

FIGURE 1 TO PARAGRAPHS (h) AND (j) OF THIS AD—PART NUMBERS AND SERIAL NUMBERS OF AFFECTED FORWARD ENGINE MOUNTS AND ATTACHMENT PINS—Continued

Serial No.	
Attachment Pin (P/N 740–2022–501)	Forward Engine Mount (P/N 745–2010–503)
1467SC	13813001
1445SC	13755001
1462SC	13789001
1464SC	13793001
1466SC	13811001
1470SC	13819001
1459SC	13783001
1463SC	13791001
1475SC	13829001
1458SC	13781001
1477SC	13833001
1474SC	13827001
1478SC	13835001
1479SC	13837001
1472SC	13823001

(i) Exception to Paragraph (g) of This AD

For airplanes with manufacturer serial numbers identified in figure 2 to paragraph (i) of this AD: If it can be conclusively determined that an engine has not been replaced after March 1, 2011 (the date of manufacture of the first airplane with affected engine mounts), the airplane is not affected by the requirements of paragraphs (g) and (h) of this AD.

FIGURE 2 TO PARAGRAPH (i) OF THIS AD—AIRPLANE MANUFACTURER SERIAL NUMBERS

Airplane manufacturer serial Nos.
4593
4602
4620
4637
4638
4642
4643
4644
4660
4677
4690
4696
4700
4701
4703
4706
4707
4710
4716
4719
4725
4726
4731
4736
4737
4741
4746
4751
4752

FIGURE 2 TO PARAGRAPH (i) OF THIS AD—AIRPLANE MANUFACTURER SERIAL NUMBERS—Continued

Airplane manufacturer serial Nos.
4753
4754
4755
4757
4761
4762
4772
4773
4774
4775
4779
4782
4783
4784
4786
4788
4790
4791
4798
4804
4813

(j) Parts Installation Prohibition

As of the effective date of this AD, no person may install on any airplane any engine mount attachment pin having P/N 740–2022–501 with a serial number identified in figure 1 to paragraphs (h) and (j) of this AD.

(k) Other FAA AD Provisions

The following provisions also apply to this AD:

(1) *Alternative Methods of Compliance (AMOCs)*: The Manager, International Branch, ANM–116, Transport Airplane Directorate, FAA, has the authority to approve AMOCs for this AD, if requested using the procedures found in 14 CFR 39.19. In accordance with 14 CFR 39.19, send your request to your principal inspector or local Flight Standards District Office, as appropriate. If sending information directly to the International Branch, send it to ATTN: Sanjay Ralhan, Aerospace Engineer, International Branch, ANM–116, Transport Airplane Directorate, FAA, 1601 Lind Avenue SW., Renton, WA 98057–3356; telephone 425–227–1405; fax 425–227–1149. Information may be emailed to: 9-ANM-116-AMOC-REQUESTS@faa.gov. 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 EASA; or Airbus's EASA DOA. If approved by the DOA, the approval must include the DOA-authorized signature.

(3) *Required for Compliance (RC)*: 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 a serviceable condition. Any substitutions or changes to procedures or tests identified as RC require approval of an AMOC.

(l) Special Flight Permits Prohibited

Special flight permits, as described in Section 21.197 and Section 21.199 of the Federal Aviation Regulations (14 CFR 21.197 and 21.199), are not allowed.

(m) Related Information

(1) Refer to Mandatory Continuing Airworthiness Information (MCAI) EASA Airworthiness Directive 2015–0004, dated January 13, 2015, for related information. This MCAI may be found in the AD docket on the Internet at <http://www.regulations.gov> by searching for and locating Docket No. FAA–2015–2963.

(2) For Airbus service information identified in this AD, contact Airbus, Airworthiness Office—EIAS, 1 Rond Point Maurice Bellonte, 31707 Blagnac Cedex, France; telephone +33 5 61 93 36 96; fax +33 5 61 93 44 51; email account.airworth-eas@airbus.com; Internet <http://www.airbus.com>. For Goodrich Aerostructures service information identified in this AD, contact UTC Aerospace Systems, ATTN: Christopher Newth—V2500 A1/A5 Project Engineer, Aftermarket—Aerostructures; 850 Lagoon Drive, Chula Vista, CA; telephone 619–498–7505; email christopher.newth@utas.utc.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 July 17, 2015.

Michael Kaszycki,

Acting Manager, Transport Airplane Directorate, Aircraft Certification Service.

[FR Doc. 2015–18533 Filed 7–29–15; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 101

[Docket No. FDA–2015–D–1839]

The Food and Drug Administration's Policy on Declaring Small Amounts of Nutrients and Dietary Ingredients on Nutrition Labels; Draft Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notification of availability.

SUMMARY: The Food and Drug Administration (FDA, we, or the Agency) is announcing the availability of a draft guidance for industry entitled “FDA’s Policy on Declaring Small Amounts of Nutrients and Dietary Ingredients on Nutrition Labels: Guidance for Industry.” The draft guidance, when finalized, will explain to manufacturers of conventional foods and dietary supplements our policy on determining the amount to declare on the nutrition label for certain nutrients and dietary ingredients that are present in a small amount.

DATES: Although you can comment on any guidance at any time (see 21 CFR 10.115(g)(5)), to ensure that the Agency considers your comment on the draft guidance before it begins work on the final version of the guidance, submit either electronic or written comments on the draft guidance by September 28, 2015.

ADDRESSES: Submit written requests for single copies of the draft guidance to the Office of Nutrition, Labeling, and Dietary Supplements, Center for Food Safety and Applied Nutrition (HFS–820), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740. Send two self-addressed adhesive labels to assist the office in processing your request. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the draft guidance.

Submit electronic comments on the draft guidance to <http://www.regulations.gov>. Submit written comments on the draft guidance to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT: Carole Adler, Center for Food Safety and Applied Nutrition (HFS–820), Food and Drug Administration, 5100 Paint Branch Pkwy., College Park, MD 20740, 240–402–2371.

SUPPLEMENTARY INFORMATION:**I. Background**

We are announcing the availability of a draft guidance for industry entitled “FDA’s Policy on Declaring Small Amounts of Nutrients and Dietary Ingredients on Nutrition Labels: Guidance for Industry.” We are issuing the draft guidance consistent with our good guidance practices regulation (21 CFR 10.115). The draft guidance represents the current thinking of FDA on our policy on declaring small amounts of nutrients and dietary ingredients on nutrition labels. 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.

The draft guidance, when finalized, will explain our nutrition labeling policy on declaring the nutrient values in conventional foods and dietary ingredient values in dietary supplements in certain cases. Specifically, declaring small amounts of nutrients and dietary ingredients in the nutrition labeling may result in a conflict between 21 CFR 101.9(c)(1) through (8) and 21 CFR 101.9(g)(4)(ii) and 21 CFR 101.9(g)(5). In such cases, we are recommending manufacturers declare nutrients and dietary ingredients in accordance with § 101.9(c)(1) through (8). If the draft guidance is finalized, we intend to consider the use of our enforcement discretion with respect to the compliance requirements in § 101.9(g)(4)(ii) and § 101.9(g)(5) when a conflict exists with § 101.9(c)(1) through (8).

We also are considering whether changes to our nutrition labeling regulations are needed, including changes to § 101.9(c) or (g), or both. If we determine that rulemaking is needed, we will consider whether to revise or withdraw the draft guidance.

II. Paperwork Reduction Act of 1995

The draft guidance refers to previously approved collections of information found in FDA regulations. These collections of information are subject to review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501–3520). The collections of information in § 101.9 have been approved under OMB control number 0910–0381.

III. Comments

Interested persons may submit either electronic comments regarding the draft guidance to <http://www.regulations.gov> or written comments regarding the draft guidance to the Division of Dockets Management (see **ADDRESSES**). It is only necessary to send one set of comments. Identify comments with the docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday, and will be posted to the docket at <http://www.regulations.gov>.

IV. Electronic Access

Persons with access to the Internet may obtain the draft guidance document

at <http://www.fda.gov/FoodGuidances> or <http://www.regulations.gov>. Use the FDA Web site listed in the previous sentence to find the most current version of the guidance.

Dated: July 24, 2015.

Leslie Kux,

Associate Commissioner for Policy.

[FR Doc. 2015–18655 Filed 7–29–15; 8:45 am]

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DEPARTMENT OF THE TREASURY**Internal Revenue Service****26 CFR Part 1**

[REG–138526–14]

RIN 1545–BM46

Issue Price Definition for Tax-Exempt Bonds; Correction

AGENCY: Internal Revenue Service (IRS), Treasury.

ACTION: Partial withdrawal of notice of proposed rulemaking, notice of proposed rulemaking, and notice of public hearing; correction.

SUMMARY: This document contains corrections to partial withdrawal of notice of proposed rulemaking, notice of proposed rulemaking, and notice of public hearing; correction (REG–138526–14) that were published in the **Federal Register** on Wednesday, June 24, 2015 (80 FR 36301). The partial withdrawal of notice of proposed rulemaking, notice of proposed rulemaking, and notice of public hearing are relating to the definition of issue price for purposes of the arbitrage restrictions under section 148 of the Internal Revenue Code (Code).

DATES: Written or electronic comments and requests for a public hearing for the notice of proposed rulemaking published at 80 FR 36301, June 24, 2015, are still being accepted and must be received by September 22, 2015.

FOR FURTHER INFORMATION CONTACT: Lewis Bell at (202) 317–6980 (not a toll-free number).

SUPPLEMENTARY INFORMATION:**Background**

The notice of proposed rulemaking that is the subject of this correction is under section 148 of the Internal Revenue Code.

Need for Correction

As published in the Wednesday, June 24, 2015 (80 FR 36301) partial withdrawal of notice of proposed rulemaking, notice of proposed