For information on accommodation options, contact conference coordinator Karen Smith or Andrea Graves at the Robert M. Kerr Food & Agricultural Products Center, Oklahoma State University, 148 FAPC, Stillwater, OK 74078–6055, 405–744–6071, FAX: 405–744–6313, or e-mail: karenl.smith@okstate.edu or andrea.graves@okstate.edu. More information is also available online at http://www.fapc.biz/fooddefense.html. (FDA has verified the Web site address, but FDA is not responsible for any subsequent changes to the Web site after this document publishes in the Federal Register.)

Registration: You are encouraged to register by October 21, 2011. The workshop has a $150 registration fee to cover the cost of facilities, materials, speakers, and breaks. Seats are limited; please submit your registration as soon as possible. The workshop 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 workshop, beginning at 8 a.m. The cost of registration at the site is $200 payable to FAPC. There is no registration fee for FDA employees.

If you need special accommodations due to a disability, please contact Karen Smith (see Contact) at least 7 days in advance.

Registration Form Instructions: To register, please complete the online registration form at http://www.fapc.biz/fooddefense.html.

Transcripts: Transcripts of the public workshop will not be available due to the format of this workshop. Course handouts may be requested after the date of the public workshop through the contact persons (see Contact) at cost plus shipping.

SUPPLEMENTARY INFORMATION: This public workshop is being held in response to the large volume of food defense inquiries from food manufacturers originating from the area covered by the FDA Dallas District Office. The SWRO presents this workshop to help achieve objectives set forth in section 406 of the Food and Drug Administration Modernization Act of 1997 (21 U.S.C. 393), which include working closely with stakeholders and maximizing the availability and clarity of information to stakeholders and the public. This is consistent with the purposes of the Southwest Regional Small Business Representative Program, which are in part to respond to industry inquiries, develop educational materials, and sponsor workshops and conferences to provide firms, particularly small businesses, with firsthand working knowledge of FDA’s regulations and compliance policies. This workshop is also consistent with the Small Business Regulatory Enforcement Fairness Act of 1996 (Pub. L. 104–121), as outreach activities by government agencies to small businesses.

The goal of this public workshop is to present information that will enable regulated industry to better comply with the regulations authorized by the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (the Bioterrorism Act) and to better understand FDA’s food defense guidance documents, especially in light of growing concerns about food protection. Information that FDA presents will be based on Agency position as articulated through regulation, guidance, and information previously made available to the public. Topics to be discussed at the workshop (both by FDA and non-FDA speakers) include: (1) Food defense awareness and definitions, (2) FDA food defense tools such as ALERT and Employees FIRST, (3) regulations issued under the Bioterrorism Act, (4) food defense guidance documents, (5) investigating food-related incidents effectively, (6) physical plant security, (7) crisis management, and other related topics. For more information, please visit http://www.fapc.biz/fooddefense.html. FDA expects that participation in this public workshop will provide regulated industry with greater understanding of the Agency’s regulatory and policy perspectives on food protection, increase compliance with FDA regulations, and heighten food defense awareness.

Dated: September 23, 2011.

Leslie Kux,
Acting Assistant Commissioner for Policy.
[FR Doc. 2011–25114 Filed 9–28–11; 8:45 am]
BILLING CODE 4160–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2011–N–0002]

Request for Notification From Industry Organizations Interested in Participating in the Selection Process and Request for Nominations for a Nonvoting Industry Representative on the Vaccines and Biological Products Advisory Committee

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is requesting that any industry organizations interested in participating in the selection of a nonvoting industry representative to serve on the Vaccines and Related Biological Products Advisory Committee for the Center for Biologics Evaluation and Research (CBER) notify FDA in writing. FDA is also requesting nominations for a nonvoting industry representative to serve the Vaccines and Related Biological Products Advisory Committee. A nominee may either be self-nominated or nominated by an organization to serve as a nonvoting industry representative. Nomination will be accepted for current vacancies effective with this notice.

DATES: Any industry organization interested in participating in the selection of an appropriate nonvoting member to represent industry interests must send a letter stating that interest to the FDA by October 31, 2011, for the vacancy listed in this document. Concurrently, nomination materials for prospective candidates should be sent to FDA by October 31, 2011.

ADDRESSES: All letters of interest and nominations should be submitted in writing to Donald Jehn (see FOR FURTHER INFORMATION CONTACT). For further information contact: Donald Jehn, Center for Biologics Evaluation and Research (HFM–71), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852–1448, 301–827–0314, FAX: 301–827–0294, e-mail: donald.jehn@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: The Agency intends to add a nonvoting industry representative on the CBER Advisory Committee.

I. Vaccines and Related Biological Products Advisory Committee

The Vaccines and Related Biological Products Advisory Committee (the Committee) advises the Commissioner
DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of Diabetes and Digestive and Kidney Diseases Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Institute of Diabetes and Digestive and Kidney Diseases Special Emphasis Panel; Digestive Diseases Core Centers.

Date: December 2, 2011.

Time: 8 a.m. to 5:30 p.m.

Agenda: To review and evaluate grant applications.

Place: Bethesda Marriott Suites, 6711 Democracy Boulevard, Bethesda, MD 20817.

Contact Person: Maria E. Davila-Bloom, PhD, Scientific Review Officer, Review Branch, DEA, NIDDK, National Institutes Of Health, Room 758, 6707 Democracy Boulevard, Bethesda, MD 20892–5452, (301) 594–7637, davilabloom@extra.niddk.nih.gov.

(Catalogue of Federal Domestic Assistance Program Nos. 93.847, Diabetes, Endocrinology and Metabolic Research; 93.848, Digestive Diseases and Nutrition Research; 93.849, Kidney Diseases, Urology and Hematology Research, National Institutes of Health, HHS)

Dated: September 23, 2011.

Jennifer S. Spaeth,
Director, Office of Federal Advisory Committee Policy.

[FR Doc. 2011–25095 Filed 9–28–11; 8:45 am]

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