Grants Management Office signs and issues the award notice.

The Commissioner will notify organizations in writing when their applications will not be funded. Every effort will be made to notify all unsuccessful applicants as soon as possible after final decisions are made.

2. Administrative and National Policy Requirements

45 CFR Part 74 and 45 CFR Part 92

Faith-based organizations that receive funding may not use Federal financial assistance, including funds, to meet any cost-sharing requirements or to support inherently religious activities, such as worship, religious instruction, or prayer.

3. Reporting

Reporting Requirements:
Programmatic Reports and Financial
Reports are required semi-annually with
final reports due 90 days after the
project end date. All required reports
will be submitted in a timely manner, in
recommended formats (to be provided),
and the final report will also be
submitted on disk or electronically
using a standard word-processing
program.

Within 90 days of project end date, the applicant will submit a copy of the final programmatic and financial reports, the evaluation report, and any program products to the National Clearinghouse on Child Abuse and Neglect, 330 C Street, SW., Washington, DC 20447. This is in addition to the standard requirement that the final program and evaluation report must also be submitted to the Grants Management Specialist and the Federal Project Officer.

VII. Agency Contacts

Program Office Contact

Marva Benjamin, 330 C St. SW., Washington, DC 20447, 202–205–8405, *mbenjamin@acf.hhs.gov*.

Grants Management Office Contact

William Wilson, 330 C St SW., Washington, DC 20447, 202–205–8913, wwilson@acf.hhs.gov.

General

The Dixon Group, ACYF Operations Center, 118 Q Street, NE., Washington, DC 20002–2132, Telephone: (866) 796– 1591.

VIII. Other Information

Additional information about this program and its purpose can be located on the following website: http://www.acf.hhs.gov/programs/cb/.

Copies of the following Forms, Assurances, and Certifications are available online at http://www.acf.hhs.gov/programs/ofs/grants/form.htm: Standard Form 424:
Application for Federal Assistance, Standard Form 424A: Budget Information, Standard Form 424B: Assurances—Non-Construction Programs, Form LLL: Disclosure of Lobbying, Certification Regarding Environmental Tobacco Smoke, Standard Form 310: Protection of Human Subjects.

The State Single Point of Contact SPOC listing is available online at http://www.whitehouse.gov/omb/grants/spoc.html.

Dated: April 23, 2004.

Joan E. Ohl,

Commissioner, Administration on Children, Youth and Families.

[FR Doc. 04–9781 Filed 4–29–04; 8:45 am] BILLING CODE 4184–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2004N-0045]

Agency Information Collection Activities; Proposed Collection; Comment Request; Health and Diet Survey—2004 Supplement; Correction

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; correction.

SUMMARY: The Food and Drug Administration is correcting a notice that appeared in the Federal Register of February 18, 2004 (69 FR 7642). The document announced an opportunity for public comment on the proposed collection of information by the agency on a voluntary consumer survey to gauge consumer understanding of dietdisease relationships, particularly those related to saturated fats, trans fatty acids, and omega-3 fatty acids, and consumer attitudes toward diet, health, and physical activity. The document was published with an incorrect docket number. This document corrects that

FOR FURTHER INFORMATION CONTACT: Joyce Strong, Office of Policy and Planning (HF–27), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827–7010. SUPPLEMENTARY INFORMATION: In FR Doc. 04–3411, appearing on page 7642 in the Federal Register of Wednesday, February 18, 2004, the following correction is made:

1. On page 7642, in the second column, in the heading of the

document, "[Docket No. 2003N-0045]" is corrected to read "[Docket No. 2004N-0045]".

Dated: April 23, 2004.

Jeffrey Shuren,

Assistant Commissioner for Policy.
[FR Doc. 04–9837 Filed 4–29–04; 8:45 am]
BILLING CODE 4160–01–8

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Arthritis 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: Arthritis 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 June 2 and 3, 2004, from 8 a.m. to 5 p.m.

Location: Food and Drug Administration, Center for Drug Evaluation and Research Advisory Committee Conference Room, 5630 Fishers Lane, rm. 1066, Rockville, MD.

Contact Person: Kimberly Littleton Topper, Center for Drug Evaluation and Research (HFD–21), Food and Drug Administration, 5600 Fishers Lane, (for express delivery, 5630 Fishers Lane, rm. 1091), Rockville, MD 20857, 301-827–7001, Fax: 301–827–6801, or e-mail: topperk@cder.fda.gov, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), code 3014512532. Please call the Information Line for up-to-date information on this meeting.

Agenda: On June 2, 2004, the committee will discuss trial design and endpoints for drugs for chronic gout, including new drug application (NDA) 21–740, oxypurinol (proposed tradename, OXIPRIM), Cardiome. On June 3, 2004, the committee will discuss trial design and endpoints for drugs for acute gout, including NDA 21–389, etoricoxib (proposed tradename, ARCOXIA), Merck.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending