4. Regulation of pesticide adjuvants. Some commenters asked the Agency to clarify the impact of this removal action on the use of tank-mix adjuvants, including with respect to tolerances and exemptions under the Federal Food, Drug and Cosmetic Act.

An adjuvant is a chemical substance separately added to a pesticide product (typically as part of a spray tank mixture). Since pesticide adjuvant products do not make pesticidal claims, they are not pesticides, and the components of adjuvants are therefore not pesticide inert ingredients. Adjuvants are not included in the inert ingredient approval process and are therefore unaffected by this policy. While adjuvants may need tolerances or tolerance exemptions in some cases, tolerances and exemptions are separate from the inert ingredient approval process.

5. No impact to the fragrance ingredient listing. One commenter noted that a few inert ingredients proposed for removal from the chemical substance list appear on the EPA Fragrance Ingredient List (FIL). The EPA FIL comprises more than 1,500 fragrance component ingredients that have undergone Agency evaluation to determine their suitability for safe use as components of fragrances in nonfood-use pesticide product formulations in accordance with the Fragrance Notification Program. Removal of an inert ingredient from the approved inert ingredient listing does not preclude use as a fragrance ingredient as part of the Fragrance Notification Program. Fragrance component ingredients are therefore unaffected by this policy.

6. Impurities. Some commenters want EPA to clarify that removing the chemical substances from the list does not prohibit the use of those chemical substances being classified as residual impurities in approved inert ingredients.

The definition of inert ingredient as given in 40 CFR 152.3 applies to chemical substances used as inert ingredients that are “intentionally included in a pesticide product” and as such the removal of a chemical substance from the approved inert ingredient list does not apply to circumstances where the chemical substance may be present as an impurity. Impurities in pesticide products are considered on a case-by-case basis as part of the Agency’s pesticide product registration process. As part of that evaluation, the Agency looks at the identity and amount of an impurity in the product manufacturing information, and the steps taken to limit or remove impurities.

7. Confirming the ingredient use in current pesticide products. Some commenters suggested that EPA provide them more time to investigate whether any of the 72 chemical substances are used in currently registered products. EPA records include no Confidential Statements of Formula for any currently registered pesticide product that list any of these chemical substances. However, if a registrant or a producer of proprietary mixtures identifies an active registration that contains one of the chemical substances that has now been removed from the approved inert ingredient listing, that registrant or producer should contact the Agency directly, using the contact for listing inquiries that is provided under FOR FURTHER INFORMATION CONTACT. If EPA confirms that the chemical substance is contained in a currently registered product, the Agency will restore the chemical substance to the list of approved inert ingredients.


James J. Jones, Assistant Administrator, Office of Chemical Safety and Pollution Prevention.

BILLCODE 6560-50-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.: 012338-001. Title: Sealand/APL Caribbean Slot Charter Agreement. Parties: Maersk Line A/S DBA Sealand; and APL Co. Pte Ltd. and American President Lines, Ltd. Filing Party: Wayne Rohde; Cozen O’Connor; 1200 Nineteenth Street NW.; Washington, DC 20036.

Synopsis: The amendment deletes Costa Rica from the geographic scope of the Agreement, reduces the amount of space chartered, and adjusts the minimum the duration of Agreement.

Agreement No.: 012446. Title: Sealand/APL Central America Slot Charter Agreement. Parties: Maersk Line A/S DBA Sealand; and APL Co. Pte Ltd. and American President Lines, Ltd. Filing Party: Wayne Rohde; Cozen O’Connor; 1200 Nineteenth Street NW.; Washington, DC 20036.

Synopsis: The Agreement authorizes Sealand to charter space to APL in the trade between the U.S. East Coast and ports in Panama, Costa Rica, and Colombia.


Synopsis: The Agreement authorizes THE Alliance and Zim to exchange slots on their respective services in the Agreement trade and to enter into cooperative working arrangements in connection therewith.


Rachel E. Dickon, Assistant Secretary.
the provisions (subpart C) of the CFPB’s Regulation V regarding other entities (“CFPB Rule”). The current clearance expires on January 31, 2017.

DATES: Comments must be filed by January 13, 2017.

ADDRESSES: Interested parties are invited to submit written comments electronically or in paper form by following the instructions in the Request for Comment part of the SUPPLEMENTARY INFORMATION section below. Write “Affiliate Marketing Disclosure Rule, PRA Comment: FTC File No. P105411” on your comment, and file your comment online at https://ftcpubliccommentworks.com/ftc/affiliatemarketingpra2, by following the instructions on the Web-based form. If you prefer to file your comment on paper, mail your comment to the following address: Federal Trade Commission, Office of the Secretary, 600 Pennsylvania Avenue NW., Suite CC–5610 (Annex J), Washington, DC 20580, or deliver your comment to the following address: Federal Trade Commission, Office of the Secretary, Constitution Center, 400 7th Street SW., 5th Floor, Suite 5610 (Annex J), Washington, DC 20024.

FOR FURTHER INFORMATION CONTACT: Requests for additional information should be addressed to Ruth Yodaiken, Attorney, Division of Privacy and Identity Protection, Bureau of Consumer Protection, Federal Trade Commission, 600 Pennsylvania Avenue NW., Room CC–8232, Washington, DC 20580, (202) 326–2127.

SUPPLEMENTARY INFORMATION: Under the PRA, 44 U.S.C. 3501–3521, federal agencies must get OMB approval for each collection of information they conduct or sponsor. “Collection of information” includes agency requests or requirements to submit reports, keep records, or provide information to a third party, 44 U.S.C. 3502(3); 5 CFR 1320.3(c). The FTC seeks clearance for its assumed share of the estimated PRA burden regarding the disclosure requirements under the FTC and CFPB Rules.

On August 15, 2016, the FTC sought public comment on the consumer notification (“disclosure”) requirements associated with the FTC Rule (August 15, 2016 Notice 1), the FTC’s shared enforcement with the CFPB of the disclosure provisions of the CFPB Rule, and the FTC’s associated PRA burden analysis. No relevant comments were received. The FTC provisionally retains its previously published PRA burden estimates subject to further public comment. For details about the FTC and CFPB Rules’ disclosure requirements, the background behind them, and the basis for the burden-related estimates stated below, see the August 15, 2016 Notice.2

Pursuant to the OMB regulations, 5 CFR part 1320, that implement the PRA, 44 U.S.C. 3501 et seq., the FTC is providing this second opportunity for public comment while seeking OMB renewed clearance regarding the FTC’s enforcement of and PRA burden estimates for the disclosure requirements at issue.

Burden Statement

A. Non-GLBA Entities
1. 894,969 annualized burden hours
2. $35,626,785 annualized labor cost
   These estimates include the start-up burden and attendant costs, such as determining compliance obligations.

B. GLBA Entities
1. 15,633 annualized burden hours
2. $818,059 annualized labor cost

C. FTC Share of Estimated PRA Burden
1. 460,205 annualized burden hours
2. $18,472,938 annualized labor cost
   The FTC’s share of total estimated burden for affected entities includes the increment apportioned to the FTC reflective of its sole jurisdiction over certain motor vehicle dealers. Capital and other non-labor costs should be minimal, at most, since the Rule has been in effect several years, with covered entities now equipped to provide the required notice.

Request for Comment

You can file a comment online or on paper. For the Commission to consider your comment, we must receive it on or before January 13, 2017. Write “Affiliate Marketing Disclosure Rule, PRA Comment: FTC File No. P105411” on your comment, and file your comment confidential treatment, and on the following address: Federal Trade Commission, Office of the Secretary, 600 Pennsylvania Avenue NW., Suite CC–5610 (Annex J), Washington, DC 20580, or deliver your comment to the following address: Federal Trade Commission, Office of the Secretary, Constitution Center, 400 7th Street SW., 5th Floor, Suite 5610 (Annex J), Washington, DC 20024.

□ If you file your comment on paper, write “Affiliate Marketing Disclosure Rule, PRA Comment: FTC File No. P105411” on your comment, and on the envelope, and mail or deliver it to the following address: Federal Trade Commission, Office of the Secretary, Constitution Center, 400 7th Street SW., 5th Floor, Suite 5610 (Annex J), Washington, DC 20024. If possible, submit your paper comment to the

1 81 FR 54086.
2 81 FR at 54089.
3 In particular, the written request for confidential treatment that accompanies the comment must include the actual and legal basis for the request, and must identify the specific portions of the comment to be withheld from the public record. See FTC Rule 4.9(c)(1), 16 CFR 4.9(c).
Commission by courier or overnight service.

Comments on the disclosure requirements subject to review under the PRA should additionally be submitted to OMB. If sent by U.S. mail, they should be addressed to Office of Information and Regulatory Affairs, Office of Management and Budget, Attention: Desk Officer for the Federal Trade Commission, New Executive Office Building, Docket Library, Room 10102, 725 17th Street NW., Washington, DC 20503. Comments sent to OMB by U.S. postal mail, however, are subject to delays due to heightened security precautions. Thus, comments instead should be sent by facsimile to (202) 395–5806.

The FTC Act and other laws that the Commission administers permit the collection of public comments to consider and use in this proceeding as appropriate. The Commission will consider all timely and responsive public comments that it receives on or before January 13, 2017. For information on the Commission’s privacy policy, including routine uses permitted by the Privacy Act, see http://www.ftc.gov/ftc/privacy.htm.

David C. Shonka,
Acting General Counsel.

[FR Doc. 2016–29946 Filed 12–13–16; 8:45 am]

BILLING CODE 6750–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier CMS–10266 and CMS–R–71]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

ACTION: Notice.

SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (the PRA), federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information (including each proposed extension or reinstatement of an existing collection of information) and to allow 60 days for public comment on the proposed action. Interested persons are invited to send comments regarding our burden estimates or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency’s functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

DATES: Comments must be received by February 13, 2017.

ADDRESSES: When commenting, please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in any one of the following ways:

1. Electronically. You may send your comments electronically to http://www.regulations.gov. Follow the instructions for “Comment or Submission” or “More Search Options” to find the information collection document(s) that are accepting comments.

2. By regular mail. You may mail written comments to the following address: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number , Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of the following:

2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov.
3. Call the Reports Clearance Office at (410) 786–1326.

FOR FURTHER INFORMATION CONTACT: Reports Clearance Office at (410) 786–1326.

SUPPLEMENTARY INFORMATION:

Contents

This notice sets out a summary of the use and burden associated with the following information collections. More detailed information can be found in each collection’s supporting statement and associated materials (see ADDRESSES). CMS–10266 Conditions of Participation: Requirements for Approval and Reapproval of Transplant Centers to Perform Organ Transplants

CMS–R–71 Quality Improvement Organization (QIO) Assumption of Responsibilities and Supporting Regulations

Under the PRA (44 U.S.C. 3501–3520), federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 3520(c)(2)(A) of the PRA requires federal agencies to publish a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice.

Information Collection

1. Type of Information Collection Request: Extension of a previously approved collection; Title of Information Collection: Conditions of Participation: Requirements for Approval and Reapproval of Transplant Centers to Perform Organ Transplants; Use: The Conditions of Participation and accompanying requirements specified in the regulations are used by our surveyors as a basis for determining whether a transplant center qualifies for approval or re-approval under Medicare. We, along with the healthcare industry, believe that the availability to the facility of the type of records and general content of records is standard medical practice and is necessary in order to ensure the well-being and safety of patients and professional treatment accountability. Form Number: CMS–10266 (OMB Control Number: 0938–1069); Frequency: Yearly; Affect ed Public: Business or other for-profits and Not-for-profit institutions; Number of Respondents: 226; Total Annual Responses: 528; Total Annual Hours: 2,523. (For policy questions regarding this collection contact Diane Corning at (410) 786–8486.)

2. Type of Information Collection Request: Extension of a previously approved collection; Title of Information Collection: Quality Improvement Organization (QIO) Assumption of Responsibilities and Supporting Regulations; Use: The Peer