

around its Internet site in order to determine whether the information, services, and materials on this web-site are presented in an appropriate technological format and whether it meets the needs, wants, and preferences of visitors or "customers" to the Internet site.

Information on the site focuses on disease prevention, health promotion,

and epidemiology. The site is designed to serve the general public, persons at risk for disease, injury, and illness, and health professionals. This research will ensure that these audiences have opportunity to provide "customer feedback" regarding the value and effectiveness of the information, services, and products of the CDC web-

site and whether these materials are easy to access, clear, and informative. The initial 60 day **Federal Register** Notice was solely for the evaluation of the National Center for HIV, STD, and TB Prevention (NCHSTP) website, but has since been modified to include the entire Agency. The total annual burden hours are 30,667.

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hrs)
Visitors to CDC Internet Site	184,000	1	0.1

Nancy Cheal,

Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention (CDC).

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

The 2000 FDA Science Forum—FDA and the Science of Safety: New Perspectives

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of meeting.

The Food and Drug Administration (FDA), Office of Science, is announcing the following meeting entitled "The 2000 FDA Science Forum—FDA and the Science of Safety: New Perspectives." The forum is devoted to the presentation and sharing of data, knowledge, and ideas among the diverse disciplines of risk management. The forum will bring FDA scientists together with industry, academia, government agencies, consumer groups, and the public to explore the scientific and practical issues related to the safety evaluation and risk management of FDA-regulated products.

Co-sponsored by FDA's Office of Science, the American Association of Pharmaceutical Scientists, FDA's Office of Women's Health, FDA's Chapter of Sigma Xi, and the Scientific Research Society.

Date and Time: The forum will be held on Monday, February 14, 2000, from 8:30 a.m. to 6 p.m., and Tuesday, February 15, 2000, from 8:30 a.m. to 5 p.m.

Location: Washington Convention Center, rms. 29 to 32 (lower level), and Hall C (upper level), 900 Ninth St. NW., Washington, DC 20001.

Contact: Susan A. Homire, Food and Drug Administration, Office of Science (HF-33), 5600 Fishers Lane, Rockville, MD 20857, 301-827-3366, e-mail "shomire@oc.fda.gov".

Registration: Registration information will be provided at a later date.

SUPPLEMENTARY INFORMATION: Speakers and panelists will address emerging issues in the safety assessment of foods, drugs, biologics, and medical devices. Plenary lectures and discussion groups will provide perspectives on the following topics: (1) Walking and Talking: The Art and Science of Risk Communication, (2) Contemporary Issues in Risk Assessment, (3) Postmarket Surveillance—Beyond Passive Surveillance, (4) The Food Safety Initiative—The Risk Perspective, (5) Risk and Gender Effects, and (6) Risk Assessment in Action.

Dated: April 26, 1999.

William K. Hubbard,

Associate Commissioner for Policy Coordination.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Microbiology Devices Panel of the Medical Devices 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). At least one portion of the meeting will be closed to the public.

Name of Committee: The Microbiology Devices Panel of the Medical Devices Advisory Committee.

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

Date and Time: The meeting will be held on May 20, 1999, 9:45 a.m. to 6:30 p.m., and May 21, 1999, 8:30 a.m. to 4:30 p.m.

Location: Corporate Bldg., conference room 020B, 9200 Corporate Blvd., Rockville, MD.

Contact Person: Freddie M. Poole, Center for Devices and Radiological Health (HFZ-440), Food and Drug Administration, 2098 Gaither Rd., Rockville, MD. 20850, 301-594-2096, or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area), code 12517. Please call the Information Line for up-to-date information on this meeting.

Agenda: On May 20, 1999, the committee will discuss and make recommendations on a premarket notification submission for a qualitative in vitro diagnostic assay intended for the detection of human cytomegalovirus (CMV) deoxyribonucleic acid (DNA) in human peripheral white blood cells and its labeling. The focus of the discussion will be the appropriate use of signal amplification terminology. The committee will also discuss, make recommendations, and vote on a premarket approval application (PMA) supplement for an in vitro diagnostic target-amplified nucleic acid probe test used for the detection of *Mycobacterium tuberculosis* complex in sediments prepared from sputum (induced or expectorated), bronchial specimens, or tracheal aspirates. The device as modified is indicated for use of acid-fast bacilli (AFB) smear negative and AFB smear positive respiratory specimens for the diagnosis of active pulmonary tuberculosis disease. On May 21, 1999, the committee will discuss, make recommendations, and vote on a PMA for an in vitro diagnostic qualitative