disseminate information regarding CMS programs, contracting opportunities, and other initiatives to faith-based and other community organizations.

19. Office of Health Insurance Portability and Accountability Act Standards (FHA)

- Develops, implements and administers the enforcement of the Health Insurance Portability and Accountability Act (HIPAA) including portability, transactions, code sets, identifiers, and security.
- Develops, implements and administers the enforcement of the Administrative Simplification Compliance Act (ASCA).
- Develops regulations to enforce the provisions of the HIPAA and the ASCA. Also develops regulations and guidance materials on HIPAA standards.
- Educates and reaches out to the public and internal CMS staff on HIPAA issues. Formulates and coordinates a public relations campaign, prepares and delivers presentations and speeches, responds to inquires on HIPAA issues, and liaisons with industry representatives.
- Works with Federal departments and agencies to identify and adopt universal messaging and clinical health data standards, and represents CMS and HHS in national projects supporting the national health enterprise architecture and the National Health Information Infrastructure.
- Provides technical assistance regarding HIPAA standards and their implementation.
- Collaborates with the Department, especially the Office for Civil Rights, on HIPAA policy issues.
- Coordinates and provides guidance on legislative and regulatory issues.
- Provides assistance and guidance for HIPAA-related budget formulation and execution activities.
- Oversees the enforcement of the insurance portability provisions of HIPAA related to non-Federal governmental health plans and States.

Dated: November 21, 2002.

Ruben J. King-Shaw, Jr.,

Deputy Administrator and Chief Operating Officer, Centers for Medicare & Medicaid Services.

[FR Doc. 03–4086 Filed 2–19–03; 8:45 am] BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 02N-0259]

Agency Information Collection
Activities; Announcement of OMB
Approval; Telephone Questionnaire
Administration to Control Subjects
Recruited into FDA Lyme Vaccine
Safety Study, "A Case-Control Study
of HLA Type and T-Cell Reactivity to
Recombinant Outer Surface Protein A
and Human Leukocyte FunctionAssociated Antigen-1"

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
Administration (FDA) is announcing
that a collection of information entitled
Telephone Questionnaire
Administration to Control Subjects
Recruited into FDA Lyme Vaccine
Safety Study, "A Case-Control Study of
HLA Type and T-Cell Reactivity to
Recombinant Outer Surface Protein A
and Human Leukocyte FunctionAssociated Antigen-1" has been
approved by the Office of Management
and Budget (OMB) under the Paperwork
Reduction Act of 1995.

FOR FURTHER INFORMATION CONTACT: JonnaLynn P. Capezzuto, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-4659.

SUPPLEMENTARY INFORMATION: In the Federal Register of October 18, 2002 (67 FR 64397), the agency announced that the proposed information collection had been submitted to OMB for review and clearance under 44 U.S.C. 3507. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. OMB has now approved the information collection and has assigned OMB control number 0910-0501. The approval expires on June 30, 2004. A copy of the supporting statement for this information collection is available on the Internet at http://www.fda.gov/ ohrms/dockets.

Dated: February 10, 2003.

Margaret M. Dotzel,

Assistant Commissioner for Policy. [FR Doc. 03–4090 Filed 2–19–03; 8:45 am] BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Contaminants and Natural Toxicants Subcommittee of the Food Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration,

HHS.

ACTION: Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

Name of Committee: Contaminants and Natural Toxicants Subcommittee of the Food Advisory Committee.

General Function of the Committee: To provide advice and recommendations to the agency on FDA's regulatory issues.

Date and Time: The meeting will be held on March 18, 2003, from 8 a.m. to 6 p.m.; and on March 19, 2003, from 8:30 a.m. to 3 p.m.

Location: U.S. Department of Agriculture, Animal and Plant Health Inspection Service, 4700 River Rd., Conference Center, Riverdale, MD, 301– 734–8010.

Contact Person: Jeanne E. Latham, Center for Food Safety and Applied Nutrition (HFS-800), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740, 301– 436–1756, or FDA Advisory Committee Information Line, 1–800–741–8138 301– 443–0572 in the Washington, DC area), code 10564. Please call the Information Line for up-to-date information on this meeting.

Agenda: The meeting's purpose is to discuss the scientific issues and principles involved in assessing and evaluating Enterobacter sakazakii contamination in powdered infant formula, risk reduction strategies based on available data, and research questions and priorities. To ensure the presence of the most relevant expertise, the membership of the subcommittee, which has expertise in contaminants, will be augmented by consultants with expertise in infant formula.

The background material for this meeting will be posted on the Internet when available or one working day before the meeting at http://www.cfsan.fda.gov/~lrd/vidtel.html.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person by March 3, 2003. Oral