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.



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.



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.

DESCAPTION

RESPRemyon Injectable Solution is a ready-ter-use stelle parenteral preparation containing tulathromyon, a semi-yon-their marville antibiotic of the solutions is a ready-ter-use stelle parenteral preparation containing tulathromyon, a semi-yon-their marville antibiotic of the solutions triamilitie. Each mil of RESPRemyon contains 100 mg of tulathromyon, 500 mg proglevel gody, 19.2 mg citric acid and 5 mg monochioplyceod. Sodium hydroxide or hydrochlaric acid may be added to adjust pit, RESPRemyon consists of an equilibrated mixture of two isomeric forms of tulathromyon in a 9:1 ratio, Structures of the isomers are shown below.





The chemical names of the isomers are (28.35.48, SR.8.108, T18, T25, T35, T48)—13[12.6-discoy-3-C-methyl-3-O-methyl-4-Ci[propylamino]
methyll-1-4-ind-hemily

Beef and Nort-Actating Dairy Cattle

Beef and Nort-Actating Dairy Cattle

BROD — RESTREMENT (In percent of boulen expiratory disease

(BRD) associated with Manufacine homolytice, Pestavella multicide, Michaphius samm, and

Appropriate boxis on the control of restrainty disease in cattle at high risk of developing BRD

associated with Manufacine homolytice, Pestavella multicide, Michaphius some, and Myraplessme boxis.

IBK — RESPIRmycin Injectable Solution is indicated for the treatment of infectious bovine keratoconjunctivitis (IBK) associated with Marazzella bavis.

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

Suckling Calves, Dairy Calves, and Veal Calves
BRD — RESPIRmydin Injectable Solution is indicated for the treatment of BRD associated with
M. hoemolytica, P. multocido, H. somni, and M. bovis.

Swine RSPRBmyrin hijectable Solution is indicated for the treatment of swine respiratory disease (SRD) associated with Admonability pleruppenamonice, Posturarilla multicidia, Bouteful bronchiseptica, Hennipolikap passaria, and Mylogolisom hippoprenominaria, and for the cantrol of \$150 associated with Admonability passaria, and Mylogolisom hippoprenominaria, and for the cantrol of \$150 associated with Admonability Posture SRD his been Bogrinoid.

DOSAGE AND ADMINISTRATION

Cattle Inject subcutaneously as a single dose in the neck at a dosage of 2.5 mg/kg (1.1 mL/100 lb) body weight (8M). Do not inject more than 10 mL per injection site. Table 1. RESPIRmyoin Cattle Dosing Guide

Animal Weight (Pounds)	Dose Volume (ml.)
100	1.1
200	2.3
300	3.4
400	4.5
500	5.7
600	6.8
700	8.0
800	9.1
900	10.2
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. RESPIRmyrin Swine Dosing Guide

Animal Weight (Pounds)	Dose Volume (mL)
15	0.2
30	0.3
50	0.6
70	0.8
90	1.0
110	1.3
130	1.5
150	1,7
170	1.9
190	2.2
210	2.4
230	2.6
250	2.8
270	3.1
290	3.3

CONTRAINDICATIONS

The use of RESPIRmycin Injectable Solution is contraindicated in animals previously found to be hypersensitive to the drug.

WARNINGS

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

RESIDUE WARNINGS

Cattle
Gattle intended for human consumption must not be slaughtered within 18 days from the last
treatment. This drug is not approved for use in female dairy cattle 20 months of age or older,
including dry dairy cows. Use in these cattle may cause drug residues in milk and/or in calves
born to these cows.

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

PRECAUTIONS

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

Swine The effects of RESPRmycin on porcine reproductive performance, pregnancy, and lactation have not been effectmented. Intramercular injection can cause a transient local lissue reaction that may result in trim loss of edible lissue as slaughter.

ADVERSE REACTIONS

Cattle In one B8D 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.

swine
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.

mus saluroun that recover a line sit to that our nature.

POST APPROVAL EXPERIENCE

I he following adverse events are based on post approval adversed may experience reporting. Not all adverse events are reported to the FBA CVM is to relaways possible to reliably estimate the adverse event frequency or establish a causal relationship to product exposure using these data. The following adverse events related in deverseing order of reporting frequency in cittle highcits on the reactions and anaphylatical/anaphylatic reactions. For a complete Isolating of adverse reactions for full attractions are related to the control of the contr

A physiological pH. Usubmervior to use twisted www.tds.gov/reportanismake.

A physiological pH. Usubmervior (a weak lobe) is approximately 50 lines more soluble in hydrophilic has hydrophile medic. This soluble with the entracellule pathogon activity typically associated with the marcindes. Markedly highe tulad mompion concentrations are observed in leangs as compared to the plasm. The enert to which upon concentrations personer five catched drug was not examined. Therefore, the clinical relevance of these elevated lung concentrations is undetermined.

unecernment. Although the relationship between buildhorpsych and the characteristics of fit antimicrobial effects has not been characteristics, as a closs, macrobides tend to be primarily bacteristatic, but may be bacteristatic, but may be bacteristatic properties. They also tend to epidiate contention independent illings the set of bacterial endication does not change once sexum drug concentrations reach. In 3 times the minimum hibilitary connection (MIQ) of the trapeley pathopes. Intel these conditions, the time that sexum concentrations reach are also the MIC becomes the major determinant of antimicrobial activity. Macrobides also entitin good-architosis reach of the conditions, the both drug and pathogen dependent. In general, by increasing the macrobide concentration and the exposure time, the PE will increase to some mazimal audious for Often work and become concentration and the opposure time, the great also to be the most powerful determinant of the duration of PAE.

Tulathromycin is eliminated from the body primarily unchanged via biliary excretion

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

²Nightingale, C.J. 1997, Pharmacokinetics and Pharmacodynamics of Newer Macrolides. Pediatr. Infect. Dis. J., 16:438—443.

Cattle
Gattle
Following subcutaneous administration into the nick of feeder calves at a disage of 2.5 mg/kg BW.
Following subcutaneous administration into the nick of feeder calves at a disage of 2.5 mg/kg BW.
Litalitationyrion is rapidly and nearly completely absorbed. Peak plasma concentrations generally occur
within 15 minutes after dissing and product relative bioavailability exceeds 90%. Itilial special extensive share distinguished 170 mil. Article. Litalitationyrion distributes extensively into body tissue, as evidenced by volume of distribution values of approximately 11 Litality animations after distribution is largely prosposable for the length animation half der distribution is largely prosposable for the length animation half der distribution is largely prosposable for the length animation half der distribution is largely prosposable for the length numation half der distribution is largely prosposable for the length of the compound versus. 8.75 days for total lang concentrations; chased on data from healthy animatiol.) Linear half described in the compound of the compound o

²Clearance and volume estimates are based on intersubject comparisons of 2.5 mg/kg BW administered by either subcutaneous or intravenous injection.

Swine Following intramuscular administration to feeder pigs at a dosage of 2.5 mg/kg BW, tulathromycin is completely and rapidly absorbed (T_{m.}—0.25 hour). Subsequently, the drug rapidly distributes into body tissues, achieving a volume of distribution exceeding 15 J. Ing. The fee drug is rapidly cleared from the systemic contained (T_{most} = 18 off Intribution, Newveet in the 3 offer periodic distribution of the system of distribution. Afficially offer all the system of distribution. Afficially administration owing to its exteriore volume of distribution. Afficially administration of the system of the system

MICRORIOLOGY

Lattle

Flathomyrich has demonstrated in vitro activity against Monoheimio haemolytica, Posteurella multocida,
Richsphillus somul, and Mycopiasma bowis, four pathogens associated with BRD, against Mononella bowis
associated with BRC, and against Fundacterium necrophorum and Prophyromonos levi associated with
bowise foot rot.

The MICs of tulathromycin against indicated BRD and IBK pathogens were determined using methods recommended by the Clinical and Laboratory Standards Institute (CLS, 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 MIXs of tulathromyoin were determined for SRD lociates obtained from calves enrolled in thespecific and a risk field studies in the U.S. in 1959. In the three-peopies crucies, solders were obtained studies and the studies of saline-treated calves that dide. If the artist studies, lociates were obtained from nasolyamped svolpts of saline-treated can be under the studies of the studies

IBK - The MKs of tulathromycin were determined for Moravella bovis isolates obtained from calves enrolled in IBK field studies in the U.S. in 2004, loolates were obtained from pretreatment conjunctival swales of calves with clinical signs of IBK enrolled in the tulathromycin injection and saline-treated groups. The results are shown in Talka 2.

Foot Rot – The IMICs of tulathromycin were determined for Fusobacterium necrophrorum and Posphyromorus Serii obtained from cattle enrolled in foot tot field studies in the U.S. and Canada in 2007. Solaties were obtained from pre-treatment interdigital biosies and owasis of cattle with clinical signs of foot rot enrolled in the tulathromycin injection and saline-treated groups. The results are shown in Table 3.

Table 3. Tulathromycin minimum inhibitory concentration (MIC) values* 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

Carlaua.							
Indicated pathogen	Date Isolated	No. of isolates	MK _{st} ** (µg/mL)	MBC ,** (pg/ml.)	WEC range (µg/mL)		
Mantheixia hoexolytica	1999	642	2	1	0.5 to 64		
Postewells multiroids	1999	221	0.5	1	0.25 to 64		
Histophilus somoi	1999	36	4	- 4	1 to 4		
Mycoplasma bovis	1999	46	0.125	1	≤ 0.063 ta > 64		
Marcuella bavis	2004	55	0.5	0.5	0.25 to 1		
Fusebacterium secrophorum	2007	116	2	64	≤ 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 pleuropneumonia
Posteurella multocida, Bardetella bronchiseptica, Haemophilus parasuis, and Mycoplosma hyopneumonia

The MMC of halathomysis against influenced SSD gathogoses were determined using methods recommended by the Clinical and Laboratory Standards Estitute (CSL, M31-A and M51-A3), MIC for Remorphism promises were determined using the control of SSD and Comparing with certain Fastistical Medium and were included up to 48 hours at 35 to 37°C in a CO-perinded atmosphere. All MiC values were determined using the 91 some ratio of this compound. budies obtained in 200 and 2002 were from lung samples from salian-treated pigs and non-treated sentinel pigs certified in Textiment of SSD field Studies in the U.S. and Canada. Description of the Comparing SSD and Comparing SSD and SSD an

Table 4. Tulathromycin minimum inhibitory concentration (MIC) values* for indicated pathogens isolated from field studies evaluating SRD in the LLS, and Canada.

Indicated pathogen	Date Isolated	No. of isolates	MK _{si} ** (µg/ml.)	MBC *** (pg/ml.)	MIC range (µg/mL)		
Actinobacillas picuropmaumonise	2000-2002 2007-2008	135 88	16 16	32 16	16 to 32 4 to 32		
Historophilus parasuis	2000-2002	31	- 1	2	0.25 to > 64		
Asterelia nultraids	2000-2002 2007-2008	55 40	1	2 2	0.5 to >64 ≤ 0.03 to 2		
Anadatolia konnekirantira	2100 2002	n	,		26.2		

*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.

FEFECTIVENESS

Cattle

Bill — In a multi-location field study, 314 cales with naturally occurring BBD were treated with
halt-toropic injection. Responses to reatment were compared to soline-treated control. A cur was
defined as a call with morned attended control, morned registration, and a restal temperature of 1647 in
Day 14. The cure rate was significantly higher (P > 0.55) in full-thromptin injection-treated calves (78%). The cure were two Bill-related deaths in the talline-treated calves (78%)
injection-treated calves (78%).

Tiffy-two talathromycin injection-treated calves and 27 saline-treated calves from the multi-location field 880 treatment study had *Misopolasma* boxis identified in cultures from pre-treatment anasopharpageal swabs. Of the 25 tubsthromycin injection-treated calves, 37 (71.2%) calves were categorized as curs and 15 (28.8%) calves were categorized as treatment failures. Of the 27 saline-treated calves, 37 (41.4%) calves were categorized as curs and 23 (85.2%) calves were treatment failures.

A Bysesian meta-inalysis was conducted to compare the BRD treatment success rate in young calves (calves weighing 250 lbs or less and feel primarily a milk-based died treated with tulathomyork injection to the success rate in older calves (calves weighing more than 250 lbs and feel primarily stroughage and grain-based died; treated with tulathomyorin injection. The analysis induded data from four BRD tereintent effectiveness subdises conducted for the approval of tulathomyorin injection in NeU. 3 and time contemporaneous studies conducted in future; The analysis showed that the BRD treatment success rate in young calves such a less at speak at the BRD treatment of BRD assistance at the Indicated with Ith hemolytics, it and/located; it many and Mr. bows in sucking when, also related in the successful of the successful with Ith hemolytics, it mollocated; it many, and Mr. bows in sucking when, also related in the successful of the successful with Ith hemolytics, it mollocated; it many, and Mr. bows in sucking when, also calves, and verall calves.

In another multi-faction field study with 390 claves a high risk of developing 880, administration to butletmorphic injection resulted in a significantly reduced incidence of 890 (11%) compared to butletmorphic injection resulted in a significantly reduced incidence of 890 (11%) compared to sindine-toted calcular solysis. Effectiveness causation was based on sorted clinical significantly are statistically continued in the significant statistical control respiration, and a restal temperature of \$100\cdot on 10\cdot on 10\cdot of 10\cd

Two induced infection model studies were conducted to confirm the effectiveness of lather topic induced infection against disciplination against disciplination against disciplination against disciplination aboves. A total of 166 calses were incutated intratactically with field strains of disciplination aboves. When calves became privace and had abnormal respiration scores, they were tested with either tatalitomiciny indiction 16.27 miles gods with earlier advanced or an equivalent two more of saline. Calves were observed for signs of 8800 for 14 days post-teamment, then were detained and necopised. To both studies, many long lesion percentages were statistically significantly lower in the lauthormytin injection-treated calves (11.3% vs. 2.85%, P = 0.0001 and 15.0% vs. 30.3%, P < 0.0007.

18K — Iwo field studies were conducted evaluating total momyorin injection for the treatment of IBK associated with Moracilla bows in 200 naturally-infected calves. The primary clinical engiporal of these studies was our early defined as as all of which no ficial signs of IBA and no cornel sides; assessed on Days 5, 9, 13, 7, and 21. Time to improvement, defined as the first day on which a call that no clinical signs of Bik inot beyes, provided that those scores were mathatized at the next day of observation, was assisted as a secondary vanish. A stall time points, in both studies, the cure rate was significantly higher (P < 0.05) for habilitationy in intercellmental cells are compared to authorise days and considerable, time to calves compared to saline-treated calves.

Foot Bot - The effectiveness of statisticomy in injection for the treatment of bovine foot rot was evaluated in 170 cattle in two field studies. Cattle diagnosed with bovine foot rot were enrolled and treated with a single subcutaneaus foos of buildstromyfin specificant (2.7 mg/g) give) or an equivalent vitume of saline. Cattle were clinically evaluated 2 days after treatment for treatment success, which was based on defined electrons in leafly success, which was based on defined electrons in leafly success, the soft studies, the breatherst success percentage was statistically significantly higher in buildningsvin injection retarted calves compared with saline-treated calves (60 Ws. 80; K. et 0000 at all 32 S. w 55%; P. et 0.080 s.

Swine
In an util sociation field study to evaluate treatment of naturally occurring 580, 266 pigs were treated
In an util slocation field study to evaluate treatment of naturally occurring 580, 266 pigs were treated
with fullathomyton injection. Responses to treatment were compared to saline-treated controls. Success
was defined as a pig with normal attitude, normal respiration, and retrail temperature of < 104° ero.

170, 180; compared to saline-treated pigs (64.18). M. Any position of the saline-treated saline-treated saline-treated seatine pigs in this study. Two induced intection model studies were
conducted to confirm the effectiveness of Insulationsystic injection apasits M. Appropriamonies. Ten days
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The effectiveness of Intalhomoral injection for the control of SSD was evaluated in a multi-basical natural infection field study. When at least 15% of the multi-conditions showed diminol separ of SRD, all the study conditions are supported by the study of SRD, and the study of SRD, a

ANIMAL SAFETY

Cattle
Safety studies were conducted in feeder calves receiving a single subcutaneous dose of 25 mg/kg BN or 3 weetly solututaneous dose of 25, 75, or 12.5 mg/kg BN or all groups, transient indications of pain after injection were seen, indicing head saking and paring at the gound linjection is serveiling, disculoration of the subcutaneous tissues at the injection site and corresponding histopathologic changes were seen in animals in all discase groups. These lesions showed group of realizing over time. No other drug-related lesions were observed macroscopically or microscopically.

An exploratory study was conducted in feeder calves receiving a single subcutaneous dose of 10, 12.5, or 15 mg/kg BW. Macroscopicilly, no lesions were observed. Microscopically, minimal to mild myocardial degeneration was seen in one of six calves administered 12.5 mg/kg BW and two of six calves administered 15 mg/kg 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 or 7.6 mg/kg BW or 8.6 mg/kg BW or 7.6 mg/kg BW or 8.6 mg/kg BW or 9.7 mg/kg B

Swine
Safety studies were conducted in pigs receiving a single intramuscular dose of 25 mg/kg BW, or 3 weekly
intramuscular doses of 25, 75, or 125 mg/kg BW, in all groups, transient indications of pain after
injection were seen, including predictiones and excessive vocaluation, fermion securiced biolity in one
injection were seen, and only a predictiones and excessive vocaluation, fermion securiced biolity in one
histopathologic changes were seen in animals at all disease, and resolved over time. Ho other
drug-related lesions were observed macroscopically or microscopically.

Store below 25°C (77°F), with excusions up to 40°C (104°F), be this product within 45 days of the first puncture and puncture a maximum of 20 times. If more than 20 punctures are antiopated, the use of aduntantic injection equipment of a protect single is recommended. When using a dame-of signific or needle with tore dameter larger than 16 gauge, discard any product remaining in the vial immediately after use.

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/reportanimales.

HOW SUPPLIED

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

Approved by FDA under ANADA # 200-730

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

PARNELL U.S. 1, Inc. 7015 College Boulevard, Level 6, Overland Park, KS, 66211

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