

with the Secretary [of Commerce] and the Secretary of Defense, and the heads of other Federal agencies, as appropriate, shall propose that any technology identified pursuant to subsection (a) [of ECRA] [which addresses the interagency process for identifying Section 1758 technologies] be added to the list of technologies controlled by the relevant multilateral export control regimes.”

Finally, BIS encourages comments addressing any other automated peptide synthesizer technology topics deemed to be relevant to this inquiry.

Comments should be submitted as described in the **ADDRESSES** section of this ANPRM and must be received no later than October 28, 2022.

This ANPRM has been designated a “significant regulatory action,” although not economically significant, under Executive Order 12866. Accordingly, this ANPRM has been reviewed by the Office of Management and Budget (OMB).

Matthew S. Borman,

Deputy Assistant Secretary for Export Administration.

[FR Doc. 2022–19430 Filed 9–12–22; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 1

[Docket No. FDA–2011–N–0179]

Prior Notice of Imported Food Questions and Answers (Edition 4); Draft Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing the availability of a draft guidance for industry entitled “Prior Notice of Imported Food Questions and Answers; Draft Guidance for Industry (Edition 4).” The draft guidance adds three additional questions. One question relates to any effect systems recognition or equivalency determinations have on prior notice requirements. The other two questions relate to FDA’s notice to a submitter of prior notice of an FDA refusal for inadequate prior notice or hold if the food article is from a foreign facility that is not registered, and address the timeframe for making requests for FDA review of such a

refusal or hold. FDA is also making other technical and editorial changes.

DATES: Submit either electronic or written comments on the draft guidance by November 14, 2022 to ensure that we consider your comment on this draft guidance before we begin work on the final version of the guidance.

ADDRESSES: You may submit comments on any guidance at any time as follows:

Electronic Submissions

Submit electronic comments in the following way:

- *Federal eRulemaking Portal:* <https://www.regulations.gov>. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to <https://www.regulations.gov> will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on <https://www.regulations.gov>.

- If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).

Written/Paper Submissions

Submit written/paper submissions as follows:

- *Mail/Hand Delivery/Courier (for written/paper submissions):* Dockets Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

- For written/paper comments submitted to the Dockets Management Staff, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

Instructions: All submissions received must include the Docket No. FDA–2011–N–0179 for “Prior Notice of Imported Food Questions and Answers; Draft Guidance for Industry (Edition 4).” Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at <https://www.regulations.gov> or at the

Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday, 240–402–7500.

- *Confidential Submissions—*To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states “THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION.” FDA will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on <https://www.regulations.gov>. Submit both copies to the Dockets Management Staff. If you do not wish your name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and you must identify this information as “confidential.” Any information marked as “confidential” will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA’s posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: <https://www.govinfo.gov/content/pkg/FR-2015-09-18/pdf/2015-23389.pdf>.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to <https://www.regulations.gov> and insert the docket number, found in brackets in the heading of this document, into the “Search” box and follow the prompts and/or go to the Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852, 240–402–7500.

You may submit comments on any guidance at any time (see 21 CFR 10.115(g)(5)). Submit written requests for single copies of the draft guidance to the Division of Operational Policy, Office of Regulatory Affairs, Food and Drug Administration, Element Building, 12420 Parklawn Dr., Rockville, MD 20852. Send one self-addressed adhesive label to assist that office in processing your requests. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the draft guidance document.

FOR FURTHER INFORMATION CONTACT: Chris Henderson, Office of Regulatory Affairs, Food and Drug Administration, Element Building, 12420 Parklawn Dr.,

Rockville, MD 20857 240-402-8186,
Christopher.Henderson@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is announcing the availability of a draft guidance for industry, entitled “Prior Notice of Imported Food Questions and Answers; Draft Guidance for Industry (Edition 4).” This draft revised guidance is being issued for public comment and has not yet been finalized. Until edition 4 is finalized, “Prior Notice of Imported Food Questions and Answers; Guidance for Industry (Edition 3),” updated most recently in 2016, remains in effect. We are issuing the draft guidance consistent with our good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the current thinking of FDA on this topic. It does not establish any rights for any person and is not binding on FDA or the public. You can use an alternate approach if it satisfies the requirements of the applicable statutes and regulations.

FDA continues to believe that it is reasonable to maintain responses to questions concerning prior notice of imported food in a single document that is periodically updated in response to additional questions or regulatory or policy changes. As in the previous editions, the following indicators are used to help users identify revisions: (1) the guidance is identified as a revision of a previously issued document; (2) the revision date appears on the cover of the guidance; (3) the edition number of the guidance is included in its title; and (4) revised or added questions and answers are identified as such in the body of the guidance.

On November 7, 2008, we published a final rule in the **Federal Register** requiring submission to FDA of prior notice of food, including food for animals, that is imported or offered for import into the United States (73 FR 66294). The rule implements section 801(m) of the Federal Food, Drug, and Cosmetic Act (the FD&C Act) (21 U.S.C. 381(m)), which was added by section 307 of the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 (the Bioterrorism Act) (Pub. L. 107-188) and requires that FDA receive prior notice of food imported or offered for import into the United States.

On December 16, 2003, FDA issued a guidance entitled “Prior Notice of Imported Food Questions and Answers (Edition 1).” FDA issued a second and third edition on May 3, 2004, and June 16, 2016, respectively. This draft will be the fourth edition of this document.

FDA is issuing this draft guidance entitled “Prior Notice of Imported Food Questions and Answers (Edition 4)” as a level 1 guidance.

The draft fourth edition guidance adds three additional questions. One question relates to any effect systems recognition or equivalency determinations have on prior notice requirements. The other two questions relate to FDA’s notice of a refusal under 801(m)(1) of the FD&C Act (in accordance with § 1.283 (21 CFR 1.283)) for inadequate prior notice or a hold under 801(l) (in accordance with § 1.285 (21 CFR 1.285)) if the food article is from a foreign facility that is not registered, as well as address the timeframe for making requests for FDA review of such a refusal or hold. The draft guidance is intended to help clarify whether food imported from a country with which FDA has a Systems Recognition Arrangement or equivalence determination is exempted from prior notice requirements. The draft guidance also intends to clarify when FDA will provide notice of the refusal or hold to the relevant party, and when the 5-calendar-day clock to request a review of the refusal or hold begins. We are also making other technical amendments to the guidance due to the expanded capabilities of the U.S. Customs and Border Protection’s Automate Broker Interface of the Automated Commercial Environment (ABI/ACE) system and FDA’s 2017 technical amendments to the prior notice rule (82 FR 15627, March 20, 2017), such as replacing references to the Automated Commercial System (ACS) and successor system with the ABI/ACE system, removing references to requirements that certain prior notice submissions be submitted in FDA’s Prior Notice Systems Interface (FDA PNSI), and updating outdated links and FDA contact information.

The draft fourth edition guidance clarifies that the existence of a Systems Recognition Arrangement with or an equivalence determination of a foreign country does not exempt imported foods from that country from FDA’s prior notice requirements.

FDA’s policy on and practice of communicating prior notice refusals and holds has changed over time. FDA previously stated that we intended to provide notice regarding refusals to carriers. Those carriers could then notify others, such as the entity that hired the carrier to transport the article of food, of a problem with the prior notice (see 73 FR 66294 at 66365). Subsequently, FDA’s Guidance for Industry “Prior Notice of Imported Food Questions and Answers (Edition 3)” was

published with the explanation that FDA will communicate the decision to examine articles of food to CBP.

The draft fourth edition clarifies that notification of these prior notice refusals and holds will be communicated to CBP and provided to the relevant party (*i.e.*, the submitter of prior notice) upon arrival of the article. FDA is clarifying its policy because providing advanced notice of a refusal or hold to a submitter could create incentives for bad actors, who may attempt to reroute their entries for the purpose of evading FDA requirements and importing unsafe food.

The draft fourth edition also clarifies the 5-calendar-day clock to request a review of these refusals and holds. Under §§ 1.283(d) and 1.285(j), certain parties may, for the enumerated reasons, request reviews of the prior notice refusals and holds within 5 calendar days of the hold or refusal. The draft fourth edition clarifies that FDA considers the 5-calendar-day clock to begin when FDA provides notice of the refusal or hold to the submitter.

Additionally, in 2016, CBP issued a notice announcing that ABI/ACE would replace ACS as the sole electronic data interchange system authorized by CBP for the processing of electronic entries of FDA-regulated products (see 81 FR 30320, May 16, 2016). ABI/ACE became the successor system to ACS. In 2017, we amended 21 CFR subpart I to replace references to ACS and successor system with ABI/ACE (see 82 FR 15627). As part of this rulemaking, we eliminated some requirements for submitting prior notice due to the expanded capabilities of ABI/ACE, such as the requirement to submit articles that have been refused under section 801(m)(1) of the FD&C Act or subpart I in FDA PNSI. Further, ABI/ACE can now accommodate entries it previously could not, such as articles of food arriving through international mail. Therefore, to reflect these changes that were implemented in the rulemaking and the expanded capabilities of ABI/ACE, we are replacing references in the guidance to ACS and successor system with ABI/ACE. In addition, we are providing clarification regarding how persons may submit prior notice for articles of food imported or offered for import by international mail.

This draft guidance is being issued consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the current thinking of FDA on “Prior Notice of Imported Food Questions and Answers (Edition 4).” It does not establish any rights for any person and is not binding on FDA or the

public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations.

II. Paperwork Reduction Act of 1995

While this guidance contains no collection of information, it does refer to previously approved FDA collections of information. Therefore, clearance by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (PRA) (44 U.S.C. 3501–3521) is not required for this guidance. The previously approved collections of information are subject to review by OMB under the PRA. The collections of information in 21 CFR 1.278 to 1.282 have been approved under OMB control number 0910–0520.

III. Electronic Access

Persons with access to the internet may obtain the draft guidance at <https://www.fda.gov/food/importing-food-products-united-states/prior-notice-imported-foods>, <https://www.fda.gov/regulatory-information/search-fda-guidance-documents>, or <https://www.regulations.gov>. Use the FDA website listed in the previous sentence to find the most current version of the guidance.

Dated: September 7, 2022.

Lauren K. Roth,

Associate Commissioner for Policy.

[FR Doc. 2022–19724 Filed 9–12–22; 8:45 am]

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DEPARTMENT OF THE TREASURY

Internal Revenue Service

26 CFR Part 301

[REG–125693–19]

RIN 1545–BP72

Resolution of Federal Tax Controversies by the Independent Office of Appeals

AGENCY: Internal Revenue Service (IRS), Treasury.

ACTION: Notice of proposed rulemaking and notice of public hearing on proposed rulemaking.

SUMMARY: This document contains proposed regulations relating to the IRS Independent Office of Appeals' resolution of Federal tax controversies without litigation and relating to requests for referral to that office following the issuance of a notice of deficiency to a taxpayer by the IRS. The proposed regulations reflect amendments to the law made by the

Taxpayer First Act of 2019. The proposed regulations apply to all persons that request to have a Federal tax controversy considered by that office. This document also provides a notice of a public hearing on these proposed regulations.

DATES: Written or electronic comments must be received by November 14, 2022. Outlines of topics to be discussed at the public hearing scheduled for November 29, 2022, must be received by November 14, 2022. If no outlines of topics are received by November 14, 2022, the public hearing will be cancelled.

ADDRESSES: Commenters are strongly encouraged to submit public comments electronically. Submit electronic submissions via the Federal eRulemaking Portal at www.regulations.gov (indicate IRS and REG–125693–19) by following the online instructions for submitting comments. Once submitted to the Federal eRulemaking Portal, comments cannot be edited or withdrawn. The Department of the Treasury (Treasury Department) and the IRS will publish for public availability any comment to its public docket. Send paper submissions to: CC:PA:LPD:PR (REG–125693–19), Room 5203, Internal Revenue Service, PO Box 7604, Ben Franklin Station, Washington, DC 20044.

FOR FURTHER INFORMATION CONTACT:

Concerning the proposed regulations, Keith L. Brau at (202) 317–5437 (not a toll-free number). Concerning submissions of comments or the public hearing, Regina Johnson, preferably at publichearings@irs.gov or (202) 317–6901 (not a toll-free number).

SUPPLEMENTARY INFORMATION:

Background

I. Overview

This document contains proposed amendments to the Procedure and Administration Regulations (26 CFR part 301) to implement section 7803(e) of the Internal Revenue Code (Code). The proposed amendments (proposed regulations) relate to the resolution by the IRS Independent Office of Appeals (Appeals) of Federal tax controversies without litigation, including guidance regarding requests for referral to Appeals following the issuance of a notice of deficiency. (References in this preamble to “Appeals” include references to the former Office of Appeals where appropriate.)

Since its establishment by the IRS in 1927, Appeals' mission has been to resolve Federal tax controversies without litigation on a basis that is fair

and impartial to both the Government and the taxpayer.¹ In doing so, Appeals has independently considered disputed administrative determinations made by the IRS in administering and enforcing the internal revenue laws arising from the IRS's examination or collection activities with respect to a particular taxpayer, and attempted to resolve those disputes without litigation. See House TFA Report, at 29. Appeals generally considers whether to resolve Federal tax controversies without litigation based on the likelihood of either the taxpayer's or the IRS's position prevailing if the Federal tax controversy was resolved before a court. When Appeals resolves a Federal tax controversy, it does so through an administrative settlement of the matter.

The IRS Restructuring and Reform Act of 1998 (RRA), Public Law 105–206 (112 Stat. 685, 689 (1998)) directed the Commissioner to restructure the IRS by establishing and implementing an organizational structure that ensured an independent appeals function within the IRS. Although the Code did not mandate the existence of an independent office within the IRS, provisions of the Code have required the independent administrative review of certain administrative determinations, such as section 6159 regarding terminating an installment agreement, sections 6320 and 6330 regarding notice and an opportunity for a hearing before a levy or upon the filing of a notice of lien, and section 7122 regarding rejections of an offer in compromise (OIC).

For decades the Internal Revenue Manual (IRM) has contained the mission statement of Appeals (Appeals Mission Statement), which is “to resolve [Federal] tax controversies, without litigation, on a basis which is fair and impartial to both the Government and the taxpayer and in a manner that will enhance voluntary compliance and public confidence in the integrity and efficiency of the Service.” See IRM

¹ See H.R. Rep. No. 39 Part 1, 116th Cong., 1st Session (House TFA Report), 28–29, fn. 4 (2019). The House TFA Report states that Appeals was established and has operated under the general authority of the Secretary of the Treasury or her delegate (Secretary) provided by section 7805 of the Code to interpret the Code, and the authority of the Commissioner of Internal Revenue (Commissioner) provided by section 7803 to, among other things, “administer, manage, conduct, direct, and supervise the execution and application of the internal revenue laws or related statutes and tax conventions to which the United States is a party,” and by section 7804 to, among other things, “employ such number of persons as the Commissioner deems proper for the administration and enforcement of the internal revenue laws, and the Commissioner shall issue all necessary directions, instructions, orders, and rules applicable to such person.” Sections 7803(a)(2)(A) and 7804(a).