necessary for the proper performance of the FDIC’s functions, including whether the information has practical utility; (b) The accuracy of the estimates of the burden of the collections of information, including the validity of the methodology and assumptions used; (c) Ways to enhance the quality, utility, and clarity of the information to be collected; and (d) Ways to minimize the burden of the collections of information on respondents, including through the use of automated collection techniques or other forms of information technology. All comments will become a matter of public record.

Dated at Washington, DC, this 31st day of March 2016.

Federal Deposit Insurance Corporation.

Robert E. Feldman,
Executive Secretary.

[Ft Doc. 2016–07819 Filed 4–5–16; 8:45 am]
BILLING CODE 6714–01–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.: 010071–044.
Title: Cruise Lines International Association Agreement.

Parties: Acromas Shipping, Ltd./Saga Shipping; Aida Cruises; AMA Waterways; American Cruise Lines, Inc.; Aqua Expeditions Pte. Ltd.; Australian Pacific Touring Pty Ltd.; Avalon Waterways; Azamara Cruises; Carnival Cruise Lines; CDF Croisières de France; Celebrity Cruises, Inc.; Celestyal Cruises; Costa Cruise Lines; Compagnie Du Ponant; Croisieurope; Cruise & Maritime Voyages; Crystal Cruises; Cunard Line; Disney Cruise Line; Emerald Waterways; Evergreen Tours; Fred.Olsen Cruise Lines Ltd.; Hapag-Lloyd Kreuzfahrten GmbH; Hebridean Island Cruises; Holland America Line; Hurtigruten, Inc.; Island Cruises; Lindblad Expeditions Pty Ltd.; Luftner Cruises; Mekong Waterways; MSC Cruises; NCL Corporation; Oceania Cruises; P & O Cruises; P & O Cruises Australia; Paul Gauguin Cruises; Pearl Seas Cruises; Phoenix Reisen Gmbh; Princess Cruises; Pullmantur Cruise Management Ltd.; Regent Seven Seas Cruises; Riviera Tours Ltd.; Royal Caribbean International; Scenic Tours UK Ltd.; Seabourn Cruise Line; SeaDream Yacht Club; Shearings Holidays Ltd.; Silversea Cruises, Ltd.; Star Cruises (HK) Limited; St. Helena Line/Andrew Weir Shipping Ltd.; Swan Hellenic; Tauck River Cruising; The River Cruise Line; Thomson Cruises; Travelmarvel; Tui Cruises Gmbh; Un-Cruises Adventures; Unilworld River Cruises, Inc.; Venice Simplon-Orient-Express Ltd./Belmond; Voyages of Discovery; Voyages to Antiquity (UK) Ltd.; and Windstar Cruises.

Filing Party: Andre Picciurro, Esq., Kaye, Rose & Partners, LLP; Emerald Plaza, 402 West Broadway, Suite 1300; San Diego, CA 92101–3542
Synopsis: The amendment would add language to clarify that the agreement can represent its members before federal and state judiciaries.

Agreement No.: 011223–052.
Title: Transpacific Stabilization Agreement.


Filing Party: David F. Smith, Esq.; Cozen O’Connor; 1200 Nineteenth Street NW.; Washington, DC 20036.
Synopsis: The amendment deletes China Shipping Container Lines (Hong Kong) Company Limited and China Shipping Container Lines Company Limited as parties to the agreement.

Agreement No.: 012288–002.
Title: Hoegh/NYK Atlantic/Pacific Space Charter Agreement.

Parties: Hoegh Autoliners AS and Nippon Yusen Kaisha.

Filing Party: Wayne Rohde, Esq.; Cozen O’Connor; 1200 Nineteenth St. NW.; Washington, DC 20036.
Synopsis: The amendment adds the trades between the U.S. West Coast, on the one hand, and Thailand, Taiwan, Indonesia, Malaysia, Brunei, Philippines, Bangladesh, Vietnam, Sri Lanka, Myanmar, Singapore, Australia and New Zealand on the other hand, to the geographic scope of the agreement.

By Order of the Federal Maritime Commission.

Dated: April 1, 2016.

Karen V. Gregory,
Secretary.

[FR Doc. 2016–07890 Filed 4–5–16; 8:45 am]
BILLING CODE 6711–AA–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–16–0469]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) has submitted the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995. The notice for the proposed information collection is published to obtain comments from the public and affected agencies.

Written comments and suggestions from the public and affected agencies concerning the proposed collection of information are encouraged. Your comments should address any of the following: (a) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility; (b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (c) Enhance the quality, utility, and clarity of the information to be collected; (d) Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses; and (e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639–7570 or send an email to omb@cdc.gov. Written comments and/or suggestions regarding the items contained in this notice should be directed to the Attention: CDC Desk Officer, Office of Management and Budget, Washington, DC 20503 or by fax to (202) 395–3806. Written comments should be received within 30 days of this notice.
Proposed Project

National Program of Cancer Registries Cancer Surveillance System (NPCR CSS), (OMB No. 0922–0469 5/31/2016). CDC plans to request OMB approval to continue collecting this information for three years. Data definitions will be updated to reflect changes in national standards for cancer diagnosis and coding, but the number of respondents and the burden per respondent will not change.

The NPCR CSS allows CDC to collect, aggregate, evaluate and disseminate cancer incidence data at the national level. The NPCR CSS is the primary source of information for United States Cancer Statistics (USCS), which CDC has published annually since 2002. The latest USCS report published in 2015 provided cancer statistics for 99% of the United States population from all cancer registries whose data met national data standards. Prior to the publication of USCS, cancer incidence data at the national level were available for only 14% of the population of the United States.

The NPCR CSS also allows CDC to monitor cancer trends over time, describe geographic variation in cancer incidence throughout the country, and provide incidence data on racial/ethnic populations and rare cancers. These activities and analyses further support CDC’s planning and evaluation efforts for state and national cancer control and prevention. In addition, datasets can be made available for secondary analysis.

Respondents are NPCR-supported central cancer registries (CCR) in 45 U.S. states, 2 territories, and the District of Columbia. Thirty-eight CCR submit data elements specified for the Standard NPCR CSS Report. Ten specialized CCR submit data elements specified for the Enhanced NPCR CSS Report, which includes additional information about treatment and follow-up for cases of breast, colorectal, and chronic myeloid leukemia cases diagnosed in 2011. Each CCR is asked to transmit two data files to CDC per year. The first file, submitted in January, is a preliminary report consisting of one year of data for the most recent year of available data. CDC evaluates the preliminary data for completeness and quality and provides a report back to the CCR. The second file, submitted by November, contains cumulative cancer incidence data from the first diagnosis year for which the cancer registry collected data with the assistance of NPCR funds (e.g., 1995) through 12 months past the close of the most recent diagnosis year (e.g., 2014). The cumulative file is used for analysis and reporting. The burden for each file transmission is estimated at two hours per response. Because cancer incidence data are already collected and aggregated at the state level the additional burden of reporting the information to CDC is small.

All information is transmitted to CDC electronically. Participation is required as a condition of the cooperative agreement with CDC. There are no costs to respondents except their time.

The total estimated annualized burden hours are 192 (152 for the Standard NPCR CSS Report, and 40 for the Enhanced NPCR CSS Report).

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
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<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
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<td>Standard NPCR CSS Report</td>
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<tr>
<td></td>
<td>Enhanced NPCR CSS Report</td>
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<td>2</td>
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</table>

**Leroy A. Richardson,**
Chief, Information Collection Review Office, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2016–07806 Filed 4–5–16; 8:45 am]
BILLING CODE 4163–18–P

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

[Docket No. FDA–2016–N–0001]

**Endocrinologic and Metabolic Drugs Advisory Committee; Amendment of Notice**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing an amendment to the notice of a meeting of the Endocrinologic and Metabolic Drugs Advisory Committee. This meeting was announced in the Federal Register of March 16, 2016. The amendment is being made to reflect a change in the Date and Time portion of the document. The Date of the meeting is changed to May 25, 2016. There are no other changes.

**FOR FURTHER INFORMATION CONTACT:** LaToya Bonner, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2417,