

commercial banks on interest rates charged on loans for new vehicles and loans for other consumer goods and personal expenses. The data are used for the analysis of household financial conditions.

The FR 2835a collects information on two measures of credit card interest rates from a sample of commercial banks with \$1 billion or more in credit card receivables and a representative group of smaller issuers. The data are used to analyze the credit card market and draw implications for the household sector.

Current Actions: On March 12, 2015 the Federal Reserve published a notice in the **Federal Register** (80 FR 13001) requesting public comment for 60 days on the extension, with revision, of the Quarterly Report of Interest Rates on Selected Direct Consumer Installment Loans and Quarterly Report of Credit Card Plans. The comment period for this notice expired on May 11, 2015. The Federal Reserve received one comment supporting the revisions. The revisions will be implemented as proposed.

3. *Report title:* Census of Finance Companies.

Agency form number: FR 3033p.

OMB control number: 7100-0277.

Frequency: Every five years.

Reporters: Domestic finance companies.

Estimated annual reporting hours: 8,000 hours.

Estimated average hours per response: 0.5 hours.

Number of respondents: 16,000.

General description of report: This information collection is authorized by law (12 U.S.C. 225a, 263, 348a, and 353-359) and is voluntary. Individual responses are exempt from disclosure pursuant to section (b)(4) of the Freedom of Information Act (5 U.S.C. 552).

Abstract: The Census of Finance Companies is a simple screening survey, which would be sent in June 2015 to all companies that meet criteria developed to identify the potential universe of domestic finance companies. An accurate census is required to form a representative sample of finance companies, to which the more detailed Survey of Finance Companies would be sent. The census would gather limited information including total assets, areas of specialization, and information on the corporate structure of the companies. The Federal Reserve has identified approximately 40,000 firms to which the census would be sent.

Current Actions: On March 12, 2015, the Federal Reserve published a notice in the **Federal Register** (80 FR 13001)

requesting public comment for 60 days on the extension, with revision, of the FR 3033p. The comment period for this notice expired on May 11, 2015. The Federal Reserve did not receive any comments. The revisions will be implemented as proposed.

Board of Governors of the Federal Reserve System, May 26, 2015.

Robert deV. Frierson,
Secretary of the Board.

[FR Doc. 2015-13005 Filed 5-28-15; 8:45 am]

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GENERAL SERVICES ADMINISTRATION

[Notice—ME—2015—01; Docket No: 2015—0002; Sequence No. 15]

Notice of Public Meeting Concerning the General Services Administration's Request for Information on Business Due Diligence

AGENCY: Office of Information Integrity, and Access; Office of Government-wide Policy; General Services Administration.

ACTION: Notice of public meeting.

SUMMARY: The purpose of this public meeting is to present information related to the government's analysis of responses to the General Services Administration's (GSA) Request for Information (RFI) on Business Due Diligence for Acquisition Involving Government Information or Information Systems, dated December 12, 2014. The meeting will focus on the problem of supply chain security, potential solution(s), and a path forward to initializing operation of the solution(s).

DATES: The meeting will be held on Tuesday, June 2, 2015 from 11:30 a.m. to 3 p.m., Eastern Standard Time, during the Software Supply Chain Assurance (SSCA) Working Groups (WGs) at MITRE. Online registration for the SSCA WGs is at <https://register.mitre.org/ssca/>. Comments are due no later than Friday, May 29, 2015.

ADDRESSES: *Meeting Location:* MITRE-1, 7525 Colshire Drive, McLean, VA 22102. If interested in speaking at the meeting, please submit a request to speak (for a maximum of five minutes during the public session) and cite Notice-ME-2015-01, in all correspondence related to this case. Submit comments in response to Notice—ME—2015—01 by any of the following methods:

- *Regulations.gov:* <http://www.regulations.gov>. Submit comments via the Federal eRulemaking portal by searching for "Notice-ME-2015-01".

Select the link "Comment Now" that corresponds with "Notice—ME—2015—01" and follow the instructions provided on the screen. Please include your name, company name (if any), and "Notice—ME—2015—01" on your attached document.

- *Mail:* General Services Administration, Office of Government-Wide Policy (ME), ATTN: Ms. Rowan Ha/Notice—ME—2015—01, 1800 F Street NW., Washington, DC 20405-0001.

Instructions: Please submit comments only and cite Notice—ME—2015—01 in all correspondence related to this case. 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: Ms. Rowan Ha, Cybersecurity Specialist, GSA Office of Government-wide Policy, at 202-219-1270, or rowan.ha@gsa.gov.

SUPPLEMENTARY INFORMATION: Federal Agencies continue to express concerns about potential risks in the products, services, and solutions they purchase. These concerns extend to all purchased items that connect in any way to a government information system and/or which contain, transmit, or process information provided by or generated for the government to support the operations and assets of a Federal agency.

Federal Agencies need better visibility into, and understanding of, how the products, services, and solutions they buy are developed, integrated, and deployed. Agencies are also interested in strengthening confidence in the processes, procedures, and practices used to improve the integrity, security, resilience, and quality of those products and services.

GSA is collaborating with its customer agencies and other stakeholders to establish a common set of risk indicators that can be used as the baseline for business due diligence research. This common core of risk indicators and risk research methodologies will be complementary to, and not a replacement for, existing government supply chain risk management activities.

Following a period of research and development to analyze and validate risk assessment processes, GSA intends to use a consensus set of common risk indicators from government and industry to enhance its current risk assessment processes. It is anticipated that the business due diligence information obtained will be used by the Federal acquisition, grant, and oversight communities to support

government risk assessments. Selection of contractors about which information may be collected during the assessment process will be a risk-based decision made at the discretion of a participating agency.

Definition: Information system in this notice means a discrete set of information resources organized expressly for the collection, processing, maintenance, use, sharing, dissemination, or disposition of information. Information systems also include specialized systems such as industrial or process controls systems, telephone switching or private branch exchange (PBX) systems, and environmental control systems (see, National Institute of Standards and Technology Special Publication 800–53 Rev. 4). Links to relevant documents can be found at: Business Due Diligence RFI: https://www.fbo.gov/index?s=opportunity&mode=form&id=230732591f542b7da9b9fc3e6c167eec&tab=core&_cview=0; Executive Order 13636, Improving Critical Infrastructure Cybersecurity: <http://www.gsa.gov/portal/content/176547>.

Dated: May 21, 2015.

Giancarlo Brizzi,

Acting Associate Administrator, Office of Government-wide Policy, General Services Administration.

[FR Doc. 2015–13016 Filed 5–28–15; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–15–1019]

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) 395–5806. Written comments should be received within 30 days of this notice.

Proposed Project

Integrating Community Pharmacists and Clinical Sites for Patient-Centered HIV Care (OMB No. 0920–1019, Expires 05/31/2017)—[Revision]—National Center for HIV, Hepatitis, STD, and TB Prevention, Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Revisions to this information collection include the addition of an Interviewer data collection worksheet, Key Informant Interviewer script, Staff communication questionnaire, Clinic cost form and Pharmacy cost form. These additions are needed in order to determine changes to clinic and pharmacy work systems, processes and outcomes in relation to the model project and how and if the model program improves patient outcomes through improved communication and collaboration between patients' clinical providers and pharmacists. In order to determine the general feasibility of the model program, the time required conducting program activities and the associated cost of program activities must be determined. Collection of data from the previously approved Initial patient information forms, Quarterly patient information forms, Pharmacy record abstraction forms, Project clinic

characteristics forms, and Project pharmacy characteristics forms is ongoing. Clinic staff will use the initial information Sheet to explain the project to patients.

CDC has entered into a partnership with Walgreen Company (a.k.a. Walgreens pharmacies, a national retail pharmacy chain) and the University of North Texas Health Science Center to develop and implement a model of HIV care that integrates community pharmacists with primary medical providers for patient-centered HIV care. The model program will be implemented at ten sites and will provide patient-centered HIV care for approximately 1,000 persons.

The patient-centered HIV care model includes the core elements of pharmacist provided Medication Therapy Management (MTM) as well as additional pharmacist services such as individualized medication adherence counseling, active monitoring of prescription refills and active collaboration between pharmacists and medical clinic providers to identify and resolve medication related treatment problems such as treatment effectiveness, adverse events and poor adherence. The expected outcomes of the model program are increased retention in HIV care, adherence to HIV medication therapy and HIV viral load suppression.

Pharmacy, laboratory and medical data are collected through abstraction of participant clients' pharmacy and medical records. These data are needed to monitor retention in care, adherence to therapy, viral load suppression and other health outcomes. Program specific data, such as the number of MTM elements completed per project site and project sites' characteristics, will be collected by project sites.

This information collection will allow CDC to conduct continuous program performance monitoring which includes identification of barriers to program implementation, solutions to those barriers, and documentation of client health outcomes. Performance monitoring will allow the model program to be adjusted, as needed, in order to develop a final implementation model that is self-sustaining and which can be used to establish similar collaborations in a variety of clinical settings. Collection of cost data will allow for the cost of the program to be estimated. There is no cost to participants other than their time. The total estimated annualized burden hours are 6,043.