DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Amended Notice of Meeting

Notice is hereby given of a change in the meeting of the Center for Scientific Review Special Emphasis Panel.

September 30, 2015, 10:00 a.m. to September 30, 2015, 11:30 a.m., National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 which was published in the Federal Register on August 27, 2015, 80 FR 49252.

The meeting will be held on October 21, 2015. The meeting location and time remain the same. The meeting is closed to the public.

Dated: August 24, 2015.

David Clary,
Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2015–22698 Filed 8–26–15; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Clinical Center; Notice of Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of a meeting of the NIH Advisory Board for Clinical Research.

The meeting will be open to the public as indicated below, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the Contact Person listed below in advance of the meeting.

The meeting will be closed to the public in accordance with the provisions set forth in section 552b(c)(9)(B), Title 5 U.S.C., as amended because the premature disclosure of to discuss personnel matters and the discussions would likely to significantly frustrate implementation of recommendations.

Name of Committee: NIH Advisory Board for Clinical Research.

Date: September 28, 2015.

Open: 10:00 a.m. to 1:00 p.m.

Agenda: Discussion of intramural clinical research operational and funding issues.

Place: National Institutes of Health, Building 10, CRC Medical Board Room 4–2551, 10 Center Drive, Bethesda, MD 20892.

Closed: 1:00 p.m. to 2:00 p.m.

Agenda: Discussion of personnel matters and/or issues of which the premature discloser may affect outcomes.

Place: National Institutes of Health, Building 10, CRC Medical Board Room 4–2551, 10 Center Drive, Bethesda, MD 20892.

Contact Person: Maureen E. Gormley, Executive Secretary, Mark O. Hatfield Clinical Research Center, National Institutes of Health, Building 10, Room 6–2551, Bethesda, MD 20892 (301) 496–2897.

Any interested person may file written comments with the committee by forwarding the statement to the Contact Person listed on this notice. The statement should include the name, address, telephone number and when applicable, the business or professional affiliation of the interested person.

In the interest of security, NIH has instituted stringent procedures for entrance onto the NIH campus. All visitor vehicles, including taxicabs, hotel, and airport shuttles will be inspected before being allowed on campus. Visitors will be asked to show one form of identification (for example, a government-issued photo ID, driver’s license, or passport) and to state the purpose of their visit.

Dated: August 21, 2015.

Michelle Trout,
Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2015–21222 Filed 8–26–15; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HOMELAND SECURITY

U.S. Customs and Border Protection

Modification of National Customs Automation Program (NCAP) Test Concerning the Submission of Certain Data Required by the Food and Drug Administration (FDA) Using the Partner Government Agency (PGA) Message Set Through the Automated Commercial Environment (ACE)


ACTION: General notice.

SUMMARY: This document announces U.S. Customs and Border Protection’s (CBP’s) plan to conduct a National Customs Automation Program (NCAP) test concerning the electronic transmission of certain import data for all Food and Drug Administration (FDA)-regulated commodities. Under the pilot, this data will be transmitted electronically through the Automated Broker Interface (ABI) for processing in CBP’s Automated Commercial Environment (ACE) system utilizing the Partnering Government Agency (PGA) Message Set.

DATES: The FDA PGA Message Set test will begin no earlier than August 27, 2015. This test will continue until concluded by way of announcement in the Federal Register. Public comments are invited and will be accepted through the duration of the test pilot.

ADDRESSES: Comments concerning this notice and any aspect of this test may be submitted at any time during the test via email to Josephine Baimonte, ACE Business Office (ABO), Office of International Trade, at josephine.baimonte@cbp.dhs.gov. In the subject line of your email, please indicate, “Comment on FDA PGA Message Set Test FRN”.

FOR FURTHER INFORMATION CONTACT: For FDA-related questions, contact Elizabeth McQueen at elizabeth.mcqueen@cbp.dhs.gov. For technical questions related to the Automated Commercial Environment (ACE) or Automated Broker Interface (ABI) transmissions, contact your assigned client representative. Interested parties without an assigned client representative should direct their questions to Steven Zaccaro at steven.j.zaccaro@cbp.dhs.gov with the subject heading “PGA Message Set FDA Test FRN-Request to Participate.” For FDA-related questions, contact Sandra Abbott at sandra.abbott@fda.hhs.gov or Max Castillo at max.castillo@fda.hhs.gov.

Any party seeking to participate in this test must provide CBP, in its request to participate, its filer code and the port(s) at which it is interested in filing the appropriate PGA Message Set information. At this time, PGA Message Set data may be submitted only for entries filed at certain ports. A current listing of those ports may be found at the following link: http://www.cbp.gov/document/guidance/list-aceids-pga-message-set-pilot-ports.

SUPPLEMENTARY INFORMATION:

I. Background

The National Customs Automation Program (NCAP) was established in Title VI—Customs Modernization (Customs Modernization Act), in the North American Free Trade Agreement Implementation Act, Public Law 103–182, 107 Stat. 2057 (19 U.S.C. 1411). Through NCAP, the initial thrust of customs modernization was on trade compliance and the development of the Automated Commercial Environment (ACE), the planned successor to the Automated Commercial System (ACS). ACE is an automated and electronic system for processing commercial trade data which is intended to streamline business processes, facilitate growth in trade, ensure cargo security, and foster participation in global commerce, while...