Tylvalosin Type B Medicated Feed

CAUTION: Federal law restricts medicated feed containing this veterinary feed directive (VFD) drug to use by or on the order of a licensed veterinarian.

Do Not Feed Undiluted

INDICATION:
Swine:
Control of porcine proliferative enteropathy (PPE) associated with *Lawsonia intracellularis* infection in groups of swine in buildings experiencing an outbreak of PPE

ACTIVE DRUG INGREDIENT: Tylvalosin* 3856 grams per ton (4250 ppm)

GUARANTEED ANALYSIS:
Crude Protein, not less than %
Lysine, not less than %
Crude Fat, not less than %
Crude Fiber, not more than %
Calcium, not less than %
Calcium, not more than %
Phosphorus, not less than %
Salt¹, not less than %
Salt¹, not more than %
Sodium², not less than %
Sodium², not more than %
Selenium, not less than ppm
Zinc, not less than ppm

¹If added
²Shall be guaranteed only when total sodium exceeds that furnished by maximum salt guarantee

INGREDIENTS:
Each ingredient must be named in accordance with the names and definitions adopted by the Association of American Feed Control Officials (AAFCO).

MIXING DIRECTIONS:
To prepare one ton of complete Type C medicated feed containing 38.6 grams per ton (42.5 ppm) tylvalosin, thoroughly mix 20 pounds of this medicated feed with 1980 pounds of non-medicated feed.

Pelleted or crumbled Type C medicated feeds must bear an expiration date of 1 week after the date of manufacture.

FEEDING DIRECTIONS:
Feed the resulting Type C medicated feed containing 38.6 grams tylvalosin/ton as the sole ration for 14 consecutive days.

CAUTION:
To assure both food safety and responsible use in swine, concurrent use of tylvalosin Type A medicated article in medicated feed and tylvalosin or another macrolide in medicated drinking water or by any other route of administration should be avoided. Not for use in swine intended for breeding. The effects of tylvalosin on swine reproductive performance, pregnancy, and lactation have not been determined. VFDs for tylvalosin shall not be refilled.
WARNINGS:

WITHDRAWAL PERIOD:

No withdrawal period is required before slaughter for human consumption.

ANTIBACTERIAL WARNINGS:
Use of antibacterial drugs in the absence of a susceptible bacterial infection is unlikely to provide benefit to treated animals and may increase the development of drug-resistant bacteria.

USER SAFETY WARNINGS:

May cause skin irritation. Tylosin has been shown to cause hypersensitivity reactions in laboratory animals. People with known hypersensitivity to tylosin should avoid contact with this product. In case of accidental ingestion, seek medical attention.

STORAGE CONDITIONS: Store in a cool dry place at or below 25°C (77°F).

NET CONTENTS: Net weight ___________ lb (___________ kg) Bag [or Bulk]

LOT NO.:

Use only as directed

For sales, technical assistance or to obtain a Safety Data Sheet, call Pharmgate Animal Health at 1-800-380-6099.

To report suspected adverse drug events, contact the ASPCA Animal Product Safety Service at 1-800-345-4735 or the FDA at 1-888-FDA-VETS.

Pharmgate Animal Health has contracted with the ASPCA Animal Product Safety Service to collect human and animal suspected adverse drug events reports for this product.

*Aivlosin® is a registered trademark of ECO Animal Health Ltd.

NADA 141-460. Approved by FDA.

Manufactured by
Blue Bird Feed Mill
Robin, IN 00000

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