1093), Rockville, MD 20857, 301–827–7001, or e-mail: cliffordj@cder.fda.gov, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), code 12529. Please call the Information Line for up-to-date information on this meeting.

Agenda: On September 9, 2003, the committee will discuss Risk Management Plans for opiate analgesic drug products with particular attention to modified-release products. On September 10, 2003, the committee will discuss the abuse liability of and Risk Management Plans for Palladone, a modified-release hydromorphone drug product indicated for the treatment of moderate to severe pain in opioid tolerant patients.

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 September 2, 2003. Oral presentations from the public will be scheduled between approximately 1:15 p.m. and 1:45 p.m. on September 9, and between 11:30 a.m. and 12 noon on September 10, 2003. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before September 2, 2003, and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation.

Persons attending FDA's advisory committee meetings are advised that the agency is not responsible for providing access to electrical outlets.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Angie Whitacre at 301–827–7001 at least 7 days in advance of the meeting.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: July 23, 2003.

Peter J. Pitts,

Associate Commissioner for External Relations.

[FR Doc. 03-19506 Filed 7-30-03; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Additives and Ingredients 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: Additives and Ingredients 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 August 26 and 27, 2003, from 8:30 a.m. to 4:30 p.m., and August 28, 2003, from 8:30 a.m. to 12 noon.

Location: St. Regis Hotel (Crystal Ballroom), 923 16th St., NW, Washington, DC 20006, 202–638–2626.

Contact Person: Richard E. Bonnette, Center for Food Safety and Applied Nutrition (HFS–255), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740, 202–418–3034, 202–418–3030, 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 upto-date information on this meeting.

Agenda: A small percentage of the population is allergic to natural rubber latex. FDA has received reports of sensitized people experiencing allergic reactions upon eating food they believed was prepared by food handlers wearing natural rubber latex gloves. The purpose of this meeting is to gather information and to provide advice and recommendations to the agency relating to reported allergic reactions to food prepared by workers wearing latex food service gloves.

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 August 20, 2003. Oral presentations from the public will be scheduled between approximately 11 a.m. and 12 noon on August 27, 2003. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before August 20,

2003, and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation.

Persons attending FDA's advisory committee meetings are advised that the agency is not responsible for providing access to electrical outlets.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Richard E. Bonnette at least 7 days in advance of the meeting.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: July 23, 2003.

Peter J. Pitts,

Associate Commissioner for External Relations.

[FR Doc. 03–19448 Filed 7–30–03; 8:45 am] **BILLING CODE 4160–01–S**

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Submission for OMB Review; Comment Request

Periodically, the Health Resources and Services Administration (HRSA) publishes abstracts of information collection requests under review by the Office of Management and Budget, in compliance with the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35). To request a copy of the clearance requests submitted to OMB for review, call the HRSA Reports Clearance Office on (301)–443–1129.

Proposed Project: Ryan White Comprehensive AIDS Resources Emergency (CARE) Act: CARE Act Data Report (CADR) Form: (OMB No. 0915– 0253)—Revision

The CARE Act Data Report (CADR) form, created in 1999 by the HIV/AIDS Bureau of the Health Resources and Services Administration (HRSA), is designed to collect information from grantees, as well as their subcontracted service providers, funded under Titles I, II, III and IV of the Ryan White (CARE) Act of 1990, as amended by the Ryan White CARE Act Amendments of 1996 and 2000 (codified under Title XXVI of

the Public Health Services Act). All Titles of the CARE Act specify HRSA's responsibilities in the administration of grant funds, the allocation of funds, the evaluation of programs for the population served, and the improvement of the quantity and quality of care. Accurate records of the providers receiving CARE Act funding, the services provided, and the clients served continue to be critical to the implementation of the legislation and thus are necessary for HRSA to fulfill its responsibilities.

CARE Act grantees are required to report aggregate data to HRSA annually. The CADR form is used by grantees and their subcontracted service providers to report data on six different areas:

Service provider information, client information, services provided/clients served, demographic information, AIDS Pharmaceutical Assistance and AIDS Drug Assistance Program, and the Health Insurance Program. The primary purposes of the CADR are to: (1) Characterize the organizations from which clients receive services; (2) provide information on the number and

characteristics of clients who receive CARE Act services; and (3) enable HAB to describe the type and amount of services a client receives. In addition to meeting the goal of accountability to Congress, clients, advocacy groups, and the general public, information collected on the CADR is critical for HRSA, State and local grantees, and individual providers to assess the status of existing HIV-related service delivery systems.

The estimated response burden for grantees is estimated as:

Title under which grantee is funded	Number of grantee respondents	Responses per grantee	Hours to coordinate receipt of data reports from	Total hour burden
Title I Only TitleII Only Title III Only	51 59 337	1 1 1	40 40 8	2,040 2,360 2,696
Title IV Only	90	1	16	1,440
Subtotal	537			8,536

The estimated response burden for service providers is estimated as:

Title under which grantee is funded	Number of respondents	Responses per provider	Hours per response	Total hour burden
Title I Only Title II Only Title III Only Title III Only Title IV Only Funded under	1,175 995 248 98 394	1 1 1 1	24 40 40 40 40 48	28,200 39,800 9,920 3,920 18,912
Subtotal	2,782			100,752
Total	3,319			109,288

Written comments and recommendations concerning the proposed information collection should be sent within 30 days of this notice to: Allison Eydt, Human Resources and Housing Branch, Office of Management and Budget, New Executive Office Building, Room 10235, Washington, DC 20503, Fax Number, 202–395–6974.

Dated: July 24, 2003.

Jane M. Harrison,

Director, Division of Policy Review and Coordination.

[FR Doc. 03–19444 Filed 7–30–03; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Submission for OMB Review; Comment Request

Periodically, the Health Resources and Services Administration (HRSA) publishes abstracts of information collection requests under review by the Office of Management and Budget, in compliance with the Paperwork Reduction Act of 1995 (44 U.S.C. Chapter 35). To request a copy of the clearance requests submitted to OMB for review, call the HRSA Reports Clearance Office on (301)-443–1129.

Proposed Project: 340B Drug Pricing Program Survey—NEW

Section 340B of the Public Health Act provides that a manufacturer that sells outpatient drugs to covered entities must agree to charge a price that will not exceed the amount determined under a statutory formula. The entities eligible to access such drug pricing (i.e., certain HHS grantees, certain disproportionate share hospitals, and other specified categories of entities) total approximately 10,000 sites. Most of these safety net providers serve the economically disadvantaged or medically uninsured.

A customer survey is being developed to collect information by mail on various aspects of the 340B Drug Pricing Program, including whether information on the program is reaching the covered entities, reasons some entities are not participating, satisfaction with the savings realized, and interest in possible