ECONOMIC IMPACT POLICY

This notice is to inform the public that the Export-Import Bank of the United States has received an application for a $19.5 million long-term guarantee to support the export of approximately $30 million worth of mining trucks to the Ukraine. The repayment term of the guarantee is 7 years. The U.S. exports will enable the Ukrainian mining company to establish a maximum production capacity of 28 million metric tons of iron ore per year. Available information indicates that all of the Ukrainian iron ore production will be sold domestically in the Ukraine. Interested parties may submit comments on this transaction by email to economic.impact@exim.gov or by mail to 811 Vermont Avenue NW., Room 947, Washington, DC 20571, within 14 days of the date this notice appears in the Federal Register.

Angela Mariana Freyre,
Senior Vice President and General Counsel.
[FR Doc 2012–8829 Filed 4–11–12; 8:45 am]
BILLING CODE 6690–01–P

FEDERAL MARITIME COMMISSION

[Docket No. 12–02]

Maher Terminal, LLC v. The Port Authority of New York and New Jersey; Notice of Filing of Complaint and Assignment

Notice is given that a complaint has been filed with the Federal Maritime Commission (Commission) by Maher Terminal, LLC, hereinafter “Complainant” against the Port Authority of New York and New Jersey (PANYNJ), hereinafter “Respondent”. Complainant asserts that it is a limited liability company registered in the State of Delaware with corporate offices and facilities located in Elizabeth, New Jersey. Complainant asserts that Respondent, PANYNJ, is a body corporate and politic created by the provision of terminal services to a common carrier.

Complainant asserts that it has sustained injuries and damages, as a result of Respondent’s actions, “including but not limited to higher costs and other undue and unreasonable payments, economic considerations, restrictions on transfers and/or changes in ownership or control interests, lost business, forgone business, and additional obligations not required of other marine terminals and other damages amounting to a sum of millions of dollars.” The full text of the complaint can be found in the Commission’s Electronic Reading Room at www.fmc.gov.

Complainant requests that the Commission require Respondent to: (1) Answer the charges in the subject complaint; (2) cease and desist from the aforementioned violations of the Shipping Act; (3) provide to Complainant the preferences provided to other marine terminal operators; (4) put in force such practices and as the Commission determines to be lawful and reasonable; and (5) pay to Complainant by way of reparations the amount of the actual injury, plus interest, cost and attorneys fees, and any other damages to be determined. Additionally, Complainant requests that the Commission order any such other relief as it determines appropriate.

This proceeding has been assigned to the Office of Administrative Law Judges. Haring in this matter, if any is held, shall commence within the time limitations prescribed in 46 CFR 502.61, and only after consideration has been given by the parties and the presiding officer to the use of alternative forms of dispute resolution. The hearing shall include oral testimony and cross-examination in the discretion of the presiding officer only upon proper showing that there are genuine issues of material fact that cannot be resolved on the basis of documents, affidavits, depositions, or other documents or that the nature of the matter in issue is such that an oral hearing and cross-examination are necessary for the development of an adequate record. Pursuant to the further terms of 46 CFR 502.61, the initial decision of the presiding officer in this proceeding shall be issued by April 8, 2013, and the final decision of the Commission shall be issued by August 6, 2013.

Karen V. Gregory,
Secretary.
[FR Doc 2012–8777 Filed 4–11–12; 8:45 am]
BILLING CODE 6730–01–P

FEDERAL RESERVE SYSTEM

Formations, Acquisitions by, and Mergers of Bank Holding Companies

The companies listed in this notice have applied to the Board for approval, pursuant to the Bank Holding Company Act of 1956 (12 U.S.C. 1841 et seq.) (BHC Act), Regulation Y (12 CFR part 225), and all other applicable statutes and regulations to become a bank holding company and/or to acquire the assets or the ownership of, control of, or the power to vote shares of a bank or bank holding company and all of the banks and nonbanking companies owned by the bank holding company, including the companies listed below.

The applications listed below, as well as other related filings required by the Board, are available for immediate inspection at the Federal Reserve Bank indicated. The applications will also be available for inspection at the offices of the Board of Governors. Interested persons may express their views in writing on the standards enumerated in the BHC Act (12 U.S.C. 1842(c)). If the proposal also involves the acquisition of a nonbanking company, the review also includes whether the acquisition of the nonbanking company complies with the standards in section 4 of the BHC Act (12 U.S.C. 1843). Unless otherwise noted, nonbanking activities will be conducted throughout the United States.

Unless otherwise noted, comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors not later than May 7, 2012.

A. Federal Reserve Bank of Boston

(Richard Walker, Community Affairs Officer) 600 Atlantic Avenue, Boston, Massachusetts 02210–2204

1. Coastway Bancorp, MHC and Coastway Bancorp, LLC, both in Cranston, Rhode Island; to become a mutual bank holding company and a stock bank holding company, respectively, by acquiring 100 percent of the voting shares of Coastway...
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2012–N–0001]

Cardiovascular and Renal Drugs Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

Name of Committee: Cardiovascular and Renal Drugs Advisory Committee.

General Function of the Committee: To provide advice and recommendations to the Agency on FDA’s regulatory issues.

Date and Time: The meeting will be held on May 23, 2012, from 8 a.m. to 5 p.m.

Location: FDA White Oak Campus, 10903 New Hampshire Ave., Building 31 Conference Center, the Great Room (rm. 1503), Silver Spring, MD 20993–0002. Information regarding special accommodations due to a disability, visitor parking, and transportation may be accessed at: http://www.fda.gov/AdvisoryCommittees/default.htm; under the heading “Resources for You,” click on “Public Meetings at the FDA White Oak Campus.” Please note that visitors to the White Oak Campus must enter through Building 1.

Contact Person: Kristina Toliver, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, rm. 2417, Silver Spring, MD 20993–0002. 301–796–9001, FAX: 301–847–8533, email: CRDAC@fda.hhs.gov, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), and follow the prompts to the desired center or product area. Please call the Information Line for up-to-date information on this meeting.

A notice in the Federal Register about last minute modifications that impact a previously announced advisory committee meeting cannot always be published quickly enough to provide timely notice. Therefore, you should always check the Agency’s Web site and call the appropriate advisory committee hot line/phone line to learn about possible modifications before coming to the meeting.

Agenda: The committee will discuss supplemental new drug application (sNDA) 202439/S–002, XARELTO (rivaroxaban), submitted by Janssen Pharmaceuticals, Inc., to reduce the risk of thrombotic cardiovascular events in patients with acute coronary syndrome (ACS) [ST elevation myocardial infarction (STEMI), non-ST elevation myocardial infarction (NSTEMI), or unstable angina (UAI)] in combination with aspirin alone or with aspirin plus clopidogrel or ticlopidine.

FDA intends to make background material available to the public no later than 2 business days before the meeting. If FDA is unable to post the background material on its Web site prior to the meeting, the background material will be made publicly available at the location of the advisory committee meeting, and the background material will be posted on FDA’s Web site after the meeting. Background material is available at http://www.fda.gov/AdvisoryCommittees(Calendar/default.htm. Scroll down to the appropriate advisory committee link.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person on or before May 9, 2012. Oral presentations from the public will be scheduled between approximately 1 p.m. and 2 p.m. Those individuals interested in making formal oral presentations should notify the contact person and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation on or before May 1, 2012. Time allotted for each presentation may be limited. If the number of registrants requesting to speak is larger than can be reasonably accommodated during the scheduled open public hearing session, FDA may conduct a lottery to determine the speakers for the scheduled open public hearing session. The contact person will notify interested persons regarding their request to speak by May 2, 2012.

Persons attending FDA’s advisory committee meetings are advised that the Agency is not responsible for providing access to electrical outlets.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Kristina Toliver at least 7 days in advance of the meeting.

FDA is committed to the orderly conduct of its advisory committee meetings. Please visit our Web site at http://www.fda.gov/AdvisoryCommittees/ucm111462.htm for procedures on public conduct during advisory committee meetings.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: April 9, 2012.

Jill Hartzler Warner,
Acting Associate Commissioner for Special Medical Programs.

[FR Doc. 2012–8824 Filed 4–11–12; 8:45 am]
BILLING CODE 4160–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2012–N–0001]

Request for Notification From Industry Organizations Interested in Participating in the Selection Process for Nonvoting Industry Representative on the Blood Products Advisory Committee, and Request for Nominations for Nonvoting Industry Representatives on the Blood Products Advisory Committee

AGENCY: Food and Drug Administration (FDA).

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is requesting that any industry organizations interested in participating in the selection of a nonvoting industry representative to serve on the Blood Products Advisory Committee for the Center for Biologics Evaluation and Research (CBER) notify FDA in writing. FDA is also requesting nominations for a nonvoting industry representative to serve on the Blood Products Advisory Committee. A nominee may either be self-nominated...