

# **Lubabegron and Monensin (Coccidiosis) and Tylosin Medicated Feedlot Cattle Feed Type C Medicated Feed**

(lubabegron and monensin and tylosin Type C 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.

## **INDICATIONS FOR USE**

For reduction of ammonia gas emissions per pound of live weight and hot carcass weight, prevention and control of coccidiosis due to *Eimeria bovis* and *Eimeria zuernii*, and reduction of incidence of liver abscesses associated with *Fusobacterium necrophorum* and *Arcanobacterium pyogenes* in beef steers and heifers fed in confinement for slaughter during the last 14 to 91 days on feed.

Effectiveness has not been demonstrated when fed for less than 14 days.

Ammonia gas emissions were measured for individual animals or small groups of animals held in environmentally controlled facilities. Based on existing information, reliable predictions of the reduction of ammonia gas emissions cannot be made on a herd, farm, or larger scale.

Increased rate of weight gain, improved feed efficiency, and increased carcass leanness have not been demonstrated with this product.

## **ACTIVE DRUG INGREDIENTS**

Lubabegron (as lubabegron fumarate) <sup>a</sup> .....	1.25 to 4.54 g/ton*
Monensin, USP <sup>b</sup> .....	10 to 40 g/ton*
Tylosin (as tylosin phosphate) <sup>c</sup> .....	8 to 10 g/ton*

## **GUARANTEED ANALYSIS**

Crude Protein, not less than.....	_____ %
Non-Protein Nitrogen (NPN) <sup>1</sup> , 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 <sup>1</sup> , not less than.....	_____ %
Salt <sup>1</sup> , not more than.....	_____ %
Sodium <sup>2</sup> , not less than.....	_____ %
Sodium <sup>2</sup> , not more than.....	_____ %
Potassium <sup>1</sup> , not less than.....	_____ %
Vitamin A <sup>1</sup> , not less than.....	_____ I.U./lb

<sup>1</sup> Guarantee required only when nutrient source is added except when the feed is intended, represented, or serves as a principal source of the nutrient.

<sup>2</sup> Sodium guarantee required only when total sodium exceeds that furnished by the 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.

## FEEDING DIRECTIONS

Feed continuously as sole ration to provide 13 to 90 mg lubabegron/hd/day, 0.14 to 0.42 mg monensin/lb body weight per day, depending upon severity of coccidiosis challenge, up to 480 mg/hd/day, and 60 to 90 mg tylosin/hd/day to beef steers and heifers fed in confinement for slaughter during the last 14 to 91 days on feed.


## CAUTION

A decrease in dry matter intake may be noticed in some animals receiving lubabegron. Lubabegron has not been approved for use in breeding animals because safety and effectiveness have not been evaluated in these animals.


Do not allow horses or other equines access to feed containing lubabegron and monensin. Ingestion of monensin by horses has been fatal. Monensin medicated cattle and goat feeds are safe for use in cattle and goats only. Consumption by unapproved species may result in toxic reactions. Feeding undiluted or mixing errors resulting in high concentrations of monensin has been fatal to cattle and could be fatal to goats. Must be thoroughly mixed in feeds before use. Do not exceed the levels of monensin recommended in the feeding directions, as reduced average daily gains may result. If feed refusals containing monensin are fed to other groups of cattle, the concentration of monensin in the refusals and amount of refusals fed should be taken into consideration to prevent monensin overdosing.

Do not use in any finished feed (supplement, concentrate or complete feed) containing in excess of 2% bentonite.

## WARNING



No withdrawal period is required when used according to labeling. A withdrawal period has not been established for this product for pre-ruminating calves. Do not use in calves to be processed for veal.



**User Safety Warning:** Not for human use. Keep out of reach of children. The active ingredient in Experior, lubabegron, is a beta-adrenergic agonist/antagonist. Individuals with cardiovascular disease should exercise special caution to avoid exposure. When mixing and handling Experior, 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 thoroughly with water; if wearing contact lenses, rinse eyes first, then remove contact lenses and continue to rinse for 5-20 minutes. If irritation persists, seek medical attention. The safety data sheet contains more detailed occupational safety information. To report adverse drug events, access medical information, or obtain additional product information, call Elanco US Inc. at

1-800-428-4441. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS or <http://www.fda.gov/AnimalVeterinary/SafetyHealth>.

**Lot Number:** \_\_\_\_\_

Approved by FDA under NADA # 141-512

**MANUFACTURED BY:**  
**BLUE BIRD FEED MILL**  
**Any town, USA 12345**

**Net Weight lb (kg) on bag or bulk**

<sup>a</sup> Sourced from Experior<sup>TM</sup> (lubabegron Type A medicated article; NADA # 141-508).

<sup>b</sup> Sourced from Rumensin<sup>TM</sup> (monensin Type A medicated article; NADA # 095-735).

<sup>c</sup> Sourced from Tylan<sup>TM</sup> (tylosin Type A medicated article; NADA # 012-491).

\*Final printed label on the formulated Type C medicated feed must bear a single concentration of each drug.

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