

**DEPARTMENT OF HEALTH AND HUMAN SERVICES****Food and Drug Administration****The Sixth Annual FDA–Orange County Regulatory Affairs (OCRA) Educational Conference “FDA and OCRA: Understanding the Changing Landscape”**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice of meeting.

The Food and Drug Administration (FDA) is announcing its sixth annual educational conference entitled “FDA and OCRA: Understanding the Changing Landscape” cosponsored with OCRA. The conference is intended to provide the drug, device and biologics industries with an opportunity to interact with FDA reviewers and compliance officers from FDA’s centers and district offices, as well as other industry experts. The main focus of this interactive conference will be product approval, compliance, and risk management in the three medical product areas. Industry speakers, interactive questions and answer, and workshop sessions will also be included to assure open exchange and dialogue on the relevant regulatory issues.

**Date and Time:** The meeting will be held on June 4 and 5, 2003, from 7:30 a.m. to 5 p.m.

**Location:** The meeting will be held at The Irvine Marriott, 18000 Von Karman Ave., Irvine, CA.

**Contact:** Ramlah Oma, Food and Drug Administration, 19900 MacArthur Blvd., suite 300, Irvine, CA 92612, 949-798-7611, FAX 949-798-7656, or OCRA, Attention to detail (ATD), 111 East Avenida San Gabriel, San Clemente, CA 92672, 949-366-1056, FAX 949-366-1057, Web site: <http://www.ocra-dg.org>. (FDA has verified the Web site address, but is not responsible for subsequent changes to the Web site after the document publishes in the **Federal Register**.)

**Registration and Meeting Information:** See OCRA Web site at <http://www.ocra-dg.org>. Contact ATD at 949-366-1056.

**Before May 20, 2003, registration fees are as follows:** \$425.00 for members, \$500.00 for nonmembers, and \$275.00 for FDA/government/full-time students with proper identification. After May 20, 2003, \$495.00 for members, \$575.00 for nonmembers, and \$325.00 for FDA/government/full-time students with proper identification.

If you need special accommodations due to a disability, please contact Ramlah Oma at least 10 days in advance.

Dated: May 2, 2003.

**Jeffrey Shuren,**

*Assistant Commissioner for Policy.*

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**BILLING CODE 4160-01-S**

**DEPARTMENT OF HEALTH AND HUMAN SERVICES****Food and Drug Administration****Psychopharmacologic Drugs 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:** Psychopharmacologic Drugs 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 16, 2003, from 8 a.m. to 5 p.m.

**Location:** Marriott Washingtonian Center, Grand Ballroom, 9751 Washingtonian Blvd., Gaithersburg, MD.

**Contact Person:** Jayne E. Peterson, Center for Drug Evaluation and Research (HFD-21), Food and Drug Administration, 5600 Fishers Lane, (for express delivery, 5630 Fishers Lane, rm. 1093), Rockville, MD 20857, 301-827-7001, e-mail: [petersonj@cder.fda.gov](mailto:petersonj@cder.fda.gov), or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area), code 12544. Please call the Information Line for up-to-date information on this meeting.

**Agenda:** On June 16, 2003, the committee will discuss the white blood cell (WBC) monitoring schedule for patients being treated long-term with clozapine. Currently, the WBC monitoring schedule is weekly for the first 6 months of continuous therapy and biweekly thereafter. The committee will consider the question of whether the frequency of WBC monitoring can be diminished further following some period of biweekly monitoring. When available, background materials for this meeting will be posted 1-business day prior to the meeting on the FDA Web site at: [www.fda.gov/ohrms/dockets/ac/acmenu.htm](http://www.fda.gov/ohrms/dockets/ac/acmenu.htm). (Click on the year 2003 and scroll down to Psychopharmacologic Drugs Advisory Committee.)

**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 June 9, 2003. Oral presentations from the public will be scheduled between approximately 1 p.m. and 2 p.m. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before June 9, 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 Jayne Peterson 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: May 5, 2003.

**Peter J. Pitts,**

*Associate Commissioner for External Relations.*

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES****Food and Drug Administration**

[Docket No. 02N-0528]

**Risk Management; Public Workshop; Reopening of Comment Period**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice of public workshop; reopening of comment period.

**SUMMARY:** The Food and Drug Administration (FDA) is reopening until May 30, 2003, the comment period for three concept papers entitled “Premarketing Risk Assessment,” “Risk Management Programs,” and “Risk Assessment of Observational Data: Good Pharmacovigilance Practices and Pharmacoepidemiologic Assessment.” The document that requested public input, review, and comments for the three concept papers was published in