Ractopamine Finishing Cattle Feed Concentrate – TD + MT Type C Medicated Top Dress Feed

For Use with Rations Containing Monensin¹ and Tylosin²

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

INDICATIONS

For increased rate of weight gain and improved feed efficiency in cattle fed in confinement for slaughter during the last 28 to 42 days on feed.

ACTIVE DRUG INGREDIENT

Ractor	pamine hydr	ochloride ^{&}		not to	exceed	300	g/to	n*
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GUARANTEED ANALYSIS

Crude Protein, not less than	%
Non-Protein Nitrogen (NPN) ⁴ , not more than	
Crude Fat, not less than	
Crude Fiber, not more than	
Calcium, not less than	
Calcium, not more than	
Phosphorus, not less than	
Salt ^s , not less than	
Salt ³ , not more than	
Sodium ⁶ , not less than	
Sodium ⁶ , not more than	
Potassium, not less than	
Vitamin A ^{(5,7} , not kess than	
When added	
⁹ If added.	
Shall be guaranteed only when total sodium exceeds that furnished by the maximum salt guarantee.	

Other than precursors of Vitamin A.

INGREDIENTS

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

DIRECTIONS FOR USE

Feed a minimum of 1.0 lb/hd/day Ractopamine Type C Top Dress TD + MT continuously to cattle fed in confinement for slaughter, to provide 70 to 400 mg/head/day ractopamine for the last 28 to 42 days on feed. Feed on top of a ration containing 10 to 40 g/ton monensin and 8 to 10 g/ton tylosin phosphate, to provide 0.14 to 0.42 mg monensin/lb body weight/day, depending on the severity of the coccidiosis challenge, up to 480 mg/hd/day and 60 to 90 mg/hd/day tylosin.

CAUTION

Ractopamine HCl is not for animals intended for breeding.

WARNING

The active ingredient in Optaflexx, ractopamine hydrochloride, is a beta-adrenergic agonist. Individuals with cardiovascular disease should exercise special caution to avoid exposure. Not for use in humans. Keep out of the reach of children. The Optaflexx 45 formulation (Type A Medicated Article) poses a low dust potential under usual conditions of handling and mixing. When mixing and handling Optaflexx, use protective clothing, impervious gloves, protective eye wear, and a NIOSH-approved dust mask. Operators should wash thoroughly with soap and water after handling. If accidental eye contact occurs, immediately rinse eyes thoroughly with water. If irritation persists, seek medical attention. The material safety data sheet contains more detailed occupational safety information. To report adverse effects, access medical information, or obtain additional product information, call 1-877-426-7765.

MANUFACTURED BY BLUE BIRD FEED MILL

Any town, USA 12345

Net Weight lb (kg) on bag or bulk

*The medicated feed label must state a single drug concentration.

¹Sourced from Rumensin[™], NADA# 95-735 ²Sourced from Tylovet[™], ANADA#200-484 [®]Sourced from Optaflexx[™], NADA# 141-221

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