

| Firm name and address                                                               | Drug labeler code |
|-------------------------------------------------------------------------------------|-------------------|
| * * *                                                                               | * * *             |
| Wellmark International, 1100 East Woodfield Rd., suite 500,<br>Schaumburg, IL 60173 | 011536            |
| * * *                                                                               | * * *             |

(2) \* \* \*

| Drug labeler code | Firm name and address                                                               |
|-------------------|-------------------------------------------------------------------------------------|
| 011536            | Wellmark International, 1100 East Woodfield Rd., suite 500,<br>Schaumburg, IL 60173 |
| * * *             | * * *                                                                               |

Dated: July 18, 2000.

**Claire M. Lathers,***Director, Office of New Animal Drug  
Evaluation, Center for Veterinary Medicine.*

[FR Doc. 00-18824 Filed 7-25-00; 8:45 am]

BILLING CODE 4160-01-F

**DEPARTMENT OF HEALTH AND  
HUMAN SERVICES****Food and Drug Administration****21 CFR Part 520****Oral Dosage Form New Animal Drugs;  
Ivermectin Sustained-Release Bolus****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 Merial  
Ltd. The supplemental NADA provides  
for changes to labeling of ivermectin  
sustained-release bolus for cattle.**DATES:** This rule is effective July 26,  
2000.**FOR FURTHER INFORMATION CONTACT:**Janis R. Messenheimer, Center for  
Veterinary Medicine (HFV-135), Food  
and Drug Administration, 7500 Standish  
Pl., Rockville, MD 20855, 301-827-  
7578.**SUPPLEMENTARY INFORMATION:** Merial  
Ltd., 2100 Ronson Rd., Iselin, NJ 08830-  
3077, filed a supplement to NADA 140-  
988 that provides for use of Ivomec®  
(ivermectin) SR bolus for cattle. The  
supplement provides for reducing the  
predicted duration of effectiveness inlabeling from approximately 135 days to  
approximately 130 days, based on bolus  
stability data. The supplement is  
approved as of June 21, 2000, and the  
regulations in 21 CFR 520.1197 are  
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 part 20 (21  
CFR part 20) and § 514.11(e)(2)(ii) (21  
CFR 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.The agency has determined under 21  
CFR 25.24(a)(1) 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 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.**§ 520.1197 [Amended]**2. Section 520.1197 *Ivermectin  
sustained-release bolus* is amended in  
paragraph (d)(2) by removing the  
parenthetical phrase "(approximately  
135 days)" and by adding in its place "  
(approximately 130 days)".

Dated: July 18, 2000.

**Claire M. Lathers,***Director, New Animal Drug Evaluation,  
Center for Veterinary Medicine.*

[FR Doc. 00-18827 Filed 7-25-00; 8:45 am]

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**DEPARTMENT OF HEALTH AND  
HUMAN SERVICES****Food and Drug Administration****21 CFR Parts 520 and 522****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 12 approved new  
animal drug applications (NADA's) from  
Merial Ltd. to Phoenix Scientific, Inc.**DATES:** This rule is effective July 26,  
2000.**FOR FURTHER INFORMATION CONTACT:**Thomas J. McKay, Center for Veterinary  
Medicine (HFV-102), Food and Drug