1. Jan Malcolm Jones, Jr., and Leslie Ann Jones, both in Jacksonville, Florida; to retain outstanding voting shares of Florida Capital Group, Inc., and thereby indirectly retain outstanding voting shares of Florida Capital Bank, both in Jacksonville, Florida.

Board of Governors of the Federal Reserve System, December 22, 2010.

Robert deV. Frierson,

Deputy Secretary of the Board.
[FR Doc. 2010–32651 Filed 12–27–10; 8:45 am]
BILLING CODE 6210–01–P

FEDERAL RESERVE SYSTEM

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

This notice corrects a notice (FR Doc. 2010–32092) published on page 80501 of the issue for Wednesday, December 22, 2010.

Under the Federal Reserve Bank of New York heading, the entry for Chuo Mitsui Trust Holding, Inc., Tokyo, Japan, is revised to read as follows:

A. Federal Reserve Bank of New York (Ivan Hurwitz, Vice President) 33 Liberty Street, New York, New York 10045–0001:

1. Chuo Mitsui Trust Holdings, Inc., Tokyo, Japan; to become a bank holding Company by acquiring The Sumitomo Trust and Banking Co., Ltd, Osaka, Japan, and thereby acquire Sumitomo Trust and Banking Co. (USA), Hoboken, New Jersey.

In connection with this application, Applicant also has applied to acquire Nikko Americas Holding Co., Inc., Nikko Asset Management Americas, Inc., and Chuo Mitsui Investments, all in New York, New York, and thereby engage in investment advisory activities, pursuant to section 225.24(b)(6) of Regulation Y.

Comments on this application must be received by January 18, 2011.

Board of Governors of the Federal Reserve System, December 22, 2010.

Robert deV. Frierson,

Deputy Secretary of the Board.
[FR Doc. 2010–32593 Filed 12–27–10; 8:45 am]
BILLING CODE 6210–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Submission for OMB Review; Comment Request; National Evaluation of the Clinical and Translational Science Awards (CTSA) Initiative

SUMMARY: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Center for Research Resources, the National Institutes of Health has submitted to the Office of Management and Budget (OMB) a request to review and approve the information collection listed below. This proposed information collection was previously published in the Federal Register on October 12, 2010, pages 62543-62544, and allowed 60-days for public comment. No comments were received. The purpose of this notice is to allow an additional 30 days for public comment. The National Institutes of Health 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. Proposed Collection:

Title: The National Evaluation of the Clinical and Translational Science Awards (CTSA) Initiative.

Type of Information Collection Request: New.

Need and Use of Information Collection: The CTSA Initiative is directed at transforming the way biomedical research is conducted nationwide by reducing the time it takes

for basic science or laboratory discoveries to become treatments for patients, and for those treatments in turn to be incorporated and disseminated throughout community practice. The primary purpose of this data collection is to provide information about the process and early outcomes associated with 46 awardees participating in the first four cohorts of CTSA awards, in order to fulfill the congressional expectations for external program evaluation. NIH will use the results to understand the extent to which the CTSA Initiative is bringing about transformational changes in clinical and translational science among academic medical centers and their research partners, increasing the efficiency of the research process, and enhancing the capacity of the field to conduct clinical and translational research. All information collected will be used to provide analytical and policy support to NCRR, assisting NIH in making decisions about current CTSA programming, future funding, and other initiatives to improve clinical and translational science. It may also provide information for NIH's Government Performance and Results Act (GPRA) report.

Frequency of Response: Biennial. Affected Public: Individuals.

Type of Respondents: Scientific researchers. The annual reporting burden is as follows:

Estimated Number of Respondents: 3,563;

Estimated Number of Responses per Respondent: 1;

Average Burden Hours Per Response: 0.13;

Estimated Total Annual Burden Hours Requested: 451.5. The annualized cost to respondents is estimated at \$14,056. There are no capital or start-up costs, and no maintenance or service cost components to report.

Respondent type	Estimated number of respondents	Estimated number of hours per respondent type	Frequency of response	Estimated total annual burden hours requested
Users survey	500	.25	.5	62.5
Nonusers survey	500	.08	.5	20.0
Trainees/scholars survey	1,213	.33	.5	200.0
Mentors survey	1,350	.25	.5	169.0
Total				451.5

Request for Comments: Written comments and/or suggestions from the public and affected agencies should address one or more of the following points: (1) Evaluate whether the proposed collection of information is necessary for the proper performance of the function of the agency, including

whether the information will have practical utility; (2) 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; (3) Enhance the quality, utility, and clarity of the information to be collected; and (4) Minimize the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

Direct Comments to OMB: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the: Office of Management and Budget, Office of Regulatory Affairs,

OIRA_submission@omb.eop.gov or by fax to 202–395–6974, Attention: Desk Officer for NIH. To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Patricia Newman, Program Analyst, Office of Science Policy, National Center for Research Resources, 6701 Democracy Boulevard, MSC 4874, Bethesda, Maryland 20892–4874, or e-mail your request, including your address to pnewman@mail.nih.gov.

Comments Due Date: Comments regarding this information collection are best assured of having their full effect if received within 30-days of the date of this publication.

Dated: December 20, 2010.

Meryl Sufian,

Supervisory Health Science Policy Analyst, Office of Science Policy, NCRR, National Institutes of Health.

[FR Doc. 2010–32659 Filed 12–27–10; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

Findings of Misconduct in Science; Correction

AGENCY: Office of the Secretary, HHS **ACTION:** Correction of notice.

SUMMARY: This document corrects errors that appeared in the notice published in the November 29, **Federal Register** entitled "Findings of Misconduct in Science."

DATES: Effective Date: December 28, 2010.

Applicability Date: The correction notice is applicable for the Findings of Misconduct in Science notice published on November 29, 2010.

FOR FURTHER INFORMATION CONTACT: Karen Gorirossi or Sheila Fleming at

Karen Gorirossi or Sheila Fleming at 240–453–8800.

SUPPLEMENTARY INFORMATION:

I. Background

In FR Doc. 2010–29867 of November 29, 2010 (75 FR 73084–73085), there was an error, which included an incorrect date of implementation of administrative actions. The error is identified and corrected in the Correction of Errors section below.

II. Correction of Errors

In FR Doc. 2010–29867 of November 29, 2010 (75 FR 73084–73085), make the following corrections:

1. On page 73084, third column, fourth paragraph, change the paragraph to read as follows: "By letter dated October 4, 2010, the Department of Health and Human Services (HHS) notified Dr. Sezen of findings of misconduct in science made by ORI and the Department's intent to debar her for a period of five (5) years pursuant to the Public Health Service Policies on Research Misconduct, 42 CFR part 50, subpart A and part 93, and HHS Implementation (2 CFR part 376) of the Office of Management and Budget (OMB) Guidelines to Agencies on Governmentwide Debarment and Suspension (2 CFR part 180). In accordance with part 93, subpart E, Dr. Sezen was afforded 30 days within which to request a hearing in this matter. As of November 4, 2010, the period of time to request a hearing expired. Thus, the following administrative actions have been implemented for a period of five (5) years, beginning on December 13, 2010."

Dated: December 17, 2010.

John Dahlberg,

Director, Division of Research Investigations, Office of Research Integrity.

[FR Doc. 2010–32555 Filed 12–27–10; 8:45 am]

BILLING CODE 4150-31-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

[Docket No. HHS-OS-2010-0033; OCIIO-

The Consumer Operated and Oriented Plan (CO-OP) Advisory Board; Office of Consumer Information and Insurance Oversight, January 13, 2011

AGENCY: Office of Consumer Information and Insurance Oversight (OCIIO), HHS. **ACTION:** Notice of meeting.

SUMMARY: This notice announces a forthcoming meeting of an advisory

committee of the Office of Consumer Information and Insurance Oversight (OCIIO) in accordance with the Federal Advisory Committee Act. The meeting is open to the public. The purpose of the meeting is to assist and advise the Secretary and Congress through the Department of Health and Human Services' Office of Consumer Information and Insurance Oversight (OCIIO) on the Department's strategy to foster the creation of qualified nonprofit health insurance issuers. Specifically, the Committee shall advise the Secretary and Congress concerning the award of grants and loans related to Section 1322 of the Affordable Care Act. In these matters, the Committee shall consult with all components of the Department, other federal entities, and non-federal organizations, as appropriate; and examine relevant data sources to assess the grant and loan award strategy to provide recommendations to OCIIO. Notice of this meeting is given under the Federal Advisory Committee Act (Pub. L. No. 92–463, 5 U.S.C., App. 2).

DATES: Meeting Date: January 13, 2011 from 8 a.m. to 5 p.m., eastern standard time (e.s.t.).

Deadline for Meeting Registration, Presentations and Comments: January 6, 2011, 5 p.m., e.s.t.

Deadline for Requesting Special Accommodations: January 6, 2011, 5 p.m., e.s.t.

ADDRESSES: Meeting Location: Jurys Hotel, 1500 New Hampshire Ave., NW., Washington, DC 20036.

Meeting Online Access: To participate in this meeting via the Internet, go to http://www.readyshow.com/ and enter participant code 78030350.

Meeting Phone Access: To participate in this meeting via phone, please dial into the toll free phone number 1–877–366–0711, and enter the phone number password 78030350#.

Meeting Registration, Presentations, and Written Comments: Brian Chiglinsky, Office of Consumer Information and Insurance Oversight, HHS, 200 Independence Avenue, SW., Washington, DC 20201, 202–260–6090, Fax: 202–260–6108, or contact by e-mail at brian.chiglinsky@hhs.gov.

Registration: The meeting is open to the public, but attendance is limited to the space available. Persons wishing to attend this meeting must register by contacting the Analyst at the address listed in the ADDRESSES section of this notice or by telephone at number listed in the FOR FURTHER INFORMATION CONTACT section of this notice, by the date listed in the DATES section of this notice.