Here to help *your* cattle breathe easier.



Veterinarians and producers know just how devastating bovine respiratory disease (BRD) can be. It compromises the productivity of an animal and overall profitability of an operation.

RESPIRmycin® (tulathromycin injection) is here to help.

This ready-to-use, trusted injectable solution contains 100 mg of tulathromycin/mL. FDA-approved, RESPIRmycin is equivalent to Draxxin[®] in:

- Active ingredient and inactive ingredients (carriers and excipients)
- Strength, dosage form and route of administration

Manufactured in Parnell's FDA-approved facility, you can count on RESPIRmycin to meet all quality and safety standards. In other words, it's the product you can count on to help your cattle breathe easier.



Available in 100 mL, 250 mL and 500 mL vials

Beef and Non-Lactating Dairy Cattle

BRD – RESPIRmycin Injectable Solution is indicated for the treatment of bovine respiratory disease (BRD) associated with *Mannheimia haemolytica, Pasteurella multocida, Histophilus* somni and Mycoplasma bovis; and for the control of respiratory disease in cattle at high risk of developing BRD associated with *M. haemolytica, P. multocida, H. somni* and *M. bovis*. **IBK** – RESPIRmycin Injectable Solution is indicated for the treatment of infectious bovine keratoconjunctivitis (IBK) associated with *Moraxella bovis*.

Foot Rot – RESPIRmycin Injectable Solution is indicated for the treatment of bovine foot rot (interdigital necrobacillosis) associated with *Fusobacterium necrophorum* and *Porphyromonas levii*.

Suckling Calves, Dairy Calves, and Veal Calves

BRD – RESPIRmycin Injectable Solution is indicated for the treatment of BRD associated with *M. haemolytica*, *P. multocida*, *H. somni* and *M. bovis*.

Swine

RESPIRmycin Injectable Solution is indicated for the treatment of swine respiratory disease (SRD) associated with *Actinobacillus pleuropneumoniae*, *Pasteurella multocida*, *Bordetella bronchiseptica*, *Haemophilus parasuis* and *Mycoplasma hyopneumoniae*; and for the control of SRD associated with *A. pleuropneumoniae*, *P. multocida* and *M. hyopneumoniae* in groups of pigs where SRD has been diagnosed.

Parnell's commitment to your success.

RESPIRmycin serves as just another example of our ongoing commitment to helping veterinarians and producers every way we can. Each product and technology in our portfolio offers customers the same quality and convenience they've come to expect from estroPLAN[®] and GONAbreed.[®] Count on it.

To learn more about RESPIRmycin or other Parnell offerings, contact your Parnell representative.



The label contains complete use information, including cautions and warnings. Always read, understand and follow the label and use directions.

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RESPIRmycin[®] (tulathromycin injection)

Injectable Solution

Antibiotic 100 mg of tulathromycin/mL

For use in beef cattle (including suckling calves), non-lactating dairy cattle (including dairy calves), veal calves, and swine. Not for use in female dairy cattle 20 months of age or older.

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

ESSPRmycin Injectable Solution is a ready-to-use stelle parenterial preparation containing tubathomycin, a semi-synthetic macrolide antibiotic of the subclass triamilite Lach mL of RESPRmycin contains 100 mg of tubathomycin, 300 mg progybere glyck), 122 mg citric add and 5 mg monorhioghcred. Sodium hydrocolde or hydrochloric caid mg/s be added to adjust pH. RESPRmycin consists of an equilitated mixture of two isomeric forms of tubathomycin in a 3:1 ratio. Structures of the isomers are shown below.

Figure 1. HJC N OH OH NICHala 0-CH3 CH3 OCH3



The chemical names of the isomers are (25,54R,58R,10R,11R,125,135,14R)-13-1[2,6-discoy-3-C,methyl-3-O,methyl-4-C ([roopJamino) methyl-3-10- territyl-3-10- territyl-3-10-

INDICATIONS

INUCAINS Beef and Non-Lactating Dairy Cattle BBD — BS/PRimycin Injectable Solution is indicated for the treatment of bovine respiratory disease (BD) associated with Mommelinia chemolytica, Pristeurella multocida, Histophilas sonni, and Alycopianna bovis: and for the control of respiratory disease in cattle at high rick of developing BDD associated with Mommelinia Internolytica, Pristeruien Multocida, Histophilas sonni, and Alycopianna associated with Mommelinia Internolytica, Pristeruien Multocida, Histophilas sonni, and Alycopianna

IBK – RESPIRmycin Injectable Solution is indicated for the treatment of infectious bovine keratoconjunctivitis (IBK) associated with Moraxella bovis.

Foot Rot – RESPIRmycin Injectable Solution is indicated for the treatment of bovine foot rot (interdigital necrobacillosis) associated with Fusobacterium necrophorum and Porohyromonos Jevii.

Suckling Calves, Dairy Calves, and Veal Calves BRD – RESPIRmycin Injectable Solution is indicated for the treatment of BRD associated with M. haemolytica, P. amiltocida, H. sonmi, and M. bovis.

Swine RESPRimprin Injectable Solution is indicated for the treatment of swine respiratory disease (SRD) associated with Actinobacillus pleuropneumoniae, Pasteurella multicida, Boutefella bronchiceptica, Heenophilus parassia, and Mycopissma hyporeumoniae, and for the control of SRD associated with Actinobacillus pleuropneumoniae, Pasteurella multicida, and Mycopissma hyponeumoniae in groups of pigs where SRD has been diagnosed.

DOSAGE AND ADMINISTRATION

linject subortaneously as a single dose in the neck at a dosage of 2.5 mg/kg (1.1 mL/100 lb) body weight (80%). Do not inject more than 10 mL per injection site. **Table 1.** RESPIRmycin Cattle Dosing Guide

Animal Weight (Pounds) Dose Volume (mL)

Annia Weight (rounus)		Dose volume (mL)		
100		1.1		
Г	200	2.3		
	300	3.4		
Γ	400	4.5		
Г	500	5.7		
Г	600	6.8		
Г	700	8.0		
	800	9.1		
	900	10.2		
Γ	1000	11.4		

Inject intramuscularly as a single dose in the neck at a dosage of 2.5 mg/kg (0.25 mL/22 lb) BW. Do not inject more than 2.5 mL per injection site. Table 2. RESPIRmycin Swine Dosing Guide

Dose Volume (mL)		
0.2		
0.3		
0.6		
0.8		
1.0		
1.3		
1.5		
1.7		
1.9		
2.2		
2.4		
2.6		
2.8		
3.1		
3.3		

CONTRAINDICATIONS ution is contraindicated in animals previously found to be The use of RESPIRmycin Injectable Solut hypersensitive to the drug. WARNINGS

FOR USE IN ANIMALS ONLY.

KEEP OUT OF REACH OF CHILDREN. NOT FOR USE IN CHICKENS OR TURKEYS.

RESIDUE WARNINGS Cattle intended for human consumption must not be slaughtered within 18 days from the last treatment. This drug is not approved for use in female dary 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. Cattle

nded for human consumption must not be slaughtered within 5 days from the last

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

Swine The effects of RESPIRmycin on porcine reproductive performance, pregnancy, and lactation have not been determined. Intramuscular injection can cause a transient local tissue reaction that may result in trim loss of edible tissue at slaughter.

ADVERSE REACTIONS

Cattle In one BRD field study, two calves treated with tulathromycin injection at 2.5 mg/kg BW exhibited transient hypersalivation. One of these calves also exhibited transient dyspnea, which may have been related to pneumonia.

iwine in one field study, one out of 40 pigs treated with tulathromycin injection at 2.5 mg/kg BW exhibited mild salivation that resolved in less than four hours.

International table Concernation and Bioth Product POT LAPPORAL EXPERIENCE The following adverse: events are abased on post approval adverse drug experience resporting, lot all adverse events are request for the FAD CMA. This is not advarys possible to reliably estimate the adverse event frequency or estabilish a causal relationship to product exposure using these data. The following adverse events and expendition devension gived or the production frequency in excitation hypothese interactions and amph/adsociation spectra to the CMA. We can experiment and according to the experiment experiment in excitation spectra to the CMA we excitation for tabilitomycin injection injecticatio sections. The other excitation and according to the excitation and advection injection between the other excitations and according to the excitation and advection injection between the other excitations and according to the excitation and advection injection between the other excitation and according to the excitation and advection injection between the other excitation and advection and advection advection between the other excitations and advection advection between the other excitation and advection advection between the other excitations and advection advectio

Comparison of the second second

unotermined. Multidupti the relationship between tuiathromycin and the characteristics of its antimicrobial effects has not been characterised; as a dars, macuillais tend to be primarily bacteristical; but may be bacterisdic against some pathogeness. They also tend to which is concentration dependent sillings the start of bacterial eradication does not change nore some drug concentrations read; 2 is 3 times the minimum inhibitory concentration (MC) of the starged pathogen. Under these conflicts, the time that are concentrations remain above the MC becomes the major determinant of antimicrobial activity. Nurolides also exhibit a post-antibicit effect. PRQ, the duration of which tends to be bed drug and pathogen dependent. In general, by increasing the macrolide concentration and the pospure time, drug concentration tends to be the most powerful determinant of affectional of PRE.

Tulathromycin is eliminated from the body primarily unchanged via biliary excretion

¹Carbon, C. 1998. Pharmacodynamics of Macrolides, Azalides, and Streptogramins: Effect on Extracellular Pathogens. Clin. Infect. Dis., **27**:28-32.

³Wightingale, C.J. 1997. Pharmacokinetics and Pharmacodynamics of Newer Macrolides. Pediatr. Infect. Dis. J., 16:438-443.

Cattle

CH-

Catle Gates Following subcutaneous administration into the seck of feeder calves at a dorage of 2.5 mg/kg BW, highborgnic is capidly and nearly completely abcoded beak plasma concentrations generally ours within 15 minutes after doraing and product relative bioavailability creeds 90%, Follar systemic devance is approximately 170 mL/r/kg. Talathomycin distributes retensively into body tissues, as develoced by volume of distribution using a poporticity FLT (Ling the PML and the DML and the DML approximately 17.20 mL/r/kg. Talathomycin distributions in the DML and the DML approximately 17.20 mL approximately 17.10 mL plasma fragma concentrations) poporticity 2.75 days for total lung concentrations (based on quantifiable terminal plasma drug concentrations) resus 825 days for total lung concentrations (based on data from healty animals)]. Linear highmanochinetics are beneved with soutcances does ranging from 127 mg/kg BWL 5.01 mg/kg BWL linear and a server of the source of the server of the cast and ender the server of the cast and the server of the server of the source of the source of the source of the server of the source o

³Clearance and volume estimates are based on intersubject comparisons of 2.5 mg/kg BW administered by either subcutaneous or intravenous injection

Suite Galowing intramuscular administration to feeder pigs at a dosage of 2.5 mg/kg BW, tutahhromycin is completely and rapidly absorbed ($T_{m_m} = 0.25$ hour). Subsequently, the drug rapidly distributes into body issues, achieving a valume of distribution exceeding 15 kJ kg. There drug is rapidly clear from the systemic circulation ($T_{m_{m_m}} = 10$ *Tm*, *thr/kg*). However, this a long terminal dimination half-life (60) concentrations are assubantably higher than a concentrations or associated in the dimination half-life (60) significance of those findings is undetermined. There are no gender differences in swine tulathromycin pharmacchinetics.

MICROBIOLOGY

Lattue Tulathromycin has demonstrated in vitro activity against Mannheimia haemolytica, Posteurella multocida, Histophilus somni, and Mycopiasma bovis, four pathogens associated with BRD; against Morazella bovis associated with IBR; and against Fusobocterium necrophorum and Porphyromonas levil associated with bovine foot not.

The MICs of tulathromycin against indicated BRD and IBK pathogens were determined using methods recommended by the Clinical and Laboratory Standards Institute (CLSI, M31-A2). The MICs against foot rot pathogens were also determined using methods recommended by the CLSI (M11-A6). All MIC values were determined using the 9:1 isomer ratio of this compound.

BRD - The MICs of tulathromycin were determined for BRD isolates obtained from calves enrolled in therapputic, and a risk field studies in the U.S. In 1999. In the therapputic studies, isolates were obtained from one-troatment costophangeal studies from all study calves, and from alm symbols cost implication saline-treated calves that died. In the at risk studies, isolates were obtained from masophanyngeal studies of saline-treated non-responders, and from lang studies in the studies of saline-treated calves that died. The results are shown in lable 3.

IBK - The MICs of tulathromycin were determined for *Monxella bovis* isolates obtained from calves enrolled in BK field studies in the U.S. in 2004, Isolates were obtained from pretreatment conjunctival sweaks of calves with cinical signs of the ronled in the tulathromycin injection and saline-treated groups. The results are shown in Table 3.

Foot Ret - The MICs of tulathromycin were determined for *Fusobacterium necopharum* and *Pophyromous levi* obtained from cattle enabled in foot rat field studies in the LIS, and Canada in 2007. Isolates were obtained from pre-treatment interdigital biopsies and swabs of cattle with chincal signs of foot ret enrolled in the tulathromychin juscica and salme-treated groups. The results arborn in Table 3:

Table 3. Tulathromycin minimum inhibitory concentration (MIC) values⁴ for indicated pathogens isolated from field studies evaluating BRD and IBK in the U.S. and from foot rot field studies in the U.S. and Canada.

Indicated pathogen	Date isolated	No. of isolates	MIC ₁₀ ** (µg/mL)	MIC ,1** (µg/mL)	MIC range (µg/mL)
Mannheimia haemolytica	1999	642	2	2	0.5 to 64
Pasteurella multocida	1999	221	0.5	1	0.25 to 64
Histophilus sonni	1999	36	4	4	1 to 4
Mycoplasma bavis	1999	43	0.125	1	< 0.063 to > 64
Moraxella bavis	2004	55	0.5	0.5	0.25 to 1
Fusobacterium necrophorum	2007	116	2	64	< 0.25 to > 128
Porphyromonos levii	2007	103	8	128	≤ 0.25 to > 128

* 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.

Swine In vitro activity of tulathromycin has been demonstrated against Actinobacillus pleuropneumo Pasteurella multocida, Bordetella bronchiseptica, Haemophilus parasuis, and Mycoplasma hyopneumo

The MIG of tubaltowing against induced SR0 pathoges were determined using methods, recommended by the Clinical and Laboratory Standards from Clinical SR0 pathoges were determined using methods, recommended by the Clinical and Laboratory Standards institute (CS), M31A and M31A3, MIG for Mempholizy pomoral were determined using theritary's Fatildowing and were included up to 84 hours at 35 to 37°C in a Cu-entrolect atmosphere. All MIC values were determined using the 91 isomer atol of this compounds basies obtained in Doo and 2002 were from Imug samples from salme-treated pigs and non-treated sentine pigs enrolled in freatment of SR0 field studies in the U.S. and Canada. Doo and 2002 were from Sunsitive atom and subthermyring in pigetion-treated pigs enrolled in the Control of SR0 field study in the US. and Canada. The results are shown in Table 4.

Table 4. Tulathromycin minimum inhibitory concentration (MIC) values[#] for indicated pathogens isolated from field studies evaluating SRD in the U.S. and Canada.

Indicated pathogen	Date isolated	No. of isolates	MIC ₃₀ ** (µg/mL)	MIC ,** (µg/mL)	MIC range (µg/mL)
Actinobacillus pleuropneumoniae	2000-2002 2007-2008	135 88	16 16	32 16	16 to 32 4 to 32
Haemophilus parasuis	2000-2002	31	1	2	0.25 to >64
Pasteurella multocida	2000-2002 2007-2008	55 40	1	2 2	0.5 to >64 ≤ 0.03 to 2
Bordetella bronchiseptica	2000-2002	42	4	8	2 to 8

* 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.

EFFECTIVENESS

Cette EFECTIVENESS BBD - in a multi-location field study, 314 calves with naturally occurring BRD were treated with tutafhomycin injection. Responses to treatment were compared to saline-treated controls. A cue was defined as a call with normal attuded activity, normal respiration, and a rectal temperature of a 1047 for Dup 11-B tocc are tax significantly higher (5 col)s in tutafhomycin injection-treated calves (2786) compared to saline-treated calves (2476). There were two BBD-related deaths in the tulafhomycin injection-treated calves compared to mis BBD-related deaths in the subintervent data calves compared to mis BBD-related deaths in the subintervent calves.

¹ High-two tulathromycin injection-treated calves and 27 saine-treated calves from the multi-location field BRD treatment study had Mopalasme havis identified in cultures from pre-treatment anapharpaged analos. Of the 52 tulathromycin injection-treated calves, 37 (12): calves were categorized as cares and 15 (28,5%) calves were categorized as treatment failures. Of the 27 saine-treated calves, 14 (145%) calves were treatogorized as treatment failures. Of the 27 saine-treated calves, 14 (145%) calves were treatogorized as cares and 25 (52%) calves were treatment failure table calves, 14 (145%) calves were treatogorized as cares and 25 (52%) calves were treatment failures that calves and the calves of the same treatment failures. Of the 27 same treatment failures are treated as the same treatment failures of the 27 same treatment failures. The same treatment failures of the 27 same treatment failures are treatment failures of the 27 same treatment failures. The 27 same treatment failures of the 27 same treat

A Bayesian meta-analysis was conducted to compare the BBD treatment success rate in young calves (calves weighing 250 lbs or less and fed primarily a mill-based diet treated with tulathromyorin injection to be success rate in older calves (calve weighing more than 250 lbs and fed primarily a roundbage and gain-based diet) treated with tulathromyoin injection. The analysis included data from four BBD treatment effectiveness suclass conducted for the approval of tulathromyoin injection in the 23, and nine tulathromyoin injections conducted effective to the approval of tulathromyoin injection in the 23, and nine in young calves was at least and you be BBD treatment success rate in older calves. K as result, tulathromyoin injections in considered effective for the treatment of BBD sasociated with M. Anemolytics P. multicolds, H. sonmi, and M. kovis in sucking calves, and veal calves.

In another multi-location field study with 29 acless at high risk of developing BRD, administration of tulathornycin injection resulted in a significantly reduced indexec of BBD (11%) compared to significantly calculated and a significantly reduced indexec of BBD (11%) compared to significantly calculated and a rectal impendance of a significantly reduced in attitude/activity, normal respiration, and a rectal impendance of <104°F no log 14. There were no BBD-reduced deaths in the tulathornycin indiction-treated calculates an non-responders in this study have the same-treated calculate. First same-transfer deaths classified as non-responders in this study have development by the BBD-relative deaths in the same-treated calculates. This study have no log to same study of same transfer in colling of provide valuation transfer and study of same transfer in the same study in the same transfer to the BBD-relative death in the same study in the same transfer and the same transfer and same study in the same study in the same transfer and the same study and same study in the same study and same study in the same study in th

The induced information and studies were considered to confine the effectiveness of tubinomycin injection against *Mycoplosmo boxis*. A total of 166 cales were inocalated instantanovally with field strates of *Mycoplosmo boxis*. When cales boxin as prese and had showing experiations cores, they were treated were indexed as a strategies of the transformation of the strategies of the strategies of the cales were takened for significant to the strategies were strategies and strategies of the cales were takened for significant to the strategies were strategies strategies and the strategies of the boxis means the strategies of BBD for 14 days post-treatment; this provides the strategies of the strategies of BBD for 14 days post-testiment to the strategies of the strategies of BBD for 14 days post-testiment to the strategies of BBD for 14 days

Beff – Two field studies were conducted evaluating tulathromycin injection for the treatment of IBK associated with *Monaclile boxis* a 200 naturally-infected calves. The primary dirical endpoint of these societies was created, defined as a alf what no clinical signs of IBK and no cornal ulace, assessed no Days 5, 91, 17, and 21. Time to improvement, defined as the first day on which a call had no clinical signs of IBK is hold by esg. provided that those scores were maintained at the next day of observation, was assessed as a secondary variable. Afail time points, in both studies, the cure rate was significantly higher (P < 0.05) for tulathromycin injection-neated calves. Additionally, time to improvement was significantly lises (P < 0.0001) in both studies for tulathromycin injection-hereated calves. Additionally, time to improve ment was significantly lises (P < 0.0001) in both studies for tulathromycin injection-hereated calves.

To Rot Rot. The effectiveness of tulathromycin injection for the treatment of bovine foot rot was evaluated in 7070 cathe in two field studies. Cathe diagnosed with bovine foot rot were enrolled and treated with a longe subscitancess does of tulathromycin (injection 12.5 mg/sgl byt) are a quivalent whome of sailnes. Cattle were chincally evaluated 7 days after treatment for treatment success, which was based on effendes decreases in leidon, welling, and lameness coses. In both studies, the treatment success statistically significantly higher in tulathromycin injection-treated calves compared with sailne-treated calves (of his-06, we C-000) and 33.¹¹ we Soly. (* 0.000) and 31.¹¹ welling the treatment success, which shall be treated calves (of his-06, welling, and lameness in Soly. * 0.000) and 31.¹¹ welling the treatment of the treatment of the treatment of the treatment success. In both studies, expension are success, which shall be treatment and the treatment of the treatment success. In both studies, expension are success, both studies, expension are success. In both studies, expension are success, both studies, expension are are success, both studies, expension are success, expension are success, expensi

Latters (MeY Hz, Shi, Pr. Cutobia and SJ:SH YL, Shi, Pr. (2006)). Swine In a multi-focation field study to evaluate the treatment of naturally occurring SR0, 266 pips were treated with haldhmorphi microlicon. Response to extrament view compared to saline-treated controls. Success word defined as a pip with normal attrudy, normal regulation, and rectal temperature of < 104° Fo BU (CSS) compared to a saline-tracted approximation (Section 2006). (CSS) compared to a latter-tracted approximation (Section 2006). Anter incoduction (Intransally and Intratarcheally with a field station (M. Intragenetized two field the tracted with mich of that compared to solution). Saline: Figure exthanted and necespised 10 disp post-treatment. The mean percentage of gross saline. They were exthanted and necespised 10 disp spost-treatment. The mean percentage of gross (Saline). The same treated pips than for saline-treated pips in this studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated pips in the studies (SaStiv Scs 2047). Saline: They than for saline-treated

The effectiveness of tulathromycin injection for the control of S40 was evaluated in a multi-location natural infection field study, When at least 15% of the study candidate showed chiral signs of S40, and transmitter of the study study of the study candidate showed chiral signs of S40, and transmitter was evaluated on Day 7. Scores was defined as 2 ang with normal altitude, normal regization, and rexta temperature of < 1047. The treatment access rate was significant 12%, so that the study insignificant resterior granured to salm-treated piss (529, 427, 428).

ANIMAL SAFETY

Cattle Staffs subscription of the set of th

An exploratory study was conducted in free draws reacting as ingle subcutaneous door of 10, 12.5, or 15 mg/hg BW. Macroscopically, no lesions were observed. Microscopically, minimal to mild myscardial dependences was need not fix calves administered 12.5 mg/hg BW and two of six calves administered 17 mg/hg BW.

A safety study was conducted in preruminant calves 13 to 27 days of age receiving 2.5 mg/kg BW or 7.5 mg/kg BW once subcutaneously. With the exception of minimal to mild injection site reactions, no drug-related clinical signs or other lesions were observed macroscopically or microscopically.

Svine Safety studies were conducted in pigs receiving a single intramuscular dose of 25 mg/hg BW, or 3 weekly intramuscular doses of 2.5, 7.5, or 12.5 mg/hg BW. In all groups, transient indications of pain after injection were seen, including rectlessness and excessive worklazion. Ternoro scurred briefly in one maint receiving 7.5 mg/s BW. Bootcations and eleman of picture site tissues and corresponding histopathologic changes were seen in annual all discages and realized even time. No other drug-skatel clicolisms were observed macroscopilly or microscopication.

Store below 25°C (77°F), with excursion up to 40°C (10°F). Use this product within 45 days of the first
 puncture an animum of 20 times. If more than 20 punctures are anticipated, the use of
 under updamentel ar a reaction of 20 times. If more than 20 punctures are anticipated, the use of
 needle with hore dameter larger than 16 gauge, discard any product remaining in the vial immediately
 after oze.

To report suspected adverse drug events, for technical assistance or to obtain a copy of the Safety Data Sheet, contact Pamell at 1-800-887-2763. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS or http://www.fda.gov/reportanimalae.

HOW SUPPLIED RESPIRmycin (tulathromycin injection) Injectable Solution is available in the following package sizes

50 mL vial 100 mL vial 250 mL vial 500 mL vial

Approved by FDA under ANADA # 200-730

Manufactured by

PARNELL TECHNOLOGIES PTY. LTD. 4/476 Gardeners Road Alexandria NSW 2015 Australia

Distributed by: PARNELL U.S. 1, Inc.

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