FEDERAL RESERVE SYSTEM

Proposed Agency Information Collection Activities; Comment Request

AGENCY: Board of Governors of the Federal Reserve System.

ACTION: Notice, request for comment.

SUMMARY: The Board of Governors of the Federal Reserve System (Board) invites comment on a proposal to extend for three years, without revision, the mandatory Reporting, Recordkeeping, and Disclosure Requirements Associated with the Guidance on Response Programs for Unauthorized Access to Customer Information (FR 4100; OMB No. 7100–0309).

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. In exercising this delegated authority, the Board is directed to take every reasonable step to solicit comment. In determining whether to approve a collection of information, the Board will consider all comments received from the public and other agencies.

DATES: Comments must be submitted on or before November 13, 2017.

ADDRESSES: You may submit comments, identified by FR 4100, by any of the following methods:

• Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments.
• Email: regs.comments@federalreserve.gov. Include OMB number in the subject line of the message.
• Fax: (202) 452–3819 or (202) 452–3102.
• Mail: Ann E. Misback, Secretary, Board of Governors of the Federal Reserve System, 20th Street and Constitution Avenue NW., Washington, DC 20551.

All public comments are available from the Board’s Web site at http://www.federalreserve.gov/apps/foia/proposedregs.aspx as submitted, unless modified for technical reasons. Accordingly, your comments will not be edited to remove any identifying or contact information. Public comments may also be viewed electronically or in paper form in Room 3515, 1801 K Street (between 18th and 19th Streets NW.) Washington, DC 20006 between 9:00 a.m. and 5:00 p.m. on weekdays.

Additionally, commenters may send a copy of their comments to the OMB Desk Officer—Shagufta Ahmed—Office of Information and Regulatory Affairs, Office of Management and Budget, New Executive Office Building, Room 10235, 725 17th Street NW., Washington, DC 20503 or by fax to (202) 395–6974.

FOR FURTHER INFORMATION CONTACT: A copy of the PRA OMB submission, including the proposed reporting form and instructions, supporting statement, and other documentation will be placed into OMB’s public docket files, once approved. These documents will also be made available on the Federal Reserve Board’s public Web site: http://www.federalreserve.gov/apps/reportforms/review.aspx or may be requested from the agency clearance officer, whose name appears below.


SUPPLEMENTARY INFORMATION:

Request for Comment on Information Collection Proposal

The Board invites public comment on the following information collection, which is being reviewed under authority delegated by the OMB under the PRA. Comments are invited on the following:

a. Whether the proposed collection of information is necessary for the proper performance of the Federal Reserve’s functions; including whether the information has practical utility;

b. The accuracy of the Federal Reserve’s estimate of the burden of the proposed 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;

d. Ways to minimize the burden of information collection on respondents, including through the use of automated collection techniques or other forms of information technology; and

e. Estimates of capital or startup costs and costs of operation, maintenance, and purchase of services to provide information.

At the end of the comment period, the comments and recommendations received will be analyzed to determine the extent to which the Federal Reserve should modify the proposal prior to giving final approval.

Proposal to approve under OMB delegated authority the extension for three years, without revision, of the following report:

Report title: Reporting, Recordkeeping, and Disclosure Requirements Associated with the Guidance on Response Programs for Unauthorized Access to Customer Information

Agency form number: FR 4100.

OMB control number: 7100–0309.

Frequency: On occasion.

Respondents: State member banks, bank holding companies, affiliates and certain non-bank subsidiaries of bank holding companies, uninsured state agencies and branches of foreign banks, commercial lending companies owned or controlled by foreign banks, and Edge and agreement corporations.

Estimated number of respondents:

Develop response program: 1; Incident notification: 412.

Estimated average hours per response:

Develop response program: 24; Incident notification: 36.

Estimated annual burden hours:

Develop response program: 24; Incident notification: 14,832.

General description of report: The ID-Theft Guidance is the information collection associated with the Interagency Guidance on Response Programs for Unauthorized Access to Customer Information and Customer Notice (security guidelines), which was published in the Federal Register in March 2005. Trends in customer information theft and the accompanying misuse of that information led to the issuance of these security guidelines applicable to financial institutions. The security guidelines are designed to facilitate timely and relevant notification to affected customers and the appropriate regulatory authority of the financial institutions. The security guidelines provide specific direction regarding the development of response programs and customer notifications.

Legal authorization and confidentiality: The Board has determined that the reporting, recordkeeping, and disclosure requirements associated with the FR 4100 are authorized by the Gramm-Leach-Bliley Act and are mandatory (15 U.S.C. 6801(b)). Since the FR 4100 provides that a financial institution regulated by the Board should notify its designated Reserve Bank upon becoming aware of an incident of

See 70 FR 15736.
unauthorized access to sensitive customer information, issues of confidentiality may arise if the Board were to obtain a copy of a customer notice during the course of an examination, a copy of a SAR, or other sensitive customer information. In such cases, the information would likely be exempt from disclosure to the public under the Freedom of Information Act (5 U.S.C. 552(b)(3), (4), (6), and (8)). Also, a federal employee is prohibited by law from disclosing a SAR or the existence of a SAR (31 U.S.C. 5318(g)).


Ann E. Misback, Secretary of the Board.

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FEDERAL RESERVE SYSTEM

Formations of, Acquisitions by, and Mergers of Bank Holding Companies

The companies listed in this notice have applied to the Board for approval, pursuant to the Bank Holding Company Act of 1956 (12 U.S.C. 1841 et seq.) (BHC Act), Regulation Y (12 CFR part 225), and all other applicable statutes and regulations to become a bank holding company and/or to acquire the assets or the ownership of, control of, or the power to vote shares of a bank or bank holding company and all of the banks and nonbanking companies owned by the bank holding company, including the companies listed below.

The applications listed below, as well as other related filings required by the Board, are available for immediate inspection at the Federal Reserve Bank indicated. The applications will also be available for inspection at the offices of the Board of Governors. Interested persons may express their views in writing on the standards enumerated in the BHC Act (12 U.S.C. 1842(c)). If the proposal also involves the acquisition of a nonbanking company, the review also includes whether the acquisition of the nonbanking company complies with the standards in section 4 of the BHC Act (12 U.S.C. 1843). Unless otherwise noted, nonbanking activities will be conducted throughout the United States.

Unless otherwise noted, comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors not later than October 5, 2017.

A. Federal Reserve Bank of Kansas City (Dennis Denney, Assistant Vice President) 1 Memorial Drive, Kansas City, Missouri 64198–0001:

1. Central Bancshares, Inc. to acquire, through its newly formed subsidiaries, CBI Midco, Inc. and CBI Merger Sub, Inc., all of Cambridge, Nebraska, up to 100 percent of the voting shares of Republic Corporation, and thereby indirectly acquire United Republic Bank, both of Omaha, Nebraska.

In connection with this application CBI Midco, Inc. and CBI Merger Sub, Inc., have applied to become bank holding companies.

B. Federal Reserve Bank of San Francisco (Gerald C. Tsai, Director, Applications and Enforcement) 101 Market Street, San Francisco, California 94105–1579:

1. Pacific Premier Bancorp, Inc.; to acquire 100 percent of Plaza Bancorp, and thereby indirectly acquire Plaza Bank, all of Irvine, California.


Yao-Chin Chao, Assistant Secretary of the Board.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA—2017–N–4765]

Center for Devices and Radiological Health Premarket Approval Application Critical to Quality Pilot Program

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration’s (FDA or Agency or we) Center for Devices and Radiological Health (CDRH or Center), Office of Compliance (OC) and Office of In Vitro Diagnostics and Radiological Health (OIR) is announcing its Premarket Approval Application Critical to Quality (PMA CtQ) pilot program. Participation in the PMA CtQ pilot program is voluntary and the program aims to evaluate device design and manufacturing process quality information early on to assist FDA in its review of the PMA manufacturing section and post-approval inspections. This voluntary pilot program is part of the FDA’s ongoing Case for Quality effort to apply innovative strategies that promote medical device quality and is a joint effort between the FDA’s CDRH and Office of Regulatory Affairs (ORA).

The pilot program is intended to provide qualifying PMA applicants with the option to engage FDA on development of CQ controls for their device and forego the standard PMA preapproval inspection. FDA would in turn, focus on the PMA applicant’s implementation of the CQ controls during a postmarket inspection.

DATES: FDA is seeking participation in the voluntary PMA CtQ pilot program starting from September 29, 2017. See the “Participation” section for instructions on how to submit a request to participate. This pilot program will run from September 29, 2017, to December 31, 2018. The voluntary PMA CtQ pilot program will accept the first nine participants with submissions that meet the acceptance criteria.

ADDRESSES: You may submit comments as follows:

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 available to the public, submit the comment as a written/paper submission 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–