# 150 mg/mL ANTIMICROBIAL

For subcutaneous injection in beef and non-lactating dairy cattle only. Not for use in female dairy cattle 20 months of age or older or in calves to be

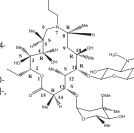
Caution: Federal (USA) law restricts this drug to use by or on the order of a READ ENTIRE BROCHURE CAREFULLY BEFORE USING THIS PRODUCT.

## DESCRIPTION

ZACTRAN® Injection for Cattle is a ready to use sterile parenteral solution containing gamithromycin, a macrolide sub-class, 7a-azalide antimicrobial. Each mL of ZACTRAN contains 150 mg of gamithromycin as the free base, 1 mg of monothioglycerol and 40 mg of succinic acid in a glycerol formal vehicle.

The chemical name of gamithromycin is 1-0xa-7-azacyclopentadecan-15-one.13-[(2.6-

dideov-3-C-methy 3-0-methyl-alpha-Lribo-hexopyranosyl) oxv1-2-ethvl-3,4,10 trihydroxy 3.5.8.10.12.14 hexamethyl-7-propyl 11-{[3,4,6-trideoxy-3-(dimethylamino)-beta-Dxvlo-hexopyranosylloxy}-[(2R\*, 3S\*,4R\*,5S\*,8R\* 10R\*.11R\*.12S\*.13S\*. 4R\*)]-and the structure is shown below.



# INDICATIONS

ZACTRAN is indicated for the treatment of bovine respiratory disease (BRD) associated with Mannheimia haemolytica, Pasteurella multocida, Histophilus somni and Mycoplasma boyis in beef and non-lactating dairy cattle. ZACTRAN is also indicated for the control of respiratory disease in beef and non-lactating dairy cattle at high risk of developing BRD associated with Mannheimia haemolytica and Pasteurella multocida.

# DOSAGE and ADMINISTRATION

Administer ZACTRAN one time as a subcutaneous injection in the neck at 6 mg/kg (2 mL/110 lb) body weight (BW). If the total dose exceeds 10 mL, divide the dose so that no more than 10 mL is administered at each injection site.

Body Weight (lb)	Dose Volume (mL)
110	2
220	4
330	6
440	8
550	10
660	12
770	14
880	16
990	18
1100	20

Animals should be appropriately restrained to achieve the proper route of administration. Use sterile equipment. Inject under the skin in front of the shoulder (see illustration).



### CONTRAINDICATIONS

As with all drugs, the use of ZACTRAN is contraindicated in animals previously found to be hypersensitive to this drug.

FOR USE IN CATTLE ONLY. NOT FOR USE IN HUMANS.

## KEEP THIS AND ALL DRUGS OUT OF REACH OF CHILDREN. NOT FOR USE IN CHICKENS OR TURKEYS.

The Safety Data Sheet (SDS) contains more detailed occupational safety information. To report suspected adverse drug events, for technical assistance, or to obtain a copy of the SDS, contact Boehringer Ingelheim Animal Health USA Inc. at 1-888-637-4251. For additional information. about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS, or online at www.fda.gov/reportanimalae.

RESIDUE WARNINGS: Do not treat cattle within 35 days of slaughter. Because a discard time in milk has not been established, do not use in female dairy cattle 20 months of age or older. A withdrawal period has not been established for this product in pre-ruminating calves. Do not use in calves to be processed for veal.

# PRECAUTIONS

The effects of ZACTRAN on bovine reproductive performance, pregnancy, and lactation have not been determined. Subcutaneous injection of ZACTRAN may cause a transient local tissue reaction in some cattle that may result in trim loss of edible tissues at slaughter.

### ADVERSE REACTIONS

Transient animal discomfort and mild to moderate injection site swelling may be seen in cattle treated with ZACTRAN.

The macrolide antimicrobials as a class are weak bases and as such

concentrate in some cells (such as pulmonary leukocytes). Prolonged

exposure of extracellular pulmonary pathogens to macrolides appears to

reflect the slow release of drug from its intracellular reservoir to the site

of action, the pulmonary epithelial lining fluid (ELF). It is the ELF that is

## CLINICAL PHARMACOLOGY

relevant to the successful treatment and control of BRD. Gamithromycin is primarily bacteriostatic at therapeutic concentrations. However, in vitro bactericidal activity has been observed at concentrations of 10 μg/mL (Mueller-Hinton broth) and after exposure to the 6-hour and 24-hour plasma samples derived from cattle dosed at 6 mg gamithromycin/kg BW. Macrolides typically exhibit substantially higher concentrations in the alveolar macrophages and ELF as compared to concentrations observed in plasma. Gamithromycin concentrations in the ELF and ELF cells exceed the concentrations observed in the plasma. Postmortem gamithromycin concentrations in ELF exceed the MIC<sub>90</sub> of M. haemolytica, H. somni and P. multocida through at least 72 hours after drug administration. Because M. haemolytica, P. multocida and H. somni are extracellular pathogens. drug concentrations in the ELF are considered to be clinically relevant. The postmortem area under the concentration-time curve (AUC) observed in lysed ELF cells (e.g., alveolar macrophages) are at least 300-times greater than that in the plasma. Although published studies suggest that inflammation can increase the release of drug from macrophages and neutrophils, these high concentrations in the alveolar macrophages should not be considered indicative of the magnitude or duration of response to the pathogens for which this product is indicated. ZACTRAN administered subcutaneously in the neck of cattle at a single dosage of 6 mg/kg BW is rapidly and completely absorbed, with peak concentrations generally occurring within 1 hour after administration. Based upon plasma and lung homogenate data, the terminal half-life (T½) of gamithromycin is approximately 3 days. In vitro plasma protein binding studies show that 26% of the gamithromycin binds to plasma protein, resulting in free drug available for rapid and extensive distribution into body tissues. The free drug is rapidly cleared from the systemic circulation with a clearance rate of 712 mL/hr/kg and a volume of distribution of 25 L/kg. Dose proportionality was established based on AUC over a range of 3 mg/kg BW to 9 mg/kg BW. Biliary excretion of the unchanged drug is the major route of elimination

# MICROBIOLOGY

The minimum inhibitory concentrations (MIC's) of gamithromycin were determined for BRD isolates obtained from calves enrolled in BRD treatment field studies in the U.S. in 2004 using methods recommended by the Clinical and Laboratory Standards Institute (M31-A2). Isolates were obtained from pre-treatment nasopharyngeal swabs from each enrolled calf and from calves removed from the study due to BRD. The results are shown above in Table 1.

Table 1. Gamithromycin minimum inhibitory concentration (MIC) values\* of indicated pathogens isolated from BRD treatment

Indicated Pathogens	Years of isolation	No. of isolates	MIC <sub>so</sub> ** (μg/mL)	MIC <sub>90</sub> ** (μg/mL)	MIC range (μg/mL)
M. haemolytica	2004	89	1	1	0.5 to >32
P. multocida	2004	79	0.5	1	0.12 to >32
H. somni	2004	32	0.5	0.5	0.25 to 1

- \* The correlation between in vitro susceptibility data and clinical effectiveness is unknown
- \*\* The lowest MIC to encompass 50% and 90% of the most susceptible isolates, respectively

#### FFFFCTIVENESS

The effectiveness of ZACTRAN for the treatment of BRD associated with Mannheimia haemolytica, Pasteurella multocida and Histophilus somni was demonstrated in a field study conducted at four geographic locations in the United States. A total of 497 cattle exhibiting clinical signs of BRD were enrolled in the study. Cattle were administered ZACTRAN (6 mg/kg BW) or an equivalent volume of sterile saline as a subcutaneous injection once on Day O. Cattle were observed daily for clinical signs of BRD and were evaluated for clinical success on Day 10. The percentage of successes in cattle treated with ZACTRAN (58%) was statistically significantly higher (p<0.05) than the percentage of successes in the cattle treated with saline (19%). The effectiveness of ZACTRAN for the treatment of BRD associated with M. bovis was demonstrated independently at two U.S. study sites. A total of 502 cattle exhibiting clinical signs of BRD were enrolled in the studies. Cattle were administered ZACTRAN (6 mg/kg BW) or an equivalent volume of sterile saline as a subcutaneous injection once on Day 0. At each site, the percentage of successes in cattle treated with ZACTRAN on Day 10 was statistically significantly higher than the percentage of successes in the cattle treated with saline (74.4% vs. 24% [p <0.001], and 67.4% vs. 46.2% [p = 0.002]). In addition, in the group of calves treated with gamithromycin that were confirmed positive for M. bovis (pre-treatment nasonharyngeal swahs), there were more calves at each site (45 of 57 calves, and 5 of 6 calves) classified as successes than as failures. The effectiveness of ZACTRAN for the control of respiratory disease in cattle at high risk of developing BRD associated with Mannheimia haemolytica and Pasteurella multocida was demonstrated in two independent studies conducted in the United States. A total of 467 crossbred beef cattle at high risk of developing BRD were enrolled in the study. ZACTRAN (6 mg/kg BW) or an equivalent volume of sterile saline was administered as a single subcutaneous iniection within one day after arrival. Cattle were observed daily for clinical signs of BRD and were evaluated for clinical success on Day 10 post-treatment. In each of the two studies, the percentage of successes in the cattle treated with ZACTRAN (86% and 78%) was statistically significantly higher (p = 0.0019 and p = 0.0016) than the percentage of successes in the cattle treated with saline (36% and 58%).

In a target animal safety study in healthy, six-month old beef cattle. ZACTRAN was administered by subcutaneous injection at 6, 18, and 30 mg/ kg bodyweight (1, 3, and 5 times the labeled dose) on Day 0, 5, and 10 (3 times the labeled administration frequency). Injection site discomfort (neck twisting, attempts to scratch or lick the injection site, and pawing at the ground) was observed in calves in the 18 mg/kg BW and 30 mg/kg BW groups at 10 minutes post-treatment following each injection. Mild to moderate injection site swelling and pathology changes consistent with inflammation were observed in the gamithromycin-treated groups. Other than injection site reactions, no clinically relevant treatment-related effects were observed.

# STORAGE CONDITIONS

Store at or below 77°F (25°C) with excursions between 59-86°F (15-30°C) Use within 18 months of first puncture

7ACTRAN is available in three ready-to-use bottle sizes. The 100, 250 and 500 mL bottles contain sufficient solution that will treat 10, 25 and 50 head of 550 lb (250 kg) cattle respectively.

# Marketed by Boehringer Ingelheim Animal Health USA Inc.

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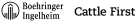
Boehringer Ingelheim



FAST. EFFECTIVE. LONG-LASTING.







# WHEN BRD STRIKES, FIGHT BACK WITH ONE POWERFUL DOSE.

<b>ACTRAN</b> °	MACROLIDES			BETA-LACTAM	FLUOROQUINOLONE	PHENICOLS		
(gamithromycin)	ZACTRAN®1	MICOTIL*2	ZUPREVO*2	DRAXXIN*2	<b>EXCEDE</b> ®2	BAYTRIL® 100°	NUFLOR*2	RESFLOR GOLD <sup>®2</sup>
Active Ingredient	gamithromycin	tilmicosin	tildipirosin	tulathromycin	ceftiofur crystalline	enrofloxacin	florfenicol	florfenicol + flunixin meglumine
BRD Pathogens Labeled for the treatment of:								
Mannheimia haemolytica	X	X	X	X	X	X	X	X
Pasteurella multocida	Χ	Х	Χ	X	X	Χ	Χ	Χ
Histophilus somni	Χ	Х	Χ	X	Χ	X	X	X
Mycoplasma bovis	Х			X		X		Χ
Time to Peak Concentration*	12 Hours	24 Hours	24 Hours	72 Hours	N/A	N/A	N/A	N/A
Challenge Study Duration of Activity	10 Days	-	-	14 Days	7 Days	-	-	-
Safety and Ease of Use Injection Site	Neck	Neck	Neck	Neck	Behind Ear	Neck	Neck	Neck
Auto Syringes	Yes	No (Injection of this drug in humans has been associated with fatalities.)	No	Yes	Yes (Injection in artery can kill cattle.)	Yes	Yes	Yes
Dose	1.8mL/100 lbs.	1.5-3.0 mL/100 lbs.	1.0 mL/100 lbs.	1.1 mL/100 lbs.	1.5 mL/100 lbs.	3.4-5.7 mL/100 lbs./ (single) 1.1-2.3 mL/100 lbs./ (multi-day)	3.0 mL/100 lbs. (IM) 2 doses required, repeat in 48 hrs; 6.0 mL/100 lbs. (SC)	6.0 mL/100 lbs.
Slaughter Withdrawal	35 Days	42 Days	21 Days	18 Days	13 Days	28 Days	38 Days (SC)	38 Days

<sup>\*</sup>Clinical relevance has not been determined



IMPORTANT SAFETY INFORMATION: Do not treat cattle within 35 days of slaughter. Because a discard time in milk has not been established, do not use in female dairy cattle 20 months of age or older, or in calves to be processed for veal. Subcutaneous injection may cause a transient local tissue reaction in some cattle that may result in trim loss of edible tissues at slaughter.

<sup>&</sup>lt;sup>2</sup> ZUPREVO, MICOTIL, EXCEDE, DRAXXIN, BAYTRIL 100, NUFLOR product labels.