comments to the Committee should be sent to Jerilyn K. Glass, M.D., PhD, Executive Secretary, ACTPCMD, at the contact information above. Individuals who plan to attend and need special assistance should notify the office at the address and phone number above at least 10 days prior to the meeting. Members of the public will have the opportunity to provide comments at the meeting.

The Advisory Committee will join the Council on Graduate Medical Education (COGME), the National Advisory Council on Nursing Education and Practice (NACNEP), and the Advisory Committee on Interdisciplinary, Community-Based Linkages (ACICBL) on Wednesday, April 21, 2010 for the third Bureau of Health Professions (BHPr) All Advisory Committee Meeting. Please refer to the Federal Register notice for the BHPr All Advisory Committee Meeting for additional details.

Dated: March 18, 2010.

Sahira Rafiullah,

Director, Division of Policy and Information Coordination.

[FR Doc. 2010-6585 Filed 3-24-10; 8:45 am]

BILLING CODE 4165-15-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Service Administration

Advisory Committee on Interdisciplinary, Community-Based Linkages; Notice of Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), notice is hereby given of the following meeting

Name: Advisory Committee on Interdisciplinary, Community-Based Linkages (ACICBL).

Dates and Times: April 22, 2010, 8:30 a.m. to 5 p.m., EST. April 23, 2010, 8:30 a.m. to 3 p.m., EST.

Place: Doubletree Hotel and Executive Meeting Center, 8120 Wisconsin Avenue, Bethesda, Maryland 20814, Telephone: 301–652–2000.

Status: The meeting will be open to the public.

Purpose: The Committee members will advance their efforts in the development of the Tenth Annual Report with the tentative working topic, Preparing the Interprofessional Workforce to Address Health Behaviors. Additionally, the Committee proposes to examine Healthy People 2010 as a strategy to identify county based data on health behavior issues. Beyond the usual health behavior foci of weight, tobacco use, stress, alcohol/substance use/ abuse, the Committee proposes studying the adherence to healthcare regimes with an emphasis on individual healthcare, systems, and community approaches. The meeting will afford Committee members with the opportunity to identify and discuss the current issues in an effort to formulate

recommendations for the Secretary of Health and Human Services and the Congress.

Agenda: The ACICBL agenda includes an overview of the Committee's general business activities, presentations by and dialogue with experts, and discussion sessions specific to the development of recommendations to be addressed in the Tenth Annual ACICBL Report. Agenda items are subject to change as dictated by the priorities of the Committee.

Supplementary Information: The ACICBL will join the Council on Graduate Medical Education (COGME), the National Advisory Council on Nurse Education and Practice (NACNEP), and the Advisory Committee on Training in Primary Care Medicine and Dentistry (ACTPCMD) on April 21, 2010, for the third Bureau of Health Professions (BHPr) All Advisory Committee Meeting. Please refer to the Federal Register notice for the BHPr All Advisory Committee Meeting for additional details. Requests to make oral comments or to provide written comments to the ACICBL should be sent to Dr. Joan Weiss, Designated Federal Official, at the contact information below. Individuals who plan to attend the meeting and need special assistance should notify Dr. Weiss at least 10 days prior to the meeting, using the address and phone number below. Members of the public will have the opportunity to provide comments at the meeting.

For Further Information Contact: Anyone requesting information regarding the ACICBL should contact Dr. Joan Weiss, Designated Federal Official with the Bureau of Health Professions, Health Resources and Services Administration, Parklawn Building, Rm 9–36, 5600 Fishers Lane, Rockville, Maryland 20857, (301) 443–6950 or jweiss@hrsa.gov. Additionally, CAPT Norma J. Hatot, Senior Nurse Consultant, can be contacted at (301) 443–2681 or nhatot@hrsa.gov.

Dated: March 18, 2010.

Sahira Rafiullah.

Director, Division of Policy and Information Coordination.

[FR Doc. 2010–6587 Filed 3–24–10; 8:45 am] BILLING CODE 4165–15–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee to the Director (ACD), Centers for Disease Control and Prevention

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC) announces the following meeting of the aforementioned committee.

Time and Date: 8:30 a.m.-3:15 p.m., April 12, 2010.

Place: CDC, Thomas R. Harkin Global Communications Center, Kent "Oz" Nelson Auditorium, 1600 Clifton Road, NE., Atlanta, GA 30333. Status: Open to the public, limited only by the space available. The meeting room accommodates approximately 150 people. The public is welcome to participate during the public comment periods. The public comment periods are tentatively scheduled for 10:50 a.m. to 11:05 a.m. and 2:30 p.m. to 2:45 p.m.

Purpose: The Advisory Committee to the Director, CDC shall advise the Secretary, HHS, and the Director, CDC on policy issues and broad strategies that will enable CDC to fulfill its mission of protecting health through health promotion, prevention, and preparedness. The committee recommends ways to prioritize CDC's activities, improve results, and address health disparities. It also provides guidance to help CDC work more effectively with its various private and public sector constituents to make health protection a practical reality.

Matters To Be Discussed: The ACD, CDC will discuss and approve recommendations by the Ethics Subcommittee on "Ethical Considerations for Decision Making Regarding Allocation of Mechanical Ventilators During a Severe Influenza Pandemic." Other agenda items will include updates from the ACD, CDC subcommittees; CDC organizational improvement; the CDC budget (including mitigating State cuts); an overview of winnable battles for CDC; and the need for establishment of additional ACD, CDC subcommittees to address agency needs and priorities.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Louis Salinas, M.P.A., Designated Federal Officer, ACD, CDC, 1600 Clifton Road, NE., M/S D–14, Atlanta, Georgia 30333. Telephone 404/639–7000. E-mail: GHickman@cdc.gov. The deadline for notification of attendance is April 7, 2010.

The Director, Management Analysis and Services Office, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Dated: March 18, 2010.

Elaine L. Baker,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention (CDC).

[FR Doc. 2010-6590 Filed 3-24-10; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Council on Graduate Medical Education; Notice of Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), notice is hereby given of the following meeting: *Name:* Council on Graduate Medical Education (COGME).

Dates and Times: April 22, 2010, 8:30 a.m.-4:15 p.m. EST; April 23, 2010, 8:30 a.m.-4:15 p.m. EST.

Place: DoubleTree Hotel Bethesda, 8120 Wisconsin Avenue, Bethesda, MD 20814, Telephone: (301) 652–2000.

Status: The meeting will be open to the public except on Friday, April 23 from 12 p.m.–1 p.m.

Agenda: On April 22, the meeting will be called to order with remarks from the COGME Chair and the Executive Secretary of COGME. There will be presentations addressing topics such as: (1) The adequacy of the pediatrician workforce physician supply; (2) the results of a recent study of primary care physician workforce projections by State; (3) the Bureau of Health Professions plans for healthcare workforce analytics; (4) a patient-centered primary care collaborative; and (5) the relationship between primary care, population health, and health care costs.

On April 23, there will be presentations on the workforce components of key health reform legislation and on challenges facing graduate medical education in the coming decade. The Council members will enter into a discussion and will formulate recommendations to the Secretary of Health and Human Services and the Congress as part of the Council's emerging report covering the primary care physician workforce.

Agenda items are subject to change as priorities dictate.

For Further Information Contact: Anyone interested in obtaining a roster of members or other relevant information should write or contact Jerald M. Katzoff, Executive Secretary, COGME, Division of Medicine and Dentistry, Bureau of Health Professions, Parklawn Building, Room 9A–27, 5600 Fishers Lane, Rockville, Maryland 20857, Telephone (301) 443–4443. The Web address for information on the Council and the April 22–23, 2010 meeting agenda is http://cogme.gov.

COGME will join the National Advisory Council on Nursing Education and Practice (NACNEP), the Advisory Committee on Training in Primary Care Medicine and Dentistry (ACTPCMD), and the Advisory Committee on Interdisciplinary, Community-Based Linkages (ACICBL) on April 21, 2010 for the third Bureau of Health Professions (BHPr) All Advisory Committee Meeting. Please refer to the Federal Register notice for the BHPr All Advisory Committee Meeting for additional details.

Supplementary Information: Requests to make oral comments or to provide written comments to the Council should be sent to Jerald M. Katzoff, Executive Secretary, COGME, at the contact information above. Individuals who plan to attend and need special assistance should notify the office at the address and phone number above at least 10 days prior to the meeting. Members of the public will have the opportunity to provide comments at the meeting.

Dated: March 18, 2010.

Sahira Rafiullah,

Director, Division of Policy and Information Coordination.

[FR Doc. 2010–6586 Filed 3–24–10; 8:45 am] BILLING CODE 4165–15–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2010-N-0001]

Food and Drug Administration Clinical Trial Requirements, Regulations, Compliance, and Good Clinical Practices; Public Workshop

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of public workshop.

SUMMARY: The Food and Drug Administration (FDA) Los Angeles District Office, in cosponsorship with the Society of Clinical Research Associates, Inc. (SoCRA) is announcing a public workshop entitled "FDA Clinical Trial Requirements, Regulations, Compliance, and Good Clinical Practices." The public workshop is intended to aid the clinical research professional's understanding of the mission, responsibilities, and authority of FDA and to facilitate interaction with FDA representatives. The program will focus on the relationships among the FDA and clinical trial staff, investigators, and ivestigational review boards (IRBs). Individual FDA representatives will discuss the informed consent process and informed consent documents, and regulations relating to drugs, devices, and biologics, as well as inspections of clinical investigators, IRBs, and research sponsors.

Date and Time: The public workshop will be held on Wednesday and Thursday, May 5 and 6, 2010, from 8 a.m. to 5 p.m.

Location: The public workshop will be held at the Hyatt Regency Newport Beach, 1107 Jamboree Rd., Newport Beach, CA 92660, 949–729–1234.

Contact: Linda Hartley, Food and Drug Administration, 19701 Fairchild, Irvine, CA 92612, 949–608–4413, FAX: 949–608–4417. Attendees are responsible for their own accommodations. To make reservations at the Hyatt Regency Newport Beach, contact the Hyatt Regency Newport Beach (see Location).

Registration: The SoCRA registration fees cover the cost of actual expenses, including refreshments, lunch, materials, and speaker expenses. Seats

are limited; please submit your registration as soon as possible. Workshop space will be filled in order of receipt of registration. Those accepted into the workshop will receive confirmation. Registration will close after the workshop is filled. Registration at the site is not guaranteed but may be possible on a space available basis on the day of the public workshop beginning at 8 a.m. The cost of registration is as follows: FDA employee (fee waived), Government employee member (\$450), Government employee nonmember (\$525), non-Government employee SoCRA member (\$575), non-Government employee non-SoCRA member (\$650).

If you need special accommodations due to a disability, please contact Linda Hartley (see *Contact*) at least 10 days in advance of the public workshop.

Extended periods of question and answer and discussion have been included in the program schedule.

Registration instructions: To register, please submit a registration form with your name, affiliation, mailing address, phone, fax number, and e-mail, along with a check or money order payable to "SoCRA." Mail to: SoCRA, 530 West Butler Ave., suite 109, Chalfont, PA 18914. To register via the Internet, go to http://www.socra.org/html/FDA_Conference.htm. (FDA has verified the Web site address, but we are not responsible for any subsequent changes to the Web site after this document publishes in the Federal Register.)

The registrar will also accept payment by major credit cards (VISA/MasterCard/AMEX only). For more information on the meeting, or for questions on registration, contact SoCRA at 800–762–7292 or 215–822–8644, FAX: 215–822–8633, or e-mail: SoCRAmail@aol.com.

SUPPLEMENTARY INFORMATION: The public workshop helps fulfill the Department of Health and Human Services' and FDA's important mission to protect the public health. The workshop will provide those engaged in FDA-regulated (human) clinical trials with information on a number of topics concerning FDA requirements related to informed consent, clinical investigation requirements, institutional review board inspections, electronic record requirements, and investigator initiated research. Topics for discussion include the following: (1) What FDA expects in a pharmaceutical clinical trial; (2) adverse event reporting science, regulation, error, and safety; (3) Part 11 Compliance—Electronic Signatures; (4) informed consent regulations; (5) IRB regulations and FDA inspections; (6)