

ITEMS TO BE DISCUSSED: Compliance matters pursuant to 52 U.S.C. 30109.

Matters relating to internal personal decisions, or internal rules and practices. Matters concerning participation in civil actions or proceedings or arbitration. Information the premature disclosure of which would be likely to have a Considerable adverse effect on the implementation of a proposed Commission action.

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PERSON TO CONTACT FOR INFORMATION:

Judith Ingram, Press Officer, Telephone: (202) 694-1220.

Shelley E. Garr,

Deputy Secretary.

[FR Doc. 2016-27328 Filed 11-8-16; 4:15 pm]

BILLING CODE 6715-01-P

FEDERAL MARITIME COMMISSION

Notice of Agreements Filed

The Commission hereby gives notice of the filing of the following agreements under the Shipping Act of 1984. Interested parties may submit comments on the agreements to the Secretary, Federal Maritime Commission, Washington, DC 20573, within twelve days of the date this notice appears in the **Federal Register**. Copies of the agreements are available through the Commission's Web site (www.fmc.gov) or by contacting the Office of Agreements at (202) 523-5793 or tradeanalysis@fmc.gov.

Agreement No.: 010099-064.

Title: International Council of Containership Operators.

Parties: Maersk Line A/S; CMA CGM, S.A.; China COSCO Shipping Corporation Limited; Crowley Maritime Corp.; Evergreen Marine Corporation (Taiwan), Ltd.; Hamburg-Süd KG; Hapag-Lloyd AG and Hapag-Lloyd USA LLC; Hyundai Merchant Marine Co., Ltd.; Kawasaki Kisen Kaisha, Ltd.; MSC Mediterranean Shipping Company S.A.; Mitsui O.S.K. Lines, Ltd.; Nippon Yusen Kaisha; Orient Overseas Container Line, Ltd.; Pacific International Lines (Pte) Ltd.; United Arab Shipping Company (S.A.G.); Wan Hai Lines Ltd.; Yang Ming Transport Marine Corp.; and Zim Integrated Shipping Services Ltd.

Filing Party: John Longstreth, Esq.; K & L Gates LLP; 1601 K Street NW., Washington, DC 20006-1600.

Synopsis: The amendment deletes Hanjin Shipping Co., Ltd. as a party to the Agreement.

Agreement No.: 012129-002.

Title: EUKOR/"K" Line Space Charter Agreement.

Parties: EUKOR Car Carriers, Inc. and Kawasaki Kisen Kaisha, Ltd.

Filing Party: John P. Meade, Esq.; Vice-President; K-Line America, Inc.; 6009 Bethlehem Road; Preston, MD 21655.

Synopsis: The amendment adds the Dominican Republic, Grand Cayman, St. Maarten, Haiti, and the Bahamas to the geographic scope of the Agreement.

Agreement No.: 012395-001.

Title: MSC/ACL Trans-Atlantic Space Charter.

Parties: Atlantic Container Line A.B. and MSC Mediterranean Shipping Company S.A.

Filing Party: Wayne R. Rohde, Esq.; Cozen O'Connor LLP; 1200 Nineteenth St. NW., Washington, DC 20036.

Synopsis: The amendment extends the duration of the Agreement for one year.

Agreement No.: 012439.

Title: THE Alliance Agreement.

Parties: Hapag-Lloyd AG and Hapag-Lloyd USA LLC (acting as one party); Kawasaki Kisen Kaisha, Ltd.; Mitsui O.S.K. Lines, Ltd.; Nippon Yusen Kaisha; and Yang Ming Marine Transport Corp.

Filing Party: David F. Smith, Esq.; Cozen O'Connor; 1200 Nineteenth Street NW., Washington, DC 20036.

Synopsis: The Agreement authorizes the Parties to charter and exchange space on one another's vessels and to rationalize, coordinate and cooperate with respect to the Parties' transportation services and operations.

By Order of the Federal Maritime Commission.

Dated: November 7, 2016.

Rachel E. Dickon,

Assistant Secretary.

[FR Doc. 2016-27185 Filed 11-9-16; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2016-D-3275]

Product Labeling for Certain Ultrasonic Surgical Aspirator Devices; Draft Guidance for Industry and Food and Drug Administration Staff; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of the draft guidance entitled "Product Labeling for Certain Ultrasonic Surgical Aspirator

Devices." FDA is providing a specific labeling recommendation in this guidance to promote the safe and effective use of ultrasonic surgical aspirator devices. The labeling recommendation is being made in light of the risk of tissue dissemination and relates to use of these devices in the removal of uterine fibroid. This draft guidance is not final nor is it in effect at this time.

DATES: Although you can comment on any guidance at any time (see 21 CFR 10.115(g)(5)), to ensure that the Agency considers your comment of this draft guidance before it begins work on the final version of the guidance, submit either electronic or written comments on the draft guidance by January 9, 2017.

ADDRESSES: You may submit comments as follows:

Electronic Submissions

Submit electronic comments in the following way:

- *Federal eRulemaking Portal:* <http://www.regulations.gov>. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to <http://www.regulations.gov> will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else's Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on <http://www.regulations.gov>.

- If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see "Written/Paper Submissions" and "Instructions").

Written/Paper Submissions

Submit written/paper submissions as follows:

- *Mail/Hand delivery/Courier (for written/paper submissions):* Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

- For written/paper comments submitted to the Division of Dockets Management, FDA will post your comment, as well as any attachments, except for information submitted,