unsupported by modern economics and at odds with the *Merger Guidelines*.

Similarly, substantiating a unilateral effects theory requires particularized evidence—also absent from the record here in some Relevant Markets—that a merger will reduce or eliminate competitive constraints, permitting the merged entity to increase prices. Without such evidence, a unilateral effects theory reduces to little more than a complaint about market structure coupled with speculation about the circumstances under which unilateral effects might occur in a post-merger world. The Merger Guidelines contemplate a more rigorous analysis.

This is not to suggest the "reason to believe" standard requires access to every piece of relevant information and a full and complete economic analysis of a proposed transaction, regardless of whether the parties wish to propose divestitures before complying with a Second Request. Rather, the standard requires only evidence sufficient to establish that competitive harm is likely. Such evidence, although quite minimal—indeed, a handful of facts in most instances—is indeed available in some Relevant Markets in this matter. and it is in those markets that I concur with the Commission's decision. While I appreciate the practical complications of requesting additional information during the course of a merger investigation, as well as the desire to conduct efficient investigations, these important pragmatic considerations do not trump the Commission's primary obligation to collect evidence sufficient to establish reason to believe the merger will harm competition before issuing a complaint and accepting a consent.

For the reasons I explain above, I find reason to believe the proposed transaction is likely to result in unilateral price effects, and thus violate the Clayton Act, in the Twin Cities, Duluth, western Wisconsin, New Orleans, western Montana, Boston/ Providence, the Mid-Atlantic region, and the western Great Lakes region. I conclude there is no reason to believe the proposed transaction will violate Section 7 in eastern Iowa, Memphis, Baton Rouge, Detroit, northern Michigan, and Grand Rapids; it follows that I believe the Commission should refrain from imposing a remedy in these markets.

[FR Doc. 2015-11724 Filed 5-14-15; 8:45 am]

BILLING CODE 6750-01-P

GENERAL SERVICES ADMINISTRATION

[OMB Control No. 3090-0205; Docket 2015-0001; Sequence 12]

General Services Administration Acquisition Regulation (GSAR; Information Collection; Environmental Conservation, Occupational Safety, and Drug-Free Workplace

AGENCY: Office of Acquisition Policy, General Services Administration (GSA). **ACTION:** Notice of request for comments regarding the extension of a previously existing OMB clearance.

SUMMARY: Under the provisions of the Paperwork Reduction Act, the General Services Administration will be submitting to the Office of Management and Budget (OMB) a request to review and approve an extension of a previously approved information collection requirement regarding Environmental Conservation, Occupational Safety, and Drug-Free Workplace.

DATES: Submit comments on or before: July 14, 2015.

ADDRESSES: Submit comments identified by Information Collection 3090–02085 by any of the following methods:

• Regulations.gov: http://www.regulations.gov.

Submit comments via the Federal eRulemaking portal by searching the OMB control number. Select the link "Submit a Comment" that corresponds with "Information Collection 3090—0205, Environmental Conservation, Occupational Safety, and Drug-Free Workplace". Follow the instructions provided at the "Submit a Comment" screen. Please include your name, company name (if any), and "Information Collection 3090—0205, Environmental Conservation, Occupational Safety, and Drug-Free Workplace" on your attached document.

• *Mail:* General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405. ATTN: Ms. Flowers/IC 3090–0205, Environmental Conservation, Occupational Safety, and Drug-Free Workplace.

Instructions: Please submit comments only and cite Information Collection 3090–0205, Environmental Conservation, Occupational Safety, and Drug-Free Workplace, in all correspondence related to this collection. All comments received will be posted without change to http://www.regulations.gov, including any personal and/or business confidential information provided.

FOR FURTHER INFORMATION CONTACT:

Kevin Funk, Procurement Analyst, General Services Acquisition Policy Division, GSA, at telephone 215–446– 4860 or via email to kevin.funk@gsa.gov.

SUPPLEMENTARY INFORMATION:

A. Purpose

The Federal Hazardous Substance Act and Hazardous Material Transportation Act prescribe standards for packaging of hazardous substances. To meet the requirements of the Acts, the General Services Administration Regulation prescribes provision 552.223–72, Hazardous Material Information, to be inserted in solicitations and contracts that provides for delivery of hazardous materials on an f.o.b. origin basis.

This information collection will be accomplished by means of the provision which requires the contractor to identify for each National Stock Number, the DOT Shipping Name, DOT Hazards Class, and whether the item requires a DOT label. Contracting Officers and technical personnel use the information to monitor and ensure contract requirements based on law and regulation.

Properly identified and labeled items of hazardous material allows for appropriate handling of such items throughout GSA's supply chain system. The information is used by GSA, stored in an NSN database and provided to GSA customers. Non-Collection and/or a less frequently conducted collection of the information resulting from provision 552.223-72 would prevent the Government from being properly notified. Government activities may be hindered from apprising their employees of; (1) All hazards to which they may be exposed; (2) Relative symptoms and appropriate emergency treatment; and (3) Proper conditions and precautions for safe use and exposure.

B. Annual Reporting Burden

Respondents: 563. Responses per Respondent: 3. Total Responses: 1689. Hours per Response: .67. Total Burden Hours: 1132.

C. Public Comments

Public Comments are particularly invited on: Whether this collection of information is necessary and whether it will have practical utility; whether our estimate of the public burden of this collection of information is accurate and based on valid assumptions and methodology; and ways to enhance the quality, utility, and clarity of the information to be collected.

Obtaining Copies of Proposals: Requesters may obtain a copy of the information collection documents from the General Services Administration, Regulatory Secretariat Division, 1800 F Street NW., Washington, DC 20405, telephone 202-501-4755.

Please cite OMB Control No. 3090-0205, Environmental Conservation, Occupational Safety, and Drug-Free Workplace, in all correspondence.

Dated: May 12, 2015.

Jeffrey A Koses,

Director, Office of Acquisition Policy, Senior Procurement Executive.

[FR Doc. 2015-11749 Filed 5-14-15; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifiers: CMS-437A & CMS-

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare &

ACTION: Notice.

Medicaid Services.

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 14, 2015.

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:

- 1. Access CMS' Web site address at http://www.cms.hhs.gov/ PaperworkReductionActof1995.
- 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-437A & CMS-437B State Agency **Sheets for Verifying Exclusions From** the Inpatient Prospective Payment **System 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 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.

Information Collection

1. Type of Information Collection Request: Revision of a currently approved collection. Title of Information Collection: State Agency Sheets for Verifying Exclusions from the Inpatient Prospective Payment System and Supporting Regulations Use: For first time verification requests for exclusion from the Inpatient Prospective Payment System (IPPS), a hospital/unit must notify the Regional Office (RO) servicing the State in which it is located that it believes it meets the criteria for exclusion from the IPPS. Currently, all new inpatient rehabilitation facilities (IRFs) must provide written certification that the inpatient population it intends to serve will meet the requirements of the IPPS exclusion criteria for IRFs. They must also complete the Form CMS-437A if they are a rehabilitation unit or complete Form CMS-437B if they are a rehabilitation hospital. This information is submitted to the State Agency (SA) no later than 5 months before the date the hospital/unit would become subject to IRF-PPS.

We propose to continue to use the Criteria Worksheets (Forms CMS-437A and CMS-437B) for verifying first-time exclusions from the IPPS, for complaint surveys, for its annual 5 percent validation sample, and for facility selfattestation. These forms are related to the survey and certification and Medicare approval of the IPPS-excluded rehabilitation units and rehabilitation hospitals.

For rehabilitation hospitals and rehabilitation units already excluded from the IPPS, annual onsite reverification surveys by the SA are not required. These hospitals and units will be provided with a copy of the appropriate CMS-437 Worksheet at least 5-months prior to the beginning of its cost reporting period, so that the hospital/unit official may complete and sign an attestation statement and complete and return the appropriate CMS-437A or CMS-437B at least 5months prior to the beginning of its cost reporting period. Fiscal Intermediaries will continue to verify, on an annual basis, compliance with the 60 percent rule (42 CFR 412.29(b)(2)) for rehabilitation hospitals and rehabilitation units through a sample of medical records and the SA will verify the medical director requirement.

The SA will maintain the documents unless instructed otherwise by the RO.