By this Public Notice, the Media Bureau extends the deadlines for filing comments and reply comments in the above-captioned proceeding. On October 6, 2022, the Commission released a Second Notice of Proposed Rulemaking (Second Notice) seeking comment on new rules to strengthen the process for identifying foreign governmental entities. The Second Notice specified comment and reply comments dates of 30 and 45 days, respectively, after Federal Register publication. That publication occurred on November 17, 2022, and on November 18, 2022, the Media Bureau released a Public Notice (Public Notice), announcing a comment filing deadline of December 19, 2022, and a reply comment filing deadline of January 3, 2023, for the Second Notice.

On December 7, 2022, the Multicultural Media, Telecom and internet Council (MMTC) and the National Association of Broadcasters (NAB) (collectively, Joint Filers) requested an extension of the comment and reply comment filing deadlines until January 9 and January 24, 2023, respectively. The Joint Filers explain how it is “challenging” under the original filing deadline “to gather relevant information from individual broadcasters and lessees affected by the proposed rules, build useful consensus around the issues in this proceeding, and draft comments and reply comments.” A coalition of religious organizations (the Religious Programmers) filed in support of the Joint Filers’ Motion, also noting the difficulties presented by the intervening holidays.

As set forth in section 1.46(a) of the Commission’s rules, the Commission’s policy is that extensions of time shall not be routinely granted. We find, however, that the Joint Filers have provided sufficient justification to warrant grant of their requested extension. As an extension should enable interested parties to present more complete and thoughtful comments to the Commission, we agree with the Joint Filers that the extension should not disadvantage any party or cause significant delay in the resolution of this proceeding.

Federal Communications Commission.

Thomas Horan, Chief of Staff, Media Bureau.

[FR Doc. 2022–28206 Filed 12–27–22; 8:45 am]

BILLING CODE 6712–01–P

FEDERAL MARITIME COMMISSION

Notice of Agreements Filed

The Commission hereby gives notice of filing of the following agreements under the Shipping Act of 1984. Interested parties may submit comments, relevant information, or documents regarding the agreements to the Secretary by email at Secretary@fmc.gov, or by mail, Federal Maritime Commission, 800 North Capitol Street, Washington, DC 20573. Comments will be most helpful to the Commission if received within 12 days of the date this notice appears in the Federal Register, and the Commission requests that comments be submitted within 7 days on agreements that request expedited review. Copies of agreements are available through the Commission’s website (www.fmc.gov) or by contacting the Office of Agreements at (202) 523–5793 or tradeanalysis@fmc.gov.

Agreement No.: 201254–002.

Agreement Name: Sealand/CMA CGM West Coast of Central America Slot Charter Agreement.

Parties: Maersk A/S DBA Sealand and CMA CGM S.A.

Filing Party: Wayne Rohde, Cozen O’Connor.

Synopsis: The amendment revises the strings and amount of space being chartered under the Agreement; adds a new Article 5.10, and updates Article 12.

Proposed Effective Date: 2/2/2023.

Location: https://www2.fmc.gov/FMC.Agreements.Web/Public/AgreementHistory/10193.

Agreement No.: 201368–001.

Agreement Name: ONE/CMA CGM Slot Exchange Agreement.

Parties: CMA CGM S.A. and Ocean Network Express Pte. Ltd.

Filing Party: Robert Magovern, Cozen O’Connor.

Synopsis: The amendment adds Malaysia, Thailand, and Vietnam to the geographic scope of the Agreement and provides for ONE to receive space on CMA CGM’s PRX and JAX service in case of slot exchange imbalance.

Proposed Effective Date: 2/4/2023.

Location: https://www2.fmc.gov/FMC.Agreements.Web/Public/AgreementHistory/49505.


JoAnne O’Bryant, Program Analyst, Secretary.

[FR Doc. 2022–28267 Filed 12–27–22; 8:45 am]

BILLING CODE P

FEDERAL RESERVE SYSTEM

Change in Bank Control Notices; Acquisitions of Shares of a Bank or Bank Holding Company

The notificants listed below have applied under the Change in Bank Control Act (Act) (12 U.S.C. 1817(j)) and §225.41 of the Board’s Regulation Y (12 CFR 225.41) to acquire shares of a bank or bank holding company. The factors that are considered in acting on the applications are set forth in paragraph 7 of the Act (12 U.S.C. 1817(j)(7)).

The public portions of the applications listed below, as well as other related filings required by the Board, if any, are available for immediate inspection at the Federal Reserve Bank(s) indicated below and at the offices of the Board of Governors. This information may also be obtained on an expedited basis, upon request, by contacting the appropriate Federal Reserve Bank and from the Board’s Freedom of Information Office at https://www.federalreserve.gov/foia/request.htm. Interested persons may express their views in writing on the standards enumerated in paragraph 7 of the Act.

Comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors, Ann E. Misback, Secretary of the Board, 20th Street and Constitution Avenue NW, Washington, DC 20551–0001, not later than January 12, 2023.

A. Federal Reserve Bank of St. Louis

(Holly A. Rieser, Senior Manager) P.O. Box 442, St. Louis, Missouri 63166–2034. Comments can also be sent electronically to Comments.applications@stls.frb.org.

1. The Steven M. Dalton 2012 Gift Trust Fund, Stephen M. Dalton, Jr., individually, and as trustee, all of Sugar Land, Texas; the Everett McCain Dalton 2012 Gift Trust Fund, Everett M. Dalton, individually, and as trustee, and William E. Dalton, Jr., all of Houston, Texas; Elizabeth McCain, Takoma Park, Maryland; James E. McCain, III, Summerfield, Florida; Marguerite M. Lloyd, individually, and as executor of the Estate of Sam Lloyd, both of Columbus, Mississippi; to join the McCain/Dalton Family Group, a group

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Agency Forms Undergoing Paperwork Reduction Act Review

In accordance with the Paperwork Reduction Act of 1995, the Centers for Disease Control and Prevention (CDC) has submitted the information collection request (ICR) titled “Distribution of Traceable Opioid Material Kits (TOM Kits) across U.S. and International Laboratories” to the Office of Management and Budget (OMB) for review and approval. CDC previously published a “Proposed Data Collection Submitted for Public Comment and Recommendations” notice on 09/16/2022 to obtain comments from the public and affected agencies. CDC received one comment related to the previous notice. This notice serves to allow an additional 30 days for public and affected agency comments.

CDC will accept all comments for this proposed information collection project. The Office of Management and Budget is particularly interested in comments that:

(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. Comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting “Currently under 30-day Review—Open for Public Comments” or by using the search function. Direct written comments and/or suggestions regarding the items contained in this notice to the Attention: CDC Desk Officer, Office of Management and Budget, 725 17th Street NW, Washington, DC 20503 or by fax to (202) 395–5806. Provide written comments within 30 days of notice publication.

Proposed Project

Distribution of Traceable Opioid Material Kits (TOM Kits) across U.S. and International Laboratories (OMB Control No. 0920–1313, expiration date 12/31/2022)—Revision—National Center for Environmental Health (NCEH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

In response to the Health and Human Services (HHS) Acting Secretary’s 2017 and ongoing public health emergency declaration on opioids, the Centers for Disease Control and Prevention (CDC) has led the development of Traceable Opioid Material Kits (TOM Kits) to support detection of emerging opioids. CDC maintains the contents of the TOM Kits based on new needs identified, in part, through the U.S. Drug Enforcement Agency (DEA) Emerging Threat Reports. For example, the DEA 2018 data indicated that fentanyl and fentanyl-related compounds accounted for approximately 76 percent of their opioid identifications.

The CDC is requesting a three-year Paperwork Reduction Act (PRA) clearance for a revision of this ICR (formerly known as “Distribution of Traceable Opioid Material Kits [TOM Kits] across U.S. Laboratories”) (OMB Control No. 0920–1313). As part of the proposed revisions, CDC will be expanding its program to include a new line of TOM Kits, the Emerging Drug Panel (EDP) Kits. For the EDP Kits, non-opioid compounds will be identified and updated by searching recent lists put out by the DEA and the Center for Forensic Science Research and Education (CFSRE). These lists provide data on all classes of drugs that were recently identified in the field and provide recommendations on which drugs should be included in testing. They are updated several times a year and keep up with the changing drug landscape in the United States. For the current round, EDP Kits will include synthetic cannabinoids, stimulants, hallucinogens, and benzodiazepines.

CDC will distribute TOM Kits through a single vendor, which will manufacture the test kits. The CDC vendor will distribute these kits to domestic laboratories, as previously approved under CDC contract. As a revision, the CDC vendor will distribute these test kits to international laboratories in partnership with the United Nations and under a separate contract with the International Narcotics Control Board (INCB) (hereafter, collectively coined the “UN”). The UN, and not the CDC, is paying the vendor to ship the kits to international requesters.

TOM Kits are not intended for diagnostic use and are free to domestic and international laboratories in the public, private, clinical, law enforcement, research, and public health domains. The CDC vendor collects both application and laboratory information on domestic laboratories when they apply for test kits. International laboratories that apply for test kits through the UN will be directed to complete and share their laboratory information with the vendor, but not with the CDC. This information is used to prioritize which laboratories will receive kits when quantities are limited. The brief web-based surveys will allow the CDC to: (1) determine what service the recipient laboratory performs; and to (2) equitably distribute test kits based on the analysis techniques and matrices used by the recipient laboratory.

Over the past three years, CDC has received 1,472 requests from interested laboratories (approximately 490 requests per year) and has distributed 3,007 TOM Kits. Based on this experience and with the addition of EDP Kits, we anticipate that up to 600 domestic laboratories will request test kits per year. Given that each application will take six minutes, the annual time burden for 600 domestic laboratories will be 60 hours. CDC will add 20 additional annual burden hours for the international distribution of test kits. We estimate that 300 international partner laboratories will apply for test kits per year with the UN, which in turn will direct these laboratories to complete a brief four-minute survey on laboratory information on the CDC vendor website.