Retirement: Kathy Hughes, Vice Chairwoman, Oneida Nation; Gail Hunt, President and Chief Executive Officer, National Alliance for Caregiving; Warren Jones, M.D., F.A.A.F.P., Executive Director, Mississippi Institute for Improvement of Geographic Minority Health; Sandy Markwood, Chief Executive Officer, National Association of Area Agencies on Aging; David W. Roberts, M.P.A., Vice President, Government Relations, Healthcare Information and Management System Society; Julie Boden Schmidt, M.S., Associate Vice President, Training and Technical Assistance, National Association of Community Health Centers; Rebecca P. Snead, Chief Executive Officer and Executive Vice President, National Alliance of State Pharmacy Associations and APME Chair; Donnna Yee, PhD, Chief Executive Officer, Asian Community Center of Sacramento Valley; Deanna Jang, Policy Director, Asian and Pacific Islander American Health Forum; Andrew Kramer, M.D., Professor of Medicine, Division of Health Care Policy and Research, University of Colorado, Denver; John Lui, PhD, M.B.A., Executive Director, Stout Vocational Rehabilitation Institute.

The agenda for the June 22, 2010 meeting will include the following:
- Recap of the previous (March 31, 2010) meeting.
- Subgroup Committee Work Summary.
- Medicare Outreach and Education Strategies.
- Public Comment.
- Listening Session with CMS Leadership.

Next Steps.

Individuals or organizations that wish to make a 5-minute oral presentation on an agenda topic should submit a written copy of the oral presentation to the DFO at the address listed in the ADDRESSES section of this notice by the date listed in the DATES section of this notice. The number of oral presentations may be limited by the space available.

Individuals not wishing to make a presentation may submit written comments to the DFO at the address listed in the ADDRESSES section of this notice by the date listed in the DATES section of this notice.

Individuals requiring sign language interpretation or other special accommodations should contact the DFO at the address listed in the ADDRESSES section of this notice by the date listed in the DATES section of this notice.


(Catalog of Federal Domestic Assistance Program No. 93.733, Medicare—Hospital Insurance Program; and Program No. 93.774, Medicare—Supplemental Medical Insurance Program)


Marilyn Taverner, Acting Administrator, Centers for Medicare & Medicaid Services.

[FR Doc. 2010–12457 Filed 5–27–10; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee on Immunization Practices (ACIP)

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:

Times and Dates:
8 a.m.–6 p.m., June 23, 2010.
8 a.m.–3 p.m., June 24, 2010.
Place: CDC, Tom Harkin Global Communications Center, 1600 Clifton Road, NE., Building 19, Kent “Oz” Nelson Auditorium, Atlanta, Georgia 30333.
Status: Open to the public, limited only by the space available.

Purpose: The committee is charged with advising the Director, CDC, on the appropriate uses of immunizing agents. In addition, under 42 U.S.C. 1396s, the committee is mandated to establish and periodically review and, as appropriate, revise the list of vaccines for administration to vaccine-eligible children through the Vaccines for Children (VFC) program, along with schedules regarding the appropriate periodicity, dosage, and contraindications applicable to the vaccines.

Matters To Be Discussed: The agenda will include discussions on: Evidence based recommendations; human papillomavirus (HPV) vaccines; 13-valent pneumococcal conjugate vaccine; meningococcal vaccine; hepatitis vaccines; a vaccine supply update; respiratory syncytial virus immunoprophylaxis vaccine; rotavirus vaccines; pertussis vaccine; and influenza vaccines. Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Antoinette Hill, Immunization Services Division, National Center for Immunization and Respiratory Diseases, CDC, 1600 Clifton Road, NE., (E–05), Atlanta, Georgia 30333, telephone 404/639–8836, fax 404/639–8905.

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 CDC and the Agency for Toxic Substance and Disease Registry.


Elaine L. Baker,
Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2010–12818 Filed 5–27–10; 8:45 am]
BILLING CODE 4160–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Partnerships To Advance the National Occupational Research Agenda (NORA)

AGENCY: The National Institute for Occupational Safety and Health (NIOSH) of the Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice of public meeting.

SUMMARY: The National Institute for Occupational Safety and Health (NIOSH) of the Centers for Disease Control and Prevention (CDC) announces the following public meeting: "Partnerships to Advance the National Occupational Research Agenda (NORA)."

Public Meeting Time and Date:
10 a.m.–4 p.m. EDT, June 16, 2010.
Place: Patriots Plaza, 395 E. Street, SW., Conference Room 9000, Washington, DC 20201.

Purpose of Meeting: The National Occupational Research Agenda (NORA) has been structured to engage partners with each other and/or with NIOSH to advance NORA priorities. The NORA Liaison Committee continues to be an opportunity for representatives from organizations with national scope to learn about NORA progress and to suggest possible partnerships based on their organization’s mission and contacts. This opportunity is now structured as a public meeting via the internet to attract participation by a larger number of organizations and to further enhance the success of NORA.

Some of the types of organizations of national scope that are especially encouraged to participate are employers, unions, trade associations, labor associations, professional associations, and foundations. Others are welcome.

This meeting will include updates from NIOSH leadership on NORA as well as updates from approximately half of the NORA Sector Councils on their progress, priorities, and implementation
plans to date, including the NORA Construction, Manufacturing, Public Safety, Services, and Wholesale and Retail Trade Councils. Updates will also be given on cross-sector activities in the areas of Healthy People 2020 and the WorkLife Initiative. After each update, there will be time to discuss partnership opportunities.

Status: The meeting is open to the public, limited only by the capacities of the conference call and conference room facilities. There is limited space available in the meeting room (capacity 34). Therefore, information to allow participation in the meeting through the internet (to see the slides) and a teleconference call (capacity 50) will be provided to registered participants. Participants are encouraged to consider attending by this method. Each participant is requested to register for the free meeting by sending an e-mail to noracoordinator@cdc.gov containing the participant’s name, organization name, contact telephone number on the day of the meeting, and preference for participation by Web meeting (requirements include: Computer, internet connection, and telephone, preferably with ‘mute’ capability) or in person. An e-mail confirming registration will include the details needed to participate in the Web meeting. Non-US citizens who do not register to attend in person on or before June 2, 2010, will not be granted access to the meeting site and will not be able to attend the meeting in person due to mandatory security clearance procedures at the Patriots Plaza facility.

Background: NORA is a partnership program to stimulate innovative research in occupational safety and health leading to improved workplace practices. Unveiled in 1996, NORA has become a research framework for the nation. Diverse parties collaborate to identify the most critical issues in workplace safety and health. Partners then work together to develop goals and objectives for addressing those needs and to move the research results into practice. The NIOSH role is facilitator of the process. For more information about NORA, see http://www.cdc.gov/niosh/nora/about.html.

Since 2006, NORA has been structured according to industrial sectors. Eight major sector groups have been defined using the North American Industrial Classification System (NAICS). After receiving public input through the web and town hall meetings, the NORA Sector Councils have been working to define sector-specific strategic plans for conducting research and moving the results into widespread practice. During 2008–2009, most of these Councils have posted draft strategic plans for public comment. Seven have posted finalized National Sector Agendas after considering comments on the drafts. For more information, see the link above and choose “Sector-based Approach,” “NORA Sector Councils,” “Sector Agendas” and “Comment on Draft Sector Agendas” from the right-side menu.

FOR FURTHER INFORMATION CONTACT: Sidney C. Soderholm, PhD, NORA Coordinator, E-mail noracoordinator@cdc.gov, telephone (202) 245–0665.

Tanja Popovic,
Deputy Associate Director for Science, Centers for Disease Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration
[Docket No. FDA–2010–N–0001]
Oncologic 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: Oncologic 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 July 20, 2010, from 8 a.m. to 3 p.m.


Contact Person: Nicole Vesely, c/o Melanie Whelan, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, rm. 6100, Silver Spring, MD 20993–0002, FAX: 301–847–8737, to reach by telephone before June 8, 2010, please call 301–827–7001; to reach by telephone after June 8, 2010, please call 301–796–9000, e-mail: nicole.vesely@fda.hhs.gov, or FDA Advisory Committee Information Line, 1–800–872–1388 (301–443–0572 in the Washington, DC area), code 3014512542. Please call the Information Line for up-to-date information on this meeting. A notice in the Federal Register about last minute modifications that impact a previously announced advisory committee meeting cannot always be published quickly enough to provide timely notice. Therefore, you should always check the agency’s Web site and call the appropriate advisory committee hot line/phone line to learn about possible modifications before coming to the meeting.

Agenda: On July 20, 2010, the committee will discuss supplemental biologics license applications (sBLAs) 125085/191 and 192 for AVASTIN (bevacizumab), manufactured by Genentech, Inc. The two proposed indications (uses) for this product are: (1) First-line treatment of a subgroup of women with metastatic breast cancer known as HER2-negative breast cancer, in combination with the chemotherapy drug docetaxel; and (2) first-line treatment of HER2-negative metastatic breast cancer in combination with one of two classes of chemotherapy drugs, known as taxanes and anthracyclines, or with the chemotherapy drug, capecitabine. In addition to the discussion of these two indications, the committee will also consider the impact of the submitted studies on the conversion from accelerated to regular approval of the indication for the treatment, in combination with the chemotherapy drug paclitaxel, of patients who have not received prior chemotherapy for their locally recurrent or metastatic HER2 negative breast cancer.

FDA intends to make background material available to the public no later than 2 business days prior to the meeting. If FDA is unable to post the background material on its Web site prior to the meeting, the background material will be made publicly available at the location of the advisory committee meeting, and the background material will be posted on FDA’s Web site after the meeting. Background material is available at http://www.fda.gov/AdvisoryCommittees/Calendar/default.htm. Scroll down to the appropriate advisory committee link.

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 on or before July 6, 2010. Oral presentations from the public will be scheduled between approximately 12:30 p.m. to 1:30 p.m. Those desiring to make formal oral presentations should