Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.fda.gov/dockets/ecomments.
Submit written requests for single copies of the SECG to Amy Green, Center for Food Safety and Applied Nutrition (CFSAN) (see FOR FURTHER INFORMATION CONTACT). Send one self-adhesive address label to assist that office in processing your request. See the SUPPLEMENTARY INFORMATION section for electronic access to the SECG.

FOR FURTHER INFORMATION CONTACT:

Amy Green, Center for Food Safety and Applied Nutrition (HFS–306), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740, 301– 436–2025, FAX 301–436–2651.

SUPPLEMENTARY INFORMATION:

I. Background

In the **Federal Register** of January 19, 2001 (66 FR 6138), FDA issued a final rule to ensure the safe and sanitary processing of fruit and vegetable juices. The regulations in part 120 (21 CFR part 120) mandate the application of HACCP principles to the processing of these foods. HACCP is a preventive system of hazard control. The effective dates of the final rule are staggered and based on the size of the business. For very small businesses (as defined in $\S 120.1(b)(1)$, the effective date is January 20, 2004. For small businesses, the effective date was January 21, 2003, and for all other size businesses the effective date was January 22, 2002.

FDA examined the economic implications of that final rule as required by the Regulatory Flexibility Act (5 U.S.C. 601–602). The agency determined that the final rule would have a significant economic impact on a substantial number of small entities.

In compliance with section 212 of the Small Business Regulatory Enforcement Fairness Act (Public Law 104–121), FDA is making available this SECG stating in plain language the requirements of the juice HAACP regulations.

FDA is issuing this SECG as level 2 guidance consistent with FDA's good guidance practices regulation (21 CFR 10.115(c)(2)). The SECG represents the agency's current thinking on the subject. 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 statutes and regulations.

II. Comments

Interested persons may submit to the Dockets Management Branch (see ADDRESSES) written or electronic

comments on the SECG entitled "Juice HACCP." Submit a single copy of electronic comments to http://www.fda.gov/dockets/ecomments or two paper copies of any mailed comments, except that individuals may submit one paper copy. Comments are to be identified with the docket number found in brackets in the heading of this document. A copy of SECG and received comments may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

III. Electronic Access

The SECG also may be viewed on a personal computer with access to the Internet at http://www.cfsan.fda.gov/guidance.html.

Dated: March 27, 2003.

Jeffrey Shuren,

Assistant Commissioner for Policy.
[FR Doc. 03–8263 Filed 4–3–03; 8:45 am]
BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 00D-1662]

"Guidance for Industry: Source Animal, Product, Preclinical, and Clinical Issues Concerning the Use of Xenotransplantation Products in Humans;" Availability

AGENCY: Food and Drug Administration,

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a document entitled "Guidance for Industry: Source Animal, Product, Preclinical, and Clinical Issues Concerning the Use of Xenotransplantation Products in Humans" dated April 2003. The document provides guidance on the production, testing, and evaluation of products intended for use in xenotransplantation. The guidance announced in this notice finalizes the draft guidance document of the same title dated February 2001.

DATES: Submit written or electronic comments on agency guidances at any time.

ADDRESSES: Submit written or electronic requests for single copies of this guidance to the Office of Communication, Training, and Manufacturers Assistance (HFM–40), Center for Biologics Evaluation and Research (CBER), Food and Drug Administration, 1401 Rockville Pike,

Rockville, MD 20852–1448. Send one self-addressed adhesive label to assist the office in processing your requests. The document may also be obtained by mail by calling the CBER Voice Information System at 1–800–835–4709 or 301–827–1800, or by fax by calling the FAX Information System at 1–888–CBER–FAX or 301–827–3844. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the guidance document.

Submit written comments on the guidance document to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.fda.gov/dockets/ecomments.

FOR FURTHER INFORMATION CONTACT: Valerie A. Butler, Center for Biologics Evaluation and Research (HFM–17), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852–1448, 301–827–6210.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is announcing the availability of a document entitled "Guidance for Industry: Source Animal, Product, Preclinical, and Clinical Issues Concerning the Use of Xenotransplantation Products in Humans" dated April 2003. The document provides guidance on the production, testing, and evaluation of products intended for use in xenotransplantation. The guidance document announced in this notice was revised based on public comments received on the draft guidance, and it finalizes the draft document of the same title dated February 2001 (66 FR 9348, February 7, 2001).

This guidance is being issued consistent with FDA's good guidance practices regulation (21 CFR 10.115). This guidance document represents the agency's current thinking on this topic. 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 requirement of the applicable statutes and regulations.

II. Comments

Interested persons may, at any time, submit written or electronic comments to the Dockets Management Branch (see ADDRESSES) regarding this guidance document. Two copies of any mailed comments are to be submitted, except individuals may submit one copy. Comments should be identified with the docket number found in the brackets in

the heading of this document. A copy of the document and received comments are available for public examination in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

III. Electronic Access

Persons with access to the Internet may obtain the document at either http://www.fda.gov/cber/guidelines.htm or http://www.fda.gov/ohrms/dockets/default.htm.

Dated: March 27, 2003.

Jeffrey Shuren,

Assistant Commissioner for Policy. [FR Doc. 03–8167 Filed 4–3–03; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HOUSING AND URBAN DEVELOPMENT

[Docket No. FR-4809-N-14]

Federal Property Suitable as Facilities To Assist the Homeless

AGENCY: Office of the Assistant Secretary for Community Planning and Development, HUD

ACTION: Notice

SUMMARY: This Notice identifies unutilized, underutilized, excess, and surplus Federal property reviewed by HUD for suitability for possible use to assist the homeless.

DATES: April 4, 2003.

FOR FURTHER INFORMATION CONTACT:

Mark Johnston, Department of Housing and Urban Development, Room 7262, 451 Seventh Street SW., Washington, DC 20410; telephone (202) 708–1234; TTY number for the hearing- and speech-impaired (202) 708–2565, (these telephone numbers are not toll-free), or call the toll-free Title V information line at 1–800–927–7588.

SUPPLEMENTARY INFORMATION: In

accordance with the December 12, 1998 court order in *National Coalition for the Homeless* v. *Veterans Administration*, No. 88–2503–OG (D.D.C.), HUD publishes a Notice, on a weekly basis, identifying unutilized, underutilized, excess and surplus Federal buildings and real property that HUD has reviewed for suitability for use to assist the homeless. Today's Notice is for the purpose of announcing that no additional properties have been determined suitable or unsuitable this week.

Dated: March 27, 2003.

John D. Garrity,

Director, Office of Special Needs Assistance Programs.

[FR Doc. 03–7857 Filed 4–3–03; 8:45 am] BILLING CODE 4210–29–M

DEPARTMENT OF THE INTERIOR

Fish and Wildlife Service

Draft Environmental Assessment and Receipt of an Application for an Incidental Take Permit for the Mayhoffer/Singletree Trail, Boulder County, CO

AGENCY: Fish and Wildlife Service, Interior.

ACTION: Notice of availability and receipt of application.

SUMMARY: Boulder County Parks and Open Space Department (Applicant) has applied to the Fish and Wildlife Service (Service) for an Incidental Take Permit pursuant to section 10(a)(1)(B) of the Endangered Species Act of 1973, as amended (Act). The permit would authorize the incidental take of Preble's meadow jumping mouse (Zapus hudsonius preblei) ("Preble's"), Federally listed as threatened, and loss and modification of its habitat associated with the construction and use of a multiple use trail on the Mayhoffer/Singletree Property, located near the Town of Superior, in unincorporated Boulder County. The permit would be in effect for 10 years from the date of issuance, to allow for construction of the proposed project and all associated mitigation activities.

We announce the receipt of the Applicant's Incidental Take Permit application that includes a combined proposed Habitat Conservation Plan (HCP) and Environmental Assessment (EA) for the Preble's on the Mayhoffer/ Singletree property. The proposed HCP/ EA is available for public comment. It fully describes the proposed project and the measures the Applicant would undertake to minimize and mitigate project impacts to the Preble's.

The Service requests comments on the HCP/EA for the proposed issuance of an Incidental Take Permit. We provide this notice pursuant to section 10(a) of the Act and National Environmental Policy Act regulations (40 CFR 1506.6). All comments on the HCP and permit application will become part of the administrative record and will be available to the public.

DATES: Written comments on the permit application, HCP, and EA should be received on or before June 3, 2003.

ADDRESSES: Comments regarding the permit application and HCP/EA should be addressed to LeRoy Carlson, Field Supervisor, U.S. Fish and Wildlife Service, Colorado Field Office, 755 Parfet Street, Suite 361, Lakewood, Colorado 80215.

FOR FURTHER INFORMATION CONTACT: Ms. Kathleen Linder, Fish and Wildlife Biologist, Colorado Field Office, telephone (303) 275–2370.

SUPPLEMENTARY INFORMATION:

Document Availability

Individuals wishing copies of the HCP/EA and associated documents for review should immediately contact the above office. Documents also will be available for public inspection, by appointment, during normal business hours at the Lakewood, Colorado, Field Office (see ADDRESSES above).

Background

Section 9 of the Act and Federal regulation prohibit the "take" of a species listed as endangered or threatened. Take is defined under the Act, in part, as to kill, harm, or harass a Federally listed species. However, the Service may issue permits to authorize "incidental take" of listed species under limited circumstances. Incidental Take is defined under the Act as take of a listed species that is incidental to, and not the purpose of, the carrying out of an otherwise lawful activity under limited circumstances. Regulations governing permits for threatened species are promulgated in 50 CFR 17.32.

The Applicant plans to develop a multiple-use trail on the Mayhoffer/ Singletree property in the vicinity of Coal Creek near Superior, Boulder County, Colorado, within portions of the property that may constitute habitat for the Preble's. Of the 32 hectares (80 acres) of potential Preble's habitat on the Mayhoffer/Singletree property, the project would impact a total of 0.27 hectare (0.67 acre) of potential Preble's habitat permanently and 0.34 hectare (0.85 acre) temporarily during construction. This reach of the Coal Creek corridor is considered to be viable Preble's habitat by the Service. Preble's have been found near this creek in 1999, approximately 0.8 kilometer (0.5 mile) upstream from the proposed project area, along the Hake Ditch running north of the creek. As discussed below, the Applicant proposes a number of measures to mitigate possible impacts of the proposed action.

Alternatives considered were—no action; alternative trail alignment, which would have taken the trail through a large prairie dog colony and