

Section 10(d) of Public Law 92-463 (5 U.S.C. Appendix I).

Substantive information may be obtained from the contact persons listed above. Summaries of the meetings and rosters of Committee members may be obtained as follows: Ms. Helen W. Garrett, Committee Management Officer, National Institute of Mental Health, Room 17C26, Parklawn Building, Rockville, Maryland 20857, (301) 443-4333.

Dated: October 26, 1982.

Sue Simons,

Committee Management Officer, Alcohol, Drug Abuse and Mental Health Administration.

[FR Doc. 82-29800 Filed 10-28-82; 8:45 am]

BILLING CODE 4160-20-M

Food and Drug Administration

[Docket No. 82F-0315]

Eastman Kodak Co.; Filing of Food Additive Petition

AGENCY: Food and Drug Administration.
ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that Eastman Kodak Co. has filed a petition proposing that the food additive regulations be amended to provide for the safe use of terephthalic acid as an alternate diacid moiety for dimethyl terephthalate as a reactant in the manufacture of ethylene-1,4-cyclohexylene dimethylene terephthalate copolymer intended for use in contact with food.

FOR FURTHER INFORMATION CONTACT: Vir Anand, Bureau of Foods (HFF-334), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202-472-5690.

SUPPLEMENTARY INFORMATION: Under the Federal Food, Drug, and Cosmetic Act (sec. 409(b)(5), 72 Stat. 1786 (21 U.S.C. 348(b)(5))), notice is given that a petition (FAP 2B3667) has been filed by Eastman Kodak Co., Eastman Chemicals Division, Kingsport, TN 37682, proposing that the food additive regulations be amended to provide for the safe use of terephthalic acid as an alternate diacid moiety for dimethyl terephthalate as a reactant in the manufacture of ethylene-1,4-cyclohexylene dimethylene terephthalate copolymer intended for use in contact with food.

The potential environmental impact of this action is being reviewed. If the agency finds that an environmental impact statement is not required and this petition results in a regulation, the

notice of availability of the agency's finding of no significant impact and the evidence supporting that finding will be published with the regulation in the Federal Register in accordance with 21 CFR 25.40(c) (proposed December 11, 1979; 44 FR 71742).

Dated: October 20, 1982.

Sanford A. Miller,
Director, Bureau of Foods.

[FR Doc. 82-29771 Filed 10-28-82; 8:45 am]

BILLING CODE 4160-01-M

[Docket No. 80N-0261]

Food Labeling Formats; Public Meeting

AGENCY: Food and Drug Administration.
ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) and the U.S. Department of Agriculture (USDA) announce the second in a series of public meetings to discuss FDA's research project on communicating food label information. Alternative nutrition labeling formats and tentative plans for consumer research to evaluate the formats will be presented and discussed.
DATES: The public meeting will be held December 2, 1982. Written comments by January 31, 1983.

ADDRESSES: The public meeting will be held in the Hubert H. Humphrey Bldg. Auditorium, 200 Independence Ave. SW., Washington, DC. Written comments to the Dockets Management Branch (HFA-305), Food and Drug Administration, Rm. 4-62, 5600 Fishers Lane, Rockville, MD 20857. Although notices of intention to attend the public meeting are not required, it would help those who are planning and conducting the meeting if the persons who are planning to attend would contact Raymond C. Stokes at the address given below.

FOR FURTHER INFORMATION CONTACT: Raymond C. Stokes, Bureau of Foods (HFF-240), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202-245-1457.

SUPPLEMENTARY INFORMATION: The Federal Register of July 8, 1980 (45 FR 45962) contained an announcement that a series of informal public meetings would be held on how to communicate label information. This reflects an interagency effort to promote public participation in food labeling formats, design, research, and evaluation.

On October 6, 1980, the first meeting was held. At the meeting, participants discussed alternative food labeling formats and work under an FDA contract on this issue. The label-design phase of the contract is now completed.

FDA intends to conduct consumer research to evaluate alternative formats for nutrition labeling on the basis of their ability to communicate the desired information.

The December 2, 1982 public meeting will provide an opportunity to review, discuss, and comment upon (1) the alternative nutrition labeling formats developed under the contract and (2) a tentative consumer research plan to evaluate the effectiveness of the formats. At the meeting, an information kit consisting of graphic and written material that will assist those who wish to submit comments will be distributed. The meeting will be held December 2, 1982, beginning at 9 a.m. in the Hubert H. Humphrey Bldg. Auditorium, 200 Independence Ave. SW., Washington, DC. The meeting will be conducted in accordance with § 10.65 *Meetings and correspondence* (21 CFR 10.65).

After the meeting the information kit also may be obtained from Raymond C. Stokes, Bureau of Foods (HFF-240), Food and Drug Administration, 200 C St. SW., Washington, DC 20204. Written comments should be submitted to the Dockets Management Branch (HFA-305), Food and Drug Administration, Rm. 4-62, 5600 Fishers Lane, Rockville, MD 20857, by January 31, 1983.

Dated: October 25, 1982.

Mark Novitch,

Acting Commissioner of Food and Drugs.

[FR Doc. 82-29770 Filed 10-28-82; 8:45 am]

BILLING CODE 4160-01-M

Anti-Infective Drugs Advisory Committee; Notice of Renewal

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: Under the Federal Advisory Committee Act of October 6, 1972 (Pub. L. 92-463, 86 Stat. 770-776 (5 U.S.C. App. I)), the Food and Drug Administration (FDA) is announcing the renewal of the Anti-Infective Drugs Advisory Committee by the Secretary, Department of Health and Human Services.

DATE: Authority for this committee will expire on October 7, 1984, unless the Secretary formally determines that renewal is in the public interest.

FOR FURTHER INFORMATION CONTACT: Richard L. Schmidt, Committee Management Office (HFA-306), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-443-2765.

Dated: October 21, 1982.
 William F. Randolph,
Acting Associate Commissioner for Regulatory Affairs.
 [FR Doc. 82-29434 Filed 10-28-82; 8:45 am]
 BILLING CODE 4160-01-M

Dermatologic Drugs Advisory Committee; Notice of Renewal

AGENCY: Food and Drug Administration.
ACTION: Notice.

SUMMARY: Under the Federal Advisory Committee Act of October 6, 1972 (Pub. L. 92-463, 88 Stat. 770-776 (5 U.S.C. App. I)), the Food and Drug Administration (FDA) is announcing the renewal of the Dermatologic Drugs Advisory Committee by the Secretary, Department of Health and Human Services.

DATE: Authority for this committee will expire on October 7, 1984, unless the Secretary formally determines that renewal is in the public interest.

FOR FURTHER INFORMATION CONTACT: Richard L. Schmidt, Committee Management Office (HFA-308), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-443-2785.

Dated: October 22, 1982.
 William F. Randolph,
Acting Associate Commissioner for Regulatory Affairs.
 [FR Doc. 82-29591 Filed 10-28-82; 8:45 am]
 BILLING CODE 4160-01-M

Consumer Participation; Notice of Open Meeting

AGENCY: Food and Drug Administration.
ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the following consumer exchange meeting: Orlando District Office, chaired by Adam J. Trujillo, District Director.

DATE: Wednesday, November 17, 1982, 9 a.m. to 11:30 a.m.

ADDRESS: Florida Cooperative Extension Service, 3406 Palm Beach Blvd., Terry Park, Ft. Myers, FL.

FOR FURTHER INFORMATION CONTACT: Lynne C. Isaacs, Consumer Affairs Officer, Food and Drug Administration, P.O. Box 118, Orlando, FL 32802, 305-885-0900.

SUPPLEMENTARY INFORMATION: The purpose of this meeting is to encourage dialogue between consumers and FDA officials, to identify and set priorities for current and future health concerns, to enhance understanding and exchange information between local consumers

and FDA's District Offices, and to contribute to the agency's policymaking decisions on vital issues.

Dated: October 22, 1982.
 William F. Randolph,
Acting Associate Commissioner for Regulatory Affairs.

[FR Doc. 82-29582 Filed 10-28-82; 8:45 am]
 BILLING CODE 4160-01-M

Consumer Participation; Notice of Open Meeting

AGENCY: Food and Drug Administration.
ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the following consumer exchange meeting, chaired by Abraham I. Kleks, District Director, Los Angeles, California..

DATE: Tuesday, November 16, 1982, 9:30 a.m.

ADDRESS: Senior NOW Programs Inc., 102 North Plumer, Tucson, AZ 85719..

FOR FURTHER INFORMATION CONTACT: Gordon L. Scott, Consumer Affairs Officer, Food and Drug Administration, 1521 West Pico Blvd., Los Angeles, CA 90015, 213-688-4395.

SUPPLEMENTARY INFORMATION: The purpose of this meeting is to encourage dialogue between consumers and FDA officials, to identify and set priorities for current and future health concerns, to enhance understanding and exchange information between local consumers and FDA's District Offices, and to contribute to the agency's policymaking decisions on vital issues.

October 22, 1982.
 William F. Randolph,
Acting Associate Commissioner for Regulatory Affairs.
 [FR Doc. 82-29594 Filed 10-28-82; 8:45 am]
 BILLING CODE 4160-01-M

Office of the Secretary

List of Agency Forms Submitted to the Office of Management and Budget for Clearance

Each Friday the Department of Health and Human Services (HHS) publishes a list of information collection packages it has submitted to the Office of Management and Budget (OMB) for clearance in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). The following are those packages submitted to OMB since the last list was published on October 22,

Public Health Service NATIONAL INSTITUTES OF HEALTH

Subject: Health Message Testing Service—Broadcast Component (0925-0046)—REINSTATEMENT.

Respondents: Individuals.
 OMB Desk Officer: Richard Eisinger.

Health Care Financing Administration

Subject: Medigap Industry Survey (HCFA-337)—NEW.

Respondents: Insurance companies which sell supplemental health insurance to Medicare beneficiaries in Calif., Fla., Miss., N.J., Wash., and Wisc.

Subject: Medicaid State Plan Preprint—Revision for Allowance of Liens and Other Recoveries (HCFA-179)—REVISED.

Respondents: State Medicaid agencies.

Subject: Medicaid State Plan Preprint—Revision for New Cost Sharing Policy (HCFA-179)—REVISED.

Respondents: State Medicaid agencies.

Subject: Medicaid State Plan Preprint—Revision for New Cost Sharing Policy (HCFA-179)—REVISED.

Respondents: State Medicaid agencies.

OMB Desk Officer: Fay S. Iudicello.

Office of the Secretary

Subject: Informal Caregiver Survey—Baseline Instrument (National Long Term Care Channeling Demonstration—NEW).

Respondents: Caregivers of elderly individuals served by the channeling demonstrations.

OMB Desk Officer: Milo Sunderhauf.

Copies of the above information collection clearance packages can be obtained by calling the HHS Reports Clearance Officer on 202-245-6511.

Written comments and recommendations for the proposed information collections should be sent directly to both the HHS Reports Clearance Officer and the appropriate OMB Desk Officer designated above at the following addresses:

J. J. Strnad, HHS Reports Clearance Officer, Hubert H. Humphrey Building, Room 524-F, Washington, D.C. 20201.
 OMB Reports Management Branch, New Executive Office Building, Room 3208, Washington, D.C. 20503.

Attn: (name of OMB Desk Officer)