FEDERAL RESERVE SYSTEM

Formations of, Acquisitions by, and Mergers of Savings and Loan Holding Companies

The companies listed in this notice have applied to the Board for approval, pursuant to the Home Owners’ Loan Act (12 U.S.C. 1461 et seq.) (HOLA), Regulation LL (12 CFR part 238), and Regulation MM (12 CFR part 239), and all other applicable statutes and regulations to become a savings and loan holding company and/or to acquire the holding of, control of, or the power to vote shares of a savings association and nonbanking companies owned by the savings and loan 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 application also will 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 HOLA (12 U.S.C. 1467a(e)). 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 10(c)(4)(B) of the HOLA (12 U.S.C. 1467a(c)(4)(B)). 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 April 19, 2013.

A. Federal Reserve Bank of Philadelphia (William Lang, Senior Vice President) 100 North 6th Street, Philadelphia, Pennsylvania 19105–1521:

1. Sunnyside Bancorp, Inc., Irvington, New York, to become a savings and loan holding company by acquiring 100 percent of the voting shares of Sunnyside Federal Savings and Loan Association of Irvington, Irvington, New York, following it conversion from a federally chartered mutual savings and loan association to a federally chartered stock savings and loan association.


Michael J. Lewandowski, Assistant Secretary of the Board.

[FR Doc. 2013–06676 Filed 3–22–13; 8:45 am]

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GENERAL SERVICES ADMINISTRATION

[OMB Control No. 3090–0288; Docket 2013–0001, Sequence 2]

Information Collection; Open Government Citizen Engagement Ratings, Rankings, and Flagging

AGENCY: Office of Citizen Services, General Services Administration (GSA).

ACTION: Notice of request for comments regarding an extension of an existing OMB information collection.

SUMMARY: Under the provisions of the Paperwork Reduction Act, the Regulatory Secretariat will be submitting to the Office of Management and Budget (OMB) a request to review and approve a previously approved information collection requirement regarding open government citizen engagement ratings, rankings, and flagging.

DATES: Comments must be submitted on or before May 24, 2013.

ADDRESSES: Submit comments identified by Information Collection 3090–0288, Open Government Citizen Engagement Ratings, Rankings, and Flagging, by any of the following methods:

• Regulations.gov: http://www.regulations.gov. Submit comments via the Federal eRulemaking portal by searching for the OMB control number. Select the link “Submit a Comment” that corresponds with “Information Collection 3090–0288, Open Government Citizen Engagement Ratings, Rankings, and Flagging”. Follow the instructions provided at the “Submit a Comment” screen. Please include your name, company name (if any), and “Information Collection 3090–0288, Open Government Citizen Engagement Ratings, Rankings, and Flagging” on your attached document.

• Mail: General Services Administration, Regulatory Secretariat (MVCB), 1275 First Street NE., Washington, DC 20417; ATTN: Hada Flowers/IC 3090–0288, Open Government Citizen Engagement Ratings, Rankings, and Flagging.

B. Public Comments Are Particularly Invited On

Pursuant to section 3506(c)(2)(A) of the PRA, GSA specifically solicits comments and information to enable it to:

(i) 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;

(ii) Evaluate the accuracy of the Agency’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;

(iii) Enhance the quality, utility, and clarity of the information to be collected; and
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2013–N–0033]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Recordkeeping Requirements for Microbiological Testing and Corrective Measures for Bottled Water

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

DATES: Fax written comments on the collection of information by April 24, 2013.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202–395–7285, or emailed to oira_submission@omb.eop.gov. All comments should be identified with the OMB control number 0910–0658.

FOR FURTHER INFORMATION CONTACT: Domini Bean, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., P150–400T, Rockville, MD 20850, 301–796–5733, domini.bean@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Recordkeeping Requirements for Microbiological Testing and Corrective Measures for Bottled Water—21 CFR 129.35(a)(3)(i), 129.80(g), and 129.80(h) (OMB Control Number 0910–0658)—Extension

The bottled water regulations in parts 129 and 165 (21 CFR parts 129 and 165) require that if any coliform organisms are detected in weekly total coliform testing of finished bottled water, followup testing must be conducted to determine whether any of the coliform organisms are Escherichia coli. The adulteration provision of the bottled water standard (§ 165.110(d)) provides that a finished product that tests positive for E. coli will be deemed adulterated under section 402(a)(3) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 342(a)(3)). In addition, the current good manufacturing practice (CGMP) regulations for bottled water in part 129 require that source water from other than a public water system (PWS) be tested at least weekly for total coliform. If any coliform organisms are detected in the source water, the bottled water manufacturers are required to determine whether any of the coliform organisms are E. coli. Source water found to contain E. coli is not considered water of a safe, sanitary quality and would be unsuitable for bottled water production. Before a bottler may use source water from a source that has tested positive for E. coli, a bottler must take appropriate measures to rectify or otherwise eliminate the cause of the contamination. A source previously found to contain E. coli will be considered negative for E. coli after five samples collected over a 24-hour period from the same sampling site are tested and found to be E. coli negative.

The respondents to this information collection are domestic and foreign bottled water manufacturers that sell bottled water in the United States.

In the Federal Register of January 18, 2013 (78 FR 4152), FDA published a 60-day notice requesting public comment on the proposed extension of this collection of information. FDA received two letters in response to the notice, which contained multiple comments.

One comment suggested that laboratory quality assurance practices should be required for the testing of bottled water. FDA’s CGMP regulations for bottled water in 21 CFR 129 do not specifically require laboratory quality assurance practices, and FDA does not have the specific statutory authority to require bottlers to use certified laboratories for water quality tests. However, the CGMP regulations for source water testing do require that “[t]est and sample methods shall be those recognized and approved by the government agency or agencies having jurisdiction over the approval of the water source, and shall be consistent with the minimum [standard of quality] requirements set forth in § 165.110(b) of this chapter” (§ 129.35(a)(3)(ii)). The CGMP regulations also state that


(iv) 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.

C. Annual Reporting Burden

The annual public reporting and recordkeeping burden for this collection of information is estimated to average up to 1,666 hours per year. Burden means the total time, effort, or financial resources expended by persons to generate, maintain, retain, or disclose or provide information to or for a Federal agency. This includes the time needed to review instructions; develop, acquire, install, and utilize technology and systems for the purposes of collecting, validating, and verifying information, processing and maintaining information, and disclosing and providing information; adjust the existing ways to comply with any previously applicable instructions and requirements which have subsequently changed; train personnel to be able to respond to a collection of information; search data sources; complete and transmit or otherwise disclose the information.

The estimated annual burden:

Respondents: 1,200,000.

Responses per Respondent: 1.

Total number of responses: 1,200,000.

Hours per Response:.0013886.

Total hours per response: 1,200,000.

Total Burden Hours: 1,666.

Obtaining Copies Of Proposals:

Requesters may obtain a copy of the information and regulatory affairs (ira) 5733, duplicating services, Regulatory Secretariat (MVCB), 1275 First Street NE., Washington, DC 20417, telephone 202–501–4755. Please cite 5733, 400T, Rockville, MD 20850, 301–796–5733, domini.bean@fda.hhs.gov.

Dated: March 19, 2013.

Casey Coleman,

Chief Information Officer.