

determined that notice and public comment are unnecessary because this amendment is nonsubstantive.

List of Subjects in 21 CFR Part 211

Drugs, Labeling, Laboratories, Packaging and containers.

Therefore, under the Federal Food, Drug, and Cosmetic Act and under authority delegated to the Commissioner of Food and Drugs, 21 CFR part 211 is amended as follows:

PART 211—CURRENT GOOD MANUFACTURING PRACTICE FOR FINISHED PHARMACEUTICALS

1. The authority citation for 21 CFR part 211 continues to read as follows:

Authority: 21 U.S.C. 321, 351, 352, 355, 356, 357, 360b, 371, 374.

§ 211.84 [Corrected]

2. Section 211.84 *Testing and approval or rejection of components, drug product containers, and closures* is amended in paragraph (c)(5) by removing the word "data" and by adding in its place the word "date".

Dated: March 16, 1998.

William K. Hubbard,

Associate Commissioner for Policy Coordination.

[FR Doc. 98-7666 Filed 3-24-98; 8:45 am]

BILLING CODE 4160-01-F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 558

New Animal Drugs For Use In Animal Feeds; Bambermycins; Technical Amendment

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule; technical amendment.

SUMMARY: The Food and Drug Administration (FDA) is amending the animal drug regulation for bambermycins to correct several cross-references in that regulation. In approving a new animal drug application (NADA) filed by Hoechst Roussel Vet, FDA failed to amend certain cross-references to conform to amendments in the approval document and to provide certain other cross-references. This document provides for those conforming amendments and cross-references.

EFFECTIVE DATE: March 25, 1998.

FOR FURTHER INFORMATION CONTACT: David L. Gordon, Center for Veterinary

Medicine (HFV-6), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301-594-1739.

SUPPLEMENTARY INFORMATION: In amending the bambermycins regulation to reflect approval of Hoechst Roussel Vet's NADA 141-034 (use of bambermycins Type A medicated articles to make Type C medicated cattle feeds), FDA amended § 558.95 (21 CFR 558.95) by redesignating paragraph (b) as paragraph (d) (see 62 FR 8373, February 25, 1997), but failed to amend the cross-references in paragraph (a). Furthermore, in approving NADA 141-034 to establish several added uses in § 558.95(b)(4) (currently § 558.95(d)(4)) (see 59 FR 15624, April 4, 1994 and 61 FR 43654, August 26, 1996), FDA failed to provide reference in paragraph (a)(5) to uses in paragraphs (b)(4)(ii) and (b)(4)(iii) (current paragraphs (d)(4)(ii) and (d)(4)(iii)). Section 558.95 is amended by revising paragraph (a), by revising the cross-references to paragraphs (d)(1), (d)(2), (d)(3), and (d)(4), as appropriate, and by expanding those references in paragraph (a)(5) to reflect all uses in paragraph (d)(4).

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:

Authority: 21 U.S.C. 360b, 371.

2. Section 558.95 is amended by revising paragraph (a) to read as follows:

§ 558.95 Bambermycins.

(a) *Approvals.* To sponsors identified by drug labeler codes in § 510.600(c) of this chapter for use of bambermycins Type A medicated articles as bambermycins activity per pound in paragraph (d) of this section as follows:

(1) To 012799: 2, 4, and 10 grams for use as in paragraphs (d)(1), (d)(2), (d)(3), and (d)(4) of this section.

(2) To 012799: 0.4 gram for use as in paragraph (d)(2) of this section.

(3) To 011490: 0.4 and 2 grams for use as in paragraph (d)(2) of this section.

(4) To 012286, 016968, and 017790: 0.4 and 2 grams for use as in paragraph (d)(2) and 2 grams for use as in paragraph (d)(3) of this section.

(5) To 012799: 10 grams to make 40 to 800 grams per ton Type B feed for use as in paragraph (d)(4) of this section.

* * * * *

Dated: March 12, 1998.

Andrew J. Beaulieu,

Acting Director, Office of New Animal Drug Evaluation, Center for Veterinary Medicine.

[FR Doc. 98-7699 Filed 3-24-98; 8:45 am]

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DEPARTMENT OF THE TREASURY

31 CFR Part 2

National Security Information

AGENCY: Department of the Treasury.

ACTION: Final rule.

SUMMARY: This rule revises regulatory text that identifies, by position title, senior Treasury officials authorized to originally or derivatively classify national security information under Executive Order 12958. These designations are now contained in Treasury Order 102-19, which is published in the **Federal Register**. This order will be updated as necessary to revise the designations of officials who have been delegated by the Secretary of the Treasury the authority to classify originally or derivatively national security information.

EFFECTIVE DATE: March 25, 1998.

FOR FURTHER INFORMATION CONTACT:

Robert A. McMenamin, Assistant Director (Information and Physical Security), Department of the Treasury, Office of Security, Room 3210 Annex, 1500 Pennsylvania Avenue, NW, Washington, D.C. 20220, (202) 622-1120.

SUPPLEMENTARY INFORMATION: This rule removes the specific designations of Treasury officials authorized to originally and derivatively classify national security information under Executive Order 12958 and previous Orders. The designation of such officials is now made by a Treasury Order that will be revised from time to time as may be necessary. This rule reduces costs by making it unnecessary to revise periodically the regulations in part 2.

Because this rule relates to agency management and personnel, notice and public procedure and a delayed effective date are not required pursuant to 5 U.S.C. 553(a)(2) and the provisions of Executive Order 12866 do not apply. Because notice and public procedure is not required, the provisions of the Regulatory Flexibility Act (5 U.S.C. 601 et seq.) do not apply.