FEDERAL RESERVE SYSTEM

Change in Bank Control Notices; Acquisitions of Shares of a Bank or Bank Holding Company Correction

This notice corrects a notice (FR DOC #2012–14579) published on page 35680 of this issue for Thursday, June 14, 2012.

Under the Federal Reserve Bank of Atlanta heading, the entry for Robert Roschman and the Robert Roschman Revocable Trust, Robert Roschman, trustee, all of Fort Lauderdale, Florida, is revised to read as follows:

A. Federal Reserve Bank of Atlanta (Chapelle Davis, Assistant Vice President) 1000 Peachtree Street NE., Atlanta, Georgia 30309:

1. Robert Roschman Revocable Trust, Robert Roschman, trustee; the Lorrie Lei Roschman Revocable Trust, Lorrie Roschman, trustee; the Revocable Trust Created by Jeffrey S. Roschman, Jeffrey Roschman, trustee; CT Foundation, Betty Roschman, Roschman Restaurant Administration, and Kerry Roschman, all of Fort Lauderdale, Florida, to collectively retain 25 percent or more of the shares and thereby control of Giant Holdings, Inc., and Landmark Bank, N.A., both of Fort Lauderdale, Florida.

Comments on this application must be received by August 1, 2012.


Robert deV. Frierson,
Deputy Secretary of the Board.

[FR Doc. 2012–16984 Filed 7–11–12; 8:45 am]
BILLING CODE 6210–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Clinical Laboratory Improvement Advisory Committee (CLIAC)

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

Times and Dates:
August 29, 2012, 8:30 a.m.–12:30 p.m.,
August 30, 2012.

Place: CDC, 1600 Clifton Road NE.,
Tom Harkin Global Communications Center, Building 19, Room 232, Auditorium B, Atlanta, Georgia 30333.

Online Registration Required: All CLIAC attendees are required to register for the meeting online at least 5 business days in advance for U.S. citizens and at least 10 business days in advance for international registrants. Register at http://www.cdc.gov/cliac/default.aspx by scrolling down and clicking the appropriate link under “Meeting Registration” (either U.S. Citizen Registration or Non-U.S. Citizen Registration) and completing all forms according to the instructions given. Please complete all the required fields before submitting your registration and submit no later than August 22, 2012, for U.S. registrants and August 15, 2012, for international registrants.

Status: Open to the public, limited only by the space available. The meeting room accommodates approximately 100 people.

Purpose: This Committee is charged with providing scientific and technical advice and guidance to the Secretary,
Department of Health and Human Services; the Assistant Secretary for Health; the Director, CDC; the Commissioner, Food and Drug Administration (FDA); and the Administrator, Centers for Medicare and Medicaid Services (CMS). The advice and guidance pertain to general issues related to improvement in clinical laboratory quality and laboratory medicine practice and specific questions related to possible revision of the CLIAC standards. Examples include providing guidance on studies designed to improve safety, efficiency, timeliness, equity, and patient-centeredness of laboratory services; revisions to the standards under which clinical laboratories are regulated; the impact of proposed revisions to the standards on medical and laboratory practice; and the modification of the standards and provision of non-regulatory guidelines to accommodate technological advances, such as new test methods and the electronic transmission of laboratory information.

Matters To Be Discussed: The agenda will include agency updates from the CDC, the CMS, and the FDA; and presentations and discussions addressing activities of the Clinical Laboratory Integration into Health Care Collaborative (CLIHC); the Laboratory Medicine Best Practices (LMBP) Initiative; the Communication in Informatics Workgroup; and the topic of usability of electronic health records. Also discussed will be the potential need for educational materials and resources for sites that test under a Provider-performed Microscopy Certificate; and the increased use of culture-independent microbiology diagnostics and the impact on public health.

Agenda items are subject to change as priorities dictate.

Providing Oral or Written Comments: It is the policy of CLIAC to accept written public comments and provide a brief period for oral public comments whenever possible.

Oral Comments: In general, each individual or group requesting to make an oral presentation will be limited to a total of five minutes (unless otherwise indicated). Speakers must also submit their comments in writing for inclusion in the meeting’s Summary Report. To assure adequate time is scheduled for public comments, individuals or groups planning to make an oral presentation should, when possible, notify the contact person below at least one week prior to the meeting date. Written Comments: For individuals or groups unable to attend the meeting, CLIAC accepts written comments until the date of the meeting (unless otherwise stated). However, it is requested that comments be submitted at least one week prior to the meeting date so that the comments may be made available to the Committee for their consideration and public distribution. Written comments, one hard copy with original signature, should be provided to the contact person below. Written comments will be included in the meeting’s Summary Report.

Availability of Meeting Materials: To support the green initiatives of the federal government, the CLIAC meeting materials will be made available to the Committee and the public in electronic format (PDF) on the internet instead of by printed copy. Refer to the CLIAC Web site on the day of the meeting for materials. http://www.cdc.gov/cliac/cliac_meeting_all_documents.aspx.

Note: If using a mobile device to access the materials, please verify the device’s browser is able to download the files from the CDC’s Web site before the meeting. Alternatively, the files can be downloaded to a computer and then emailed to the portable device. An internet connection, power source and limited hard copies may be available at the meeting location, but cannot be guaranteed.

Contact Person for Additional Information: Nancy Anderson, Chief, Laboratory Practice Standards Branch, Division of Laboratory Science and Standards, Laboratory Science, Policy and Practice Program Office, Office of Surveillance, Epidemiology and Laboratory Services, Centers for Disease Control and Prevention, 1600 Clifton Road NE., Mailstop F–11, Atlanta, Georgia 30333; telephone (404) 498–2741; fax (404) 498–2219; or via email at NAnderson@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.

Dated: July 2, 2012.

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

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and Prevention

Subcommittee for Dose Reconstruction Reviews (SDRR), Advisory Board on Radiation and Worker Health (ABRW or the Advisory Board), National Institute for Occupational Safety and Health (NIOSH)

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

Time and Date: 8:30 a.m.–5 p.m., Eastern Time, August 6, 2012.

Place: Cincinnati Airport Marriott, 2395 Progress Drive, Hebron, Kentucky 41018, Telephone: (859) 334–4611, Fax: (859) 334–4619.

Status: Open to the public, but without an oral public comment period. To access by conference call dial the following information: 1 (866) 659–0537, Participant Pass Code 9933701.

Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Key functions of the Advisory Board include providing advice on the development of probability of causation guidelines that have been promulgated by the Department of Health and Human Services (HHS) as a final rule; advice on methods of dose reconstruction, which have also been promulgated by HHS as a final rule; advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program; and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC).

In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to CDC. NIOSH implements this responsibility for CDC. The charter was issued on August 3, 2001, renewed at appropriate intervals, and will expire on August 3, 2013.

Purpose: The Advisory Board is charged with (a) Providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the scientific...