Date of approval.

FREEDOM OF INFORMATION SUMMARY

Micotil® 300 Injection (tilmicosin phosphate)

Supplement to NADA 140-929

"...for the treatment of ovine respiratory disease (ORD) associated with Mannheimia (Pasteurella) haemolytica."

> SUBMITTED BY: ELANCO ANIMAL HEALTH



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1. GENERAL INFORMATION:

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a.	NADA Number:	140-929			
b.	Sponsor:	Elanco Animal Health A Division of Eli Lilly & Co., Lilly Corporate Center, Indianapolis, IN 46285			
		Drug Labeler Code: 000986			
c.	Established Name:	Tilmicosin phosphate			
d.	Proprietary Name:	Micotil® 300 Injection			
e.	Dosage Form:	Ready-to-use injectable solution			
f.	How Supplied:	50 mL, 100 mL, and 250 mL multidose amber glass bottles			
g.	How Dispensed:	R _x			
h.	Amount of Active Ingredient:	300 mg tilmicosin as tilmicosin phosphate per mL			
i.	Route of Administration:	subcutaneous injection			
j.	Species/Class:	Ovine/sheep			
k.	Recommended Dosage:	a single subcutaneous injection of 10 mg/kg body weight (1 mL/30 kg or 1.5 mL /100 lb body weight)			
1.	Pharmacological Category:	antimicrobial			
m.	Indications:	Micotil [®] 300 Injection is indicated for the treatment of ovine respiratory disease (ORD) associated with Mannheimia (Pasteurella) haemolytica.			
n.	n. Effect of Supplement: Provides for the use of tilmicosin phosphate (Micotil® 300) ir a new animal species, sheep.				

2. EFFECTIVENESS:

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Section 514.1(d) of Title 21 of the Code of Federal Regulations (CFR) permits extrapolation of data from a major species to a minor species to satisfy the requirements of section 512 of the Federal Food, Drug, and Cosmetic Act with respect to the effectiveness of a new animal drug. A combination of data from sheep (a minor species) and a closely-related approved major species (cattle) were used to support the determination of effectiveness, consistent with the Guideline for Industry – FDA Approval of New Animal Drugs for Minor Uses and minor Species (Guideline #61 FDA/CVM April 1999).

For the purposes of this supplement for use in sheep, a determination of medical equivalence was based on a pharmacokinetic comparison demonstrating that serum concentrations of tilmicosin in sheep and cattle are comparable when tilmicosin is administered as a single subcutaneous injection at a rate of 10 mg/kg. The data for the effectiveness study was generated under INAD 9693 and submitted in a Public Master File (PMF) 5673. A notice of availability of this PMF is published in the Federal Register (65 FR 47992, August 4, 2000)..

3. TARGET ANIMAL SAFETY:

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The comparative study of tilmicosin pharmacokinetics, conducted under INAD 9693 and published under PMF 5673, in sheep and cattle indicates that the target animal safety of tilmicosin should be comparable in sheep and cattle when administered as a single subcutaneous injection at 10 mg/kg. None of the animals died during the study and no adverse effects of tilmicosin were observed on blood pressure, heart rate, or respiration rate of sheep. The results indicate that tilmicosin can be used safely in sheep at the recommended dose for sheep with a minimum body weight of 15 kg. The availability of this data in PMF 5673 is published in the Federal Register (65 FR 47992, August 4, 2000.

4. HUMAN SAFETY:

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a. Toxicology

See the Freedom of Information (FOI) Summary for the approval of the original application for MICOTIL® 300 (NADA 140-929), approved March 24, 1992. A copy of this FOI summary can be obtained from Mrs. Marylin H. Broderick, HFV-12, Center for Veterinary Medicine, 7500 Standish Place, Rockville, MD 20855.

b. Safe Concentration of Total Residues:

- Acceptable Daily Intake (ADI): The ADI previously established for tilmicosin is 1.5 mg/person/day or (25 µg/kg body weight/day for a 60 kg person). Ref. 21 CFR 556.735.
- 2) Safe Tissue Concentrations (STC):

The STC is calculated by dividing the ADI per person by the tissue consumption factors (300 g/day for muscle, 50 g/day for kidney or fat, and 100 g/day for liver.

 $STC_{(muscle)} = 1.5 \text{ mg/day} \div 300 \text{ g/day} = 5 \text{ }\mu\text{g/g} = 5 \text{ }ppm$ $STC_{(kidney \text{ or fat})} = 1.5 \text{ }m\text{g/day} \div 50 \text{ }g\text{/day} = 30 \text{ }\mu\text{g/g} = 30 \text{ }ppm$ $STC_{(liver)} = 1.5 \text{ }m\text{g/day} \div 100 \text{ }g\text{/day} = 15 \text{ }\mu\text{g/g} = 15 \text{ }ppm$

c. Total Residue and Metabolism Studies

Total residues of ¹⁴C-tilmicosin in sheep tissues were evaluated in two studies:

1) Study HRC/LLY36/930447

The Metabolism and Residues of ¹⁴C Tilmicosin Following Subcutaneous Administration to Sheep. L.F. Elsom, D.R. Hawkins, D.H. Dighton, A. Kaur, D.M. Cameron, 1993.

Huntingdon Research Centre Ltd., Huntingdon, Cambridgeshire, U.K.

The purpose of this study was to evaluate the excretion profile of ¹⁴C tilmicosin; to assess the depletion of total radioactive residues in edible tissues; to establish the metabolic profiles in edible tissues and compare them with the metabolic profile in urine and to identify and quantify the major metabolites of tilmicosin in sheep.

Fourteen, 10 to 11 week old Beulah-cross lambs, 16 to 23 kg body weight were administered a single subcutaneous injection of ¹⁴C tilmicosin at 20 mg/kg body weight. Plasma samples were collected at 1, 2, 4, 6, 8, 12, 24, 36, 48, 72, 96, 120, 144 and 168 hours after dosing. Urine and feces were collected at pre-dose and in 24-hour intervals for up to 7 days from time of dosing. Tissue samples of liver, kidney, lung, skeletal muscle, fat and injection site were collected for radioactivity assay and TLC and HPLC analyses.

A mean total of 85.2% of the radioactivity dosed was excreted in the 7 days after dosing. The majority of the radioactivity was excreted in the feces (a mean of 71.9%). The urine contained a mean of 13.2% of the total dose in the 7 days. Mean concentrations of radioactivity in tissues of treated sheep are summarized in Table 4.1.

Parent tilmicosin accounted for approximately 75% of the urinary radioactivity with metabolites T-1 and T-2 accounting for <1% of the dose. In the liver and kidney, parent tilmicosin and metabolite T-2 were the major components. The concentration of parent tilmicosin in the liver declined while T-2 correspondingly increased as withdrawal time increased.

Table 4.1: Concentrations (ppm) (Mean \pm SD) of radioactivity in tissues of sheep following a single subcutaneous dose of ¹⁴C-tilmicosin at a dose of 20 mg/kg body weight

Tissue		Sacrifie	e (days)	
	3	7	21	28
Liver	9.98±0.78	5.77±0.30	3.67±0.72	2.70±0.60
Kidneys	21.09±4.49	4.07±1.35	1.42±0.89	0.55±0.13
Omental Fat	<1.36	<1.32	<1.32	<1.35
Renal Fat	<1.24	<1.15	<1.17	<1.20
Muscle	1.26±0.18	0.57	< 0.26	<0.26
Lungs	5.11±0.08	1.53±0.07	NS*	NS
Inj. Site Skin	63.02±25.80	18.74±5.50	32.91±28.83	6.51±5.69
Inj. Site Muscle	43.19±0.33	14.38±2.11	5.32±6.00	1.32±0.51
*Not compled		11.502.2.11	5.52±0.00	1.52±(

*Not sampled

2) Study CVLS4/92

Tilmicosin: Metabolism and Residues of ¹⁴C Tilmicosin Following Subcutaneous Administration to Sheep. R. M. Parker and A.M. Walker, 1993.

Central Veterinary Laboratory, Weybridge, Addlestone, KT15 3NB, U.K.

The depletion and quantification of ¹⁴C tilmicosin in plasma and tissues was studied in 16 sheep after subcutaneous administration of 20 mg/kg body weight tilmicosin. Plasma was collected at 1, 2, 3, 4, 6, 8, 12, 24, 36, 48, 72, 96, 120, 144 and 168 hours after dosing. Kidney, liver, fat, muscle, injection site and lung tissues were collected at days 3, 7, 21 and 28 post-injection. The plasma and tissue samples were assayed by validated HPLC methods.

Values for the mean plasma concentration of tilmicosin ranged from 958.8 ng/mL (T_{max}) at 4 hours post-injection to below the validated limit of determination (50 ng/mL or 0.05 ppm) at 48 hours post-injection. Between 6 and 24 hours, t_{y} was approximately 7 hours and between 48 and 96 hours, t_{y} was approximately 41 hours.

Concentrations of tilmicosin were highest in the injection site and kidney at Day 3 post-injection. At Day 28, mean tilmicosin concentration was 160.0 ng/g (0.16 μ g/g) in the liver, 122 ng/g (0.12 μ g/g) in the injection site and 63 ng/g (0.63 μ g/g) in the kidney. Tilmicosin was not detected in muscle and fat 21 and 28 days after dosing above the limit of detection of the method (50 ppb). Mean concentrations (ppb) of tilmicosin in the tissues of treated sheep are summarized in Table 4.2.

Tissue				
	3	7	21	28
Liver	2444.0±278.6*	733.0±43.8	310.0±162.7	160.0±116.2
Kidney	12414.0±5226.5	1286.0±500.6	467.3±352.9	73.0±29.9
Muscle	478.5±186.0	193.5±202.9	ND†	ND
Renal Fat	73.0±14.1	ND	ND	ND
Injection Site	20352±3369.7	7067.3±1019.1	3626.5±3544.6	121.8±28.4
Lung	2810.3±871.0	322.5±111.0	NS††	NS

Table 4.2: Concentrations (mean \pm SD) of tilmicosin in sheep tissues

*Values are reported as ppb

†Not determined

††Not sampled

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Sheep injected subcutaneously with ¹⁴C tilmicosin at 20 mg/kg body weight excrete 85% of the dose within the first 7 days. Plasma concentrations of tilmicosin decline close to the limits of detection by 36 hours. Tissue concentrations of tilmicosin in all tissues are less than the safe concentration for total residues at all sampling times studied and well below the safe concentrations 28 days after dosing.

Because parent tilmicosin accounts for a substantial portion of the dose in urine and of the residue in kidney and liver, it is selected as the marker residue. As in cattle, liver is selected as the target tissue.

Although a separate comparative metabolism report was not provided, a review of work previously conducted to support NADA 140-929 along with the metabolism data provided for sheep demonstrate that the metabolic profiles for sheep and the toxicological species, the rat, are comparable. Therefore, the toxicological species has been autoexposed to the metabolites of tilmicosin present in the edible tissues of treated sheep.

d. Tolerance for the Marker Residue

The total residue and metabolism study in sheep demonstrates that the marker residue, parent tilmicosin, represents approximately 20% of the total residue in liver. When this percentage is applied to the calculated safe concentrations, a tolerance of 3 ppm is calculated for residues of tilmicosin in liver. Applying the 20% to the muscle safe concentration results in a calculated muscle tolerance of 1 ppm. However, consistent with the requested withdrawal period, we are assigning tolerances of 1.2 ppm for sheep liver and 0.1 ppm for sheep muscle. A liver tolerance of 1.2 ppm is the same tolerance value currently assigned to cattle. A muscle tolerance of 0.1 ppm for residues of parent tilmicosin in sheep muscle is the same as the muscle tolerance assigned to cattle and consistent with the Maximum Residue Limit recommended by the 47th meeting of the Joint FAO/WHO Expert Committee on Food Additives (JECFA), Rome 1996. Additionally, the analytical method has an LOO of half the proposed muscle tolerance and can serve as a monitoring method for residues of tilmicosin in sheep tissues. We propose to assign a withdrawal period of 28 days for the use of tilmicosin, as Micotil® 300, in sheep as a single subcutaneous injection at a dose of up to 10 mg/kg body weight.

e. Withdrawal Period

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Depletion of residues in the edible tissue of sheep was evaluated in two studies:

1) Study CVLS6/91

Tilmicosin: Residues in Sheep. R.K.P. Patel, R.M. Parker and H.A. Simmons, 1992.

Analytical Chemistry Unit, Central Veterinary Laboratory, New Haw, Weybridge, Surrey KT15 3NB.

The purpose of this study was to measure residues of tilmicosin in sheep after subcutaneous administration to establish an approximate withdrawal period for the parent drug.

Twenty-four 6 month old sheep, Scottish Blackface, 30 to 40 kg body weight were administered single subcutaneous doses of 30 mg tilmicosin/kg body weight into the left dorsolateral chest wall. Liver, kidney, muscle, fat and injection site tissues were collected at Days 14, 21, 28, 35, 42 and 56 post-injection and analyzed by a validated HPLC method with a limit of determination of 50 ng/g (50 ppb).

The highest mean concentration of tilmicosin was in liver at Day 14 (1554 ng/g) and the minimum concentration of tilmicosin was obtained in muscle and fat (all values were below the limit of determination). At Day 28, the mean concentration of tilmicosin in liver was 418.0 ng/g, at injection site, 150.3 ng/g, and in kidney, 82.0 ng/g. At Day 28, residues in muscle and fat residues fell below the limit of determination (50 ppb). Mean concentrations (ppb) of tilmicosin in sheep tissues are summarized in Table 4.3.

Tissue Withdrawal (days) 14 21 28 35 42 56 Liver 1554±160 1170±213 418±208 310±126 199±106 81±19 Kidney 478±178 148±65 93±6 68±4 <100 51 Muscle <LOQ <LOO <LOO <LOQ <LOQ <LOQ Fat <LOO <LOQ <LOQ <LOO <LOO <LOQ Injection 506±310 641±469 150±95 159±103 161±84 81±29 Site

Table 4.3: Concentration (ppb) of tilmicosin residues in sheep tissues following subcutaneous administration of Micotil® 300 at a dose of 30 mg/kg body weight

LOQ=50 ppb

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Tilmicosin was metabolized and/or excreted to a great extent during the study interval. Because the dose was three fold higher than the proposed therapeutic dose, residues would be expected to be lower in animals treated with the proposed dose, 10 mg/kg.

2) Study CVLS/23/95

Tilmicosin: Residues in Sheep after its subcutaneous administration. R.K.P. Patel and R.M. Parker, 1995.

Analytical Chemistry Unit, Central Veterinary Laboratory, New Haw, Addlestone, Surrey KT15 3NB.

The purpose of this study was to determine the residues of tilmicosin in tissues at various withdrawal time points after its subcutaneous administration to sheep at the intended use level.

Twenty-eight Swaledale sheep in the weight range 26.2 to 51.2 kg body weight were divided into seven groups of four sheep per group, consisting of 2 males and 2 females. Six of the seven groups were administered single subcutaneous doses of 10 mg tilmicosin/kg body weight into the left dorsolateral chest wall. The seventh group was designated as control and received no injection. Each group was designated for slaughter at one of the following times post-treatment: 14, 21, 28, 35, 42, or 49 days. (The four sheep in the control group were designated for slaughter at 14 days.)

Liver, kidney, thigh muscle, renal fat and injection site tissues were collected at slaughter and analyzed by a validated HPLC method. Mean concentrations (ppb) of tilmicosin in sheep tissues are summarized in Table 4.4.

Tissue			Withdra	awal (days)	
	14	21	28	35	42	49
Liver	107±11	80±43	<loq< td=""><td>59</td><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<>	59	<loq< td=""><td><loq< td=""></loq<></td></loq<>	<loq< td=""></loq<>
Kidney	162±76	73±16	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""></loq<></td></loq<>	<loq< td=""></loq<>
Muscle	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""></loq<></td></loq<>	<loq< td=""></loq<>
Fat	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""></loq<></td></loq<>	<loq< td=""></loq<>
Injection Site	1527±504	143±55	80±30	<loq< td=""><td><loq< td=""><td><loq< td=""></loq<></td></loq<></td></loq<>	<loq< td=""><td><loq< td=""></loq<></td></loq<>	<loq< td=""></loq<>

Table 4.4: Concentration (ppb) of tilmicosin residues in sheep tissues following subcutaneous administration of Micotil® 300 at a dose of 10 mg/kg body weight

LOQ=50 ppb

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The highest mean concentration of tilmicosin was in injection site tissue at Day 14 (1527 ng/g for the average of the four sheep) and the minimum concentration of tilmicosin was obtained in muscle and fat (all values were below the LOQ for the method).

The mean residue concentration in the liver tissue was 107 ng/g at Day 14. Residues in liver were less than the LOQ by Day 49.

Summary of the two residue studies is presented in Table 4.5.

Study no.	Dose	Liver	Kidney	Muscle	Fat	Injection Site
HRC/LLY36/ 930447	¹⁴ C 20 mg/kg	2.70±0.60	0.55±0.13	<0.26*	<1.20*	1.32±0.51
CVLS4/92	¹⁴ C 20 mg/kg	0.16±0.12	0.07±0.03	ND†	ND	0.12±0.03
CVLS6/91	cold 30 mg/kg	0.42±0.21	0.09±0.01	ND	ND	0.16±0.1
CVLS/23/95	cold 10 mg/kg	<loq< td=""><td><loq< td=""><td>ND</td><td>ND</td><td>.080±0.03</td></loq<></td></loq<>	<loq< td=""><td>ND</td><td>ND</td><td>.080±0.03</td></loq<>	ND	ND	.080±0.03

Table 4.5. Summary of residues detected at Day 28 post-injection for the two studies

*Below limits of detection †not detected Limit of quantitation (LOQ) = 0.05 μg/g

These studies confirm that residues of tilmicosin resulting from a single subcutaneous injection of Micotil® 300 to sheep at a dose of 10 mg/kg body weight are below the assigned liver and muscle tolerance levels 28 days after treatment, the assigned withdrawal period.

f. Analytical Methods for Residues

The regulatory analytical method for residues is a reverse phase HPLC method with UV detection. The method is available from Residue Chemistry Team (HFV-151), Center for Veterinary Medicine, 7500 Standish Place, Rockville, MD 20855.

g. User Safety Concerns

User safety concerns associated with direct contact have been satisfactorily addressed by establishing label warnings. In addition, a toll-free number is provided on the label for additional information and reporting adverse events.

5. AGENCY CONCLUSIONS:

The data submitted in support of this supplemental NADA satisfy the requirements of Section 512 of the Food, Drug, and Cosmetic Act and 21 CFR 514.1 of the implementing regulations. The data demonstrate that Micotil® 300 Injection (tilmicosin phosphate), when used under labeled conditions of use is safe and effective for the treatment of ovine respiratory disease (ORD) associated with *Mannheimia* (*Pasteurella*) haemolytica.

The human food safety data demonstrate that residues resulting from a single subcutaneous injection of tilmicosin to sheep at a dose of 10 mg/kg body weight are below the assigned liver and muscle tolerance levels 28 days after treatment, the assigned withdrawal period.

The product remains a prescription drug for safe and effective use by a veterinarian for the treatment of properly diagnosed pneumonia in sheep.

This approval does not qualify for marketing exclusivity under Section 512(c)(2)(F)(iii) of the Federal Food, Drug, and Cosmetic Act.

In accordance with 21 CFR 514.106(b)(2)(vii), this is a Category II change. This supplement provides for the use of tilmicosin in sheep, a new animal species. The approval of this change is not expected to have any adverse effect on the safety or effectiveness of this new animal drug. Accordingly, this approval did not require a reevaluation of the safety and effectiveness data in the parent application.

Tilmicosin is under U.S. patent number 4,820,695 expiring April 11, 2006.

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ATTACHMENTS: 6.

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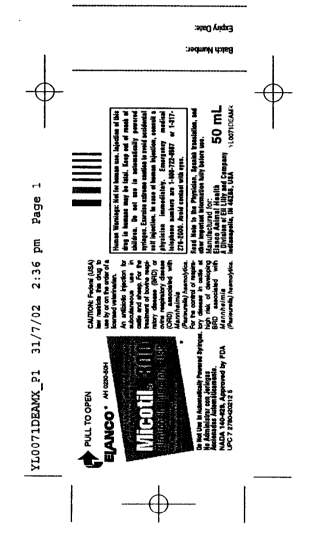
A. Facsimile R_x labeling for 50 mL, 100 mL, and 250 mL bottles. B. Facsimile package insert

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DOTTED LINES INDICATE OVERPRINT AREA ONLY, AND ARE NOT TO BE PRINTED.

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CAUTION: Do Not Administer to Swine. Injection in Swine Has Been Shown to be Fatal.



WARNINGS: Animals intended for human consumption must not be slaughtered within 28 days of the last treatment. Do not use in female dairy cattle 20 months of age or older. Use of tilmicosin in this class of cattle may cause milk residues. Do not use in lactating ewes if the milk is intended for human consumption.

CAUTION: The safety of tilmicosin has not been established in pregnant cattle and in animals used for breeding purposes. Intramuscular injection will cause a local reaction which may result in trim loss. The safety of tilmicosin 12

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NADA 140-929, Approved by FDA Micotil® 300 Injection

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Tilmicosin injection, USP CAUTION: Federal (USA) law restricts this drug to use by or on the orde of a licensed veterinarian.

Human Warnings: Not for human use. Injection of this drug in huma may be fatal. Keep out of reach of children. Do not use in automatica powered syringes. Exercise extreme caution to avoid accidental sinjection. In case of human injection, consult a physician immediate Emergency medical telephone numbers are 1-800-722-0987 1-317-276-2000. Avoid contact with eyes.



NOTE TO THE PHYSICIAN: The cardiovascular system appears to be the target of toxicity. This antibiotic persists in tissues for several days. The cardiovascular system should be monitored closely and supportive treatment provided. Dobutamine partially offset the negative inotropic effects induced by Micotillo in dogs. β -adrenergic antagonists, such as propranoiol, exacerbated the negative inotropy of Micotil-induced tachycardia in dogs. Epinephrine potentiated lethality of Micotillo in pigs.

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For cattle, injection under the skin behind the shoulders and over the rit suggested.

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For sheep, injection in a skin fold behind the shoulders and over the ric suggested.

Note-Swelling at the subcutaneous site of injection may be observed by transient and usually mild.

CONTRAINDICATION: Do not use in automatically powered syringes. Do administer intravenously to cattle or sheep. Intravenous injection in catt! sheep will be fatal. Do not administer to animals other than cattle or sh Injection of this antibiotic has been shown to be fatal in swine and non-hur primates, and it may be fatal in horses and goats.

Mannheimia (Pasteurelia) haemolytica for at least 3 days following the single injection. Serum tilmicosin levels are a poor indicator of total body tilmicosin. The lung/serum tilmicosin ratio in favor of lung tissue appeared to equilibrate by 3 days post injection at approximately 60. In a study with radioactive tilmicosin, 24% and 68% of the dose was recovered from urine and feces, respectively, over 21 days.

Directions—Inject Subcutaneously in Cattle and Sheep Only. Administer a single subcutaneous dose of 10 mg/kg of body weight (1 mL/30 kg or 1.5 mL per 100 ibs).

Do not inject more than 15 mL per injection site. Do not use in lambs less than 15 kg body weight. If no improvement is noted within 48 hours, the diagnosis should be reevaluated.

ADVERTENCIAS PARA HUMANOS: No usarlo en humanos. 1 inyección de esta droga en humanos puede ser fatal. Conservar fuera del alcance de los niños. No administrar con jaringaccionadas automáticamente. Ejercer precauciones extremas pa evitar inyectarse uno mismo. En casos de inyección a humane consultar con un médico inmediatamente. Los números telefónica para emergencias médicas son 1-800-722-0987 o 1-317-276-20C Evitar contacto con ojos.

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NOTA AL MEDICO: EL SISTEMA CARDIOVASCULAR PARECE SER EL BLANCO DE LA TOXICIDAD DE ESTE PRODUCTO. ESTE ANTIBIÓTICO PERSISTE EN LOS TEJIDOS POR VARIOS DIAS. EL SISTEMA CARDIOVASCULAR DEBERA OBSERVARSE CUIDADOSAMENTE Y TRATAMIENTO DE SOPORTE DEBERA PROPORCIONARSE. DOBUTAMINA PARCIALMENTE BLOQUEA LOS EFECTOS INOTROPICOS NEGATIVOS INDUCIDOS POR MICOTIL EN PERROS. β-ADRENERGICOS ANTAGO-NISTAS, COMO EL PROPRANOLOL, EXCERBARON LA INOTROPIA NEG-ATIVA DE LA TAQUICARDIA INDUCIDA POR MICOTIL EN PERROS. LA EPINEFRINA POTENCIALIZA LA LETALIDAD DE MICOTIL EN CERDOS.

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minimal myocardial necrosis in some animals in the 50 mg/kg gr-Subcutaneous doses of 150 mg/kg injected at 72-hour intervals resulted in de. Edema was marked at the site of injection. Minimal myocardial necrosis was only lesion observed at necropsy. Deaths of cattle have been observed w: single intravenous dose of 5 mg/kg of body weight.

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In sheep, single subcutaneous injections of 10 mg/kg dose did not cause deaths and no adverse effects of tilmicosin were observed on blood press heart rate, or respiratory rate.

Pharmacology: A single subcutaneous injection of Micotil® at 10 mg/kg of the weight dose in cattle resulted in peak tilmicosin levels within one hour detectable levels (0.07 µg/mL) in serum beyond 3 days. However, lung concertions of tilmicosin remained above the tilmicosin MIC 95% of 3.12 µg/mI

In monkeys, a single intramuscular dose of 10 mg/kg caused no signs of toxicity. A single dose of 20 mg/kg caused vomiting and 30 mg/kg caused the death of the only monkey tested.

In swine, intramuscular injection of 10 mg/kg caused increased respiration, emesis, and a convulsion, 20 mg/kg resulted in mortality in 3 of 4 pigs, and 30 mg/kg caused the death of all 4 pigs tested.

Results of genetic toxicology studies were all negative. Results of teratology and reproduction studies in rats were negative. The no effect level in dogs after daily oral doses for up to one year is 4 mg/kg of body weight.

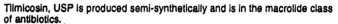
In cattle, subcutaneous doses of 10, 30, and 50 mg/kg of body weight, each injected 3 times at 72 hour intervals did not cause any deaths. As expected, edema at the site of injection was noted. The only lesion observed at necropsy was 8

For Subcutaneous Use in Cattle and Sheep Only. Do Not Use in Automatically Powered Syringes.

Indications: Micotile 300 is indicated for the treatment of bovine respire disease (BRD) and ovine respiratory disease (ORD) associated Mannheimia (Pasteureila) haemolytica. Miccuile 300 is indicated for control of respiratory disease in cattle at high risk of developing t associated with Mannheimia (Pasteurella) haemolytica,

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Description: Micotile 300 is a solution of the antibiotic tilmicosin. Each contains 300 mg of tilmicosin, USP as tilmicosin phosphate in 2 propylene glycol, phosphoric acid as needed to adjust pH and water injection, q.s.



Actions: Activity—Tilmicosin has an *in-vitro** antibacterial spectrum that is predominantly gram-positive with activity against certain gram-negative microorganisms. Activity against several mycoplasma species has also been detected.



Ninety-five percent of the Mannheimia (Pasteurella) haemolytica isolates were inhibited by 3.12 µg/mL or less.

Microorganism	MIC* (ug/mL)
Mannheimia (Pasteurella) haemolytica	3.12
Pasteurella multocida	6.25
Haemophilus somnus	6.25
6	

Mycoplasma dispar	0.097
M. bovirhinis	0.024
M. bovoculi	0.048

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*The clinical significance of this in vitro data in cattle has not been demonstrated.

Toxicology: The heart is the target of toxicity in laboratory and domestic anii given Micotile 300 by oral or parenteral routes. The primary cardiac effects increased heart rate (tachycardia) and decreased contractility (neginotropy).

inotropy). Upon injection subcutaneously, the acute median lethal dose of tilmicosin in r is 97 mg per kg, and in rats is 185 mg/kg of body weight. Given orally, median lethal dose is 800 mg/kg and 2250 mg/kg in fasted and nonfasted r respectively. No compound-related lesions were bound at necropsy.

Mannheimia

EANCO * AH 0230-50H



De Not Use in Automatically Powered Syrin No Administrar con Jeringas Accionadas Automáticamente, NADA 140-929, Approved by FDA UPC 7 2780420212 5

212 5	·		BÃO	associated theimia	with	indianapolis, IN 4
	50	mL	(Paste	urella) haemo	dytica.	YL.00710EAM

CAUTION: Federal (USA) law restricts this drug to use by or on the order of a Human Warnings: Not for human use. Injection of this drug in humans may be tatal. Koop out of reach of licensed veterinarian. children. Do not use in automatically powered An antibiotic injection for syringer. Exercise extreme coution to avoid accidsubcutaneous use in catle and sheep. For the ental self injection. In same of human injection, consult a physician immediately. Emergency treatment of bovine respimedical telephone numbers are 1-500-722-0987 or 1-317-275-2000. Avoid contact with eyes. ratory disease (BRD) or ovine respiratory disease (ORD) associated with Read Note to the Physician, Spanish translation, and other important information fully before use. (Pasteurella) haemolytica.

For the control of respira-tory disease in cattle at Manufactured for: Elanco Animal Health A Division of El Lilly and Con Indianapolis, IN 46285, USA i iterature for the high risk of developing

product shi be attached. is missing

has not been established for sheep with a body weight of less than 15 k in pregnant sheep or sheep used for breeding purposes.

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How Supplied: Micotil is supplied in 50 mL, 100 mL and 250 mL multid amber glass bottles.

Storage: Store at room temperature, 86°F (30°C) or below. Protect fr direct sunlight. Conservar a 86°F (30°C). Protejer de la directa luz solar.

*Elanco®, Micotil®, and the diagonal color bar are trademarks of Eli Lilly and Comp Text revised April (), 2002

Manufactured for:

Elanco Animal Health, A Division of Eli Lilly and Company Indianapolis, IN 46285, USA YL0071DEA

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ELANCO Micotil / Label / Arr	erica	Product code: AH 0230	Artwork Ref. No: Y	L0071DEAMX_P2	Affiliate Proof No.: 2	BLACK
Created at/by: Speke/MIKE	Date Crea	ited: 24.04.2002	Last amended: 31.07	.2002 Speke/ MIKE	Lilly	BLACK
Label size: 50 mL	Artwork si	ze: 90 mm W x 40 mm H	Studio approval:		WARNING THIS MATERIAL IS COPYRIGHT AND	P298
Barcode: 419 (B) (L0L00L00)	File Type: QuarkXPress 4.0		- Date: (_OK matches approved proof / file / ISDN)		Eli Liliy and Company Limited, Spoke Operations UNAUTHORISED POSSESSION, USE OR	BLUE
Primary Pack: GPG080	Reason fo	r Revision: PROOFS FOR			REPRODUCTION WILL ATTRACT CRIMINAL PENALTIES AND LABILITY FOR DAMAGES AND LEGAL COSTS. ALL FORTS SUPPLIED ARE FOR THE	100%
Approved by:	Date:	Q.C. approved by	/:	Date:	GENERATION OF THIS WORK ONLY. IN LINE WITH COPYNIGHT REGULATIONS IT IS PROHIMITED TO MAKE COPIES OF THE FONTS.	PRINT

Process colour reproduction may not match PANTONE-Identified solid colour standards. Refer to current PANTONE Colour Publications for the accurate colour.

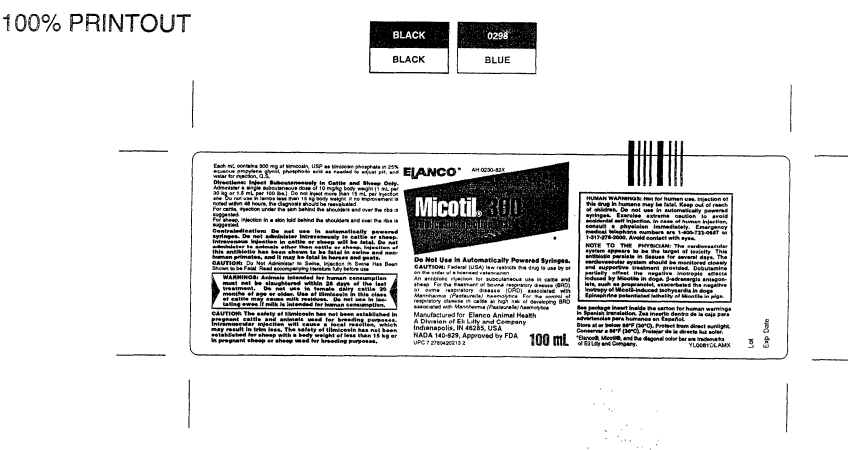
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ELANCO MICOTIL 300 / LABEL / AM		L / AMERICA	Product code: AH0230	Artwork Reference No: Y	L0081DEAMX_P3	Affiliate Proof No.: 3
Created at/by: Speke / MIKE		Date created: 22.04.2002		Last amendment: 26.04.2002 Speke / MIKE		Lilly
Pack size: 100 ml		Artwork size: 157	mm x 54 mm	Studio approval: Date:		
Pharma Code	: 391 (B) (L000L000)	File Type: Adobe	Illustrator 8.01	(/OK matches approved proof / file / ISDN)		AND PROPRIETY TO Eli Lilly and Company Limited, Speke Operations UNAUTHORISED POSSESSION, USE OR
Base Spec.: (GPG021	Reason for Revisio	n: REVISED PROOFS FOR	R FDA (No text approval code).		ARPRODUCTION WILL ATTRACT CRIMINAL PENALTES AND LIABILITY FOR DAMAGES AND LEGAL COSTS ALL FONTS SUPPLIED ARE FOR THE
Approved by:	Approved by:		Q.C. approved by:		Date:	GENERATION OF THIS WORK ONLY IN LINE WITH COPYRIGHT REGULATIONS IT IS PROHIBITED TO MAKE COPIES OF THE FONTS

Process colour reproduction may not match PANTONE-Identified solid colour standards. refer to current PANTONE Colour Publications for the accurate colour.

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EANCO	Micotil 300 / Inse	ort / America	Product code: AH 02			
Created at	/by: Speke/MIKE	Date Created: 22.04.2002				
Label size:	100 ml	Artwork size: A5 148 mm x 210 n				
Barcode: 39	91 (B) (L000L000)	File Type: QuarkXPress 4.0				
Base Spec	: MXG410	Reason for F	Revision: REVISED PI			
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Process colour reproduction may not match PANTONE-Identified solid colour standards, refer to

100% PRINTOUT

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		Pharmacode Bar Height =	12 mm
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AH 0230	NADA 140-929, Approved by FDA	PA9041DEAMP	
	Micotil® 300 Tlimicosin Injection, USP		
CAUTION: Federal (USA) law	restricts this drug to use by or on the order of	of a licensed veterinarian.	
of reach of children. Do not to avoid accidental self inje Emergency medical telepho with eyes. NOTE TO PHYSICIAN: The	uman use. Injection of this drug in humans use in automatically powered syringes. Ex ction. In case of human injection, consult a one numbers are 1-800- 722-0987 or 1-317-2 e cardiovascular system appears to be the	vercise extreme caution physician immediately. 276-2000. Avoid contact a target of toxicity. This	
closely and supportive trea effects induced by Micotile	s for several days. The cardiovascular syste tment provided. Dobutamine partially offse n dogs. Badrenergic antagonists, such as p otil-induced tachycardia in dogs. Epinephrine	t the negative inotropic or the negative inotropic or the negative inotropic of the negative ino	
humanos puede ser fatal. Co accionadas automáticament En casos de inyección a h telefónicos para emergencia los ojos. NOTA AL MEDICO: EL Si TOXICIDAD DE ESTE PRO VARIOS DIAS. EL SISTEMA TRATAMIENTO DE SOPOR BLOQUEA LOS EFECTOS ROS. BETA ADRENERGICO	UMANOS: No usarlo en humanos. La inyec nservario fuera del alcance de los niños. No e. Ejercer precauciones extremas para evitau umanos, consultar con un médico inmedia a médicas son 1-800-722-0987 o 1-317-276-20 STEMA CARDIOVASCULAR PARECE SER DOUCTO. ESTE ANTIBIOTICO PERSISTE E CARDIOVASCULAR DEBERA OBSERVARSI TE DEBERA PROPORCIONARSE. DOBUTA INOTROPICOS NEGATIVOS INDUCIDOS P S ANTAGONISTAS, COMO EL PROPRANOL LA TAQUICARDIA INDUCIDA POR MIC	administrar con jeringas r inyectarse uno mismo. atamente. Los números 000. Evitar contacto con R EL BLANCO DE LA N LOS TEJIDOS POR E CUIDADOSAMENTE Y MINA PARCIALMENTE OR MICOTIL EN PER- OL, EXACERBARON LA	PHARMAG AREA
For Subcutaneous Use in Ca Indications: Micotile 300 is ovine respiratory disease Micotile 300 is indicated for t BRD associated with Mannh Description: Micotile 300 is	A LA LETALIDAD DE MICOTIL EN CERDO ttle and Sheep Only. Do Not Use in Automat indicated for the treatment of bovine respir (ORD) associated with Mannheimia (P he control of respiratory disease in cattle a eimia (Pasteurella) haemolytica. a solution of the antibiotic tilmicosin. Each	tically Powered Syringes. atory disease (BRD) and <i>asteurella</i>) haemolytica. at high risk of developing h mL contains 300 mg of	
adjust pH and water for injec	•		–
Actions: Activity—Tilmicosir positive with activity agains mycoplasma species has als		at is predominantly gram- Activity against several	BARCODE READER
Ninety-five percent of the 3.12 µg/mL or less.	Mannheimia (Pasteurella) haemolytica iso	plates were inhibited by	DEF
Microorganism	MIC* (μg/mL)		0
Mannheimia (Pasteurella) ha Pasteurella multocida	emolytica 3.12 6.25		ARC
Haemophilus somnus	6.25		B
Mycoplasma dispar	0.097		
M. bovirhinis			
M. bovoculi	0.024 0.048		

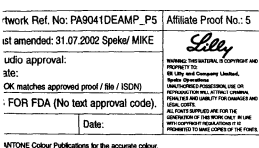
The clinical significance of this *in vitro* data in cattle has not been demonstrated. **Toxicology:** The heart is the target of toxicity in laboratory and domestic animals given Micotile 300 by oral or parenteral routes. The primary cardiac effects are increased heart rate (tachycardia) and decreased contractility (negative inotropy). Upon injection subcutaneously, the acute median lethal dose of tilmicosin in mice is 97 mg/per kg, and in rats is 185 mg/kg of body weight. Given orally, the median lethal dose is 800 mg/kg and

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2250 mg/kg in fasted and nonfasted rats, respectively. No compound-related lesions were found at necropsy.

In monkeys, a single intramuscular dose of 10 mg/kg caused no signs of toxicity. A single dose of 20 mg/kg caused vomiting and 30 mg/kg caused the death of the only monkey tested.

In swine, intramuscular injection of 10 mg/kg caused increased respiration, emesis, and a convulsion; 20 mg/kg resulted in mortality in 3 of 4 pigs; and 30 mg/kg caused the death of all 4 pigs tested.

Results of genetic toxicology studies were all negative. Results of teratology and reproduction studies in rats were negative. The no effect level in dogs after daily oral doses for up to one year is 4 mg/kg of body weight.

In cattle, subcutaneous doses of 10, 30, and 50 mg/kg of body weight, each injected 3 times at 72 hour intervals did not cause any deaths. As expected, edema at the site of injection was noted. The only lesion observed at necropsy was minimal myocardial necrosis in some animals in the 50 mg/kg group. Subcutaneous doses of 150 mg/kg injected at 72-hour intervals resulted in deaths. Edema was marked at the site of injection. Minimal myocardial necrosis was the only lesion observed at necropsy. Deaths of cattle have been observed with a single intravenous dose of 5 mg/kg of body weight.

5 mg/kg of body weight. In sheep, single subcutaneous injections of 10 mg/kg dose did not cause any deaths and no adverse effects of tilmicosin were observed on blood pressure, heart rate, or respiratory rate. **Pharmacology:** A single subcutaneous injection of Micotile at 10 mg/kg of body weight dose in cattle resulted in peak tilmicosin levels within one hour and detectable levels (0.07 µg/mL) in serum beyond 3 days. However, lung concentrations of tilmicosin remained above the tilmicosin MiC 95% of 3.12 µg/mL for *Mannheimia (Pasteurella) haemolytica* for at least 3 days following the single injection. Serum tilmicosin levels are a poor indicator of total body tilmicosin. The lung/serum tilmicosin ratio in favor of lung tissue appeared to equilibrate by 3 days post injection at approximately 60. In a study with radioactive tilmicosin, 24% and 68% of the dose was recovered from urine and feces, respectively, over 21 days. **Directions—Inject** Subcutaneously in **Cattle and Sheep Only**. Administer a single

Directions—Inject Subcutaneously in Cattle and Sheep Only. Administer a single subcutaneous dose of 10 mg/kg of body weight (1 mL/30 kg or 1.5 mL/100 lbs). Do not inject more than 15 mL per injection site. Do not use in lambs less than 15 kg body weight.

If no improvement is noted within 48 hours, the diagnosis should be reevaluated.

For cattle, injection under the skin behind the shoulders and over the ribs is suggested.

For sheep, injection in a skin fold behind the shoulders and over the ribs is suggested.

Note-Swelling at the subcutaneous site of injection may be observed but is transient and usually mild.

CONTRAINDICATION: Do not use in automatically powered syringes. Do not administer intravenously to cattle or sheep. Intravenous injection in cattle or sheep will be fatal. Do not administer to animals other than cattle or sheep. Injection of this antibiotic has been shown to be fatal in swine and non-human primates, and it may be fatal in horses and goats.

CAUTION: Do Not Administer to Swine. Injection in Swine Has Been Shown to be Fatal.

WARNINGS: Animals intended for human consumption must not be slaughtered within 28 days of the last treatment. Do not use in female dairy cattle 20 months of age or older. Use of tilmicosin in this class of cattle may cause milk residues. Do not use in lactating ewes if the milk is intended for human consumption.

CAUTION: The safety of tilmicosin has not been established in pregnant cattle and in animals used for breeding purposes. Intramuscular injection will cause a local reaction which may result in trim loss. The safety of tilmicosin has not been established for sheep with a body weight of less than 15 kg or in pregnant sheep or sheep used for breeding purposes.

How Supplied: Micotil is supplied in 50 mL, 100 mL and 250 mL multidose amber glass bottles. Storage: Store at room temperature, 86°F (30°C) or below. Protect from direct sunlight.

Conservar a 86°F (30°C). Protejer de la directa luz solar.

*Elanco® and Micotil® are trademarks of Eli Lilly and Company.

Text Revised April (), 2002

Manufactured for: Elanco Animal Health A Division of Eli Lilly and Company Indianapolis, IN 46285, USA

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PULL TO OPEN

FOLL TO OFERV Each mL contains 300 mg of tilmicosin, USP as tilmicosin phosphate in 25% equeous proprierus giyool, phosphoric acid as needed to adjust pH, and water for injection, Q.S.

as measure to aquat pH, and water for rejection, Q.S. Diversions: hipsets Suboutbasewark in Cattle and Shawe Only. Administer a single suboutbaseous does of 10 mg/kg body weight. (I mU20 kg or 15 mL per 100 ba). Do not hipset mase than 15 mL per hipsetion sits. Do not use in tembs teas than 15 kg body weight. In or tropowenen it a noted within 44 hours, the diagnosts should

callie, injection under the skin behind the shoulders and over For case, injection under the stoh behind the shoulders and over the ribe is suggested. For sheep, injection in a skin fold behind the shoulders and over the ribe is suggested.

Araindication: Do not use in auto inges. De not administer interim causer, us not use in automatically powered o not administre infravously to cattle or sheep, b injection in cattle or chasp will be fatal. Do not to animale other than cattle or chasp, injection of the hase been shown to be fatal in surine and non-tation, and it may be fatal in horses and guals. fals and CAUTION: Do Not Administer to Swine, Injection in Swi - Man Rea on to be Fatal. Read accompanying iterature fully before use

WARNINGS: Animals intended for human consump-tion must not be staughtered within 22 days of the last treatment. Do not use in themale dairy catiliti 20 months of age or older. Use of Windowin in this class of eatthe may cause milk residues. Do not use in lactating swee if the milk indended for insuma consumption.

CAUTION: The safety of Windoowin has not been satublished in program cattle and in animals used for branding purposes. Intramuscular injection will cause a load reaction which may result in tim loss. The safety of Windowin has not been satublished for sheep with a body weight of less than 15 to or in pregnant sheep or since used to fee than

Store at or below &6°F (30°C) Protect from direct sunlight. Conservar a 86°F (30°C). Protect de la directa luz aclar



Do Not Use in Automatically Powered Syringes. No Administrar con Jeringas

Accionadas Automáticamente. CAUTION: Federal (USA) law restricts this drug to use by or on the order of a licensed veterinarian. use by or on the order of a licensed veterinarian. An antibiotic injection for subcutaneous use in cattle and sheep. For the treatment of bowine respiratory disease (BAD) or ovine respiratory disease (ORD) associated with Mannheimia (Pasteurella) haemotytica. For the control of respiratory disease in cattle at high risk of developing BRD associated with Mannheimia (Pasteurella) haemotytica.

NADA 140-929, Approved by FDA UPC 7 2780420214 9 250 ml

HUMAN WARNINGS: Not for human use, injection of this drug in humans may be fatal. Keep out of reach of children. Do not use in sufomalically powered syringes. Exercise extreme caution to avoid accidential self injection. In case of human injection, consult a physicia Immediately. Emergency medical telephone numbers an 1-800-722-0987 or 1-317-276-2000, Avoid contact with eyes. 1-600-722-0687 or 1-317-276-2000, Avoid contact with sysa. NOTE TO THE PHYSHCIAN: The cardiovascular system appears to be the target of tockicly. This ambiotic persists in dissues for several days. The cardiovascular system should be monitored closely and supportive trashment provided. Dobutamine particity offiset the negative instropic effects induced by Micolable in dogs. Featurengic antagonists, such as programoid, excercisited the negative inducey of Micolable in dogs. Epinephrine potentiaset industry of Micolable in dogs. ADVERTENCIAS PARA HUMANOS: No usario en

AUTENTICIECICUS PARA HUBANDOS: No usano en humanos. La hiprocición de está droga en humanos puede ser fatal. Conservario fuera del alcance de los miños. No administrar: con jeringas accionadas automáticamento. Epircer precesciones extremas para evitar inyectarse uno mismo. En caso de inyección a humanos, consultar con un midicio immediatamente. Los nómorous telefónicos para amergiencias midicas aon 1.400-722-0081 o 1.317/271 2000. Evitar contacto con los ojos.

Manufactured for: Elance Animal Health A Division of Eli Lilly and Company Indianapolis, IN 45285, USA "Encode Necesita, and the diagonal core bar are trade El Lilly and Company. YL0061DEAMX

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NADA 140-929, Approved by FDA

Micotil® 300 Tilmicosin Injection, USP

CAUTION: Federal (USA) law restricts this drug to use by or on the order of a licensed veterinarian.

Human Warnings: Not for human use. Injection of this drug in humans may be fatal. Keep out of reach of children. Do not use in automatically powered syringes. Exercise extreme caution to avoid accidental self injection. In case of human injection, consult a physician immediately. Emergency medical telephone numbers are 1-800-722-0987 or 1-317-276-2000. Avoid contact with eyes.

NOTE TO PHYSICIAN: The cardiovascular system appears to be the target of toxicity. This antibiotic persists in tissues for several days. The cardiovascular system should be monitored closely and supportive treatment provided. Dobutamine partially offset the negative inotropic effects induced by Micotil@ in dogs. β-adrenergic antagonists, such as propranolol, exacerbated the negative inotropy of Micotil-induced tachycardia in dogs. Epinephrine potentiated lethality of Micotil@ in pigs.

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YL0061DEAMX_P4 31/7/02 3:09 pm Page 5

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ADVERTENCIAS PARA HUMANOS: No usarlo en humanos. La inyección de esta droga en humanos puede ser fatal. Conservarlo fuera del alcance de los niños. No administrar con jeringas accionadas automáticamente. Ejercer precauciones extremas para evitar inyectarse uno mismo. En casos de inyección a humanos, consultar con un médico inmediatamente. Los números telefónicos para emergencias médicas son 1-800-722-0987 o 1-317-276-2000. Evitar contacto con los olos.

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NOTA AL MEDICO: EL SISTEMA CARDIOVASCULAR PARECE SER EL BLANCO DE LA TOXICI-DAD DE ESTE PRODUCTO. ESTE ANTIBIOTICO PERSISTE EN LOS TEJIDOS POR VARIOS DIAS. EL SISTEMA CARDIOVASCULAR DEBERA OBSERVARSE CUIDADOSAMENTE Y TRATAMIENTO DE SOPORTE DEBERA PROPORCIONARSE. DOBUTAMINA PARCIALMENTE BLOQUEA LOS EFECTOS INOTROPICOS NEGATIVOS INDUCIDOS POR MICOTIL EN PERROS. BETA ADRENERGICOS ANTAGONISTAS, COMO EL PROPRANOLOL, EXACERBARON LA INOTROPIA NEGATIVA DE LA TAQUICARDIA INDUCIDA POR MICOTIL EN PERROS. LA EPINEFRINA POTENCIALIZA LA LETAL-IDAD DE MICOTIL EN CERDOS.

For Subcutaneous Use in Cattle and Sheep Only. Do Not Use in Automatically Powered Syringes.

Indications: Micotil® 300 is indicated for the treatment of bovine respiratory disease (BRD) and ovine respiratory disease (ORD) associated with Mannheimia (Pasteurella) haemolytica. Micotil® 300 is indicated for the control of respiratory disease in cattle at high risk of developing BRD associated with Mannheimia (Pasteurella) haemolytica.

Storage: Store at room temperature, 86°F (30°C) or below. Protect from direct sunlight. Conservar a 86°F (30°C). Protejer de la directa luz solar.

*Elanco® and Micotil® are trademarks of Eli Lilly and Company

Text Revised April (), 2002

Manufactured for: Elanco Animal Health A Division of Eli Lilly and Company Indianapolis, IN 46285, USA

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escription: Micotile 300 is a solution of the antibiotic tilmicosin. Each mL contains 300 mg of tilmicosin, USP as micosin phosphate in 25% propylene glycol, phosphoric acid as needed to adjust pH and water for injection, q.s. imicosin, USP is produced semi-synthetically and is in the macrolide class of antibiotics.

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ctions: Activity—Tilmicosin has an *in-vitro** antibacterial spectrum that is predominantly gram-positive with tivity against certain gram-negative microorganisms. Activity against several mycoplasma species has also been stected.

inety-five percent of the Mannheimia (Pasteurella) haemolytica isolates were inhibited by 3.12 µg/mL or less.

Microorganism	MIC* (µg/mL)
Mannheimia (Pasteurella) haemolytica	3.12
Pasteurella multocida	6.25
Haemophilus somnus	6.25
Mycoplasma dispar	0.097
M. bovirhinis	0.024
M. bovoculi 🔋	0.048

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*The clinical significance of this in vitro data in cattle has not been demonstrated.

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YL0061DEAMX_P4 31/7/02 3:09 pm Page 9

Toxicology: The heart is the target of toxicity in laboratory and domestic animals given Micotil® 300 by oral or parenteral routes. The primary cardiac effects are increased heart rate (tachycardia) and decreased contractility (negative inotropy).

Upon injection subcutaneously, the acute median lethal dose of tilmicosin in mice is 97 mg per kg, and in rats is 185 mg/kg of body weight. Given orally, the median lethal dose is 800 mg/kg and 2250 mg/kg in fasted and nonfasted rats, respectively. No compound-related lesions were found at necropsy.

in monkeys, a single intramuscular dose of 10 mg/kg caused no signs of toxicity. A single dose of 20 mg/kg caused vomiting and 30 mg/kg caused the death of the only monkey tested.

In swine, intramuscular injection of 10 mg/kg caused increased respiration, emesis, and a convulsion, 20 mg/kg resulted in mortality in 3 of 4 pigs, and 30 mg/kg caused the death of all 4 pigs tested.

Results of genetic toxicology studies were all negative. Results of teratology and reproduction studies in rats were negative. The no effect level in dogs after daily oral doses for up to one year is 4 mg/kg of body weight.

In cattle, subcutaneous doses of 10, 30, and 50 mg/kg of body weight, each injected 3 times at 72 hour intervals did not cause any deaths. As expected, edema at the site of injection was noted. The only lesion observed at necropsy was minimal myocardial necrosis in some animals in the 50 mg/kg group. Subcutaneous doses of 150 mg/kg injected at 72-hour intervals resulted in deaths. Edema was marked at the site of injection. Minimal myocardial necrosis was the only lesion observed at necropsy. Deaths of cattle have been observed with a single P4

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intravenous dose of 5 mg/kg of body weight.

In sheep, single subcutaneous injections of 10 mg/kg dose did not cause any deaths and no adverse effects of tilmicosin were observed on blood pressure, heart rate, or respiratory rate.

Pharmacology: A single subcutaneous injection of Micotike at 10 mg/kg of body weight dose in cattle resulted in peak tilmicosin levels within one hour and detectable levels (0.07 µg/mL) in serum beyond 3 days. However, lung concentrations of tilmicosin remained above the tilmicosin MIC 95% of 3.12 µg/mL for *Mannheimia (Pasteurella) haemolytica* for at least 3 days following the single injection. Serum tilmicosin levels are a poor indicator of total body tilmicosin. The lung/serum tilmicosin ratio in favor of lung tissue appeared to equilibrate by 3 days post injection at approximately 60. In a study with radioactive tilmicosin, 24% and 68% of the dose was recovered from urine and feces respectively over 21 days.

Directions—Inject Subcutaneously in Cattle and Sheep Only. Administer a single subcutaneous dose of 10 mg/kg of body weight (1 mL/30 kg or 1.5 mL per 100 lbs).

Bo not inject more than 15 mL per injection site.

Do not use in lambs less than 15 kg body weight.

If no improvement is noted within 48 hours, the diagnosis should be reevaluated.

For cattle, injection under the skin behind the shoulders and over the ribs is suggested.

For sheep, injection in a skin fold behind the shoulders and over the ribs is suggested.

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YL0061DEAMX_P4 31/7/02 3:09 pm Page 7

CONTRAINDICATION: Do not use in automatically powered syringes. Do not administer intravenously to cattle or sheep. Intravenous injection in cattle or sheep will be fatal. Do not administer to animals other than cattle or sheep. Injection of this antibiotic has been shown to be fatal in swine and non-human primates, and it may be fatal in horses and goats.

CAUTION: Do Not Administer to Swine, Injection in Swine Has Been Shown to be Fatal.

WARNINGS: Animals intended for human consumption must not be slaughtered within 28 days of the last treatment.

Do not use in female dairy cattle 20 months of age or older. Use of tilmicosin in this class of cattle may cause milk residues. Do not use in lactating ewes if the milk is intended for human consumption.

CAUTION: The safety of tilmicosin has not been established in pregnant cattle and in animals used for breeding purposes. Intramuscular injection will cause a local reaction which may result in trim loss. The safety of tilmicosin has not been established for sheep with a body weight of less than 15 kg or in pregnant sheep or sheep used for breeding purposes.

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How Supplied: Micotil is supplied in 50 mL, 100 mL and 250 mL multidose amber glass bottles.