (17) coordinates all relevant Division activities with the Office of the Director.

Information Resources Branch (HCC42). (1) Operates the Institute's libraries for occupational safety and health information for use by occupational safety and health professionals; (2) acquires, disseminates, and coordinates scientific and technical information relating to occupational safety and health in support of Division activities and NIOSH research programs; (3) plans, implements, and coordinates dissemination activities for all NIOSH publications (printed and electronic); (4) verifies printing clearance for NIOSH publications within NIOSH/CDC procedures; (5) develops and manages the NIOSH exhibit program for professional meetings and conferees; (6) develops and maintains electronic data systems for the Institute to assess information; and (7) establishes and maintains the NIOSH archives.

Risk Analysis and Document Development Branch (HCC43). (1) From existing scientific literature develops documents containing (a) criteria for recommended occupational safety and health standards, and (b) technical and scientific information relevant to a variety of occupational safety and health issues for the U.S. Department of Labor and other Federal agencies; (2) coordinates testimony in response to the Department of Labor, Environmental Protection Agency, and other Federal and State agencies' rulemaking; (3) incorporates recommended work practices, engineering controls, and available evidence of technological feasibility into documents and testimony; (4) analyzes the economics of occupational safety and health

interventions; (5) maintains the NIOSH Docket Office; (6) coordinates scientific review of NIOSH policy documents and testimony; (7) conducts risk analyses and develops risk profiles; (8) researches and develops new quantitative risk assessment methodologies; (9) assists the Director of NIOSH in establishing a priority system for surveillance, research, document development, recommended standards, and training; (10) identifies information on worker exposures, hazard severity, potential for intervention, and advances in new technology; and (11) coordinates requests for policy and/or scientific review of internationally produced documents.

Training and Educational Systems Branch (HCC44). (1) Develops, through research and evaluation, training resources in industrial hygiene, safety, occupational medicine, nursing, and allied professions; (2) collaborates on cooperative training programs with qualified outside organizations; (3) determines strategies for and advises on occupational safety and health workforce needs on a nationwide basis; (4) defines and evaluates selected workforce certification/accreditation programs; (5) establishes career development guidelines for training of employers and employees in the prevention of injuries and diseases; (6) provides graphic design and audio-visual standards and support for the Institute; (7) consults and advises NIOSH professionals on presentation techniques and selection of media; and (8) consults on workforce development.

Delete in their entirety the titles and functional statements for the *Division of Training and Manpower Development (HCC9)* and the *Division of Standards Development and Technology Transfer (HCCC)*.

Dated: June 9, 1995.

David Satcher,
Director.

[FR Doc. 95-15033 Filed 6-19-95; 8:45 am]

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DEPARTMENT OF THE INTERIOR

Fish and Wildlife Service

Extension of the Public Comment Period—Availability of an Environmental Assessment and Receipt of an Application for an Incidental Take Permit From Aronov Realty and Management Inc., in Baldwin County, AL

AGENCY: Fish and Wildlife Service, Interior.

ACTION: Notice of extension of the public comment period.

SUMMARY: The Fish and Wildlife Service (Service) gives notice that the public comment period on the environmental assessment/habitat conservation plan for the Aronov Realty and Management Incorporated (Applicant) application for an incidental take permit pursuant to Section 10(a) of the Endangered Species Act (Act) is being extended. The Applicant has been assigned permit number PRT-802986. The original 30-day comment period was to end on June 30, 1995 (**Federal Register** 60:28428). In the intervening period, the Applicant has proposed certain additional mitigation and minimization measures to fully address the potential impacts to the Bon Secour National Wildlife Refuge and the endangered Alabama Beach mouse (ABM).

DATES: The public comment period for this proposal, which originally closed on June 30, 1995, is now extended until July 14, 1995.

ADDRESSES: Persons wishing to review the application, Environmental Assessment, or Habitat Conservation Plan may obtain a copy *by writing* the Service's Southeast Regional Office, Atlanta, Georgia. Documents will also be available for public inspection, by appointment, during normal business hours at the Regional Office, or the field office. Written data or comments concerning the application, Environmental Assessment, or Habitat Conservation Plan should be submitted to the Regional Office. Please reference permit under **PRT-802986** in such comments.

Regional Permit Coordinator (TE), U.S. Fish and Wildlife Service, 1875 Century Boulevard, Suite 210, Atlanta, Georgia 30345, (telephone 404/679-7110, fax 404/679-7280).
Field Supervisor, U.S. Fish and Wildlife Service, 6578 Dogwood View Parkway, Suite A, Jackson, Mississippi 39213 (telephone 601/965-4900, fax 601/965-4340).

FOR FURTHER INFORMATION CONTACT: Rick G. Gooch at the Atlanta, Georgia, Regional Office.

SUPPLEMENTARY INFORMATION: The ABM, *Peromyscus polionotus ammobates*, is a subspecies of the common oldfield mouse *Peromyscus polionotus* and is restricted to the dune systems of the Gulf Coast of Alabama. The known current range of ABM extends from Fort Morgan eastward to the western terminus of Alabama Highway 182, including the Perdue Unit on the Bon Secour National Wildlife Refuge (BSNWR). The sand dune systems