The annual burden for this information collection is estimated to be 7,806 hours. This represents an increase of 5,472 hours from the current burden estimate of 2,334 hours. This increase is not due to any new requirements imposed by the FDIC. Rather, it is due to FDIC’s reassessment of the burden hours associated with the contracting process and to better account for the burdens associated with requests for proposals and price quotations as well as RFIs.

Request for Comment

Comments are invited on: (a) Whether the collection of information is necessary for the proper performance of the FDIC’s functions, including whether the information has practical utility; (b) the accuracy of the estimates of the burden of the information collection, including the validity of the methodology and assumptions used; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. All comments will become a matter of public record.

Dated at Washington, DC, on August 13, 2018.

Federal Deposit Insurance Corporation.

Robert E. Feldman,
Executive Secretary.

[FR Doc. 2018–17647 Filed 8–15–18; 8:45 am]

BILLING CODE 6714–01–P

FEDERAL RESERVE SYSTEM

Agency Information Collection Activities: Announcement of Board Approval Under Delegated Authority and Submission to OMB

AGENCY: Board of Governors of the Federal Reserve System.

SUMMARY: The Board of Governors of the Federal Reserve System (Board) is adopting a proposal to extend for three years, with revision, the Report of Selected Money Market Rates (FR 2420; OMB No. 7100–0357). The revisions are applicable as of October 1, 2018.


SUPPLEMENTARY INFORMATION:

On June 15, 1984, the Office of Management and Budget (OMB) delegated to the Board authority under the Paperwork Reduction Act (PRA) to approve of and assign OMB control numbers to collection of information requests and requirements conducted or sponsored by the Board. Board-approved collections of information are incorporated into the official OMB inventory of currently approved collections of information. Copies of the Paperwork Reduction Act Submission, supporting statements and approved collection of information instrument(s) are placed into OMB’s public docket files. The Board may not conduct or sponsor, and the respondent is not required to respond to, an information collection that has been extended, revised, or implemented on or after October 1, 1995, unless it displays a currently valid OMB control number.

Final approval under OMB delegated authority of the extension for three years, with revision, of the following report:


Agency form number: FR 2420. OMB control number: 7100–0357.

Effective Date: October 1, 2018.

Frequency: Daily.

Respondents: Commercial banks, savings associations, U.S. branches and agencies of foreign banks, International Banking Facilities, and significant banking organizations representing entities actively participating in the federal funds and/or other money markets.

Estimated number of respondents: 99 commercial banks and savings associations, 84 U.S. branches and agencies of foreign banks, 2 International Banking Facilities, and 1 significant banking organization.

Estimated average hours per response: 1.8 commercial banks and savings associations, 1.8 U.S. branches and agencies of foreign banks, 1.0 International Banking Facilities, and 1.8 significant banking organization.

Estimated annual burden hours: 44,550 commercial banks and savings associations, 37,800 U.S. branches and agencies of foreign banks, 2,050 International Banking Facilities, and 450 significant banking organization.

General description of report: The FR 2420 is a transaction-based report that collects daily liability data on federal funds purchased, selected borrowings from non-exempt entities, Eurodollar transactions, and time deposits and certificates of deposits (CDs) from (1) domestically chartered commercial banks and savings associations that have $18 billion or more in total assets as well as those that have total assets above $5 billion but less than $18 billion and meet the activity threshold, (2) U.S. branches and agencies of foreign banks with total third-party assets of $2.5 billion or more, and (3) significant banking organizations that are active participants in money markets.¹ The FR 2420 also collects daily data on Eurodollar transactions from International Banking Facilities (IBFs) of the above-referenced institutions. The FR 2420 data are used in the publication of the effective federal funds rate (EFFR) and overnight bank funding rate (OBFR) and in analysis of current money market conditions.

Legal authorization and confidentiality: The FR 2420 is authorized by section 11(a)(2) of the Federal Reserve Act, which authorizes the Board to require depository institutions to make such reports of their liabilities and assets as the Board may determine to be necessary or desirable to enable the Board to discharge its responsibility to monitor and control monetary and credit aggregates (12 U.S.C. 248(a)(2)). The FR 2420 is also authorized pursuant to section 7(c)(2) of the International Banking Act (IBA), which provides that Federal branches and agencies of foreign banks are subject to section 11(a) of the Federal Reserve Act as if they were a state member bank (12 U.S.C. 3105(c)(2)). Section 7(c)(2) of the IBA also provides that state-licensed branches and agencies of foreign banks are subject to the requirement in section 9 of the Federal Reserve Act that they file reports of condition with the appropriate Federal Reserve Bank (12 U.S.C. 324). The obligation to comply with the reporting requirements of FR 2420 is mandatory.

The individual financial institution information provided by each respondent would not be otherwise available to the public. The proposed revisions, as well as information currently collected, would be accorded confidential treatment under the

¹ A selected borrowing from a non-exempt entity is an unsecured borrowing (an unsecured primary obligation undertaken by the reporting institution as a means of obtaining funds) in U.S. dollars from a counterparty that is a non-exempt entity as derived from Regulation D, Section 204.2(a)(vii).
authority of exemption 4 of the Freedom of Information Act (5 U.S.C. 552(b)(4)). Exemption 4 protects from disclosure trade secrets and privileged or confidential commercial or financial information.

Current actions: On May 18, 2018, the Board published a notice in the Federal Register (83 FR 23276) requesting public comment for 60 days on the extension, with revision, of the FR 2420. The Board proposes to revise the FR 2420 by adding Selected Deposits (Part D) and removing Selected Borrowings from Non-Exempt Entities (Part AA). Other minor edits in the reporting instructions are proposed to improve clarity. The first report for the proposed revisions to FR 2420 would be as of October 1, 2018. The comment period for this notice expired on July 17, 2018. The Board received one comment from a government entity supporting the continued collection of data on the FR 2420. The revisions will be implemented as proposed.


Ann Misback,
Secretary of the Board.

[FR Doc. 2018–17670 Filed 8–15–18; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA—2018–N–2970]

Agency Information Collection Activities; Proposed Collection; Comment Request; Surveys and Interviews With Investigational New Drug Sponsors To Assess Current Communication Practices With Food and Drug Administration Review Staff Under the Sixth Authorization of the Prescription Drug User Fee Act

AGENCY: Food and Drug Administration, HHS.

ACTIONS: Notice.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (PRA), Federal Agencies are required to publish notice in the Federal Register concerning each proposed collection of information and to allow 60 days for public comment in response to the notice. This notice solicits comments on a proposed information collection involving surveys and interviews of sponsors of commercial investigational new drugs (INDs) to obtain feedback about communication practices with FDA review staff.

DATES: Submit either electronic or written comments on the collection of information by October 15, 2018.

ADDRESSES: You may submit comments as follows. Please note that late, untimely filed comments will not be considered. Electronic comments must be submitted on or before October 15, 2018. The https://www.regulations.gov electronic filing system will accept comments until midnight Eastern Time at the end of October 15, 2018. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are postmarked or the delivery service acceptance receipt is on or before that date.

Electronic Submissions

Submit electronic comments in the following way:

• Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on https://www.regulations.gov.

• If you want to submit a comment with confidential information that you do not wish to be made public, you must identify this information as “confidential” and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).

Written/Paper Submissions

Submit written/paper submissions as follows:

• Mail/Hand Delivery/Courier (for written/paper submissions): Dockets Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

• For written/paper comments submitted to the Dockets Management Staff, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

Instructions: All submissions received must include the Docket No. FDA—2018–N–2970 for “Surveys and Interviews with Investigational New Drug (IND) Sponsors to Assess Current Communication Practices with FDA Review Staff under the Sixth Authorization of the Prescription Drug User Fee Act (PDUFA VI).” Received comments, those filed in a timely manner (see ADDRESSES), will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at https://www.regulations.gov or at the Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday.

• Confidential Submissions—To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states “THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION.” The Agency will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on https://www.regulations.gov. Submit both copies to the Dockets Management Staff. If you do not wish your name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and you must identify this information as “confidential.” Any information marked as “confidential” will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA’s posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: https://www.gpo.gov/fdsys/pkg/FR-2015-09-18/pdf/2015-23389.pdf.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to https://www.regulations.gov and insert the docket number, found in brackets in the heading of this document, into the “Search” box and follow the prompts and/or go to the Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.