The Proposed Amendment

Accordingly, under the authority delegated to me by the Administrator, the FAA proposes to amend 14 CFR part 39 as follows:

PART 39—AIRWORTHINESS DIRECTIVES

1. The authority citation for part 39 continues to read as follows:

Authority: 49 U.S.C. 106(g), 40113, 44701.

§ 39.13 [Amended]

2. The FAA amends § 39.13 by adding the following new airworthiness directive (AD):


(a) Comments Due Date

We must receive comments by December 30, 2013.

(b) Affected ADs

None.

(c) Applicability

This AD applies to all Pratt & Whitney Canada Corp. (P&WC) PT6A–114 and PT6A–114A turboprop engines.

(d) Reason

This AD was prompted by several incidents of compressor turbine (CT) blade failure, including two fatalities, resulting in power loss and in-flight shutdown (IFSD) of the engine. We are issuing this AD to prevent CT blade failure, which could lead to damage to the engine or to the airplane.

(e) Actions and Compliance

Comply with this AD within the compliance times specified, unless already done.

(1) Within 150 operating hours after the effective date of this AD, perform a borescope inspection (BSI) of CT blades for engines with 500 or more hours since new (TSN) that have not been previously inspected, or more than 500 flight hours since last inspection (TSLI).

(2) Thereafter, repeat the inspection in paragraph (e)(1) of this AD within every additional 500 flight hours TSLI.

(3) During the next hot section inspection (HSI) after the effective date of this AD, replace the complete set of CT blades with blades eligible for installation.

(4) If CT blades listed in paragraphs (g)(1) or (g)(2) of this AD, are installed to comply with paragraph (e)(3) of this AD, you must still comply with the 500-hour TSLI repetitive inspection requirement of paragraph (e)(2) of this AD.

(f) Optional Terminating Action

Replacing all CT blades with new CT blades, P/N 3072791–01, and Disk Balance Assembly, P/N 3072801–01; or with new CT blades, P/N 3072791–02, and Disk Balance Assembly, P/N 3072801–02; is terminating action for this AD.

(g) Definition

CT blades eligible for installation are:

(1) New CT blades, other than those listed in paragraphs (g)(3) and (g)(4) of this AD.

(2) CT blades, other than those listed in paragraphs (g)(3) and (g)(4) of this AD, that have met the inspection requirements of paragraphs (e)(1) and (e)(2) of this AD;

(3) CT blade, P/N 3072791–01, and Disk Balance Assembly, P/N 3072801–01; and

(4) CT blade, P/N 3072791–02, and Disk Balance Assembly, P/N 3072801–02.

(h) Credit for Previous Actions

If you performed P&WC Service Bulletin (SB) No. PT6A–72–1669, Revision 9, dated June 28, 2013, or earlier versions, you have met the initial inspection requirements of this AD. However, you must still comply with the 500-hour TSLI repetitive inspection requirement of paragraph (e)(2) of this AD.

(i) Alternative Methods of Compliance (AMOCs)

The Manager, Engine Certification Office, FAA, may approve AMOCs to this AD. Use the procedures found in 14 CFR 39.19 to make your request.

(j) Related Information


(3) For guidance on the initial and repetitive BSIs mandated by this AD, refer to P&W SB No. PT6A–72–1669 and P&W SB No. PT6A–72–1727, which are not incorporated by reference in this AD. The BSIs can be obtained from Pratt & Whitney Canada Corp. using the contact information in paragraph (j)(4) of this AD.

(4) For service information identified in this AD, contact Pratt & Whitney Canada Corp., 1000 Marie-Victorin, Longueuil, Quebec, Canada, J4G 1A1; phone: 800–266–8000; fax: 450–647–2888; Internet: www.pwc.ca.

(5) You may view this service information at the FAA, Engine & Propeller Directorate, 12 New England Executive Park, Burlington, MA. For information on the availability of this material at the FAA, call 781–238–7125.

Issued in Burlington, Massachusetts, on October 7, 2013.

Colleen M. D’Alessandro,
Assistant Directorate Manager, Engine & Propeller Directorate, Aircraft Certification Service.

BILING CODE 4910–13–P

FEDERAL TRADE COMMISSION

16 CFR Part 312

RIN 3084–AB20

Children’s Online Privacy Protection Rule Applications for Approval of Proposed Parental Consent Methods by AssertID, Inc., Imperium LLC, and iVeriFly, Inc.; Application for Approval of Safe Harbor Program by kidSAFE Seal Program

AGENCY: Federal Trade Commission (FTC or Commission).

ACTION: Notice of extension of Commission determination and public comment deadlines.

SUMMARY: The FTC is extending the deadlines for Commission determination of applications for approval of proposed parental consent methods by AssertID, Inc. (“AssertID”), Imperium LLC (“Imperium”), and iVeriFly, Inc. (“iVeriFly”) pursuant to the Children’s Online Privacy Protection Rule. In addition, the FTC is extending the deadline for filing public comments concerning Imperium’s application for approval of a parental consent method and the proposed self-regulatory guidelines submitted by the kidSAFE Seal Program (“kidSAFE”).

DATES: Written comments must be received by November 4, 2013.

ADDRESSES: Interested parties may file comments online or on paper, by following the instructions in the Request for Comment part of the SUPPLEMENTARY INFORMATION section below. For comments concerning Imperium, write “Imperium Application for Parental Consent Method, Project No. P–135419” on your comment and file your comment online at https://ftcpublic.commentworks.com/ftc/ pncoppaimperiumapp, by following the instructions on the web-based form. For comments concerning kidSAFE, write “kidSAFE Application for Safe Harbor, Project No. P–135418” on your comment and file your comment online at https://ftcpublic.commentworks.com/ftc/ coppakidsafeapp, by following the instructions on the web-based form. If you prefer to file your comment on paper, mail or deliver your comment to the following address: Federal Trade Commission, Office of the Secretary, Room H–113 (Annex E), 600 Pennsylvania Avenue NW., Washington, DC 20580.
determinations on the applications. The federal government re-opened on October 17, 2013. In order to ensure that it can give full consideration to the applications submitted by AssertID, Imperium, and iVeriFly, the Commission has determined to extend the timetable laid out in 16 CFR 312.12(a) by sixteen days in order to account for the time period in which the government was shut down. Accordingly, the Commission will issue its determination for AssertID by November 13, 2013; for Imperium by December 26, 2013; and for iVeriFly by February 4, 2014.

In addition, during the time when the government was shut down, interested parties were unable to submit comments on Imperium’s application for approval of a parental consent method or on kidSAFE’s proposed self-regulatory guidelines. The Commission has decided to extend the comment period for both matters until November 4, 2013.

Section B. Invitation to Comment
You can file a comment online or on paper. For the Commission to consider your comment, we must receive it on or before November 4, 2013. For comments concerning Imperium, write “Imperium Application for Parental Consent Method, Project No. P–135419” on your comment. For comments concerning kidSAFE, write “kidSAFE Application for Safe Harbor, Project No. P–135418” on your comment. Your comment—including your name and your state—will be placed on the public record of this proceeding, including, to the extent practicable, on the Commission Web site, at http://www.ftc.gov/os/publiccomments.shtm. As a matter of discretion, the Commission tries to remove individuals’ home contact information from comments before placing them on the Commission Web site.

Because your comment will be made public, you are solely responsible for making sure that your comment doesn’t include any sensitive personal information, such as Social Security number, date of birth, driver’s license number or other state identification number or foreign country equivalent, passport number, financial account number, or credit or debit card number. You are also solely responsible for making sure that your comment doesn’t include any sensitive health information, including medical records or other individually identifiable health information. In addition, do not include any “[t]rade secret or any commercial or financial information which is obtained from any person and which is privileged or confidential,” as provided in Section 6(f) of the FTC Act, 15 U.S.C. 46(f), and FTC Rule 4.10(a)(2), 16 CFR 4.10(a)(2).

In particular, do not include competitively sensitive information such as costs, sales statistics, inventories, formulas, patterns, devices, manufacturing processes, or customer names.

If you want the Commission to give your comment confidential treatment, you must file it in paper form, with a request for confidential treatment, and follow the procedure explained in FTC Rule 4.9(c), 16 CFR 4.9(c). Your comment will be kept confidential only if the FTC General Counsel, in his or her sole discretion, grants your request in accordance with the law and the public interest.

Postal mail addressed to the Commission is subject to delay due to heightened security screening. As a result, we encourage you to submit your comments online. To make sure that the Commission considers your online comment, you must file your comment concerning Imperium at https://ftcpublic.commentworks.com/ftc/pncoppaimperiumapp, and your comment concerning kidSAFE at https://ftcpublic.commentworks.com/ftc/coppakidsafeapp, by following the instructions on the web-based form. If this Notice appears at http://www.regulations.gov/#/home, you also may file a comment through that Web site.

If you file your comment concerning Imperium on paper, write “Imperium Application for Parental Consent Method, Project No. P–135419” on your comment and on the envelope, and mail or deliver it to the following address: Federal Trade Commission, Office of the Secretary, Room H–113 (Annex E), 600 Pennsylvania Avenue NW., Washington, DC 20580. If you file your comment concerning kidSAFE on paper, write “kidSAFE Application for Safe Harbor, Project No. P–135418” on your comment and on the envelope, and mail or deliver it to the following address: Federal Trade Commission, Office of the Secretary, Room H–113 (Annex E), 600 Pennsylvania Avenue NW., Washington, DC 20580. If possible, submit your paper comment to the Commission by courier or overnight service.

Visit the Commission Web site at http://www.ftc.gov to read this Notice and the news release describing it. The FTC Act and other laws that the Commission administers permit the

1 64 FR 59888 (1999).
2 16 CFR part 312.
3 78 FR 3972 (2013).
4 16 CFR 312.12(a); 78 FR at 3991–3992, 4013.
5 16 CFR 312.11; 78 FR at 3995–96, 4012–13.
6 16 CFR 312.12(a).

In particular, the written request for confidential treatment that accompanies the comment must include the factual and legal basis for the request, and must identify the specific portions of the comment to be withheld from the public record. See FTC Rule 4.9(c), 16 CFR 4.9(c).
collection of public comments to consider and use in this proceeding as appropriate. The Commission will consider all timely and responsive public comments that it receives on or before November 4, 2013. You can find more information, including routine uses permitted by the Privacy Act, in the Commission’s privacy policy, at http://www.ftc.gov/privacy.htm.

By direction of the Commission.

Donald S. Clark, Secretary.

[FR Doc. 2013–25452 Filed 10–28–13; 8:45 am]

BILLING CODE 6750–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Parts 12, 225, 500, 507, and 579

[Docket No. FDA–2011–N–0922]

Current Good Manufacturing Practice and Hazard Analysis and Risk-Based Preventive Controls for Food for Animals; Public Meeting on Proposed Rule

AGENCY: Food and Drug Administration, HHS.

ACTION: Notification of public meeting.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing three public meetings to discuss the proposed rule to establish requirements for current good manufacturing practice and hazard analysis and risk-based preventive controls for animal food. This proposed rule is one of several proposed rules that will establish the foundation of, and central framework for, the modern food safety system envisioned by Congress in the FDA Food Safety Modernization Act (FSMA). The purpose of the public meetings is to inform the public of the provisions of the proposed rule and the rulemaking process (including how to submit comments, data, and other information to the rulemaking docket) as well as solicit oral stakeholder and public comments on the proposed rule and to respond to questions about the proposed rule.

DATES: See section II, “How to Participate in the Public Meetings,” in the SUPPLEMENTARY INFORMATION section.

ADDITIONAL INFORMATION: See section II, “How to Participate in the Public Meetings,” in the SUPPLEMENTARY INFORMATION section.

FOR FURTHER INFORMATION CONTACT: For general questions about the meeting, for assistance to register for the meeting, to request an opportunity to make an oral presentation, or to request special accommodations due to a disability, contact: Aleta Sindelar, Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Pl., Rm. 133, Rockville, MD 20855, 240–276–9230, FAX: 240–276–9241, email: aleta.sindelar@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

FSMA (Pub. L. 111–353) was signed into law by President Obama on January 4, 2011, to better protect public health by helping to ensure the safety and security of the food supply. FSMA amends the Federal Food, Drug, and Cosmetic Act (the FD&C Act) to establish the foundation of a modernized, prevention-based food safety system. Among other things, FSMA requires FDA to issue regulations requiring preventive controls for human food and animal food, set standards for produce safety, and require importers to have a program to verify that the food products they bring into the United States are produced in a manner consistent with applicable FDA food safety requirements.

FSMA was the first major legislative reform of FDA’s food safety authorities in more than 70 years, even though FDA has increased the focus of its food safety efforts on prevention over the past several years. The proposed rule for preventive controls for food for animals can be found elsewhere in this issue of the Federal Register, and it establishes a docket so that the public can review the proposed rule and submit comments to FDA. This proposed rulemaking is one of several key proposals in furtherance of FSMA’s food safety mandate.

The proposed rule would establish regulations regarding the manufacturing, processing, packing, or holding of animal food in two ways. First, it would create new current good manufacturing practice (CGMP) regulations that specifically address the manufacturing, processing, packing, and holding of animal food. Second, it would include new preventive control provisions intended to implement section 103 of FSMA for animal food. In general, with some exceptions the new preventive control provisions would apply to animal food facilities that are required to register with FDA under the FD&C Act. These preventive controls would include requirements for covered facilities to maintain a food safety plan, perform a hazard analysis, and institute preventive controls for the mitigation of those hazards. Facilities would also be required to monitor their controls, verify that they were effective, take appropriate corrective actions, and maintain records documenting these actions.

For information on the proposed rule for preventive controls for food for animals and related fact sheets, see FDA’s FSMA Web page located at www.fda.gov/FSMA.

II. How To Participate in the Public Meetings

FDA is holding the public meetings on the proposed rule for preventive controls for food for animals to inform the public about the proposed rule and the rulemaking process, including how to submit comments, data, and other information to the rulemaking docket; to respond to questions about the proposed rule; and to provide an opportunity for interested persons to make oral presentations. Due to limited space and time, FDA encourages all persons who wish to attend the meetings to register in advance. There is no fee to register for the public meetings, and registration will be on a first-come, first-served basis. Early registration is recommended because seating is limited. Onsite registration will be accepted, as space permits, after all preregistered attendees are seated.

Those requesting an opportunity to make an oral presentation during the time allotted for public comment at the meeting are asked to submit a request and to provide the specific topic or issue to be addressed. Due to the anticipated high level of interest in presenting public comment and limited time available, FDA is allocating 3 minutes to each speaker to make an oral presentation. Speakers will be limited to making oral remarks; there will not be an opportunity to display materials such as slide shows, videos, or other media during the meeting. If time permits, individuals or organizations that did not register in advance may be granted the opportunity to make an oral presentation. FDA would like to maximize the number of individuals who make a presentation at the meeting and will do our best to accommodate all persons who wish to make a presentation or express their opinions at the meeting.

FDA encourages persons and groups who have similar interests to consolidate their information for presentation by a single representative. After reviewing the presentation requests, FDA will notify each participant before the meeting of the approximate time their presentation is scheduled to be held and remind them of the presentation format (i.e., 3-minute oral presentation without visual media).