- El Paso, TX, El Paso Intl, RNAV (GPS) RWY 26L, Orig
- El Paso, TX, El Paso Intl, RNAV (GPS) RWY 26R, Orig
- El Paso, TX, El Paso Intl, GPS RWY 26L, Orig, CANCELLED
- Houston, TX, West Houston, RNAV (GPS) Y RWY 33, Orig
- Temple, TX, Draughon-Miller Central Texas Regional, LOC/DME BC RWY 33, Amdt 3A, CANCELLED
- Wichita Falls, TX, Sheppard AFB/ Wichita Falls Muni, NDB RWY 33L, Amdt 11
- Wichita Falls, TX, Sheppard AFB/ Wichita Falls Muni, VOR–D, Amdt 14
- Wichita Falls, TX, Sheppard AFB/ Wichita Falls Muni, RNAV (GPS) RWY 15R, Orig
- Wichita Falls, TX, Sheppard AFB/ Wichita Falls Muni, RNAV (GPS) RWY 33L, Orig
- Evanston, WY, Evanston-Uinta County Burns Field, VOR/DME RWY 23, Amdt 2B
- Evanston, WY, Evanston-Uinta County Burns Field, RNAV (GPS) RWY 5, Orig
- Evanston, WY, Evanston-Uinta County Burns Field, RNAV (GPS) RWY 23, Orig

The FAA published the following procedure in Docket No. 30339; Amdt. No. 3031 to Part 97 of the Federal Aviation Regulations (Vol. 67, FR No. 225, Page 70155; dated Thursday, November 21, 2002) under section 97.29 effective November 28, 2002 which is hereby amended to be effective December 26, 2002:

Sacramento, CA, Sacramento Mather, ILS RWY 22L, Amdt 3

[FR Doc. 02–30441 Filed 12–2–02; 8:45 am] BILLING CODE 4910–13–M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Parts 520

Oral Dosage Form New Animal Drugs; Lincomycin Hydrochloride Soluble Powder

AGENCY: Food and Drug Administration, HHS.

ACTION: Technical amendment.

SUMMARY: The Food and Drug Administration (FDA) is amending the animal drug regulations for preslaughter withdrawal time for lincomycin soluble powder products used to make medicated drinking water for swine to correct inadvertant editorial errors. This action is being taken to ensure accuracy and clarity in the agency's regulations.

DATES: This rule is effective December 3, 2002.

FOR FURTHER INFORMATION CONTACT:

George K. Haibel, Center for Veterinary Medicine (HFV–6), Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855, 301–827–4567, email: ghaibel@cvm.fda.gov.

SUPPLEMENTARY INFORMATION: FDA has found that § 520.1263c (21 CFR 520.1263c) does not reflect the approved preslaughter withdrawal time for three lincomycin soluble powder products used to make medicated drinking water for swine. The 6-day withdrawal time was inadvertently removed for a generic product approved under ANADA 200-189 at the time it was being removed for the pioneer product approved under NADA 111-636 (64 FR 13341, March 18, 1999). The conditions of use for two other products approved February 4, 1999, under ANADA 200-241 (64 FR 13508, March 19, 1999) and September 22, 1999, under ANADA 200-233 (64 FR 66382, November 26, 1999) were subsequently codified without a withdrawal period. At this time, the regulations are being amended in § 520.1263c to correct these errors.

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 520

Animal drugs.

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 520 is amended as follows:

PART 520—ORAL DOSAGE FORM NEW ANIMAL DRUGS

1. The authority citation for 21 CFR part 520 continues to read as follows:

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

§520.1263c [Amended]

2. Section 520.1263c *Lincomycin hydrochloride soluble powder* is amended in paragraph (d)(1)(iii) by adding at the end the sentence "For Nos. 046573 and 051259: Do not slaughter swine for 6 days following last treatment." Dated: November 8, 2002. **Steven D. Vaughn,** *Director, Office of New Animal Drug Evaluation, Center for Veterinary Medicine.* [FR Doc. 02–30639 Filed 12–2–02; 8:45 am] **BILLING CODE 4160–01–S**

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 520

Oral Dosage Form New Animal Drugs; Ivermectin Paste

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 an abbreviated new animal drug application (ANADA) filed by Virbac AH, Inc. The ANADA provides for oral use of ivermectin paste in horses for treatment and control of various internal parasites or parasitic conditions.

DATES: This rule is effective December 3, 2002.

FOR FURTHER INFORMATION CONTACT: Lonnie W. Luther, Center for Veterinary Medicine (HFV–104), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301–827–8549, email: lluther@cvm.fda.gov.

SUPPLEMENTARY INFORMATION: Virbac AH, Inc., 3200 Meacham Blvd., Ft. Worth, TX 76137, filed ANADA 200-320 for EQUELL (ivermectin) Paste. The application provides for oral use of 1.87 percent ivermectin paste in horses for the treatment and control of various species of internal parasites or parasitic conditions. Virbac's EQUELL Paste is approved as a generic copy of Merial Limited's EQUALEN Paste, approved under NADA 134-314. ANADA 200-320 is approved as of August 9, 2002, and 21 CFR 520.1192 is amended to reflect the approval. The basis of approval is discussed in the freedom of information summary.

In accordance with the freedom of information provisions of 21 CFR part 20 and 514.11(e)(2)(ii), a summary of safety and effectiveness data and information submitted to support approval of this application may be seen in the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852, between 9 a.m. and 4 p.m., Monday through Friday.