

application was received and filed in the Center for Drug Evaluation and Research on December 27, 1994, which shall be considered the filing date for purposes of the act.

Interested persons may submit relevant information on the application to the Dockets Management Branch (address above) in two copies (except that individuals may submit single copies) and identified with the docket number found in brackets in the heading of this document. These submissions may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

The agency encourages any person who submits relevant information on the application to do so by March 13, 1995, and to provide an additional copy of the submission directly to the contact person identified above, to facilitate consideration of the information during the 30-day review period.

This notice is issued under the Federal Food, Drug, and Cosmetic Act (sec. 802 (21 U.S.C. 382)) and under authority delegated to the Commissioner of Food and Drugs (21 CFR 5.10) and redelegated to the Center for Drug Evaluation and Research (21 CFR 5.44).

Dated: February 9, 1995.

Edward Miracco,

Acting Deputy Director, Office of Compliance, Center for Drug Evaluation and Research.

[FR Doc. 95-5059 Filed 2-28-95; 8:45 am]

BILLING CODE 4160-01-F

[Docket No. 95N-0049]

Drug Export; Zyluprim (Allopurinol Sodium) for Injection Equivalent to 500 Milligrams Allopurinol Sterile Lyophilized Powder

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that Burroughs Wellcome Co. has filed an application requesting approval for the export of the human drug Zyluprim (allopurinol sodium) for Injection equivalent to 500 milligrams (mg) allopurinol to Canada.

ADDRESSES: Relevant information on this application may be directed to the Dockets Management Branch (HFA-305), Food and Drug Administration, rm. 1-23, 12420 Parklawn Dr., Rockville, MD 20857, and to the contact person identified below. Any future inquiries concerning the export of human drugs under the Drug Export Amendments Act of 1986 should also be directed to the contact person.

FOR FURTHER INFORMATION CONTACT: James E. Hamilton, Center for Drug Evaluation and Research (HFD-310), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-594-2073.

SUPPLEMENTARY INFORMATION: The drug export provisions in section 802 of the Federal Food, Drug, and Cosmetic Act (the act) (21 U.S.C. 382) provide that FDA may approve applications for the export of drugs that are not currently approved in the United States. Section 802(b)(3)(B) of the act sets forth the requirements that must be met in an application for approval. Section 802(b)(3)(C) of the act requires that the agency review the application within 30 days of its filing to determine whether the requirements of section 802(b)(3)(B) have been satisfied. Section 802(b)(3)(A) of the act requires that the agency publish a notice in the **Federal Register** within 10 days of the filing of an application for export to facilitate public participation in its review of the application. To meet this requirement, the agency is providing notice that Burroughs Wellcome Co., 3030 Cornwallis Rd., P.O. Box 12700, Research Triangle Park, NC 27709-2700, has filed an application requesting approval for the export of the human drug Zyluprim (allopurinol sodium) for Injection equivalent to 500 mg allopurinol to Canada. This product is primarily indicated for its prophylactic usage in patients with leukemia, lymphomas, or other malignancies, receiving antineoplastic treatment (radiation or cytotoxic drugs) which might induce increased uric acid levels. The firm does have new drug application approval for Zyluprim (allopurinol) Tablets in two dosage strengths. The application was received and filed in the Center for Drug Evaluation and Research on January 25, 1995, which shall be considered the filing date for purposes of the act.

Interested persons may submit relevant information on the application to the Dockets Management Branch (address above) in two copies (except that individuals may submit single copies) and identified with the docket number found in brackets in the heading of this document. These submissions may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

The agency encourages any person who submits relevant information on the application to do so by March 13, 1995, and to provide an additional copy of the submission directly to the contact person identified above, to facilitate

consideration of the information during the 30-day review period.

This notice is issued under the Federal Food, Drug, and Cosmetic Act (sec. 802 (21 U.S.C. 382)) and under authority delegated to the Commissioner of Food and Drugs (21 CFR 5.10) and redelegated to the Center for Drug Evaluation and Research (21 CFR 5.44).

Dated: February 8, 1995.

Edward Miracco,

Acting Deputy Director, Office of Compliance, Center for Drug Evaluation and Research.

[FR Doc. 95-5062 Filed 2-28-95; 8:45 am]

BILLING CODE 4160-01-F

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[NM-070-5101-G014; NMNM93652]

Notice of intent, and Notice of Scoping Meetings and Comment Period; New Mexico

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of intent, and notice of scoping meetings with a public comment period.

SUMMARY: In accordance with the National Environmental Policy Act, the Bureau of Land Management is directing the preparation of an environmental document for the construction, operation, and maintenance of a pipeline that would be 12 or 16 inches in diameter and approximately 400 miles in length. The proposed project is known as the Mid-American Pipeline Company Four Corners Loop Project. The environmental document is being prepared as an environmental assessment, but will be advanced to the environmental impact statement level if this is indicated by scoping or by a determination of significant impacts in the environmental assessment. Public meetings will be held for the proposed pipeline project with a public comment period.

DATES: Public scoping meetings are planned in Albuquerque, New Mexico at the Albuquerque Convention Center, San Miguel Room, 401 Second Street, NW., on March 15, 1995 at 3:00 p.m. to 9:00 p.m. and in Roswell, New Mexico at the Roswell Inn, Berrendo Room, 1815 North Main at 3:00 p.m. to 9:00 p.m. on March 16, 1995, respectively. The meeting agenda will be to conduct an open-house to receive interested parties from 3:00 p.m. to 5:30 p.m. with a formal presentation starting at 7:00 p.m., followed by a workshop to receive comments, and ending at 9:00 p.m.

Written comments on the proposed project accepted until March 31, 1995.

ADDRESSES: Any comments should be sent to the Bureau of Land Management, Farmington District Office, Attention: Jerry Crockford, 1235 LaPlata Highway, Farmington, NM 87401.

FOR FURTHER INFORMATION CONTACT: Jerry Crockford, (505) 599-6333.

SUPPLEMENTARY INFORMATION: Pursuant to Section 28 of the Mineral Leasing Act of 1920 (30 U.S.C.), as amended by the Act of November 16, 1973 (37 Stat. 567), the Mid-America Pipeline Company has applied for a right-of-way, serial number NMNM93652, for the construction, operation, and maintenance of a pipeline. The proposed project crosses Federal, State, Indian, and private land. The pipeline starts at the Chaco Pump Station approximately 12 miles south of Farmington, New Mexico and ends at the Mid-American Pipeline Company Hobbs Pump Station located in Texas southeast of Hobbs, New Mexico. The proposed project will parallel existing pipelines for most of its length except as dictated by resource conflicts. The pipeline is designed to transport 50,000 barrels of natural gas liquids per day. Maps of the Mid-America Pipeline Company proposed route are available for viewing at the Bureau of Land Management, Albuquerque District Office, 435 Montano NE., Albuquerque, New Mexico; Farmington District Office, 1235 LaPlata Highway, Farmington, New Mexico; Roswell District Office, 1717 West Second, Roswell, New Mexico; Carlsbad Resource Area Office, 620 East Greene, Carlsbad, New Mexico.

Dated: February 23, 1995.

Joel E. Farrell,

Assistant District Manager for Lands and Renewable Resources.

[FR Doc. 95-5046 Filed 2-28-95; 8:45 am]

BILLING CODE 4310-FB-M

Fish and Wildlife Service

Notice of Receipt of Applications for Permit

The following applicants have applied for a permit to conduct certain activities with endangered species. This notice is provided pursuant to Section 10(c) of the Endangered Species Act of 1973, as amended (16 U.S.C. 1531, et seq.):

Applicant: S.O.S. Care, Escondido, CA, PRT-796422

The applicant requests a permit to import and reexport newborn-captive-born tigrina (*Felis tigrina*) from the Conservation Project at the Sao Paulo

Zoo, Sao Paulo, Brazil for medical treatment for the purpose of enhancement of survival of the species.

Applicant: Zoological Society of San Diego, San Diego, CA, PRT-799314

The applicant requests a permit to export DNA samples from captive-born lowland tapir (*Tapirus terrestris*), Turkmanian kulan (*Equus hemionus kulan*), Somali wild ass (*Equus africanus somalicus*) and Przewalski's wild horse (*Equus przewalskii*) for the purpose of scientific research.

Applicant: International Crane Foundation, Baraboo, WI, PRT-799313

The applicant requests a permit to export 8 captive-produced eggs of red-crowned crane (*Grus japonensis*) and 8 captive-produced eggs of white-naped crane (*Grus vipio*) to V.A. Andronov, Khinganski Nature Reserve, Amurskaja Reg., Russia for the purpose of enhancement and propagation of the species through rearing and reintroduction to the wild.

Applicant: International Crane Foundation, Baraboo, WI, PRT-799312

The applicant requests a permit to export up to 10 captive-produced eggs of Siberian crane (*Grus leucogeranus*) to Oka State Nature Reserve, Spasski Region, Rjazan Oblast, Russia for the purpose of enhancement and propagation of the species through rearing and reintroduction to the wild.

Applicant: International Fisheries, Inc., Hialeah, FL, PRT-798217

The applicant requests a permit to import up to 2,000 captive-bred Asian bonytongue (*Scleropages formosus*) from the Rainbow Aquarium in Singapore for the purpose of enhancement of survival of the species.

Applicant: National Zoological Park, Washington, D.C., PRT-700309

The applicant requests a permit to take, import, export, reexport and purchase in interstate and foreign commerce, blood, hair and other tissue samples, and salvaged carcasses from any endangered wildlife exotic to the United States for the purpose of scientific research to enhance the survival of endangered species in the wild. Tissues are to be obtained from animals in the wild and in zoos. Tissues collected from animals in the wild are to be collected opportunistically during immobilization of the animals by local wildlife management officials. Wild animals will be immobilized, but not harmed, for collection of tissues.

Written data or comments should be submitted to the Director, U.S. Fish and

Wildlife Service, Office of Management Authority, 4401 North Fairfax Drive, Room 420(c), Arlington, Virginia 22203 and must be received by the Director within 30 days of the date of this publication.

Documents and other information submitted with these applications are available for review, subject to the requirements of the Privacy Act and Freedom of Information Act, by any party who submits a written request for a copy of such documents to the following office within 30 days of the date of publication of this notice: U.S. Fish and Wildlife Service, Office of Management Authority, 4401 North Fairfax Drive, Room 420(c), Arlington, Virginia 22203. Phone: (703/358-2104); FAX: (703/358-2281).

Dated: February 24, 1995.

Caroline Anderson,

Acting Chief, Branch of Permits, Office of Management Authority.

[FR Doc. 95-5045 Filed 2-28-95; 8:45 am]

BILLING CODE 4310-55-P

National Park Service

Notice of Intent To Repatriate a Cultural Item in the Possession of the Field Museum of Natural History, Chicago, IL

AGENCY: National Park Service, Interior.

ACTION: Notice.

Notice is hereby given under the Native American Graves Protection and Repatriation Act of 1990 of the intent to repatriate an item in the possession of the Field Museum of Natural History, Chicago, IL, under a compromise of repatriation claim.

The item is a wampum belt consisting of purple and white shell beads woven into a 32" long by 5" wide rectangular panel. The beads are placed to form a series of diamond-shaped figures inside oblongs. The belt is bound with buckskin with buckskin fringe attached at the ends. The belt was purchased by the Field Museum from Walter C. Wyman in December 1900 (FM# 68566). Museum records indicate that the belt was originally purchased by Wyman from the grandson of Chief Skenandoa on May 8, 1898, approximately one year after the chief's death.

Authorized representatives of the Oneida Indian Nation of New York and the Oneida Tribe of Indians of Wisconsin have been provided with copies of museum records and photographs of the belt. In a letter dated February 7, 1994, the Oneida Indian