FEDERAL MARITIME COMMISSION
(DOCKET NO. 16–11)

Notice of Filing of Complaint and Assignment—Correction

LANDERS BROTHERS AUTO GROUP, INC., D/B/A LANDERS HONDA (JONESBORO), LANDERS BROTHERS AUTO NO. 4, LLC, D/B/A/LANDERS HONDA (PINE BLUFF), INDIVIDUALLY AND ON BEHALF OF OTHERS SIMILARLY SITUATED.

V.

NIPPON YUSEN KABUSHIKI KAISHA, NYK LINE (NORTH AMERICA) INC., MITSUI O.S.K. LINES, LTD., MITSUI O.S.K. BULK SHIPPING (USA), INC., WORLD LOGISTICS SERVICE (USA) INC., HOEGH AUTOLINERS AS, HOEGH AUTOLINERS, INC., NISSAN MOTOR CAR CARRIERS CO. LTD., KAWASAKI KISEN KAISHA, LTD., “K” LINE AMERICA, INC., WALLENIUS WILHELMSEN LOGISTICS AS, WALLENIUS WILHELMSEN LOGISTICS AMERICAS LLC, EUKOR CAR CARRIERS INC., COMPANIA SUD AMERICANA DE VAPORES S.A., AND CSAV AGENCY NORTH AMERICA, LLC.

In a Notice of Filing of Complaint and Assignment published on Wednesday, May 4, 2016, 81 FR 26793, the last sentence stated that “[t]he initial decision of the presiding officer in this proceeding shall be issued by April 28, 2017 and the final decision of the Commission shall be issued by November 13, 2017.” Due to a clerical error, the date for the initial decision was incorrect. That sentence is corrected to read as follows: “The initial decision of the presiding officer in this proceeding shall be issued by May 12, 2017 and the final decision of the Commission shall be issued by November 13, 2017.”

Karen V. Gregory,
Secretary.

ORDER

The amendment deletes Agreement No.: 012408.
Title: WWL/Grimaldi Euromed SPA Space Charter Agreement.
Parties: Wallenius Wilhelmsen Logitics AS and Grimaldi Euromed SPA.
Filing Party: Wayne R. Rohde, Esq.; Cozen O’Connor LLP; 1200 Nineteenth St. NW.; Washington, DC 20036.
Synopsis: The Agreement authorizes the parties to charter space to/from one another on an “as needed/as available” basis in the trade between ports on the Atlantic Coast of the United States and ports in North Europe and on the Mediterranean Sea.
Agreement No.: 012409.
Title: CMA CGM/COSCON Slot Exchange Agreement Asia—U.S. West Coast.
Synopsis: The Agreement authorizes the parties to exchange slots on their respective services in the trade between the United States West Coast and the People’s Republic of China (including Hong Kong), Singapore, Malaysia, Vietnam, and Canada.
Agreement No.: 201202–009.
Title: Oakland MTO Agreement.
Parties: Everport Terminal Services, Inc.; SSA Terminals, LLC; SSA Terminals (Oakland), LLC; and Trapac, Inc.
Filing Party: Wayne R. Rohde, Esq.; Cozen O’Connor; 1200 19th Street NW.; Washington, DC 20036.
Synopsis: The amendment deletes Ports America Outer Harbor Terminal, LLC as a party to the Agreement.
By Order of the Federal Maritime Commission.
Dated: May 6, 2016.
Rachel E. Dickon,
Assistant Secretary.

[FR Doc. 2016–11084 Filed 5–10–16; 8:45 am]
BILLING CODE 6731–AA–P

FEDERAL MINE SAFETY AND HEALTH REVIEW COMMISSION
(BAC 6735–01)

Sunshine Act Notice

May 9, 2016.
TIME AND DATE: 10:00 a.m., Wednesday, May 18, 2016.
STATUS: Closed.

MATTERS TO BE CONSIDERED: The Commission will consider and act upon the following in closed session as a continuation of the meeting held on May 4, 2016: Secretary of Labor v. Newtown Energy, Inc., Docket No. WEVA 2011–283 (Issues include whether the Administrative Law Judge erred by concluding that the violation in question was not significant and substantial and was not the result of an unwarrantable failure to comply.)

Any person attending this meeting who requires special accessibility features and/or auxiliary aids, such as sign language interpreters, must inform the Commission in advance of those needs. Subject to 29 CFR 2706.150(a)(3) and § 2706.160(d).


Sarah Stewart,
Deputy General Counsel.

[FR Doc. 2016–11199 Filed 5–9–16; 4:15 pm]
BILLING CODE 6731–AA–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and Prevention
[30Day–16–0822]
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
acquiescence 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) 395–5806. Written comments should be received within 30 days of this notice.

Proposed Project
The National Intimate Partner and Sexual Violence Survey (NISVS) (OMB Control No. 0920–0822, expiration date 6/30/2016)—Revision — National Center for Injury Prevention and Control (NCIPC), Centers for Disease Control and Prevention (CDC).

Background and Brief Description
This is a revision request for the currently approved National Intimate Partner and Sexual Violence Survey (NISVS). OMB approval is requested for three years.

In 2010, NISVS reported that approximately 6.9 million women and 5.6 million men experienced rape, physical violence and/or stalking by an intimate partner within the last year. The health care costs of Intimate Partner Violence (IPV) exceed $5.8 billion each year, nearly $3.9 billion of which is for direct medical and mental health care services.

In order to address this important public health problem, CDC implemented, beginning in 2010, the National Intimate Partner and Sexual Violence Surveillance System that produces national and state level estimates of IPV, Sexual Violence (SV) and stalking on an annual basis. CDC is requesting a continuation of data collection among non-institutionalized adult men and women aged 18 years or older in the United States assessing lifetime experiences of IPV, SV and stalking with a new and improved data collection tool. The revisions to the survey are aimed at reducing the time and complexity of the instrument, thus reducing the burden on the respondent. The simplified structure of the instrument will also reduce the complexity of the data set, making it more assessable for public use.

Additionally, in collaboration with the Department of Defense (DoD), NISVS will collect information regarding the experiences of IPV, SV and stalking among active duty women and men in the military and wives of active duty men.

Data collected are used by local, state and national governments and organizations to inform prevention programs and policy making related to intimate partner violence, sexual violence and stalking. This data collection will take place during the first three months of data collection. Data are analyzed using appropriate statistical software to account for the complexity of the survey design to compute weighted counts, percentages, confidence intervals using both national and state level data.

To comply with OMB requirements, CDC is developing an expert panel to address methodological issues with the NISVS survey. The panel will meet multiple times over the course of the next year. The members of this panel will provide guidance on how to improve both survey design (methods, sampling frame, recruitment, mode of administration) and content/question wording with the goals of increasing response rates, reducing non-response bias, and maximizing the opportunities across Federal surveys for covering populations of interest. The survey will be conducted among English or Spanish speaking male and female adults (18 years and older) living in the United States. The estimated annual burden hours are 27,106. There are no extra costs to respondents.

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<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>
</tr>
</thead>
<tbody>
<tr>
<td>Non-Participating Household (Screened)</td>
<td>NISVS Survey Instrument. First section non-participating.</td>
<td>170,000</td>
<td>1</td>
<td>3/60</td>
</tr>
<tr>
<td>Eligible Household (Completes Survey)</td>
<td>NISVS Survey Instrument. Section for participating.</td>
<td>25,000</td>
<td>1</td>
<td>25/60</td>
</tr>
<tr>
<td>Non-Participating DoD Household (Screened)</td>
<td>NISVS Survey Instrument. Section for DoD participating.</td>
<td>73,800</td>
<td>1</td>
<td>3/60</td>
</tr>
<tr>
<td>Eligible DoD Household (Completes Survey)</td>
<td>NISVS Survey Instrument. Section for participating.</td>
<td>10,800</td>
<td>1</td>
<td>25/60</td>
</tr>
</tbody>
</table>
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services


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 July 11, 2016.

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 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–10261 Part C Medicare Advantage Reporting Requirements and Supporting Regulations in 42 CFR 422.516(a)


CMS–10463 Cooperative Agreement To Support Navigators in Federally-Facilitated and State Partnership Exchanges

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 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(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.

1. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Part C Medicare Advantage Reporting Requirements and Supporting Regulations in 42 CFR 422.516(a); Use: Medicare Advantage Organizations (MAOs) must have an effective procedure to develop, compile, evaluate, and report to CMS, to its enrollees, and to the general public, at the times and in the manner that CMS requires, and while safeguarding the confidentiality of the doctor-patient relationship, statistics and other information with respect to: the cost of its operations; the patterns of service utilization; the availability, accessibility, and acceptability of its services; to the extent practical, developments in the health status of its enrollees; information demonstrating that the MAO has a fiscally sound operation; and other matters that CMS may require. CMS also has oversight authority over cost plans which includes establishment of reporting requirements. This revision would add five new data elements to the reporting section: Organization Determinations and Reconsiderations. These new data elements are needed to obtain more information about case reopenings. The revision would also suspend the Sponsor Oversight of Agents reporting section beginning 2017 so that the reporting section can be reassessed based on burden and usage. Form Number: CMS–10261 (OMB control number: 0938–1054); Frequency: Yearly and Semi-annually; Affected Public: Private sector (Business or other For-profits); Number of Respondents: 544; Total Annual Responses: 3,508; Total Annual Hours: 160,215. (For policy questions regarding this collection contact Terry Lied at 410–786–8973).

2. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Reporting Requirements for States Under Transitional Medical Assistance (TMA) Provisions; Use: The HHS Secretary is required to submit annual reports to Congress with information collected from states in accordance with section 5004(d) of the American Recovery and Reinvestment Act of 2009. Medicaid agencies in 50 states complete the reports while we review the information to determine if each state has met all of the reporting requirements specified under section 5004(d). Form Number: