NADA No.	Drug Name
139–913	Equron Sulka-S-Bolus

Accordingly, the agency is amending the regulations in 21 CFR 510.600(c)(1) and (c)(2) to reflect the change of sponsor. The drug labeler code assigned to Solvay Animal Health is being retained as the drug labeler code for the new sponsor.

List of Subjects in 21 CFR Part 510

Administrative practice and procedure, Animal drugs, Labeling, Reporting and recordkeeping requirements.

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 510 is amended as follows:

PART 510—NEW ANIMAL DRUGS

1. The authority citation for 21 CFR part 510 continues to read as follows:

Authority: Secs. 201, 301, 501, 502, 503, 512, 701, 721 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321, 331, 351, 352, 353, 360b, 371, 379e).

§510.600 [Amended]

2. Section 510.600 Names, addresses, and drug labeler codes of sponsors of approved applications is amended in the table in paragraph (c)(1) by removing the entry for "Solvay Animal Health, Inc." and by alphabetically adding a new entry for "Fort Dodge Animal Health, A Division of American Cyanamid Co., P.O. Box 1339, Fort Dodge, IA 50501" and in the table in paragraph (c)(2) in the entry for "053501" by removing the sponsor name and address "Solvay Animal Health, Inc., 1201 Northland Dr., Mendota Heights, MN 55120" and adding in its place "Fort Dodge Animal Health, A Division of American Cyanamid Co., P.O. Box 1339, Fort Dodge, IA 50501".

Dated: July 22, 1997.

Robert C. Livingston,

Director, Office of New Animal Drug Evaluation, Center for Veterinary Medicine. [FR Doc. 97–20249 Filed 7-30-97; 8:45 am] BILLING CODE 4160–01–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Parts 510 and 524

New Animal Drugs; Change of Sponsor

AGENCY: Food and Drug Administration,

HHS.

ACTION: Final rule.

SUMMARY: The Food and Drug Administration (FDA) is amending the animal drug regulations to reflect a change of sponsor for three approved new animal drug applications (NADA's) from Syntex Animal Health, Inc., Division of Syntex Agribusiness, Inc., to Medicis Dermatologics, Inc.

EFFECTIVE DATE: July 31, 1997.

FOR FURTHER INFORMATION CONTACT:

Thomas J. McKay, Center for Veterinary Medicine (HFV–102), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301–827–0213.

SUPPLEMENTARY INFORMATION: Syntex Animal Health, Inc., Division of Syntex Agribusiness, Inc., 3401 Hillview Ave., P.O. Box 10850, Palo Alto, CA 94303, has informed FDA that it has transferred ownership of, and all rights and interests in NADA's 15–151 (fluocinolone acetonide, neomycin sulfate cream), 15–152 (fluocinolone acetonide cream), and 15–298 (fluocinolone acetonide solution) to Medicis Dermatologics, Inc., 4343 East Camelback Rd., suite 250, Phoenix, AZ 85018–2700. Accordingly, the agency is

amending the regulations in 21 CFR 524.981a, 524.981b, and 524.981c to reflect the transfer of ownership. The agency is also amending the regulations in 21 CFR 510.600(c)(1) and (c)(2) by alphabetically adding a new listing for Medicis Dermatologics, Inc.

List of Subjects

21 CFR Part 510

Administrative practice and procedure, Animal drugs, Labeling, Reporting and recordkeeping requirements.

21 CFR Part 524

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 parts 510 and 524 are amended as follows:

PART 510—NEW ANIMAL DRUGS

- 1. The authority citation for 21 CFR part 510 continues to read as follows: **Authority:** Secs. 201, 301, 501, 502, 503, 512, 701, 721 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 321, 331, 351, 352, 353, 360b, 371, 379e).
- 2. Section 510.600 is amended in the table in paragraph (c)(1) by alphabetically adding a new entry for "Medicis Dermatologics, Inc." and in the table in paragraph (c)(2) by numerically adding a new entry for "099207" to read as follows:

§ 510.600 Names, addresses, and drug labeler codes of sponsors of approved applications.

(c) * * *

(1) * * *

Firm name and address				Drug labeler code			
*	*	*	*	*	*	*	
		t Camelback Rd., suite 25	0, 099207				
Phoenix, AZ 8	35018–2700.						
*	*	*	*	*	*	*	

Drug labeler code				Firm Name and address			
*	*	*	*	*	*	*	
099207				Medicis Dermatologics, Inc., 4343 East Camelback Rd., suite 250, Phoenix, AZ 85018–2700.			
*	*	*	*	*	*	*	

PART 524—OPHTHALMIC AND TOPICAL DOSAGE FORM NEW ANIMAL DRUGS

3. The authority citation for 21 CFR part 524 continues to read as follows:

Authority: Sec. 512 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360b).

§ 524.981a [Amended]

4. Section 524.981a Fluocinolone acetonide cream is amended in paragraph (b) by removing "000033" and adding in its place "099207".

§ 524.981b [Amended]

5. Section 524.981b *Fluocinolone acetonide solution* is amended in paragraph (b) by removing "000033" and adding in its place "099207".

§ 524.981c [Amended]

6. Section 524.981c Fluocinolone acetonide, neomycin sulfate cream is amended in paragraph (b) by removing "000033" and adding in its place "099207".

Dated: July 23, 1997.

Robert C. Livingston,

Director, Office of New Animal Drug Evaluation, Center for Veterinary Medicine. [FR Doc. 97–20248 Filed 7-30-97; 8:45 am] BILLING CODE 4160–01–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 520

Animal Drugs, Feeds, and Related Products; Change of Sponsor; Corrections

AGENCY: Food and Drug Administration, HHS.

ACTION: Final rule; correction.

SUMMARY: The Food and Drug Administration (FDA) is correcting a document that appeared in the **Federal Register** of June 30, 1997 (62 FR 35075 at 35076). The document amended the animal drug regulations to reflect the change of sponsor for 52 approved new animal drug applications (NADA's) from Fermenta Animal Health Co. to Boehringer Ingelheim Animal Health, Inc. The document was published with

two inadvertent errors. This document corrects those errors.

EFFECTIVE DATE: July 31, 1997. FOR FURTHER INFORMATION CONTACT: Thomas J. McKay, Center for Veterinary Medicine (HFV–102), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301–827–0213.

In FR Doc. 97–16967, appearing on page 35075, in the **Federal Register** of Monday, June 30, 1997, the following corrections are made: On page 35076, in the first column, in amendment 11, in the third line, "(a)(6)" is corrected to read "(b)(6)"; and on the same page, in the second column, in amendment 19, beginning in the fourth line, "000069, 054273, and 057561" is corrected to read "000069, 054273, 057561, and 059130".

Dated: July 21, 1997.

Robert C. Livingston,

Director, Office of New Animal Drug Evaluation, Center for Veterinary Medicine. [FR Doc. 97–20250 Filed 7-30-97; 8:45 am] BILLING CODE 4160–01–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 556

Tolerances for Residues of New Animal Drugs in Food; Apramycin

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 two supplemental new animal drug applications (NADA's) filed by Elanco Animal Health, A Division of Eli Lilly & Co. The supplemental NADA's provide for revised tolerances for total residues of apramycin (i.e., the safe concentration) in edible swine tissues.

EFFECTIVE DATE: July 31, 1997. **FOR FURTHER INFORMATION CONTACT:** George K. Haibel, Center for Veterinary Medicine (HFV–133), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301–594–1644. SUPPLEMENTARY INFORMATION: Elanco Animal Health, A Division of Eli Lilly & Co., Lilly Corporate Center, Indianapolis, IN 46285, is sponsor of supplemental NADA 106–964 that provides for the use of Apralan® (apramycin sulfate) soluble powder in swine drinking water and supplemental NADA 126–050 that provides for the use of Apralan® (apramycin sulfate) Type A medicated article in swine feed, both for control of porcine colibacillosis (weanling pig scours) caused by strains of Escherichia coli sensitive to apramycin. These supplemental NADA's provide for a change in the tolerance for total residues of apramycin (i.e., the safe concentration) in edible swine tissues as provided in § 556.52 (21 CFR 556.52). Review of these supplements involved a review of new toxicology studies and information in the original approvals.

In evaluating these supplements, FDA's Center for Veterinary Medicine also considered that the proof of human food safety for antimicrobial animal drug residues includes a determination of their antimicrobial activity for all antimicrobial new animal drug products. In the absence of studies to determine the microbiological safety of antimicrobial drug residues, the acceptable daily intake (ADI) for apramycin is limited to 25 micrograms per kilogram (µg/kg) of body weight per day (for appropriate studies see "Guidance: Microbial Testing of Antimicrobial Drug Residues in Food," January, 1996). As indicated in the freedom of information summaries, the safe concentration for total apramycin residues is established at 5 parts per million (ppm) for muscle, 15 ppm for liver, and 30 ppm for fat and kidney. These revised safe concentrations warrant removal of the existing tolerances for total residues in § 556.52, because those tolerances are now incorrect. Because this approval does not result in a different tolerance than that currently codified for marker residue in swine kidney, and because the sponsor did not petition FDA to change the tolerance, the tolerance of 0.1 ppm in swine kidney remains codified. FDA is also codifying the ADI for apramycin of 25 μg/kg of body weight per day. The supplement is