

Assistance; 83.543, Individual and Family Grant (IFG) Program; 83.544, Public Assistance Grants; 83.545, Disaster Housing Program; 83.548, Hazard Mitigation Grant Program.)

Joe M. Allbaugh,

Director.

[FR Doc. 02-804 Filed 1-11-02; 8:45 am]

BILLING CODE 6718-02-P

FEDERAL MARITIME COMMISSION

Notice of Agreement(s) Filed

The Commission hereby gives notice of the filing of the following agreement(s) under the Shipping Act of 1984. Interested parties can review or obtain copies of agreements at the Washington, DC offices of the Commission, 800 North Capitol Street, NW., Room 962. Interested parties may submit comments on an agreement to the Secretary, Federal Maritime Commission, Washington, DC 20573, within 10 days of the date this notice appears in the **Federal Register**.

Agreement No.: 011742-001.

Title: P&O Nedlloyd-Farrell/Hapag-Lloyd/Zim Mediterranean Space Charter Agreement.

Parties: Farrell Lines, Inc. Hapag-Lloyd Container Linie GmbH. P&O Nedlloyd Limited. P&O Nedlloyd B.V. Zim Israel Navigation Co., Ltd.

Synopsis: The proposed agreement modification adds authority for the parties to discuss and agree on the phasing-in and phasing-out of vessels for maintenance and to discuss and agree on criteria to measure adherence to any agreed-upon schedule, as well as any remedial action in the event of non-adherence. The modification also clarifies the parties' authority to use common terminals and adds provisions dealing with force majeure situations, notices, and enforceability. The parties request expedited review.

Dated: January 9, 2000.

By Order of the Federal Maritime Commission.

Bryant L. VanBrakle,

Secretary.

[FR Doc. 02-845 Filed 1-11-02; 8:45 am]

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FEDERAL RESERVE SYSTEM

Change in Bank Control Notices; Acquisition of Shares of Bank or Bank Holding Companies

The notificants listed below have applied under the Change in Bank Control Act (12 U.S.C. 1817(j)) and § 225.41 of the Board's Regulation Y (12

CFR 225.41) to acquire a bank or bank holding company. The factors that are considered in acting on the notices are set forth in paragraph 7 of the Act (12 U.S.C. 1817(j)(7)).

The notices are available for immediate inspection at the Federal Reserve Bank indicated. The notices also will be available for inspection at the office of the Board of Governors. Interested persons may express their views in writing to the Reserve Bank indicated for that notice or to the offices of the Board of Governors. Comments must be received not later than January 28, 2002.

A. Federal Reserve Bank of Minneapolis (JoAnne F. Lewellen, Assistant Vice President) 90 Hennepin Avenue, Minneapolis, Minnesota 55480-0291:

1. *Edward T. Christian, trustee of Edward T. Christian Revocable Trust*, Albert Lea, Minnesota; to acquire voting shares of Kiester Investments, Inc., Kiester, Minnesota, and thereby indirectly acquire voting shares of First National Bank of Kiester, Kiester, Minnesota.

Board of Governors of the Federal Reserve System, January 8, 2002.

Robert deV. Frierson,

Deputy Secretary of the Board.

[FR Doc. 02-795 Filed 1-11-02; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 01N-0402]

Agency Information Collection Activities; Submission for OMB Review; Comment Request; Medical Devices; Third-Party Premarket Submission Review and Quality System Inspections Under United States/European Community Mutual Recognition Agreement

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that the proposed collection of information listed below has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995 (the PRA).

DATES: Submit written comments on the collection of information by February 13, 2002.

ADDRESSES: Submit written comments on the collection of information to the Office of Information and Regulatory Affairs, OMB, New Executive Office Bldg., 725 17th St. NW., rm. 10235, Washington, DC 20503, Attn: Stuart Shapiro, Desk Officer for FDA.

FOR FURTHER INFORMATION CONTACT: Peggy Schlosburg, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-1223.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Medical Devices; Third-Party Premarket Submission Review and Quality System Inspections Under United States/European Community Mutual Recognition Agreement (OMB Control No. 0910-0378)—Extension

The third-party program under the United States/European (U.S./EC) Community/Mutual Recognition Agreement (MRA) is intended to implement that part of the U.S./EC MRA that covers the exchange of quality system evaluation reports for all medical devices and premarket evaluation reports for selected low-to-moderate risk devices. Under the MRA, firms may apply to become designated as a U.S. Conformity Assessment Body (CAB). Firms who are designated will be qualified to conduct quality system evaluations for all classes of devices and product type examinations and verifications for selected devices based on EC requirements under the voluntary third-party program authorized by MRA. Firms designated as European Union (EU) CABs could conduct quality system evaluations for all classes of devices and premarket 510(k) evaluations for selected devices based on FDA requirements. Under the voluntary third-party program, reports of these evaluations would be submitted by the EU CABs to FDA. The EU CABs would also be required to maintain copies of their evaluation reports.

FDA requests approval of the following collection of information:

Requests for Designation as U.S. CABs—Under this program, U.S. companies were allowed to apply for designation as a U.S. CAB. Such designation enabled the company to perform third-party reviews of U.S. products for export to the EU and third-party audits of quality systems established by manufacturers of medical devices manufactured for export to the EU. Third-party review of U.S. products