
(2) For affected fuel pumps that have a part number or combination of part numbers that are specified in paragraphs (h)(2)(i) through (h)(2)(iii) of this AD: Do the replacement within 90 months after the effective date of this AD.

(i) All of the affected fuel pumps have P/N 568–1–28300–100.

(ii) All of the affected fuel pumps have P/N 568–1–28300–101.

(iii) The affected fuel pumps have a combination of P/Ns 568–1–28300–100 and 568–1–28300–101.

(i) Definitions

(1) For the purpose of this AD, an “affected fuel pump” is defined as any pump having P/N 568–1–28300–001, 568–1–28300–002, 568–1–28300–100, or 568–1–28300–101.

(2) For the purpose of this AD, a “serviceable fuel pump” is a pump having a part number not listed in paragraph (i)(1) of this AD.

(j) No Reporting Requirement

Although Airbus Service Bulletin A330–28–3127, Revision 01, dated September 24, 2015; Airbus Service Bulletin A340–28–4138, Revision 01, dated September 24, 2015; or Airbus Service Bulletin A340–28–5060, Revision 01, dated September 24, 2015, specifies to submit certain information to the manufacturer, and specifies that action as “RC” (Required for Compliance), this AD does not include that requirement.

(k) Parts Installation Prohibition

After the identification of the fuel pump part numbers as required by paragraph (g) of this AD, comply with the prohibition required by paragraph (k)(1) or (k)(2) of this AD, as applicable.

(1) For an airplane that does not have an affected fuel pump installed: After the identification of the fuel pump part numbers as required by paragraph (g) of this AD, do not install an affected fuel pump.

(2) For an airplane that has an affected fuel pump installed: After modification of an airplane as required by paragraph (b) of this AD, no person may install an affected fuel pump on any airplane.

(l) Credit for Previous Actions

This paragraph provides credit for actions required by paragraphs (g) and (h) of this AD, if those actions were performed before the effective date of this AD using service information included in paragraphs (l)(1), (l)(2), and (l)(3) of this AD, which are not incorporated by reference in this AD.


(m) Other FAA AD Provisions

The following provisions also apply to this AD:


Before using any approved AMOC, notify your appropriate principal inspector, or lacking a principal inspector, the manager of the local flight standards district office/certificate holding district office. The AMOC approval letter must specifically reference this AD.

(2) Contacting the Manufacturer: For any requirement in this AD to obtain corrective actions from a manufacturer, the action must be accomplished using a method approved by the Manager, International Branch, ANM–116, Transport Airplane Directorate, FAA; or the European Aviation Safety Agency (EASA); or Airbus’s EASA Design Organization Approval (DOA). If approved by the DOA, the approval must include the DOA-authorized signature.

(3) Required for Compliance (RC): Except as provide by paragraph (j) of this AD, if any service information contains procedures or tests that are identified as RC, those procedures and tests must be done to comply with this AD; any procedures or tests that are not identified as RC are recommended. Those procedures and tests that are not identified as RC may be deviated from using accepted methods in accordance with the operator’s maintenance or inspection program without obtaining approval of an AMOC, provided the procedures and tests identified as RC can be done and the airplane can be put back in an airworthy condition. Any substitutions or changes to procedures or tests identified as RC require approval of an AMOC.

(n) Related Information


(2) For service information identified in this AD, contact Airbus SAS, Airworthiness Office—EAL, 1 Rond Point Maurice Bellonte, 31707 Blagnac Cedex, France; telephone: +33 5 61 93 36 96; fax: +33 5 61 93 45 80; email: airworthiness.A330-A340@airbus.com; Internet: http://www.airbus.com. You may view this service information at the FAA, Transport Airplane Directorate, 1601 Lind Avenue SW., Renton, WA. For information on the availability of this material at the FAA, call 425–227–1221.

Issued in Renton, Washington, on April 28, 2016.

Dionne Palermo,
Acting Manager, Transport Airplane Directorate, Aircraft Certification Service.

[FR Doc. 2016–10633 Filed 5–6–16; 8:45 am]
BILLING CODE 4910–13–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Parts 1, 11, 16, and 111

[Docket No. FDA–2015–N–0797]

The Food and Drug Administration (FDA or we) is announcing three one-day public meetings in different regions throughout the United States to provide importers and other interested persons an opportunity to have an in-depth discussion on the implementation of the FDA Food Safety Modernization Act (FSMA) Import safety programs (i.e., foreign supplier verification programs (FSVPs) for importers of food for humans and animals, accreditation of third-party certification bodies, and FDA’s Voluntary Qualified Importer Program (VQIP)). During these meetings, participants and key FDA subject matter experts will discuss the next phase of FSMA implementation related to import safety programs, which includes establishing the operational framework for these programs and plans for guidance documents, training, education, and technical assistance. The purpose of the regional outreach public meetings is to continue the dialogue with the importer community on FSMA and eliciting ideas that will help to inform FDA and our stakeholders on how to continue to work together to successfully comply with FSMA mandates and regulations.

DATES: See section III for dates and times of the regional outreach meetings, closing dates for advance registration, and requests for special accommodations due to disability.

ADDRESSES: See section III for meeting locations.

FOR FURTHER INFORMATION CONTACT:
For questions about registering for the meeting, or to register by phone: Peggy Walker, Planning Professionals Ltd., 1210 West McDermott St., Suite 111, Allen, TX 75013, 214–384–0667, FAX: 469–854–0692, email: pwalker@planningprofessionals.com.

For general questions about the meeting or for special accommodations due to a disability: Juanita Yates, Center for Food Safety and Applied Nutrition (HFS–009), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740, 240–402–1731, email: juanita.yates@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

On May 2, 2014, we released our “Operational Strategy for Implementing the FDA Food Safety Modernization Act (FSMA),” electronically at http://www.fda.gov/Food/GuidanceRegulation/FSMA/ucm395105.htm, to guide the next phase of FSMA implementation following the establishment of regulations and relevant programs. Within the “Operational Strategy for Implementing FSMA,” there is an appendix that outlines guiding principles for how the operational strategy can be implemented with respect to food and feed facilities, produce safety standards, and import oversight. The guiding principles include the following: Expanding inspection and surveillance; administering new administrative enforcement tools; developing guidance, education, and technical assistance tools; and building a prevention-oriented import system.

On April 23, 2015, FDA hosted a public meeting as an opportunity to share views concerning how FDA should address the operational aspects of FSMA implementation as suggested by the guiding principles. We provided an update on current planning efforts and received input from the public to inform the development of operational work plans in the areas of produce safety, preventive controls for foods for humans and animals, measures to address intentional adulteration, FSVP, and the FDA third-party accreditation program. In addition, we established a docket to obtain comments on a range of operational issues that we might consider in our FSMA implementation approach.

On March 21, 2016, FDA hosted a kick-off public meeting to brief participants on the key components of the FSVP and third-party certification final rules; brief participants on the status of the VQIP; discuss the plans for guidance documents related to import safety, as well as training, education, and technical assistance; provide an update on the development of a risk-based industry oversight framework that is at the core of FSMA; and answer questions about these import programs. The public meeting was an opportunity for FDA to share its current thinking on implementation plans for programs related to import safety. During that public meeting, we mentioned plans to continue dialogue on implementation of these import safety programs with a series of regional meeting across the United States.

The agendas, recordings, and transcripts for the FSMA implementation and prevention-oriented import system public meetings are accessible on our FSMA Web site at http://www.fda.gov/FSMA.

II. Purpose and Format of the Regional Outreach Meetings

FDA plans to hold three one-day public meetings in different regions throughout the United States to provide importers and other interested persons an opportunity to have an in-depth discussion on the implementation of FSMA import safety programs (i.e., FSVPs for importers of food for humans and animals, accreditation of third-party certification bodies, and FDA’s VQIP). We invite the public to provide information, share experiences, and raise issues on implementation topics related to import safety including (but not limited to): Increasing awareness/reaching the regulated community, potential partners on outreach and implementation, state of readiness, barriers to implementation, training and education for industry and regulators, guidance needs, promotion of best practices, technical assistance, compliance and enforcement issues, and long-term implementation success. The purpose of the regional outreach meetings is to continue the dialogue with the importer community and elicit ideas that will help to inform FDA and the regulated population on how to continue to work together to successfully comply with FSMA mandates and regulations.

III. How To Participate in the Public Meeting

We are holding three one-day public meetings in different regions throughout the United States.

Due to limited space and time, we encourage all persons who wish to attend the meeting to register in advance. There is no fee to register for the regional outreach meetings, and registration will be on a first-come, first-served basis. Early registration is recommended because seating is very limited.

Table 1 provides information on participation in the regional outreach meetings.

<table>
<thead>
<tr>
<th>Regional outreach meetings</th>
<th>Date</th>
<th>Address</th>
<th>Preregister</th>
<th>Electronic address</th>
<th>Special accommodations</th>
<th>Other information</th>
</tr>
</thead>
<tbody>
<tr>
<td>California Regional Outreach Meeting</td>
<td>June 7, 2016, from 8:30 a.m. to 3 p.m. PDT.</td>
<td>The Hilton Costa Mesa, 3560 Bristol Street, Costa Mesa, CA 92626.</td>
<td>May 26, 2016: Closing date for Registration.</td>
<td>Please preregister at <a href="http://www.fda.gov/NewsEvents/WorkshopsMeetingsConferences/default.htm">http://www.fda.gov/NewsEvents/WorkshopsMeetingsConferences/default.htm</a>.</td>
<td>May 25, 2016: Closing date to request special accommodations due to a disability.</td>
<td>Registration check-in begins at 8 a.m.</td>
</tr>
<tr>
<td>New Jersey Regional Outreach Meeting</td>
<td>June 15, 2016, from 8:30 a.m. to 3 p.m. EDT.</td>
<td>Renaissance Meadowlands Hotel, 801 Rutherford Avenue, Rutherford, NJ 07070.</td>
<td>June 3, 2016: Closing date for Registration.</td>
<td>Please preregister at <a href="http://www.fda.gov/NewsEvents/WorkshopsMeetingsConferences/default.htm">http://www.fda.gov/NewsEvents/WorkshopsMeetingsConferences/default.htm</a>.</td>
<td>June 2, 2016: Closing date to request special accommodations due to a disability.</td>
<td>Registration check-in begins at 8 a.m.</td>
</tr>
</tbody>
</table>
I. Background

On February 3, 2012, HUD published in the Federal Register, at 77 FR 5662, a final rule titled “Equal Access to Housing in HUD Programs Regardless of Sexual Orientation or Gender Identity” (the Equal Access Rule) in order to address evidence that lesbian, gay, bisexual, and transgender (LGBT) individuals and families do not have equal access to housing, and to promote the federal goal of providing decent housing and a suitable living environment for all.

II. Definition of Terms

A. Definition of Terms

1. "Sexual Orientation or Gender Identity": Sexual orientation or gender identity means being or not being gay, lesbian, bisexual, or transgender (LGBT) or has a record of, or is perceived or believed to have had, a sexual orientation or gender identity. It is not limited to persons who self-identify as LGBT.

2. "Equal Access to Housing in HUD Programs Regardless of Sexual Orientation or Gender Identity Rule": The Department’s final rule titled “Equal Access to Housing in HUD Programs Regardless of Sexual Orientation or Gender Identity” published in the Federal Register on February 3, 2012, at 77 FR 5662, which incorporated the requirements of the Equal Access Rule.

III. Purpose and Need

The purpose of the proposed rule is to ensure equal access to decent, safe, and sanitary housing and a suitable living environment for all. The proposed rule would revise regulations to ensure that HUD-assisted and HUD-insured housing is available to all eligible individuals and families without regard to sex, sexual orientation, gender identity, or marital status. The proposed rule is necessary to remove any barriers to obtaining decent housing and a suitable living environment for all.

TABLE 1—INFORMATION ON PARTICIPATION IN THE MEETING—Continued

<table>
<thead>
<tr>
<th>REGIONAL OUTREACH MEETINGS</th>
<th>DATE</th>
<th>ADDRESS</th>
<th>PREREGISTER</th>
<th>ELECTRONIC ADDRESS</th>
<th>SPECIAL ACCOMMODATIONS</th>
<th>OTHER INFORMATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Michigan Regional Outreach Meeting</td>
<td>June 21, 2016, from 8:30 a.m. to 3 p.m. EDT.</td>
<td>Double Tree Suites by Hilton Hotel Detroit—Downtown Fort Shelby, 525 W Lafayette Blvd., Detroit, MI 48226.</td>
<td>June 10, 2016: Closing date for Registration.</td>
<td>Please preregister at <a href="http://www.fda.gov/Food/NewsEvents/WorkshopsMeetingsConferences/default.htm">http://www.fda.gov/Food/NewsEvents/WorkshopsMeetingsConferences/default.htm</a>.</td>
<td>June 9, 2016: Closing date to request special accommodations due to a disability.</td>
<td>Registration check-in begins at 8 a.m.</td>
</tr>
</tbody>
</table>

1. You may also register via email, mail, or fax. Please include your name, title, firm name, address, and phone and fax numbers in your registration information and send to: Peggy Walker, Planning Professionals Ltd., 1210 West McDermott St., Suite 111, Allen, TX 75013, 214–384–0667, FAX: 469–854–6992, email: pwalker@planningprofessionals.com.