Board of Governors of the Federal Reserve System, September 1, 1999.

Robert deV. Frierson,

Associate Secretary of the Board. [FR Doc. 99–23290 Filed 9–7–99; 8:45 am] BILLING CODE 6210–01–F

FEDERAL RESERVE SYSTEM

Government in the Sunshine Meeting Notice

AGENCY HOLDING THE MEETING: Board of Governors of the Federal Reserve System.

TIME AND DATE: 10:00 a.m., Monday, September 13, 1999.

PLACE: Marriner S. Eccles Federal Reserve Board Building, 20th and C Streets, N.W., Washington, D.C. 20551.

STATUS: Closed.

MATTERS TO BE CONSIDERED:

1. Personnel actions (appointments, promotions, assignments, reassignments, and salary actions) involving individual Federal Reserve System employees.

2. Any items carried forward from a previously announced meeting.

CONTACT PERSON FOR MORE INFORMATION:

Lynn S. Fox, Assistant to the Board; 202–452–3204.

SUPPLEMENTARY INFORMATION: You may call 202–452–3206 beginning at approximately 5 p.m. two business days before the meeting for a recorded announcement of bank and bank holding company applications scheduled for the meeting; or you may contact the Board's Web site at http://www.federalreserve.gov for an electronic announcement that not only lists applications, but also indicates procedural and other information about the meeting.

Dated: September 3, 1999.

Robert deV. Frierson,

Associate Secretary of the Board. [FR Doc. 99–23469 Filed 9–3–99; 3:35 am] BILLING CODE 6210-01-P

GENERAL SERVICES ADMINISTRATION

Office of Communications

Standard and Optional Forms Management Office Cancellation of a Standard Form

AGENCY: General Services Administration.

ACTION: Notice.

SUMMARY: The Office of Personnel Management cancelled the need for

Standard Form 66B, Caution Personnel Record—Restricted Usage because of low usage.

FOR FURTHER INFORMATION CONTACT: Ms. Barbara Williams (202) 501–0581. DATES: Effective September 8, 1999.

Dated: August 30, 1999.

Barbara M. Williams,

Deputy Standard and Optional Forms Management Officer.

[FR Doc. 99–23256 Filed 9–7–99; 8:45 am]

GENERAL SERVICES ADMINISTRATION

Office of Communications

Stocking change of a Standard Form

AGENCY: General Services Administration.

ACTION: Notice.

SUMMARY: Because of low usage, the Department of the Treasury is not stocking the following Standard Form: SF 1198, Request by Employee for Allotment of Pay for Credit to Savings Account with a Financial Organization.

You can get this form from: Department of the Treasury—FMS, Ardmore Industrial Center, 3361–L 75th Avenue, Landover, MD 10785.

The form is also available on the internet. Address: http://www.gsa.gov/forms/forms.htm.

FOR FURTHER INFORMATION CONTACT: Mr. Irv. Wilson (202) 622–1575. This contact is for information about completing the form only.

DATES: Effective on September 8, 1999.

Dated: August 23, 1999.

Barbara M. Williams,

Deputy Standard and Optional Forms Management Officer.

[FR Doc. 99–23257 Filed 9–7–99; 8:45 am] BILLING CODE 6820–34–M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[INFO-99-35]

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 the CDC Reports Clearance Officer on (404) 639–7090.

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 for other forms of information technology. Send comments to Seleda Perryman, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: Vibrio Illness Investigation Report Form—(0920-0322)—Reinstatement—The National Center for Infectious Disease (NCID)-The purpose of the Cholera and other Vibrio Illness Investigation Report Form is to collect information on illnesses occurring as a result of infection with Vibrio species. Vibrios are important pathogens in the United States, primary septicemia, gastroenteritis, and wound infections have been associated with various species. Gastroenteritis and primary septicemia have been associated with the consumption of undercooked shellfish, particularly with raw, Gulf Coast oysters. Associations have also been linked to wound infections with exposure of broken skin to seawater. Most importantly, Vibrio cholerae 01 is the organism responsible for cholera, a severe, dehydrating diarrheal illness. Although infections with Vibrio cholerae 01 are notifiable in all states, an official report form for this illness did not previously exist. The Vibrio Illness Investigation Report Form is used to record information on all Vibrio-related illnesses, as well as more detailed information on cholera illness, which is currently a reportable disease in all states. The form has a separate, optional Seafood Investigation section to be completed when applicable. The form provides a consolidated, systematic method by which health departments can report such information, and is then used to gain a better understanding of the incidence, etiology, and epidemiology of all Vibriorelated illnesses occurring in the United States.

There is no change in the frequency of reporting or projected reporting. Most

respondents are epidemiologists or nurses in the local health department, but in some instances, infection control nurses or physicians might complete the form. The total cost per respondent is estimated at \$11.00. This is primarily salary but also includes postage and telephone calls.

Respondents	No. of respondents	No. of responses/ respondent	Avg. burden of response (in hrs.)	Total burden (in hrs.)
Local health department staff	90 45 15	1 1 1	.33 .33 .33	30 15 5
Total				50

Dated: September 1, 1999.

Nancy Cheal,

Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention (CDC).

[FR Doc. 99–23282 Filed 9–7–99; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Clinical Laboratory Improvement Advisory Committee (CLIAC) Meeting: Correction

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) announced the following committee meeting in the **Federal Register** on August 23, 1999, Volume 64, Number 162, Page 45971.

Name: Clinical Laboratory Improvement Advisory Committee (CLIAC).

Times and Dates: 8:30 a.m.–5 p.m., September 22, 1999. 8:30 a.m.–3:30 p.m., September 23, 1999.

Correction: Please note, "potential rulemaking for genetic testing" should be added to the previously published agenda.

Contact Person for Additional Information: John C. Ridderhof, Dr.P.H., Division of Laboratory Systems, Public Health Practice Program Office, CDC, 4770 Buford Highway, NE, M/S G–25, Atlanta, Georgia 30341–3724, telephone 770/488–8076, FAX 770/488–8282.

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 the both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

John C. Burckhardt,

Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 99–23382 Filed 9–3–99; 10:00 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket Nos. 91N-0101, 91N-0098, 91N-0103, and 91N-100H]

Food Labeling; Health Claims and Label Statements; Request for Scientific Data and Information

AGENCY: Food and Drug Administration, HHS

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is requesting scientific data, research study results, and other related information on four substance-disease relationships in order to reevaluate the scientific evidence for these relationships. The agency is taking this action to comply with a recent court decision in which FDA was instructed to reconsider whether to authorize health claims for these relationships in dietary supplement labeling. The four health claims to be reconsidered are: "Consumption of antioxidant vitamins may reduce the risk of certain kinds of cancer," "Consumption of fiber may reduce the risk of colorectal cancer. "Consumption of omega-3 fatty acids may reduce the risk of coronary heart disease," and "0.8 mg of folic acid in a dietary supplement is more effective in reducing the risk of neural tube defects than a lower amount in foods in common form." The agency will use the data and information to determine, for each substance-disease relationship, if an appropriate scientific basis exists to support the issuance of a proposed rule to authorize a health claim for the relationship.

DATES: Written comments by November 22, 1999.

ADDRESSES: Written comments to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT:

Christine J. Lewis, Center for Food Safety and Applied Nutrition (HFS– 451), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202–205–4168.

SUPPLEMENTARY INFORMATION: The Nutrition Labeling and Education Act of 1990 (the 1990 amendments), which amended the Federal Food, Drug, and Cosmetic Act (the act), directed the Secretary of Health and Human Services, among other things, to evaluate the scientific evidence on 10 substance-disease relationships to determine their scientific validity as the basis for health claims in food labeling. For conventional foods, the 1990 amendments state that a health claim is permitted only if FDA determines that there is significant scientific agreement among qualified experts that the claim is supported by the totality of publicly available scientific evidence, including evidence from well-designed studies conducted in a manner that is consistent with generally recognized scientific procedures and principles (section 403(r)(3)(B)(i) of the act (21 U.S.C. 343(r)(3)(B))). While the 1990 amendments allowed FDA to consider a different scientific standard for health claims for dietary supplements (section 403(r)(5)(D) of the act (21 U.S.C. 343(r)(5)(D))), FDA issued regulations in 21 CFR 101.14(c) in 1994 that applied the same standard as that used for health claims for conventional foods (59 FR 395, January 4, 1994).

FDA conducted rulemakings in which it reviewed the scientific evidence for all 10 substance-disease relationships. Although the agency issued regulations authorizing health claims for most of these relationships, it concluded that there was insufficient scientific agreement regarding the scientific validity of the four health claims listed in the **Summary** section of this document. Therefore, the agency issued regulations providing that these claims were not authorized. (See § 101.71(a), (c), (e) (21 CFR 101.79(c)(2)(i)(G)).