



100 mg/mL Antimicrobial Injectable Solution For Subcutaneous Use In Beef Cattle And Non-Lacfating Dairy Cattle For Intramuscular Of Subcutaneous Use In Swine Not For Use In Female Dairy Cattle 20 Months Of Age Or Older Or In Calves To Be Processed For Veal

CAUTION

Federal (U.S.A.) law restricts this drug to use by or on the order of a licensed veterinarian. Federal (U.S.A.) law prohibits the extra-label use of this drug in food-producing animals. To assure responsible antimicrobial drug use, enrofloxacin should only be used as a second-line drug for colibacillosis in swine following consideration of other therapeutic options.

PRODUCT DESCRIPTION: Baytril® 100 is a sterile, ready-to-use injectable antimicrobial solution that contains enrofloxacin, a broad-spectrum fluoroquinolone antimicrobial agent.

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1-cyclopropyl-r (-4-emyl-r-piperaziny)-o-fludro-1,4-dinydro-4-oxo-3-duinolimearboxylic a INDICATIONS: Cattle - Single-Dose Therapy: Baytril® 100 is indicated for the treatment of bovine respiratory disease (BRD) associated with Mannherima haemolytica, Pasteurella multocida, Histophilus sommi and Mycoplasma bovis in beef and non-lactating dairy cattle; and for the control of BRD in beef and non-lactating dairy cattle at high risk of developing BRD associated with M. haemolytica, P. multocida, H. sommi and M. bovis.

associated with *M. naemolytica, P. Muticica, H. Somm and M. Dows.* **Cattle - Mutiple-Day Therapy:** Baytril® 100 is indicated for the treatment of bovine respiratory disease (BRD) associated with *Mannheimia haemolytica, Pasteurella multocida* and *Histophilus somni* in beef and non-lactating dairy cattle. **Swine:** Baytril® 100 is indicated for the treatment and control of swine respiratory disease (SRD) associated with *Actinobacillus pleuropneumoniae, Pasteurella multocida, Haemophilus parasuis, Streptococcus suis, Bordetella bronchiseptica* and *Mycoplasma hyopneumoniae.* Baytril® 100 is indicated for the control of colibacillosis in groups or pens of weaned pigs where colibacillosis associated with *Escherichia coli* has been diagnosed. DECARG AMD ADMINICTATION:

Baytife 100 provides flexible dosages and durations of therapy. Baytife 100 provides flexible dosages and durations of therapy. Baytife 100 may be administered as a single dose for one day for treatment and control of BRD (cattle), for treatment and control of SRD or for control of collabacillosis (swine), or for multiple days for BRD treatment (cattle). Selection of the appropriate dose and duration of therapy for BRD treatment in cattle should be based on an assessment of the severity of the disease, pathogen susceptibility and clinical response. Cattle:

Cattle

Single-Dose Therapy (BRD Treatment): Administer, by subcutaneous injection, a single dose of 7.5-12.5 mg/kg of body weight (3.4-5.7 mL/100 lb).

Multiple-Day Therapy (BRD Treatment): Administer daily, a subcutaneous dose of 2.5-5 mg/kg of body weight (1.1-2.3 mL/100 lb). Treatment should be repeated at 24-hour intervals for three days. Additional treatments may be given on Days 4 and 5 to animals that have shown clinical improvement but not total recovery.

Single-Dose Therapy (BRD Control): Administer, by subcutaneous injection, a single dose of 7.5 mg/kg of body weight (3.4 mL/100 lb). Examples of conditions that may contribute to calves being at high risk of developing BRD include, but are not limited to.

the following:

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Transportation with animals from two or more farm origins.
An extended transport time with few to no rest stops.
An environmental temperature change of 230°F during transportation.
A 330°F range in temperature fluctuation within a 24-hour period.
Exposure to wet or cold weather conditions.
Excessive shrink (more than would be expected with a normal load of cattle).
Stressful arrival processing procedures (e.g., castration or dehorning).
Exposure within the prior 72 hours to animals showing clinical signs of BRD.
Administered dose volume should not exceed 20 mL per injection site.

	Treatment		Control
Weight (Ib)	Single-Dose Therapy 7.5 - 12.5 mg/kg Dose Volume (mL)	Multiple-Day Therapy 2.5 - 5.0 mg/kg Dose Volume (mL)	Single-Dose Therapy 7.5 mg/kg Dose Volume (mL)
100	3.5 - 5.5	1.5 - 2.0	3.5
200	7.0 - 11.0	2.5 - 4.5	7.0
300	10.5 - 17.0	3.5 - 6.5	10.5
400	14.0 - 22.5	4.5 - 9.0	14.0
500	17.0 - 28.5	5.5 - 11.5	17.0
600	20.5 - 34.0	7.0 - 13.5	20.5
700	24.0 - 39.5	8.0 - 16.0	24.0
800	27.5 - 45.5	9.0 - 18.0	27.5
900	31.0 - 51.0	10.0 - 20.5	31.0
1000	34.0 - 57.0	11.0 - 23.0	34.0
1100	37.5 - 62.5	12.5 - 25.0	37.5

*Dose volumes have been rounded to the nearest 0.5 mL within the dose range. Swine: Administer, either by intramuscular or subcutaneous (behind the ear) injection, a single dose of 7.5 mg/kg of body weight (3.4 mL/100 lb). Administered dose volume should not exceed 5 mL per injection site. For the control of colibacillosis, administration should be initiated within the first 60 days post-weaning when clinical signs are present in at least 2% of the animals in the group. If no improvement is noted within 48 hours, the diagnosis

should be reevaluated

Table 2 - Baytril® 100 Dose Schedule for Swine

Weight (lb)	Dose Volume (mL)	
15	0.5	
30	1.0	
50	1.7	
100	3.4	
150	5.1	
200	6.8	
250	85	

Dilution of Baytril® 100: Baytril® 100 may be diluted with sterile water prior to injection. The diluted product should be used within 24 hours. Store diluted solution in amber glass bottles between 4-40°C (36-104°F). Table 3 – Dilution Schedule*

Swine Weight	mL of Baytril [®] 100	mL of sterile water	Number of doses
10 lb	34 mL	66 mL	100
15 lb	51 mL	49 mL	100
20 lb	68 mL	32 mL	100
25 lb	85 ml	15 ml	100

*For 1 mL dose volume from diluted solution

Use within 30 days of first puncture and puncture a maximum of 30 times with a needle or 4 times with a dosage delivery device. Any product remaining beyond these parameters should be discarded.



RESIDUE WARNINGS: Cattle: Animals intended for human consumption must not be slaughtered within 28 days from the last treatment. This product is not approved for female dairy cattle 20 months of age or older, including dry dairy cows. Use in these cattle may cause drug residues in milk and/or in calves born to these cows. A withdrawal period has not been established for this product in per-uminating calves. Do not use in calves to be processed for yeal. **Swine:** Animals intended for human consumption must not be slaughtered within 5 days of receiving a single-injection dose.

HUMAN WARNINGS:

Not for use in humans. Keep out of reach of children. Avoid contact with eyes. In case of contact, immediately flush eyes with copious amounts of water for 15 minutes. In case of dermal contact, wash skin with soap and water. Consult a physician if irritation persists following ocular or dermal exposures. Individuals with a history of hypersensitivity to quinolones should avoid this product. In humans, there is a risk of user photosensitization within a few hours after excessive exposure to quinolones. If excessive accidental exposure occurs, avoid idrect sunlight. For customer service or to obtain product information, including a Safety Data Sheet, call 1-800-633-3796. For medical emergencies or to report adverse reactions, call 1-800-422-9874.

PRECAUTIONS:

The effects of enrolloxacin on cattle or swine reproductive performance, pregnancy and lactation have not been adequately determined. The long-term effects on articular joint cartilage have not been determined in pigs above market weight. Subcutaneous injection in cattle and swine, or intramuscular injection in swine, can cause a transient local tissue reaction that may result in trim loss of edible tissue at slaughter.

may result in trim loss of edible tissue at slaughter. Baytrile "IO contains different excipients than other Baytrile" products. The safety and efficacy of this formulation in species other than cattle and swine have not been determined. Ouinolone-class drugs should be used with caution in animals with known or suspected Central Nervous System (CNS) disorders. In such animals, quinolones have, in rare instances, been associated with CNS stimulation which may lead to convulsive seizures. Quinolone-class drugs have been shown to produce erosions of cartilage of weight-bearing this and other signs of arthropathy in immature animals of various species. See Animal Safety section for additional information.

ADVERSE REACTIONS: No adverse reactions were observed during clinical trials.

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

MICROBIOLOGY: Errofloxacin is bactericidal and exerts its antibacterial effect by inhibiting bacterial DNA gyrase (a type II topoisomerase) thereby preventing DNA supercoiling and replication which leads to cell death.¹ Enrofloxacin is active against Gram-negative and Gram-positive bacteria.

EFFECTIVENESS: Cattle: A total of 845 calves with naturally-occurring BRD were treated with Baytril® 100 in eight field trials located in five cattle-feeding states. Response to treatment was compared to non-treated controls. Single-dose and multiple-day therapy regimens were evaluated. BRD and mortality were significantly reduced in enrofloxacin-treated calves. No adverse reactions were reported in treated animals.

in treated animals. The effectiveness of Baytril[®] 100 for the control of respiratory disease in cattle at high risk of developing BRD was evaluated in a six-location study in the U.S. and Canada. A total of 1,150 crossbred beef calves at high risk of developing BRD were enrolled in the study. Baytin[®] 100 (7.5 mg/Kg BW) or an equivalent volume of sterile saline was administered as a single subcutaneous injection within two days after arrival. Cattle were observed daily for clinical signs of BRD and were evaluated for success on Day 14 post-treatment. Treatment success in the Baytril[®] 100 group (457/573, 87,83%) was significantly higher (P = 0.0013) than success in the saline control group (455/571, 80.92%). In addition, there were more treatment successes (n = 3) than failures (n = 3) in the group of animals positive for *M. bovis* on Day 0 that were treated with Baytril[®] 100. No product-related adverse reactions were reported.

Subcess in the same Collind (reginer) (450/87), 00.92%). In addition, there were indire treatment of December 2014 that hardings a positive of *M. boxis* on Day 0 that were treated with Baytril® 100. No product-related adverse reactions were reported. **Swine:** A total of 590 pigs were treated with Baytril® 100 or saline in two separate natural infection SRD field trials. For the treatment of SRD, the success rate of enrofloxacin-treated pigs that were defined as "sick and febrile" (increased respiratory rate, abored or dyspineic breathing, depressed attitude and a rectal temperature $\ge 104^{\circ}$) was statistically significantly (nee trial) and morbidity were statistically significantly (new for enrofloxacin-treated pigs in pens containing a percentage of "sick and febrile" pigs compared to saline-treated pigs. The effectiveness of Baytril® 100 administered as a single SC dose of 7.5 mg/kg BW for the treatment and control of SRD associated with *M. hyopneumoniae* was demonstrated using an induced infection model study and three single-site natural infection field studies. In the model study, 72 healthy pigs were challenged with a representative *M. hyopneumoniae* usa demonstrated using an induced infection model study and three single-site natural infection field studies. In the model study, 72 healthy pigs were challenged with a representative *M. hyopneumoniae* usa demonstrated using an induced infection model study and three baytril® 100 or saline. A statistically significant (+ 0.0001) in the saline-treated with Baytril® 100 erasine. At 7 days post-treatment, the cure rate was statistically significantly (higher at each site (+ 0.0001) in the Baytril® 100 or saline. At 7 days post-treatment of SRD, and cure as statistically significantly (higher at each site (+ 0.0001) in the Baytril® 100 or saline. At 7 days post-treatment, the cure rate was statistically significantly (higher at each site (+ 0.0001) in the Baytril® 100 or saline.

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Endet in tabitation and the second were observed in the tabits at does of 25 mg/kg or in rats at 50 mg/kg.
ANIMAL SAFETY:
Cattles: Safety studies were conducted in feeder calves using single doese of 25 mg/kg for 15 consecutive days and 50 mg/kg for 5 consecutive days. No clinical signs of toxicity were observed when a does of 5 mg/kg was administered for 15 days. Clinical signs of toxicity were observed in the second does of 15 mg/kg for 15 consecutive days. No clinical signs of toxicity were observed when a does of 5 mg/kg was administered for 3 days. No during the second does of 5.15 and 25 mg/kg for 15 consecutive days. No clinical signs of depression, incoordination and muscle fasciculation were observed in calves when doese of 15 or 25 mg/kg were administered for 10 to 15 days. Clinical signs of depression, incoordination and muscle fasciculation were observed in calves when doese of 15 or 25 mg/kg for 15 days. A were identified. No articular cartilage lesions were observed at the reaximation of stifle joints from animals administered 25 mg/kg for 15 days. A safety study was conducted in 23-day-old calves using doese of 5, 15 and 25 mg/kg for 15 consecutive days. No clinical signs of toxicity or changes in clinical pathology parameters were observed. No articular cartilage lesions were observed. No articular cartilage lesions were observed in the stifle joints at any dose level at 2 days and 9 days following 15 days of drug administration. An injection site study conducted in feeder calves demonstrated that the formulation may induce a transient reaction in the subcutaneous stately. A safety study was conducted in 16 consecutive days. Including the safine transient controls. Musculoskelat stiffnees was observed following the 5 and 25 mg/kg for 15 consecutive days. Including the safine transitient controls usous observed stiffnees was observed in all groups, including the safine transitient controls. Musculoskelat stiffnees was observed in leador was observed after treatment. Clinical sign

ceased and most animals were clinically normal at necropsy. A second study was conducted in two pigs weighing approximately 23 kg (50 lb), treated with 50 mg/kg for 5 consecutive days. There were no clinical signs of toxicity or pathological changes. An injection site study conducted in pigs demonstrated that the formulation may induce a transient reaction in the subcutaneous tissue. No painful responses to administration were observed. Intramuscular Safety: Asafety study was conducted in 48 weaned, 20- to 22-day-old pigs. Pigs were administered Baytril® 100, at 7.5, 22.5 and 37.5 mg/kg BW by IM injection into the neck once weekly for 3 consecutive weeks. All pigs remained clinically normal throughout the study. Transient decreases in feed and water consumption were observed after each ingtamation was found on post-mortem examination in all enrofloxacin-treated groups. STORAGE CONDITIONS: Protect from direct suplicity. Do not refire the study are a to 37.5 mg/kg BW by BM box.

STORAGE CONDITIONS: Protect from direct sunlight. Do not refrigerate or freeze. Store at 20-30°C (68-86°F), excursions permitted up to 40°C (104°F). Precipitation may occur due to cold temperature. To redissolve, warm and then shake the vial. HOW SUPPLIED:

- Baytril[®] 100: 100 mg/mL 100 mg/mL 100 mg/mL 100 mL Bottle 250 mL Bottle 500 mL Bottle

REFERENCES:

Hooper, D. C., Wolfson, J. S., Quinolone Antimicrobial Agents, 2nd ed, 59 - 75, 1993.
 For customer service or to obtain product information, including a Safety Data Sheet, call 1-800-633-3796.

For medical emergencies or to report adverse reactions, call 1-800-422-9874. Baytril® 100

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