governing section 18 require publication of a notice of receipt of an application for a specific exemption proposing the first food use of a registered chemical. Such notice provides for opportunity for public comment on the application. Accordingly, interested persons may submit written views on this subject to the Field Operations Division at the address above.

The Agency, accordingly, will review and consider all comments received during the comment period in determining whether to issue the emergency exemption requested by the Florida Department of Agriculture and Consumer Services.

Dated: August 15, 1991.

Anne E. Lindsay,

Director, Registration Division, Office of Pesticide Programs.

[FR Doc. 91-20910 Filed 8-29-91; 8:45 am] BILLING CODE 6560-50-F

[ER-FRL-3991-5]

Environmental Impact Statements; Availability

Responsible Agency: Office of Federal Activities, General Information (202) 382-5073 or (202) 382-5075.

Availability of Environmental Impact Statements Filed August 19, 1991 Through August 23, 1991 Pursuant to 40 CFR 1506.9.

EIS No. 910291, Final EIS, FHW, NC, NC-228/Spruce Pine Bypass Construction, US 19E southwest of Spruce Pine to NC-226 northwest of Minpro, Funding, COE Section 404 Permit and TVA Section 26A Permit, North Toe River, Mitchell County, NC. Due: September 30, 1991, Contact:

Nicholas Graf, P.E. (919) 856-4346. EIS No. 910292, Final EIS, FHW, WV, Chelyan Bridge Replacement and Upgrading, Reconstruction, US 60 from Diamond to WV61 Hugheston, COE 404 Permit and Coast Guard Bridge Permit, Chelyan Cabin Creek and Kanawha River, Kanawha County, WV Due: September 30, 1991. Contact: Billy R. Higginbotham (304) 348-3093.

EIS No. 910293, Final EIS, FHW, PR, PR-3 Relocation, between the Municipalities of Fajardo, and Humacao, COE 404 Permit, NPDES Permit and Funding, PR, Due: September 30, 1991, Contact: Juan O. Cruz (809) 766-5600.

EIS No. 910294, Final EIS, FHW, CA, CA-198 Freeway Improvements, Plaza Road to Mooney Boulevard, Funding and Possible Section 404 Permit, City of Visalia, Tulare County, CA, Due:

September 30, 1991, Contact: John R. Schultz (916) 551-1140.

EIS No. 910295, Draft EIS, AFS, AZ, Mt. Lemmon Ski Valley Area, Development and Management, Special Use Permit, Santa Catalina District, Coronado National Forest, Pima County, AZ, Due: October 29, 1991, Contact: James R. Abbott (602) 749-8700.

EIS No. 910296, Final EIS, AFS, ID, Accelerated Engelmann Spruce Harvest and Reforestation in Brush Creek, Hendricks Creek, and Copet Creek Salvage Timber Sales, Implementation, McCall Ranger District, Payette National Forest, Adams and Idaho Counties, ID, Due: September 30, 1991, Contact: Linda Fitch (208) 634-1401.

EIS No. 910297, Final EIS, USA, MD, VA, MA. Cameron Station Comprehensive Base Closure and Realignment of Fort Belvoir, Fort Myer and Fort McNair, Implementation, Fairfax and Arlington Counties, VA and Washington, DC, Due: September 30, 1991, Contact: Keith Harris (301) 962-4999.

Amended Notices

EIS No. 910134, Draft EIS, AFS, MT, East Boulder Mine Project, Platinum and Palladium Mining, Construction and Operation, Plan of Operations Approval and COE Section 404 Permit, Gallatin National Forest, Sweet Grass County, MT, Due: October 15, 1991, Contact: Leonard L. Lucero (406) 587-6701. Published FR 05-10-91-Reveiw period reopened and extended.

EIS No. 910273, Draft Supplement, AFS, MT, East Boulder Mine Project, Platinum and Palladium Mining. Construction and Operation. Additional Alternative, Plan of Operations Approval and COE Section 404 Permit, Gallatin National Forest, Sweet Grass County, MT, Due: October 15, 1991, Contact: Sherm Sollid (406) 587-6701. Published FR 08-23-91-Review period reestablished.

Dated: August 27, 1991.

William D. Dickerson,

Deputy Director, Office of Federal Activities. [FR Doc. 91-20913 Filed 8-29-91; 8:45 am] BILLING CODE 6560-50-M

FEDERAL MARITIME COMMISSION

[Docket No. 88-27]

Ariel Maritime Group, Inc., et al. v. New York Shipping Association, Inc., et al.; Order To Show Cause Why Certain Claims Should Not be Dismissed With Prejudice

August 23, 1991.

Notice is given that, at the request of two complainants in this proceeding. AMG Services, Inc., formerly Ariel Maritime Group, Inc., and Ariel Maritime (USA) Inc. (the two Ariel complainants), a procedure has been established to determine whether the two Ariel complainants are the real parties in interest to assert claims arising out of implementation of the socalled Rules on Containers and whether certain other named complainants purportedly associated with the two Ariel complainants and their alleged claims should not be dismissed with prejudice.

Notice of the filing of the complaint in this proceeding was published in the Federal Register, 54 FR 185, on January 4, 1989. In addition to the two Ariel complainants and to seven other complainants who are not associated with the two Ariel complainants, the following ten companies were identified as trade names of complainant Ariel Maritime (USA) Inc.: TransAfrica Line, Oasis Express Line, Javelin Line, Coast Container Line, Interlink Lines, Buccaneer Line, Union Exportadora Lines, Canbel Line, Cedar Star Line, and

Liberty Lines.

The two Ariel complainants have asserted that they are the real parties in interest to assert any claims relating to the operations of the Ariel group of companies, including the ten companies named above, and that these ten companies and their alleged claims should be dismissed with prejudice. However, because these ten companies may not in fact have been represented, before any adverse action is taken against them, the Ariel complainants have asked that suitable notice be provided the ten companies by publication in the Federal Register and by mail where feasible.

Accordingly, the ten complainant companies named above are notified that they are ordered to show cause in writing why they and their alleged claims should not be dismissed with prejudice from the proceeding. Responses to this order shall be mailed so as to reach the undersigned on or before September 30, 1991. The responses shall be addressed to: Norman D. Kline, Administrative Law

Judge, Office of Administrative Law Judges, Federal Maritime Commission, Washington, DC 20573.

Failure to comply with this order will result in a dismissal of the complainant and its alleged claims with prejudice.

Norman D. Kline.

Administrative Law Judge.

[FR Doc. 91-20777 Filed 8-29-91; 8:45 am]

BILLING CODE 6730-01-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

Agency Forms Submitted to the Office of Management and Budget for Clearance

On Fridays, the Department of Health and Human Services, Office of the Secretary publishes a list of information collections it has submitted to the Office of Management and Budget (OMB) for clearance in compliance with the Paperwork Reduction Act (44 U.S.C. chapter 35). The following are those information collections recently submitted to OMB.

1. Social Security Client Satisfaction Survey-0990-0171-revision-This survey of Social Security beneficiaries will obtain information on client satisfaction with Social Security services as part of the Inspector General's on-going monitoring of SSA. The information will be used to identify areas where improvements in service delivery are necessary. In particular, some program indicators used in implementing the Chief Financial Officer Act are data elements from this survey. Respondents: Individuals: Annual Number of Respondents: 975; Frequency of Response: One time; Average Burden per Response: 20 minutes; Estimated Total Annual Burden: 325 hours.

OMB Desk Officer: Allison Eydt.
Copies of the information collection
packages listed above can be obtained
by calling the OS Reports Clearance
Officer on (202) 619–0511. Written
comments and recommendation for the
proposed information collection should
be sent directly to the OMB desk officer
designated above at the following
address: OMB Reports Management
Branch, New Executive Office Building,
room 3208, Washington, DC 20503.

Dated: August 20, 1991.

James F. Trickett,

Deputy Assistant Secretary for Management and Acquisition.

[FR Doc. 91-20447 Filed 8-29-91; 8:45 am]
BILLING CODE 4150-04-M

Administration for Children and Families

Forms Submitted to the Office of Management and Budget for Clearance

The Administration for Children and Families will publish on Fridays information collection packages submitted to the Office of Management and Budget (OMB) for clearance, in compliance with the Paperwork Reduction Act (44 U.S.C. chapter 35). Following is the package submitted to OMB since the last publication. (For a copy of a package, call the FSA, Report Clearance Officer 202–401–5604.)

Tribal JOBS Program Quarterly
Report—Form ACF-114-New—The
information collected on Form ACF-114
will provide the database to properly
monitor, analyze, and assess JOBS
program administrated by Tribal
grantees nationwide. Also, Tribal JOBS
program information will be used to
address Congressional inquiries.
Respondents: State or local
governments/Tribal grantees; Number
of Respondents: 76; Frequency of
Response: Quarterly; Average Burden
per Response: 8 hours; Estimated
Annual Burden: 2,432 hours.

OMB Desk Clearance Officer: Laura Oliven.

Written comments and recommendations for the proposed information collection should be sent directly to the appropriate OMB Desk Officers designated above at the following address: OMB Reports Management Branch, New Executive Office Building, Room 3201, 725 17th Street, NW., Washington, DC 20503.

Dated: August 13, 1991.

Naomi B. Marr,

Associate Administrator, Office of Management & Information Systems. [FR Doc. 91–20382 Filed 8–29–91; 8:45 am] BILLING CODE 4150–04–M

U.S. Advisory Board on Child Abuse and Neglect; Meeting

AGENCY HOLDING THE MEETING: Administration for Children and Families, HHS.

TIMES AND DATES: 2 p.m. September 11, 1991-6 p.m., September 15, 1991.

PLACE

- 2 p.m.-6 p.m., September 11, 1991, Hotel Radisson, 1550 Court Place, Denver, Colorado.
- 8 a.m.—8 p.m., September 12, 1991, Ganow Tower, University of Colorado, Boulder, Colorado.

8:30 a.m. September 13, 1991–1 p.m. September 14, 1991, Hotel Radisson, 1550 Court Place, Denver, Colorado. 3:30 p.m.–6 p.m. September 15, 1991, Colorado Convention Center, Denver,

STATUS: The meeting is closed to public observation from 2 p.m. on September 11, 1991 until 8 p.m. on September 12, 1991, and open to public observation at all other times.

MATTERS TO BE CONSIDERED: During portions of this meeting the Advisory Board will review the first two years of Board operations and plan the next two years of Board operations. Therefore, those portions of the meeting will be closed in order to protect the free exchange of internal views among Board members and to avoid undue interference with the operation of the Board.

During the open portions of this meeting the Advisory Board will: Be briefed on plans for the release of its 1991 report; review the workplan for the development of its 1993 report on a new national child protection strategy; decide on the subject of its 1992 report; discuss plans for a topical report on the Federal child protection research effort; review National Center developments since the May, 1991 meeting and Inter-Agency Task Force developments as well as receive an update on the DHHS initiative on Child Abuse and Neglect; discuss with Congressional staff the implications of the 1991 report for current and future reauthorizations of the Child Abuse Prevention and Treatment Act; elect a Chairperson and Vice-Chairperson to serve until the end of the 1993 Board Program Year; hear a presentation on the role of religious institutions in the new national strategy; discuss with four experts on sexual abuse the implications for the prevention and treatment of sexual abuse of the new national strategy; and participate in a Hearing of the House Select Committee on Children, Youth and Families.

CONTACT PERSON FOR MORE INFORMATION: Eileen H. Lohr, Program Assistant, U.S. Advisory Board on Child Abuse and Neglect, room 2433, Switzer

Abuse and Neglect, room 2433, Switzer Building, Washington, DC 20201, (202) 445–6670.

Dated: August 23, 1991.

Byron D. Metrikin-Gold,

Executive Director, U.S. Advisory Board on Child Abuse and Neglect.

[FR Doc. 91-20839 Filed 8-29-91; 8:45 am]

BILLING CODE 4130-01-M

Food and Drug Administration

[Docket No. 78N-301H]

RIN 0905-AA06

Hydrocortisone; Marketing Status as an External Analgesic Drug Product for Over-the-Counter Human Use; Notice of Enforcement Policy

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an enforcement policy allowing over-thecounter (OTC) marketing of external analgesic drug products containing above 0.5 percent up to 1 percent hydrocortisone or its hydrocortisone acetate equivalent. The OTC marketing of such drug products is being permitted pending establishment under the OTC drug review of a final monograph covering external analgesic drug products. FDA anticipates that external analgesic drug products containing above 0.5 up to 1 percent hydrocortisone or its hydrocortisone acetate equivalent will be determined to be generally recognized as safe and effective and not misbranded.

EFFECTIVE DATE: The enforcement policy is effective August 30, 1991.

ADDRESSES: Written comments to the Dockets Management Branch (HFA–305), Food and Drug Administration, rm. 1–23, 12420 Parklawn Dr., Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: William E. Gilbertson, Center for Drug Evaluation and Research (HFD-210), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–295–8000.

SUPPLEMENTARY INFORMATION: In an amendment of the tentative final monograph for OTC external analgesic drug products, published in the Federal Register of February 27, 1990 (55 FR 6932), FDA proposed conditions under which products containing hydrocortisone or its hydrocortisone acetate equivalent above 0.5 percent up to 1 percent could be marketed OTC. This proposal was based on an evaluation of available data and submitted studies supporting general recognition of the safety and effectiveness of topical hydrocortisone for such use. The studies have been placed in the Dockets Management Branch (address above) and may be seen there by interested persons.

The agency also invited public comment on the proposed change in marketing status that would switch hydrocortisone above 0.5 percent up to 1 percent from its current status as a prescription drug to OTC status. The agency proposal did not allow OTC marketing to begin at the time of publication of the amendment of the tentative final monograph. The agency referred to the Federal Register of June 3, 1983 (48 FR 24925), in which FDA explained the enforcement policy for drugs that were originally on prescription status but which were being proposed for OTC marketing under the OTC drug review. As noted there, 21 CFR 330.13 permits OTC marketing of a drug previously limited to prescription use prior to publication of a final monograph provided that certain conditions are met. To qualify for such treatment, the drug must at a minimum have been considered by an OTC drug advisory review panel and either been recommended for OTC marketing by the panel or subsequently determined by FDA to be suitable for OTC marketing. Hydrocortisone was evaluated by the Advisory Review Panel on OTC Topical Analgesic, Antirheumatic, Otic, Burn, and Sunburn Prevention and Treatment Drug Products (the Panel) in its consideration of the prescription-to-OTC switch of hydrocortisone preparations, but the Panel recommended limiting the concentration for OTC use to 0.25 to 0.5 percent (December 4, 1979, 44 FR 69768 at 69813 through 69824).

In response to the proposal to switch above 0.5 up to 1 percent hydrocortisone from prescription to OTC status, eight drug manufacturers, numerous health professionals, one manufacturer's association, one law firm, and three health professional associations submitted comments. There was one request for an oral hearing from the American Academy of Dermatology (Ref. 1). In a subsequent letter, the Academy withdrew its request for a hearing, stating that its Board of Directors had taken definitive action not to oppose the switch (Ref. 2). Copies of the comments are on public display in the Dockets Management Branch

After carefully reviewing all of the comments received, the agency is issuing a notice of enforcement policy permitting OTC marketing of above 0.5 up to 1 percent hydrocortisone or its hydrocortisone acetate equivalent for topical use prior to publication of the final monograph for OTC external analgesic drug products. This decision is based on extensive supportive safety and effectiveness data, and the following facts: (1) The majority of the comments, both for and against the proposal, were of a testimonial nature without substantive data; (2) many of the comments opposed to the proposal

misunderstood that the proposed indication for OTC use was limited to temporary use to relieve the itching associated with minor skin irritation and rashes due to specific limited causes listed in the proposed monograph and the drug was not to be labeled for the treatment or cure of any skin disease with symptoms of itching; (3) no information not previously known by the agency was provided by the comments; and (4) the objections and concerns regarding the current proposal are the same or similar to those raised when 0.25 to 0.5 percent hydrocortisone was originally proposed for OTC use. Those objections and concerns have been disproven by the available scientific and medical evidence and a history of safe marketing of 0.25 to 0.5 percent hydrocortisone during 9 years of OTC use as well as years of safe useexperience of 1 percent hydrocortisone as a prescription drug.

The agency addressed the safety. effectiveness, and labeling concerns expressed by the comments in the amendment of the external analgesic tentative final monograph proposing OTC status for above 0.5 up to 1 percent hydrocortisone (55 FR 6932). Based on the comments received in response to the proposal, the agency is revising the proposed label warning on the 7-day use limitation from "Do not use this or any other * * "" to read. "* * " stop use of this product and do not begin use * (see below). With this revision, the agency believes there are no unresolved safety or effectiveness issues relating to the OTC use of above 0.5 up to 1 percent hydrocortisone as an antipruritic (antiitch) external analgesic. Accordingly, the agency has determined that it would be inappropriate to continue to bar the interim marketing of such products. The agency's enforcement policy, which is set out in § 330.13, relating to OTC marketing of drug products containing certain ingredients that are under consideration in FDA's review of OTC drugs makes it clear that FDA may by notice in the Federal Register permit interim marketing of products such as hydrocortisone above 0.5 up to 1 percent. The agency advises that any drug product intended for OTC use as an antipruritic external analgesic that contains above 0.5 up to 1 percent hydrocortisone or its hydrocortisone acetate equivalent may be marketed pending issuance of the final monograph, subject to the risk that the agency may, in the final monograph, adopt a different position that could require relabeling, recall, or other regulatory action. Marketing of such products with labeling not in accord

with the labeling proposed in the amended tentative final monograph and this notice also may result in regulatory action against the product, the marketer, or both.

The labeling for OTC hydrocortisone products proposed in the amended tentative final monograph, as revised, is stated below. This labeling is required for marketing any OTC drug product containing above 0.5 up to 1 percent hydrocortisone or its hydrocortisone acetate equivalent. Also, as indicated in the amendment (55 FR 6932 at 6945 and 6946), the same labeling should apply to all OTC concentrations of hydrocortisone, ranging from 0.25 to 1.0 percent. The agency encourages manufacturers to revise the labeling of the currently marketed lower concentrations (0.25 to 0.5 percent) as soon as possible. The following labeling is to be used for all OTC drug products containing hydrocortisone or its

hydrocortisone acetate equivalent: Statement of Identity: "Antipruritic (anti-itch)," "anti-itch," "antipruritic (anti-itch) (insert dosage form, e.g., cream, lotion, ointment, or spray.)" or "anti-itch (insert dosage form, e.g.

cream, lotion, or spray)."

Indications: One of the following should be used: (1) "For the temporary relief of itching associated with minor skin irritations and rashes" [which may be followed by "due to" (select one or more of the following: "eczema," "insect bites," "poison ivy, poison oak, or poison sumac," "soaps," "detergents,"
"cosmetics," "jewelry," "seborrheic
dermatitis," "psoriasis,") and/or ("and
for external" (select one or more of the following: "genital," "feminine," and "anal") "itching"); or (2) "For the temporary relief of itching associated with minor skin irritations, inflammation, and rashes due to" (select one or more of the following: "eczema," "insect bites," "poison ivy, poison oak, or poison sumac," "soaps," "detergents," "cosmetics," "jewelry," "seborrheic dermatitis," "psoriasis,") and/or ("and for external" (select one or more of the following: "gential," "feminine," and "anal") "itching").

In addition, the indications section must include the following statement: "Other uses of this product should be only under the advice and supervision of a" (select one of the following:

"physician" or "doctor").

Warnings: "For external use only. Avoid contact with the eyes. If condition worsens, or if symptoms persist for more than 7 days or clear up and occur again within a few days, stop use of this product and do not begin use of any other hydrocortisone product unless you have consulted a" (select one of the

following: "physician" or "doctor"). "Do not use for the treatment of diaper rash. Consult a" (select one of the following: "physician" or "doctor").

If the product is labeled with the

indications "for external genital itching" or "for external feminine itching," the warnings must include the statement "Do not use if you have a vaginal discharge. Consult a" (select one of the following: "physician" or "doctor"). If the product is labeled with the indication "for external anal itching," the warnings must include the following statements: "Do not exceed the recommended daily dosage unless directed by a doctor. In case of bleeding, consult a doctor promptly. Do not put this product into the rectum by using fingers or any mechanical device or applicator." (The word "physician" may be substituted for the word "doctor" in these statements.)

Directions: Adults and children 2 years of age and older: Apply to affected area not more than 3 to 4 times daily. Children under 2 years of age: Do not use, consult a (select one of the following: physician or doctor).

If the product is labeled with the indication "for external anal itching." the directions must include the following statements: "Adults: When practical, cleanse the affected area" (selected one or both of the following: "with mild soap and warm water and rinse thoroughly" or "by patting or blotting with an appropriate cleansing pad"). "Gently dry by patting or blotting with toilet tissue or a soft cloth before application of this product." (Other appropriate directions in this section may be inserted here.) "Children under 12 years of age: consult a" (select one of the following: "physician" or "doctor"

The final monograph for OTC external analgesic drug products, when published, will establish the final labeling that will be required for all OTC drug products that contain hydrocortisone.

References

(1) Comment No. HER1, Docket No. 78N-301H, Dockets Management Branch. (2) Comment No. WDL1, Docket No. 78N-301H, Dockets Management Branch.

Interested persons may submit written comments to the Dockets Management Branch (address above). Such comments will be considered in determining whether further amendments to or revisions of this policy are warranted. Three copies of all comments shall be submitted, except that individuals may submit single copies. 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 Dockets Management Brabetween 9 a.m. and 4 p.m., Monday through Friday.

Dated: August 14, 1991. Michael R. Taylor, Deputy Commissioner for Policy. [FR Doc. 91-20834 Filed 8-29-91; 8:45 am] BILLING CODE 4160-01-M

[Docket No. 91N-0318]

Human Organ and Tissue Transplantation; Public Hearing

AGENCY: Food and Drug Administration. HHS

ACTION: Notice of public hearing: requests for comments.

SUMMARY: The Food and Drug Administration(FDA), with the concurrence of the Health Resources and Services Administration (HRSA), the National Institutes of Health (NIH), the Centers for Disease Control (CDC), and the Health Care Financing Administration (HCFA), is announcing a public hearing to solicit information and views of interested persons on the need to expand the Federal regulation of organ and tissue transplantation. The information from this public hearing will be used to evaluate whether and how the Federal government should develop a new regulatory program to address aspects of this industry.

DATES: Written notices of participation should be filed by September 30, 1991. The hearing will begin at 9 a.m. on October 16, 1991. The record will remain open for 15 days following the hearing, by which time any additional written material must be submitted.

ADDRESSES: The public hearing will be held at the Jack Masur Auditorium, Warren Grant Magnusun Clinical Center, Bldg. 10, National Institutes of Health, 9000 Rockville Pike, Bethesda, MD 20892. Written notices of participation and any comments are to be sent to the Dockets Management Branch (HFA-305), Food and Drug Administration, rm. 1-23, 12420 Parklawn Dr., Rockville, MD 20857 Transcripts of the hearing, copies of data and information submitted during the hearing, and any written comments will be available for review at the Dockets Management Branch (address above).

FOR FURTHER INFORMATION CONTACT: Nicholas P. Reuter, Office of the Associate Commissioner for Health Affairs (HFY-20), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-443-1382.