Date of Approval: Aug. 9, 2004

FREEDOM OF INFORMATION SUMMARY

NADA 134-314

EQVALAN Paste 1.87%

ivermectin

This supplement amends the EQVALAN (ivermectin) Paste 1.87% labeling to reflect a change in the indications section. Specifically, under the sub-heading Small Strongyles, the labeling has been revised to separate the listing of adult species from the fourth-stage larvae.

Sponsored by:

Merial Ltd.

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1. GENERAL INFORMATION:

a. File Number: NADA 134-314

b. Sponsor: Merial Ltd.

3239 Satellite Blvd.

Building 500

Duluth, GA 30096-4640

Drug Labeler Code: 050604

c. Established Name: Ivermectin

d. Proprietary Name: EQVALAN 1.87% Paste For Horses

e. Dosage Form: Paste containing 1.87 % ivermectin

f. How Supplied: Individual dose syringe contains sufficient paste to

treat one 1250 lb horse orally. Each weight marking on the syringe plunger delivers enough

paste to treat 250 lb of body weight

g. How Dispensed: OTC

h. Amount of Active Ingredients: Each syringe contains 113 mg of ivermectin

i. Route of Administration: Oral

j. Species/Class: Equine

k. Recommended Dosage: 91 mcg ivermectin per lb (200 mcg/kg)

1. Pharmacological Category: Anthelmintic

m. Indications: For treatment and control of the following parasites

in horses: **Large strongyles** (adults)-*Strongylus vulgaris* (also early forms in blood vessels), *S. edentatus* (also tissue stages), *S. equinus*,

Triodontophorus spp. including T. brevicauda and T. serratus and Craterostomum acuticaudatum; Small Strongyles (adults, including those resistant

to some benzimidazole class compounds)-Coronocyclus spp. including C. coronatus, C. labiatus, and C. labratus, Cyathostomum spp. including C. catinatum and C. pateratum, Cylicocyclus spp. including C. insigne, C.

leptostomum, C. nassatus, and C. brevicapsulatus,

Cylicodontophorus spp., Cylicostephanus spp. including C. calicatus, C. goldi, C. longibursatus, and C. minutus, and Petrovinema poculatum; Small Strongyles-fourth stage larvae; Pinworms (adults and fourth stage larvae)-Oxyuris equi; Ascarids (adults and third and fourth stage larvae)-Parascaris equorum; Hairworms (adults)-Trichostrongylus axei; Large-mouth Stomach Worms (adults)-Habronema muscae; Bots (oral and gastric stages)-Gasterophilus spp. including G. intestinalis and G. nasalis; Lungworms (adults and fourth stage larvae)-Dictyocaulus arnfieldi; **Intestinal Threadworms** (adults)-Strongyloides westeri; Summer Sores caused by Habronema and Draschia spp. cutaneous third stage larvae; Dermatitis caused by neck threadworm microfilariae, Onchocerca sp.

n. Effect of Supplement:

This supplement amends the EQVALAN (ivermectin) Paste 1.87% labeling to reflect a change in the indications section. Specifically, under the sub-heading Small Strongyles, the labeling has been revised to separate the listing of adult species from the fourth-stage larvae.

2. EFFECTIVENESS:

This supplement revises the indications to better reflect the original effectiveness data. Specifically, the new language separates out the unspeciated fourth-stage larvae from the speciated adult small strongyles. The new language reads as follows: "Small Strongyles (adults, including those resistant to some benzimidazole class compounds)-Coronocyclus spp. including C. coronatus, C. labiatus, and C. labratus, Cyathostomum spp. including C. catinatum and C. pateratum, Cylicocyclus spp. including C. insigne, C. leptostomum, C. nassatus, and C. brevicapsulatus, Cylicodontophorus spp., Cylicostephanus spp. including C. calicatus, C. goldi, C. longibursatus, and C. minutus, and Petrovinema poculatum; Small Strongyles-fourth stage larvae"

No new data was required for this supplement to NADA 134-314 (original approval May 29, 1984) and this supplement did not require review of the original effectiveness data for this product.

3. TARGET ANIMAL SAFETY:

No new data was required for this supplement to NADA 134-314 (original approval May 29, 1984) and this supplement did not require review of the original target animal safety data for this product.

4. HUMAN SAFETY:

This drug is intended for use in horses, which are non-food animals. Because this new animal drug is not intended for use in food-producing animals, data on human safety pertaining to drug residues in food were not required for approval of this NADA.

Human Warnings are provided on the product label as follows: "Not for use in humans. Keep this and all drugs out of the reach of children." "Refrain from smoking and eating when handling. Wash hands after use. Avoid contact with eyes." "Do not use in horses intended for human consumption."

5. AGENCY CONCLUSIONS:

The data submitted in support of this NADA satisfy the requirements of section 512 of the Federal Food, Drug, and Cosmetic Act and 21 CFR Part 514 of the implementing regulations. The data demonstrates that EQVALAN Paste 1.87% is safe and effective for the labeled claim.

The drug is available over-the-counter for lay use. Routine deworming of horses is a widely accepted and recommended practice performed by the layperson. A diagnosis of parasite infection prior to deworming is not necessary. The syringe carton and package outsert contain detailed directions for use.

This approval does not qualify for marketing exclusivity under section 512(c)(2)(F)(iii) of the Federal Food, Drug, and Cosmetic Act.

According to the Center's supplemental approval policy (21 CFR 514.106), this is a Category I change. Therefore, this action did not require a reevaluation of the safety and effectiveness data in the parent application.

6. ATTACHMENTS:

Facsimile labeling is attached as indicated below:

Carton

Package Outsert

Syringe Label