This section of the FEDERAL REGISTER contains regulatory documents having general applicability and legal effect, most of which are keyed to and codified in the Code of Federal Regulations, which is published under 50 titles pursuant to 44 U.S.C. 1510.

The Code of Federal Regulations is sold by the Superintendent of Documents. Prices of new books are listed in the first FEDERAL REGISTER issue of each week.

DEPARTMENT OF TRANSPORTATION

Federal Aviation Administration

14 CFR Part 39


RIN 2120–AA64

Airworthiness Directives; Airbus Airplanes

AGENCY: Federal Aviation Administration (FAA), DOT.

ACTION: Final rule; correction.

SUMMARY: The FAA is correcting an airworthiness directive (AD) that published in the Federal Register. That AD applies to all Airbus Model A300 B4–600, B4–600R, and F4–600R series airplanes, Model A300 C4–605R Variant F airplanes (collectively called A300–600 series airplanes), and Model A310 series airplanes. The agency docket number specified throughout the AD is incorrect. No other part of the preamble or regulatory information has been changed; therefore, only the changed portion of the final rule is being published in the Federal Register.

The effective date of this AD remains April 25, 2012.

Correction of Non-Regulatory Text

In the Federal Register of March 21, 2012, AD 2012–06–02, Amendment 39–16983 (77 FR 16430), is corrected as follows:

On page 16430, in the second column, change the docket number to read as follows:


Correction of Regulatory Text

§ 39.13 [Corrected]

In the Federal Register of March 21, 2012, AD 2012–06–02, Amendment 39–16983 (77 FR 16430), on page 16431, in the third column, the product identification line of AD 2012–06–02 is corrected to read as follows:

* * * * *


* * * * *

Issued in Renton, Washington, on April 13, 2012.

John P. Piccola,
Acting Manager, Transport Airplane Directorate, Aircraft Certification Service.

BILLING CODE 4910–13–P
78020, December 19, 2008 (the current regulation), must be sent to the Office of Foreign Labor Certification’s (OFLC’s) Chicago National Processing Center (CNPC) and postmarked no later than midnight April 22, 2012, the last day before the effective date of the H–2B Final Rule. The guidance also provides that applications postmarked on or after April 23, 2012 will be adjudicated in accordance with the requirements described in the Final Rule.

The Department is revising its guidance to clarify that the Final Rule will not be operative until April 27, 2012. In accordance with the Congressional Review Act, (CRA), 5 U.S.C. 801, et seq., April 27, 2012 is 60 days after February 27, 2012, the date on which the rule was reported to Congress, and the earliest date on which the rule can become operative under the CRA. See 5 U.S.C. 801(a)(3). While section 801(a)(3) does not alter the date a rule goes into effect, it prevents an agency from enforcing the rule for 60 days after the rule is reported to Congress.

Accordingly, applications filed under the current regulation must be sent to the CNPC and postmarked no later than midnight April 26, 2012, and applications postmarked on or after April 27, 2012 will be adjudicated in accordance with the requirements described in the Final Rule. Any application filed under the current regulation that is postmarked on or after April 27, 2012 will be returned, and the employer (and its agent or attorney) informed of the need to file a new application in accordance with the provisions of the new H–2B Final Rule. Please note that, as provided in the March 20th guidance, employers who file H–2B applications with a start date of need before October 1, 2013 will not be required to obtain the pre-approved H–2B registration under 20 CFR 655.15, and the Department will continue to adjudicate temporary need during the processing of applications by reviewing the employer’s statement of temporary need in Section B of the ETA Form 9142. Employers with H–2B applications postmarked on or after April 27, 2012 with a start date of need on or after October 1, 2013, must comply with all the requirements contained in the registration process unless the OFLC publishes additional guidance in the Federal Register.

Employers with questions are encouraged to submit their questions to H-2B.Regulation@dol.gov. The Department will provide responses in the form of Frequently Asked Questions (FAQs) on its Web site.

Signed in Washington, this 17th day of April 2012.

Jane Oates, Assistant Secretary, Employment and Training Administration.

[FR Doc. 2012–9612 Filed 4–20–12; 8:45 am]

BILLING CODE 4510–FP–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

21 CFR Part 558
[Docket No. FDA–2012–N–0002]
New Animal Drugs for Use in Animal Feeds; Tiamulin

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule.

SUMMARY: The Food and Drug Administration (FDA) is amending the animal drug regulations to reflect approval of a supplemental new animal drug application (NADA) filed by Novartis Animal Health US, Inc. The supplemental NADA provides for approval of a new concentration of a Type A medicated article.

DATES: This rule is effective April 23, 2012.

FOR FURTHER INFORMATION CONTACT: Cindy L. Burnsteel, Center for Veterinary Medicine (HFV–130), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 240–276–8341, email: cindy.burnsteel@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: Novartis Animal Health US, Inc. (Novartis), 3200 Northline Ave., Suite 300, Greensboro, NC 27408, filed a supplement to NADA 139–472 for DENAGARD (tiamulin hydrogen fumarate) Type A medicated articles for use of a new product formulation in medicated swine feed. The supplemental NADA is approved as of January 6, 2012, and the regulations in 21 CFR 558.4 and 558.60 are amended to reflect the approval.

The Agency has determined under 21 CFR 25.33 that this action is of a type that does not individually or cumulatively have a significant effect on the human environment. Therefore, neither an environmental assessment nor an environmental impact statement is required.

This rule does not meet the definition of “rule” in 5 U.S.C. 804(3)(A) because it is a rule of “particular applicability.” Therefore, it is not subject to the congressional review requirements in 5 U.S.C. 801–808.

List of Subjects in 21 CFR Part 558

Animal drugs, Animal feeds.

Therefore, under the Federal Food, Drug, and Cosmetic Act and under authority delegated to the Commissioner of Food and Drugs and redelegated to the Center for Veterinary Medicine, 21 CFR part 558 is amended as follows:

PART 558—NEW ANIMAL DRUGS FOR USE IN ANIMAL FEEDS

1. The authority citation for 21 CFR part 558 continues to read as follows:


2. In paragraph (d) of § 558.4, in the “Category II” table, revise the entries for “Tiamulin” to read as follows:

§ 558.4 Requirement of a medicated feed mill license.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Assay limits percent ¹ Type A</th>
<th>Type B maximum (100x)</th>
<th>Assay limits percent ¹ Type B/C ²</th>
</tr>
</thead>
<tbody>
<tr>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Tiamulin hydrogen fumarate</td>
<td>90–115</td>
<td>10 g/lb</td>
<td>90–115/70–130</td>
</tr>
</tbody>
</table>

¹ Percent of labeled amount.

² Values given represent ranges for either Type B or Type C medicated feeds. For those drugs that have two range limit, the first set is for a Type B medicated feed and the second set is for a Type C medicated feed. These values (ranges) have been assigned in order to provide for the possibility of dilution of a Type B medicated feed with lower assay limits to make a Type C medicated feed.