Synopsis: The amendment would add Evergreen Line Joint Service Agreement as a non-shareholder party to the agreement.

Agreement No.: 012048.

Title: The Container Trades Statistics Agreement.


Synopsis: The agreement authorizes the parties to gather, compile, aggregate, exchange, and disseminate demand and supply forecasts, a volume database, and a price index relating to the trade in containers. It also establishes a forum for the parties to meet and discuss such data. By Order of the Federal Maritime Commission.


Karen V. Gregory,
Assistant Secretary.
[FR Doc. E8–20958 Filed 9–9–08; 8:45 am]
BILLING CODE 6730–01–P

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 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 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. Additional information on all bank holding companies may be obtained from the National Information Center website at www.ffiec.gov/nic/

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 3, 2008.

A. Federal Reserve Bank of St. Louis

(1) Heritage Bancorp, Inc., Mason, Tennessee; to directly acquire 2.12 percent, for total direct and indirect ownership of 51.77 percent, of Mason Bancorp, Inc., Mason, Tennessee, and thereby indirectly acquire The Bank of Mason, Mason, Tennessee.


Robert deV. Frierson,
Deputy Secretary of the Board.
[FR Doc. E8–20958 Filed 9–9–08; 8:45 am]
BILLING CODE 6210–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day–08–0494]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call 404–439–5960 and send comments to Maryam I. Daneshvar, CDC Acting Reports Clearance Officer, 1600 Clifton Road, MS–D74, Atlanta, GA 30333 or send an e-mail to omb@cdc.gov.

Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency’s estimate of the burden of the proposed collection of information; (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. Written comments should be received within 60 days of this notice.

Proposed Project

Exposure to Aerosolized Brevetoxins during Red Tide Events (OMB No. 0920–0494)—Revision—National Center for Environmental Health (NCEH), Centers for Disease Control and Prevention (CDC.)

Background and Brief Description

Karenia brevis (formerly Gymnodinium breve) is the marine dinoflagellate responsible for extensive blooms (called Florida red tides) that forms in the Gulf of Mexico. K. brevis produces potent toxins, called brevetoxins, which have been responsible for killing millions of fish and other marine organisms. The biochemical activity of brevetoxins is not completely understood and there is still little information regarding human health effects from environmental exposures, such as inhaling brevetoxin that has been aerosolized and swept onto the coast by offshore winds. The National Center for Environmental Health (NCEH), Centers for Disease Control and Prevention (CDC) has recruited people who work along the coast of Florida and who are periodically occupationally exposed to aerosolized red tide toxins.

NCEH administered a baseline respiratory health survey and conducted pre- and post-shift pulmonary function tests (PFTs) during a time when there is no red tide reported near the area. When a red tide developed, NCEH administered a symptom survey and conducted PFTs. NCEH compared symptoms reported before the shift with symptoms reported after the shift. NCEH also examined changes in PFT test results (post-shift values compared to pre-shift values). NCEH did these comparisons during a time when there was no red tide and during a time when
there was a red tide and then examined the data to see if red tide exposure had an effect on symptom reports or PFT results.

NCEH requests a revision of data collection procedures for the currently approved project and an additional three year extension. Unfortunately, the exposures experienced by the study cohort have been minimal, and NCEH plans to conduct another study (using the same symptom surveys and PFTs) during a more severe red tide event.

First, NCEH wants to quantify the levels of cytokines in nasal exudates to assess whether they can be used to verify exposure and to demonstrate a biological effect (i.e., allergic response) following inhalation of aerosolized brevetoxins. NCEH will collect nasal exudates at the same time the PFTs are done.

NCEH plans to include the study subjects who have been involved in the earlier studies and any new individuals (n=25) who have been hired to work at the relevant beaches.

There is no cost to respondents other than their time.

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**ESTIMATED ANNUALIZED BURDEN HOURS**

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Disease Control and Prevention**

**Ethics Subcommittee, Advisory Committee to the Director, Centers for Disease Control and Prevention (CDC); Meeting**

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention announce the following Subcommittee meeting.

**Name:** Ethics Subcommittee, Advisory Committee to the Director.

**Time and Date:** 12 p.m.–1:30 p.m., EDT, Thursday, September 25, 2008.

**Place:** This meeting will be held by conference call. The call in number is (866) 919–3560 and entering code 4168828.

**Status:** Open to the public. The public is welcome to comment during the public comment period which is tentatively scheduled from 1 p.m.–1:15 p.m.

**Purpose:** The Ethics Subcommittee will provide counsel to the ACD, CDC regarding a broad range of public health ethics questions and issues arising from programs, scientists and practitioners.

**Matters To Be Discussed:** Agenda items will include review of ethics guidance for public health emergency preparedness and response.

**For Further Information Contact:** For more information about this meeting contact Drue Barrett, PhD, Designated Federal Official, Ethics Subcommittee, CDC, 1600 Clifton Road, NE., M/S D–50, Atlanta, Georgia 30333. Telephone (404)639–4690, e-mail: dbarrett@cdc.gov.

The Director, Management Analysis and Services Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.


Elaine L. Baker, Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. E8–20967 Filed 9–9–08; 8:45 am]

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

**[Docket No. FDA–2008–N–0239]**

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Regulations for In Vivo Radiopharmaceuticals Used for Diagnosis and Monitoring

**AGENCY:** Food and Drug Administration, HHSS.

**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 October 10, 2008.

**ADDRESS:** 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: 202–395–6974, or e-mailed to bagula@omb.eop.gov. All comments should be identified with the OMB control number 0910–0409. Also include the FDA docket number found in brackets in the heading of this document.

**FOR FURTHER INFORMATION CONTACT:**

Elizabeth Berbakos, Office of Information Management (HFA–710), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–796–3792.

**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.

**Regulations for In Vivo Radiopharmaceuticals Used for Diagnosis and Monitoring (OMB Control Number 0910–0409)—Extension**

FDA is requesting OMB approval of the information collection requirements contained in 21 CFR 315.4, 315.5, and 315.6. These regulations require manufacturers of diagnostic radiopharmaceuticals to submit information that demonstrates the safety and effectiveness of a new diagnostic radiopharmaceutical or of a new indication for use of an approved diagnostic radiopharmaceutical.

In response to the requirements of section 122 of the Food and Drug Administration Modernization Act of 1997 (Public Law 105–115), FDA published a final rule in the *Federal Register*...