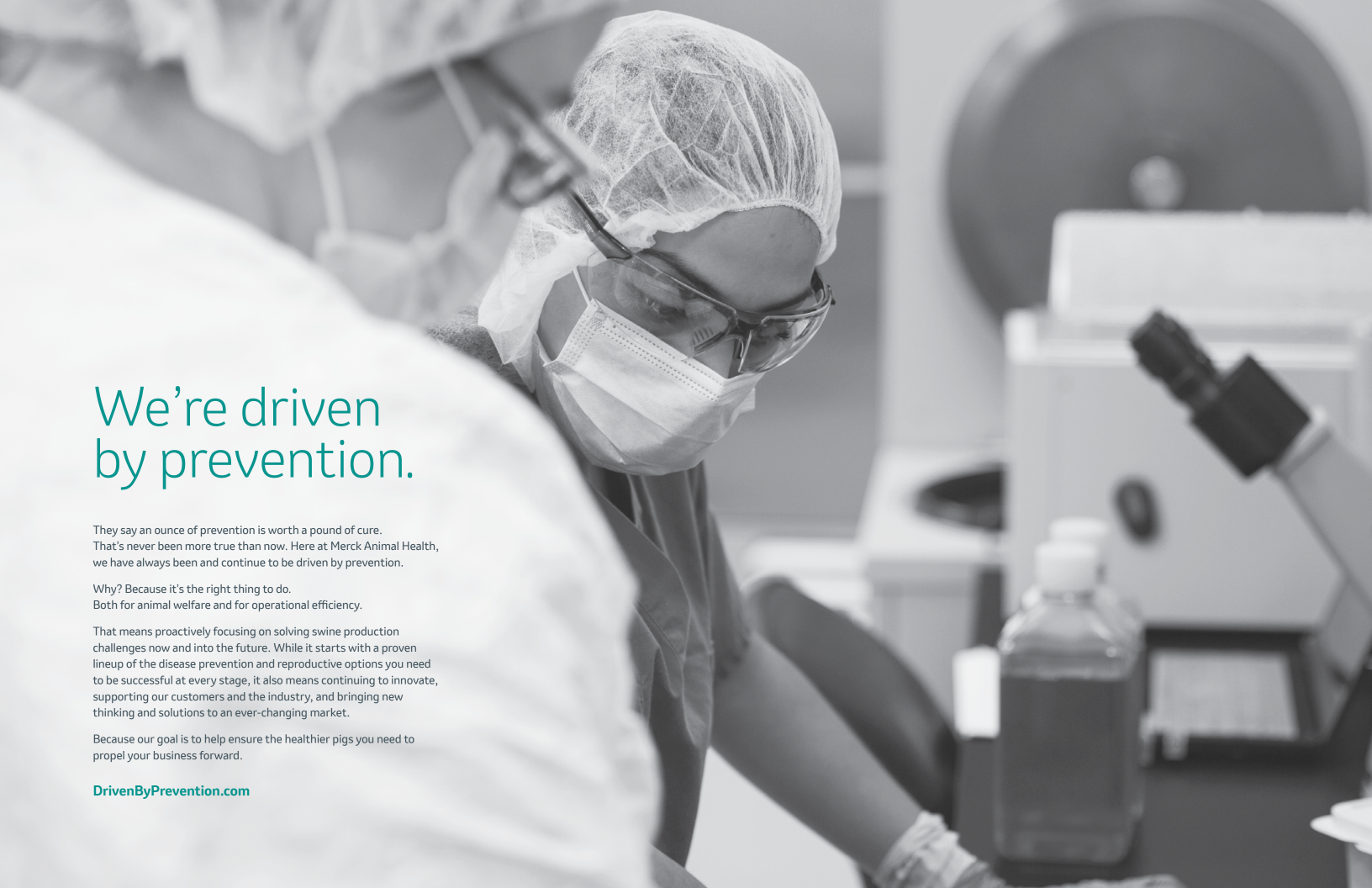


SWINE PRODUCT GUIDE







CIRCUMVENT® CML

With MICROSOL DILUVAC FORTE® - 50 dose, 250 dose (NEW 500 dose available soon)

This product has been shown effective for the vaccination of healthy swine 3 weeks of age or older against disease caused by porcine circovirus types 2a and 2d, Mycoplasma hyopneumoniae and Lawsonia intracellularis. Duration of immunity (DOI) for porcine circovirus type 2d is at least 16 weeks, and for Mycoplasma hyopneumoniae is at least 10 weeks. Duration of immunity for ileitis due to Lawsonia intracellularis is at least 20 weeks. Duration of immunity for porcine circovirus type 2a has not been established.



CIRCUMVENT® PCV G2

With MICROSOL DILUVAC FORTE® - 50 dose, 250 dose (NEW 500 dose available soon)

An aid in the prevention of viremia, an aid in the reduction of virus shedding and an aid in the reduction of lymphoid infection caused by porcine circovirus type 2. Convenient dosing options (one x 2 mL or two x1mL) for one-dose and two-dose programs. (See complete label instructions.) The only PCV2 vaccine approved for use in pigs as early as 3 days of age (two-dose option). Five-month PCV2 duration of immunity.



Two-Dose Option:

1 mL at 3 days of age or older, followed by second 1 mL 3 weeks later.



One-Dose Option:

2 mL once at 3 weeks of age or older.



CIRCUMVENT® PCV-M G2

With MICROSOL DILUVAC FORTE® - 50 dose, 250 dose (NEW 500 dose available soon)

An aid in the prevention of viremia, an aid in the reduction of virus shedding, an aid in the reduction of lymphoid infection caused by porcine circovirus type 2 and an aid in the reduction of lung lesions caused by Mycoplasma hyopneumoniae. Convenient dosing options (one x 2 mL or two x 1 mL) for one-dose and two-dose programs. (See complete label instructions.) The only PCV2 vaccine approved for use in pigs as early as 3 days of age (two-dose option). Five-month PCV2 DOI



Two-Dose Option:

1 mL at 3 days of age or older, followed by second 1 mL 3 weeks later.



One-Dose Option:

2 mL once at 3 weeks of age or older.



M+PAC®

With EMUNADE® - 250 dose

An aid in the prevention of pneumonia caused by Mycoplasma hyopneumoniae infection in swine. Convenient one- or two-dose options (one x 2 mL and two x 1 mL) for one-dose and two-dose programs. (See complete label instructions.) Unique patented dual-emulsion adjuvant. Duration of immunity of at least 4 months has been established for the Intramuscular route with 6 week or older swine.





Two-Dose Option:

1 mL at 7-10 days of age or older, followed by second 1 mL 2 weeks later.



One-Dose Option:

2 mL once at 6 weeks of age or older.

WEEK 5 OR OLDER



MYCO SILENCER® ONCE

With MICROSOL DILUVAC FORTE® - 250 dose

An aid in the prevention of pneumonia caused by Mycoplasma hyopneumoniae infection in swine. Convenient one- or two-dose options (one x 2 mL and two x 1 mL) for one-dose and two-dose programs. (See complete label instructions.) Unique patented dual-emulsion adjuvant. Up to six months DOI with a single shot.



Two-Dose Option:

Second 1 mL shot given 2-3 weeks after first 1 mL shot. (Second shot not required if 2 mL shot is given at week 3.)





PORCILIS® ILEITIS

With MICROSOL DILUVAC FORTE° - 50 dose, 250 dose (NEW 500 dose available soon)

An aid in the control of ileitis caused by Lawsonia intracellularis, an aid in the reduction of colonization by Lawsonia and an aid in the reduction of duration of fecal shedding. Convenient dosing options (one x 2 mL or two x 1 mL) for one-dose and two-dose programs as early as 3 days of age (two-dose option). (See complete label instructions.) DOI for at least 20 weeks has been demonstrated.



Two-Dose Option:

1 mL at 3 days of age or older, followed by second 1 mL 3 weeks later.



One-Dose Option:

2 mL once at 3 weeks of age or older.



PRIME PAC® PRRS RR

Modified Live Virus (MLV) - 100 dose

This product has been shown to be effective for the vaccination of healthy swine 3 weeks of age or older against respiratory form of the disease caused by Porcine Reproductive and Respiratory Syndrome (PRRS) virus and for the vaccination of female breeding age swine against reproductive disease caused by PRRS virus. Duration of immunity against reproductive form of the disease is at least 20 weeks. Duration of immunity against respiratory form of the disease is at least 23 weeks. For sows and gilts, inject a single 1mL dose intramuscularly eight weeks prior to breeding, and for piglets, inject a single 1 mL dose IM at 3 weeks of age or older. (See complete label instructions.) Freeze-dried MLV vaccine with diluent.





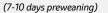
PROSYSTEM® ROTA

Modified Live Virus (MLV) - 50 dose

An aid in the prevention of rotaviral diarrhea in young pigs; a 1-mL oral dose and 1-mL IM dose to pigs preweaning. (See complete label instructions.) Unique rotavirus protection includes two major serotypes - G4 and G5 - of serogroup A. Freeze-dried MLV vaccine with diluent.









INJECTION ADMINISTRATION



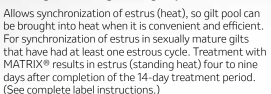
KEY





MATRIX®

(altrenogest) - 2.2 mg altrenogest per mL (0.22%)



IMPORTANT SAFETY INFORMATION: Gilts must not be slaughtered for human consumption for 21 days after the last treatment. Do not use in gilts having a previous or current history of uterine inflammation. Underfeeding of MATRIX® (altrenogest) Solution 0.22% may lead to the occurrence of cystic follicles. Always use the MATRIX® Dosing Device to administer this product. Skin contact must be avoided as MATRIX® is readily absorbed through unbroken skin, and exposure may result in serious side effects to both women and men. Always wear vinyl, neoprene, or nitrile protective gloves when handling MATRIX® or when in contact with equipment or surfaces contaminated by this product. Latex gloves are not protective. PREGNANT WOMEN OR WOMEN WHO MAY BE PREGNANT SHOULD NOT HANDLE MATRIX® (altrenogest). WOMEN OF CHILDBEARING AGE SHOULD EXERCISE EXTREME CAUTION WHEN HANDLING THIS PRODUCT. Accidental absorption, such as absorption through the skin, could lead to a disruption of the menstrual cycle or prolongation of pregnancy. Wash off accidental spillage on the skin immediately with soap and water. Any equipment or surfaces that come in contact with MATRIX® should be adequately cleaned and decontaminated to prevent human exposure. Other people who should not handle this product include those with thrombophlebitis, thromboembolic disorders, or with a history of these events, cerebral-vascular or coronary-artery disease, suspected estrogen-dependent neoplasia, benign or malignant tumors which developed during the use of oral contraceptives or other estrogen containing products, liver dysfunction or disease, and women with known or suspected carcinoma of the breast or undiagnosed vaginal bleeding. For complete safety information, refer to the product label



P.G. 600°

(serum gonadotropin and chorionic gonadotropin for injection)

Maximizes pig flow by helping more gilts and weaned sows cycle, particularly in summer, producing more pigs when market prices are high.^{1,2} For induction of fertile estrus (heat) in healthy prepuberal (noncycling) gilts over 5½ months of age and weighing at least 187 lbs. For induction of estrus in healthy weaned sows experiencing delayed return to estrus. (See complete label instructions.)

Induction and Synchronization of Estrus in Prepuberal Gilts and Anestrous Sows by a PMSG/ HCG-Compound Technical Report No. 9.

²The Attainment of Estrus in Sows Administered with 400 I.U. Pregnant Mare Serum Gonadotropin and 200 I.U. Human Chorionic Gonadotropin at Weaning.

IMPORTANT SAFETY INFORMATION: Treatment will not induce estrus in gilts that have already reached puberty (begun to cycle). Gilts that are less than five and one-half months of age or that weigh less than 85 kg (187 lb.) may not be mature enough to continue normal estrus cycles or maintain a normal pregnancy to full term after treatment. Treatment will not induce estrus in sows that are returning to estrus normally three to seven days after weaning. Delayed return to estrus is most prevalent after the first litter; the effectiveness of P.G. 600 has not been established after later litters. Delayed return to estrus often occurs during periods of adverse environmental conditions, and sows mated under such conditions may farrow smaller than normal litters. For complete safety information, refer to the product label.

PRE-FARROW
PRODUCTS FOR REPRODUCTION
EFFICIENCY AND HEALTH



PROSYSTEM® RCE

Modified Live Virus (MLV) - 25 dose

An aid in prevention of rotaviral diarrhea and enterotoxemia colibacillosis in nursing pigs of vaccinated sows/gilts. Unique rotavirus and seven-way scours protection. Includes two major rotavirus serotypes – G4 and G5 – of serogroup A. Freeze-dried MLV vaccine with bacterin/toxoid diluent. (See complete label instructions.)

SAFE-GUARD® (fenbendazole) DEWORMER

20% Type A Medicated Feed Article

For the treatment and control of: **Lungworms:** Adult *Metastrongylus apri*, adult *Metastrongylus pudendotectus*; **Gastrointestinal worms:** Adult and larvae (L3, L4 stages, liver, lung, intestinal forms) large roundworms (*Ascaris suum*), Adult nodular worms (*Oesophagostomum dentatum*, *O. quadrispinulatum*), Adult small stomach worms (*Hyostrongylus rubidus*), Adult and larvae (L2, L3, L4 stages – intestinal mucosal forms) whipworms (*Trichuris suis*); and **Kidney worms:** Adult and larvae *Stephanurus dentatus*. The SAFE-GUARD Dewormer 20% Type A Medicated Feed article can only be purchased by an approved medicated feed mill. (See complete label instructions.) Ask your Merck Animal Health representative for the various SAFE-GUARD presentations available from our distributor partners.

IMPORTANT SAFETY INFORMATION: Swine must not be slaughtered for human consumption within 4 days following last treatment with this drug product. Consult your veterinarian for assistance in the diagnosis, treatment and control of parasitism.

SAFE-GUARD® AQUASOL

(fenbendazole oral suspension) - 200 mg fenbendazole/mL

Indicated for swine, except for nursing piglets, for the treatment and control of: **Lungworms:** Adult *Metastrongylus apri*, Adult *Metastrongylus pudendotectus*; **Gastrointestinal worms:** Adult and larvae (L3, L4 stages, liver, lung, intestinal forms) large roundworms (*Ascaris suum*); Adult nodular worms (*Oesophagostomum dentatum*, *O. quadrispinulatum*); Adult small stomach worms (*Hyostrongylus rubidus*); Adult and larvae (L2, L3, L4 stages – intestinal mucosal forms) whipworms (*Trichuris suis*); and **Kidney worms:** Adult and larvae *Stephanurus dentatus*. See Product Information for complete directions and warnings.

IMPORTANT SAFETY INFORMATION: Swine intended for human consumption must not be slaughtered within 2 days of the last treatment. Consult your veterinarian for assistance in the diagnosis, treatment and control of parasitism.

SYMPTOMATIC CARE PRODUCTS FOR PIG HEALTH



BANAMINE®-S

(flunixin meglumine injection) - 50 mg/mL

For control of pyrexia (fever) associated with swine respiratory disease. (See complete label instructions.)

IMPORTANT SAFETY INFORMATION: Swine must not be slaughtered for human consumption within 12 days of last treatment. Do not use in animals showing hypersensitivity to flunixin meglumine. Use judiciously, when renal impairment or gastric ulceration is suspected. Not for use in breeding swine. For complete information on Banamine-S Injectable Solution use, contraindications and warnings, see accompanying product package insert.

BIOSECURITY PRODUCTS FOR PIG HEALTH



ARMATREX® ANTIMICROBIAL

A Silane Quaternary Ammonium Salt

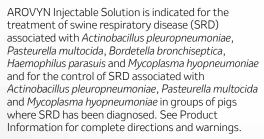
EPA-registered for use in livestock and companion animal facilities as a final bacteriostatic finish to impart fungistatic (mold and mildew) or algistatic activity that provides freshness and reduces surface deterioration or microbiologically induced corrosion.

R

AROVYN™

18181118888

(tulathromycin injection) - 100 mg/mL



IMPORTANT SAFETY INFORMATION: The pre-slaughter withdrawal time for AROVYN in swine is five days. AROVYN should not be used in animals known to be hypersensitive to the product.

NUFLOR®-S

For the vaccination of healthy swine as a control against disease caused by

Porcine Epidemic Diarrhea Virus.

(florfenicol) Injectable Solution - 300 mg/mL



The first and only injectable florfenicol approved for use in U.S. swine. For treatment of swine respiratory disease associated with Actinobacillus pleuropneumoniae, Pasteurella multocida, Salmonella Choleraesuis, Streptococcus suis, Bordetella bronchiseptica and Glaesserella (Haemophilus) parasuis in swine except for nursing piglets and swine of reproductive age intended for breeding. See Product Information for complete directions and warnings.

IMPORTANT SAFETY INFORMATION: Do not use in animals intended for breeding purposes. Perianal inflammation, rectal eversion, rectal prolapse and diarrhea may occur transiently following treatment. Swine intended for human consumption must not be slaughtered within 11 days of the last intramuscular treatment. Intramuscular injection may result in trim loss of edible tissue at slaughter. The effects of florfenicol on porcine reproductive performance, pregnancy and lactation have not been determined. Not for human use and keep away from children. Avoid direct contact with skin, eyes, and clothing. Pregnant women should wear gloves and exercise caution or avoid handling this product. See Product Information for Full Prescribing Information.

RNA PARTICLE TECHNOLOGY PRODUCTS FOR PIG HEALTH

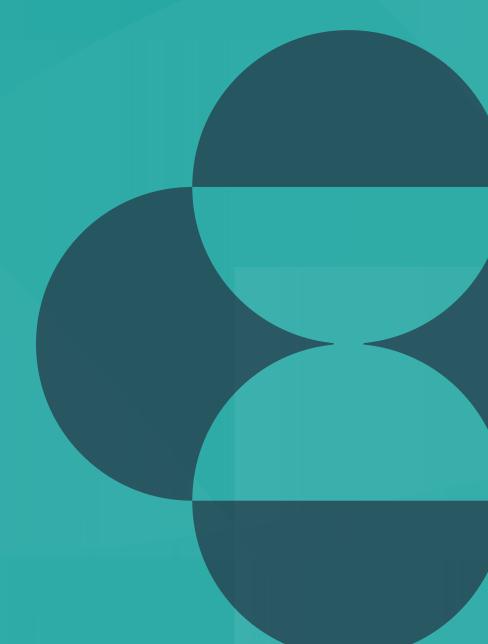


MERCK		Protection														
Animal Health				F41						<u> </u>						
VACCINE			ype C	K99, 987P, I	a		iae			Porcine Epidemic Diarrhea (PED)						
ANTIGEN			ns T	K99,	Swin	żi	mor	pe 2	/pe 3	arrhe					sinis	in
CHART			ringe	,88	s of !	llular	naudo	us Ty	us Ty	ic Di					eraes	imur
			perf	coli k	Viru	trace	a hyc	covir	covir	dem					Chol	Typh
• = Indicated Use			dium	ichia	ıza A	nia in	lasm	e Cir	e Cir	e Epi		us A	us B	us C	nella	rella
■ = Modified Live	Sows	Pigs	Clostridium perfringens Type	Escherichia coli K88,	nfluenza A Virus of Swine	_awsonia intracellularis	Mycoplasma hyopneumoniae	Porcine Circovirus Type 2	Porcine Circovirus Type 3	orcin(PRRSV	Rotavirus A	Rotavirus B	Rotavirus C	Salmonella Choleraesuis	Salmonella Typhimurium
Circonium Vancium	Š	P	Ü	Ë	=	7	Z,	<u>A</u>	Ğ	<u>A</u>	<u>a</u>	R	R	R	SS	S
Circovirus Vaccines CIRCUMVENT® CML		•				•	•	•								
CIRCUMVENT® PCV G2		•				_	•	•								\vdash
CIRCUMVENT® PCV-M G2		•					•	•								
Respiratory Vaccines																
MYCO SILENCER® ONCE		•					•									
M+PAC [®]		•					•									
PRIME PAC® PRRS RR	•	•									•					
Enteric Vaccines																
PORCILIS® ILEITIS		•				•										
ARGUS® SC/ST		•													•	•
PROSYSTEM® RCE	•		•	•								•				
PROSYSTEM® ROTA		•										•				
RNA Particle Technology Va	accin	es														
SEQUIVITY®*	•				•			•	•	•		•	•	•		

 $^{{}^{\}star}\mathsf{Talk}\ \mathsf{to}\ \mathsf{a}\ \mathsf{Merck}\ \mathsf{Animal}\ \mathsf{Health}\ \mathsf{representative}\ \mathsf{to}\ \mathsf{learn}\ \mathsf{more}\ \mathsf{about}\ \mathsf{all}\ \mathsf{SEQUIVITY}\ \mathsf{antigens}.$

RECOMMENDED NEEDLE SIZES AND LENGTHS:					
GAUGE	LENGTH				
18 or 20	5/8" or 1/2"				
16 or 18	3/4" or 5/8"				
16	1"				
14 or 16	1" or 11/2"				
GAUGE	LENGTH				
16 or 18	1/2"				
16	3/4"				
14 or 16	1"				
	GAUGE 18 or 20 16 or 18 16 14 or 16 GAUGE 16 or 18 16				

PRODUCT INFORMATION AND LABEL INSTRUCTIONS



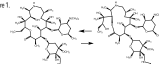


100 mg of tulathromycin/mL

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. DESCRIPTION

AROWN Injectable Solution is a ready-to-use sterile parenteral preparation containing tulathromycia a semi-synthetic macrolide antihiotic of the subclass triamilide. Each mL of AROVYN contains 100 m nycin, 500 mg propylene glycol, 19.2 mg citric acid and 5 mg monothioglycerol. Sodiun or hydrochloric acid may be added to adjust pH. AROWN consists of an equiliblated nixture of two isomeric forms of tulathromycin in a 9:1 ratio. Structures of the isomers are shown



The chemical names of the isomers are (2R.3S,4R.5R.8R,10R,11R,12S,13S,14R).13.[12.6 dideoxy-3-C-methyl-3-C-met 3-(dimethylamino)-β-D-xylohexopyranosyl]oxy]-1-oxa-4- azacyclotridecan-13-one, respectively

Beef and Non-Lactating Dairy Cattle
BRD - AROVYN 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 Mannheimia haemolytica, Pasteurella multocida, Histophilus somni, and Mycoplasma bovis. IBK – AROWYN Injectable Solution is indicated for the treatment of infectious bovine keratoconjunctivitis (IBK) associated with Moraxella bovis.

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

necrobacillosis) associated with Fusobactenium necrophorum and Porphyromonas levii. Sudding Calves, Dainy Calves, and Veal Calves BRD - AROYM Injectable Solution is indicated for the treatment of BRD associated with M. haemofytica, P. mulkocida, H. sommi, and M. bovis. Swine AROYM Injectable Solution is indicated for the treatment of swine respiratory disease (SRD) received with Articohacillies relaxeneeumonics. Between the swine respiratory disease (SRD)

associated with Actinobacillus pleuropneumoniae, Pasteurella multocida, Bordetella bronchiseptica, Haemophilus parasuis, and Mycoplasma hyponeumoniae; and for the control of SRD associated with Actinobacillus pleuropneumoniae, Pasteurella multocida, and Mycoplasma hyponeumoniae in groups of pigs where SRD has been diagnosed.

DOSAGE AND ADMINISTRATION

Inject subcutaneously as a single dose in the neck at a dosage of 2.5 mg/kg (1.1 mL/100 lb) body weight (BW). Do not inject more than 10 mL per injection site.

Table 1. AROVYN Cattle Dosing Guide

	Table 1. ANOVIN Calle Dosing Guide						
	Animal Weight (Pounds)	Dose Volume (mL)	Animal Weight (Pounds)	Dose Volume (mL)			
ı	100	1.1	600	6.8			
1	200	2.3	700	8.0			
	300	3.4	800	9.1			
	400	4.5	900	10.2			
١	500	5.7	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

Table 2. AKOVYN SWINE D	osing Guide		
Animal Weight (Pounds)	Dose Volume (mL)	Animal Weight (Pounds)	Dose Volume (mL)
15	0.2	170	1.9
30	0.3	190	2.2
50	0.6	210	2.4
70	0.8	230	2.6
90	1.0	250	2.8
110	1.3	270	3.1
130	1.5	290	3.3

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

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

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 dairy cattle 20 months of age or older, including dry dairy cows. Use in these attle may cause drug residues in milk and/or in calves born to these cows

ine intended for human consumption must not be slaughtered within

PRECAUTIONS

Cattle
The effects of AROYYN 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 AROVYN 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

caute 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 b transient hypersalivation. One of these calves also exhibited transient dyspnea, which may have beer related to pneumonia. Swine In one field study, one out of 40 pigs treated with fullathromycin injection at 2.5 mg/kg BW exhibited and a filter to the translated in beauthon four hours.

POST APPROVAL EXPERIENCE

POST APPROVAL EXPERIENCE
The following adverse events are based on post approval adverse drug experience reporting. Not all adverse events are reported to the FDA CVM. It is not always possible to reliably estimate the adverse event frequency or establish a causal relationship to product exposure using these data. The following adverse events are listed in decreasing order of reporting frequency in cattle: Injection site reactions and anaphylaxids-anaphylaction feations. For a complete listing of adverse reactions for tulathromycin injectable solution reported to the CVM see: http://www.fda.gov/reportanimalae.

CLINICAL PHARMACOLOGY

CLINICAL PHARMACOLOGY
At physiological pH, tulathromycin (a weak base) is approximately 50 times more soluble in hydrophiloit than hydropholoit media. This solubility profile is consistent with the extracellular pathogen activity hydrophiloit media. This solubility profile is consistent with the extracellular pathogen activity hydrophiloit solubility as of the foliation or concentrations are observed in the lungs as compared to the plasma. The extent to which lung concentrations represent free factive) drug was not examined. Therefore, the clinical relevance of these elevated lung concentrations is undetermined. Although the relationship between tulathromycin and the characteristics of its antimicrobial effects has not been characterized, as a class, macrolifes tend to be primarily bacteriostatic, but may be bacterioidal against some pathogens.²⁷ They also tend to exhibit concentration independent killing; the rate of bacterial eradication does not change once serum drug concentrations reach 2 to 3 times the minimum inhibitory concentration (MIC) of the targeted pathogen. Under these conditions, the time that serum concentrations remain above the MIC becomes the major determinant of antimicrobial activity. Macrolides also exhibit a post-antibiotic effect (PAE), the duration of which thends to be both drug and pathogen dependent in queneal, by MIL Decomes the major determinant or animinuous activity, reactivities as which the period process makes a feefect (PAE), the duration of which thanks to be both droug and pathogen dependent in la peneral, by increasing the macrolide concentration and the exposure time, the PAE will increase to some maximum and control the process of the process of

the most powerful determinant of the duration of PAE. Iulathromycin is eliminated from the body primarily unchanged via biliany exception.

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

'Mightingale, C. J. 1997. Pharmacokinetics and P harmacodynamics of Newer Macrolides. Pediatr. Infect. Dis. J., 16:438-443.

Cattle
Following subcutaneous administration into the neck of feeder calves at a dosage of 2.5 mg/kg BW, tulathromycin is rapidly and nearly completely absorbed. Peak plasma concentrations generally occur within 15 minutes after dosing and product relative biovavailability exceeds 90%. Total systemic dearance is approximately 170 mUrkur, Tulathromycin distributes extensively into body tissues, as evidenced by volume of distribution values of approximately 11 L/kg in healthy ruminating calves.³ evidenced by volume of distribution values of approximately 1 LVsg in neathy ruminating caives. This extensive volume of distribution is largely responsible for the long elimination half-life of this compound [approximately 2.75 days in the plasma (based on quantifiable terminal plasma drug concentrations) versus 8.75 days for total lung concentrations (based on data from healthy animals). Linear pharmacokinetic are observed with subcutaneous doses ranging from 1.27 mg/kg BW to 5.0 mg/kg BW. No pharmacokinetic differences are observed in castrated male versus female calves. "Clearance and volume estimates are based on intersubject comparisons of 2.5 mg/kg BW administered by either subcutaneous or intravenous injection.

Swine Pollowing intramuscular administration to feeder pigs at a dosage of 2.5 mg/kg BW, tulathromydin is completely and rapidly absorbed (T_{max} "0.25 hour). Subsequently, the drug rapidly distributes into body itssues, achieving a volume of distribution exceeding 15 L/kg. The free drug is rapidly deared from the systemic circulation (C_{0,5 femic} = 187 mL/hr/kg.) However, it has a long terminal elimination half-life (60 to 90 hours) owing to its extensive volume of distribution. Although pulmonary tulathromyric noncentrations are usubstantially higher than concentrations observed in the plasma, the clinical significance of these findings is undetermined. There are no gender differences in swine bubbbrowing a hourse are lossed. tulathromycin pharmacokinetics.

ulathromycin has demonstrated *in vitro* activity against *Mannheimia haemolytica, Pasteurella* multocida, Histophilus somni, and Mycoplasma bovis, four pathogens associated with BRD; against Moraxella bovis associated with IBK; and against Fusobacterium necrophorum and Porphyromonas levii associated with boyine foot rot.

Moraveila bow's associated with IBK, and against rusobactenium necrophorum and Prophyromonas lewi's associated with bowine foot indicated BRD and IBK pathogens were determined using methods recommended by the Clinical and Laboratory Standards Institute (CLSI, MS1 + A2). The MICs against foot tot pathogens were also determined using methods recommended by the CLSI (M11+A6). BIMC values were determined using methods recommended by the CLSI (M11+A6). BIMC values were determined using the 9-1 isomer ratio of this compound. BRD—The MICs of tulathromycin were determined for BRD isolates obtained from cakes enrolled in the appetute and at-risk field busides in the U.S. in 1999. In the therapeutic studies, isolates were obtained from pretreatment nasopharyngeal swabs from all study calves, and from lung swabs or lung tissue of saline-treated calves that died. In the at-risk studies, isolates were obtained from nasopharyngeal swabs of saline-treated anon-responders, and from lung swabs or lung tissue of saline-treated calves that died. The results are shown in Table 3. IBK—The MICs of fullathromycin were determined for Moravella bow's isolates obtained from calves enrolled in IBK field studies in the U.S. in 2004. Isolates were obtained from pre-treatment conjunctual swabs of calves with dirical signs of IBK enrolled in the tulathromycin injection-treated and saline-treated groups. The results are shown in Table 3.

Foot Rot – The MICs of tulathromycin were determined for Fusobacterium necrophorum and Porphyromonas levii obtained from cattle enrolled in foot rot field studies in the U.S. and Canada in Prophylomonas levi obtained from cattle enrolled in foot for field studies in the U.S. and Canada in 2007. Isolates were obtained from pretreatment interdigital biopsies and swabs of cattle with clinical signs of foot rot enrolled in the tulathromycin injection-treated 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

0.5. una canada.					
Indicated pathogen	Date isolated	No. of isolates	MIC ₅₀ ** (ug/mL)	MIC ₉₀ ** (ug/mL)	MIC range (ug/mL)
Mannheimia haemolytica	1999	642	2	2	0.5 to 64
Pasteurella multocida	1999	221	0.5	1	0.25 to 64
Histophilu somni	1999	36	4	4	1 to 4
Mycoplasma bovis	1999	43	0.125	1	≤ 0.063 to > 64
Moraxella bovis	2004	55	0.5	0.5	0.25 to 1
Fusobacterium necrophorum	2007	116	2	64	≤ 0.25 to > 128
Porphyromonas 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.

In vitro activity of tulathromycin has been demonstrated against Actinobacillus pleuropne In vitro activity of tulathromycin has been demonstrated against Actinobacillus pleuropneumoniae, Pasteurella mulhocida, Bordetella bronchiseptica, Haemophis parasuis, and Mycoplasma hypopneumoniae. The MICs of tulathromycin against indicated SRD pathogens were determined using methods recommended by the Clinical and Laboratory Standards Institute (CLSI, M31 A and M31-A3) MICs of Haemophilus parasuis were determined using Veterianary Fasticious Medium and were incubated up to 48 hours at 35 to 37°C in a CO2-enriched atmosphere. All MIC values were determined using the 9:1 isomer ratio of this compound. Isobates obtained in 2000 and 2002 were from lung samples from saline-treated and the rated set entire lips enrolled in Teatment of SRD field studies in the U.S. and Canada. Isolates obtained in 2007 and 2008 were from lung samples from saline-treated and halthromycin incircin treated entire promiled in the Control of SRD. samples from saline-treated and tulathromycin injection-treated pigs enrolled in the Control of SRD field study in the U.S. 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 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 ₅₀ ** (ug/mL)	MIC ₉₀ ** (ug/mL)	MIC range (ug/mL)	
Actinobacillus pleuropneumonia	2002-2002 2007-2008	135 88	16 16	32 16	16 to 32 4 to 32	l
Haemophilus parasuis	2000-2002	31	1	2	0.25 to > 64	ı
Pasteurella multocida	2002-2002 2007-2008	55 40	1	2 2	0.5 to > 64 ≤ 0.03 to > 2	
Bordetella bronchiseptica	2002-2002	42	4	8	2 to 8	ı

*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

Cattle BRD – In a multi-location field study, 314 calves with naturally occurring BRD were treated with tulathromycin injection. Responses to treatment were compared to saline-treated controls. A cure was defined as a calf with normal studied/activity, normal respiration, and a rectal temperature of $\leq 104^{\circ}$ fon Day 14. The cure rate was significantly higher ($f \leq 0.05$) in tulathromycin injection-treated calves (248). There were two BRD related deaths in the tulathromycin injection-treated calves (248). There were two BRD related deaths in the saline-treated calves calves.

Fifty-two tulathromycin injection-treated calves and 27 saline-treated calves from the multi-location Field BRD treatment study had Mycoplasma bowis identified in cultures from pre-treatment nasopharyngeal swabs. Of the 52 tulathromycin injection-treated calves, 37 (71.2%) calves were categorized as cures and 15 (28.8%) calves were categorized as treatment failures. Of the 27 saline ated calves, 4 (14.8%) calves were categorized as cures and 23 (85.2%) calves were treatment

an meta-analysis was conducted to compare the BRD treatment success rate in young calve A Bayesian meta-analysis was conducted to compare the BRD treatment success rate in young claves (calesweighing 250 lbs or less and fled primarily a milk based dielt pruted with tulathromycin injection to the success rate in older calves (calves weighing more than 250 lbs and fed primarily a roughage and grain-based dielt breated with tulathromycin injection. The analysis induced data from four BRD treatment effectiveness studies conducted for the approval of tulathromycin injection in the U.S. and nine contemporaneous studies conducted in Europe. The analysis showed that the BRD treatment success rate in young calves was at least as good as the BRD treatment success rate in older calves. East a result in taldbrowning in injection is; conscienced effective for the treatment of BRD. in older calves. As a result, tulathromycin injection is considered effective for the treatment of BRD associated with M. haemolytica, P. multocida, H. somni, and M. bovis in suckling calves, dairy calves, and year caives. In another multi-location field study with 399 calves at high risk of developing BRD, administration of

In another multi-location field study with 399 calves at high risk of developing BRD, administration of tulathromycin injection resulted in a significantly reduced incidence of BRD (11%) compared to saline-treated calves (59%). Effectiveness evaluation was based on scored clinical signs of normal attitude activity, normal respiration, and a retal temperature of ≤ 104°F on Day 14. There were no BRD-related deaths in the ulathromycin injection-treated calves compared to two BRD-related deaths in the saline-treated calves. Fifly saline-treated calves calves described in cultures of post treatment nasopharyngeal swabs or lurg tissue. Two induced infection model studies were conducted to confirm the effectiveness of fullathromycin injection against Mycop/sarab box's. Attoal of 164 calves were inculated intratracheally with field staries of Mycoplasma box's. When calves became pyrexic and had abnormal respiration scores, they were treated with either tulathromycin injection (25 mg/kg MV) subcutaneously or an equivalent volume of saline. Calves were observed for signs of BRD for 14 days post-treatment, then were eurhanized and necropsied. In both studies, mean jung lesion percentages were statistically significantly lower in the tulathromycin injection-treated calves compared with staller-treated calves (11.3% vs. 28.7%, P = 0.0001 and 15.0% vs. 30.7%, P < 0.0001).

(11.3% vs. 28.9%, P = 0.0001 and 15.0% vs. 30.7%, P < 0.0001).
BIK – Two field studies were conducted evaluating ulathromycin injection for the treatment of IBK associated with Monavella bows in 200 naturally-infected calves. The primary clinical endpoint of these studies was cure rate, defined as a calf with no clinical signs of IBK and no comeal ulcer, assessed on Days 5, 9, 13, 17, and 21. Time to improvement, defined as the first day on which a calf had no clinical signs of IBK in both eyes, provided that those scores were maintained at the next day of

clinical signs of IBK in both eyes, provided that those scores were maintained at the next day of obsenation, was assessed as a scorday variable. All time points, in both studies, the cure rate was significantly higher (P < 0.05) for tulathromycin injection-treated calves. Additionally, time to improvement was significantly less (P < 0.0001) in both studies for tulathromycin injection injection injection injection injection injection for the treatment of sovine foot rot was evaluated in 170 cattle in two field studies. Cattle diagnosed with bovine foot rot were enrolled and treated with a single subcutaneous dose of tulathromycin injection (2.5 mg/kg BM) or an equivalent olium of saline treatment of the saline studies. Cattle diagnosed with bovine foot rot were enrolled and treated with a single subcutaneous dose of tulathromycin injection (2.5 mg/kg BM) or an equivalent oliume of saline. Cattle were dinicially evaluated of 3/84 after treatment for treatment success, which was based on defined decreases in lesion, swelling, and lameness scores. In both studies, the treatment success perentage was scatistically significantly higher in tulathromycin injection-treated calves compared with saline-treated calves (60% vs. 8%, P < 0.0001 and 83.3% vs. 50%, P = 0.0088). Swine

In a multi-location field study to evaluate the treatment of naturally occurring SRD, 266 pigs were In a multi-location field study to evaluate the treatment of naturally occurring SRD, 266 pigs were treated with fulshromyrin injection. Responses to teatment were compared to saline-treated controls. Success was defined as a pig with normal attitude, normal respiration, and rectal temperature of < 104°F on Day 7. The treatment success rate was significantly greater (P ≤ 0.05) in tulathromyrin injection-treated pigs (70°Fs) oppraced to saline-treated pigs (46°Fs). My popneumoniae was isolated from 106 saline-treated and non-treated sentinel pigs in this study. Two induced infection model studies were conducted to confirm the effectiveness of tulathromyrin injection-agains IM. hypopneumoniae. Ind aps, after inoculation intranasally and intratancheally with a field starion of My popneumoniae. 144 pigs were treated with either tulathromyrin injection 12 5 mg/kg BW) intramuscularly or an equivalent volume of saline. Pigs were euthanized and necopsied 10 days post-treatment. The mean percentage of gross preumonic lung lesions was statistically significantly lower (P < 0.0001) for fulathromyrin injection-treated pigs than for saline treated pigs in the for saline treated pigs in injection of the control of SRD was evaluated in a multi-location natural infection field study. When at least 15% of the study candidates showed clinical signs of SRD, all pigs were emolled and treated with fulsathromyrin injection (22° pigs) osaline (22° pigs) osaline (22° pigs).

SRD, all pigs were enrolled and treated with tulathromycin injection (226 pigs) or saline (227 pigs). Responses to treatment were evaluated on Day 7. Success was defined as a pig with normal attitude normal respiration, and rectal temperature of < 104 °E. The treatment success rate was significantly greater (P < 0.05) in tulathromycin injection-treated pigs compared to saline-treated pigs (59.2%)

ANIMAL SAFETY

Cattle Safety studies were conducted in feeder calves receiving a single subcutaneous dose of 25 mg/kg BW or 3 weekly subcutaneous doses of 25, 7.5, or 12.5 mg/kg BW. In all groups, transient indication of pain after injection were seen, including head shaking and pawing at the ground. Injection ste swelling, discoloration of the subcutaneous tissues at the injection site and corresponding histopathologic changes were seen in animals in all dosage groups. These lesions showed signs of resolving over time. No other drug-related lesions were observed microsopically or incircosopically. An exploratory study was conducted in feeder calves receiving a single subcutaneous dose of 10, 125, or 15 mg/kg BW. Macrosopically, no lesions were observed. Microsopically minimal to mild myocardial degeneration was seen in one of six calves administered 12.5 mg/kg BW and two of six calves administered 12.5 mg/kg BW and two of six calves administered 15 mg/kg BW.

calves administered 13 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 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.

Swine Safety studies were conducted in pigs receiving a single intramuscular dose of 25 mg/kg BW, or 3 weekly intramuscular doses of 2.5, 7.5, or 12.5 mg/kg BW. In all groups, transient indications of pain after injection were seen, including neaflessness and excessive vocalization. Temos occurred briefly in one animal receiving 7.5 mg/kg BW. Discoloration and edema of injection site tissues and corresponding histopathologic changes were seen in animals at all dosages and resolved over time. No other drug-related lesions were observed macroscopically or microscopically.

tore below 30°C (86°F), with excursions up to 40°C (104°F). Use this product within 84 days of the Some Deums 30 Clop 7, with exclusions up to 40 C (104 + 1) see this product within 64 bays of the first puncture and puncture a maximum of 20 times. If more than 20 punctures are anticipated, the use of automatic injection equipment or a repeater syringe is recommended. When using a draw-of spike or needle with bowe diameter larger than 16 gauge, discard any product remaining in the vial immediately after use.

HOW SUPPLIED VYN Injectable Solution is available in the following package sizes

STORAGE CONDITIONS

500 mL vial Approved by FDA under ANADA # 200-715 Tulathromycin (active ingred.) made in China. Formulated in Germany.

Intervet Inc. (d/b/a Merck Animal Health), Madison, NJ 07940 menera, include merck united readint, Madiston, RUD/Y940 To report suspected adverse events, for technical assistance or to obtain a copy of the Safety Data Sheet (SDS), contact Merck Animal Health at 1-800-211-3573. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA VETS or online at http://www.rub.nc/







BANAMINE®-S

VIXIN MEGLUMINE INJECTION (50 MG/ML))

For intramuscular use in swine.

Veterinary Not for use in breeding swine.

NADA #101-479, Approved by FDA.

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

DESCRIPTION: Each milliliter of BANAMINE-S (flunixin meglumine injection) contains 50 mg flunixin (equivalent to 83 mg flunixin meglumine), 0.1 mg edetate disodium, 2.5 mg sodium formaldehyde sulfoxylate, 4.0 mg diethanolamine, 207.2 mg propylene glycol; 5.0 mg phenol as preservative, hydrochloric acid, water for injection a.s.

CLINICAL PHARMACOLOGY: Flunixin meglumine is a potent non-narcotic, nonsteroidal, analgesic agent with anti-inflammatory and antipyretic activity. It is significantly more potent than pentazocine, meperidine, and codeine as an analgesic in the rat yeast paw test.

Flunixin is known to persist in inflammatory tissues¹ and is associated with anti-inflammatory properties which extend well beyond the period associated with detectable plasma drug concentrations². Therefore, prediction of drug concentrations based upon estimated plasma terminal elimination half-life will likely underestimate both the duration of drug action and the concentration of drug remaining at the site of activity.

The pharmacokinetic profiles were found to follow a 2-compartmental model, although a deep (third) compartment was observed in some animals. The mean terminal elimination half-life (ß half-life) of flunixin after a single intramuscular injection of Banamine (2.2 mg/kg) to pigs was between 3 and 4 hours. The mean observed maximum plasma concentration was 2944 ng/mL, achieved at a mean time of approximately 0.4 hours. The mean AUC, was 6431 ng*hr/mL. Following IM administration of flunixin, quantifiable drug concentration could be measured up to 18 hours post dose. The mean volume of distribution was 2003 mL kg and the mean total clearance was 390 mL/hr/kg. The mean absolute bioavailability of flunixin following an intramuscular injection in the neck was 87%.

 $\textbf{INDICATION:} \ \textbf{BANAMINE-S (flunixin meglumine injection) is indicated for the control of pyrexia associated$ with swine respiratory disease

DOSE AND ADMINISTRATION: The recommended dose for swine is 2.2 mg/kg (1 mg/lb; 2 mL per 100 lbs) body weight given by a single intramuscular administration. The injection should be given only in the neck musculature with a maximum of 10 mL per site.

USE WITHIN 28 DAYS OF FIRST PUNCTURE AND PUNCTURE A MAXIMUM OF 10 TIMES. WHEN USING A DRAW-OFF SPIKE OR NEEDLE WITH BORE DIAMETER LARGER THAN 18 GAUGE, DISCARD ANY PRODUCT REMAINING IN THE VIAL IMMEDIATELY AFTER USE.

Note: Intramuscular injection may cause local tissue irritation and damage. In an injection-site irritation study, the tissue damage did not resolve in all animals by Day 28 post-injection. This may result in trim loss of edible tissue at slaughter

CONTRAINDICATIONS: There are no known contraindications to this drug in swine when used as directed. Do not use in animals showing hypersensitivity to flunixin meglumine. Use judiciously when renal impairment or gastric ulceration is suspected.



RESIDUE WARNINGS: SWINE MUST NOT BE SLAUGHTERED FOR HUMAN CONSUMPTION WITHIN 12 DAYS OF THE LAST TREATMENT.



PRECAUTIONS: As a class, cyclo-oxygenase inhibitory NSAIDs may be associated with gastrointestinal, renal and hepatic toxicity. Sensitivity to drug-associated adverse events varies with the individual patient. Patients at greatest risk for adverse events are those that are dehydrated, on concomitant diuretic therapy, or those with existing renal, cardiovascular and/or henatic dysfunction. Concurrent administration of potentially nephrotoxic drugs should be carefully approached. NSAIDs may inhibit the prostaglandins that maintain normal homeostatic function. Such prostaglandin effects may result in clinically significant disease in patients with underlying or pre-existing disease that has not been previously diagnosed

Since many NSAIDs possess the potential to produce gastrointestinal ulceration, concomitant use of flunixin meglumine with other anti-inflammatory drugs, such as other NSAIDs and corticosteroids, should be avoided.

Not for use in breeding swine. The reproductive effects of BANAMINE-S (flunixin meglumine injection) have

Intramuscular injection may cause local tissue irritation and damage. In an injection site irritation study, the tissue damage did not resolve in all animals by Day 28 post-injection. This may result in trim loss of edible tissue at slaughter.

ADVERSE REACTIONS: Flunixin was mildly irritating at the injection sites. No other flunixin-related changes (adverse reactions) were noted in swine administered a 1X (2.2 mg/kg; 1.0 mg/lb) dose for 9 days.

ANIMAL SAFETY: Minimal toxicity manifested itself as statistically significant increased spleen weight at elevated doses (5X or higher daily for 9 days) with no change in normal microscopic architecture.

HOW SUPPLIED: BANAMINE-S (flunixin meglumine injection), 50 mg/mL is available in 100-mL (NDC #

Store at or below 25°C (77°F). Do not freeze.

See the In-Use statement as provided in the Dose and Administration section.

REFERENCES:

- 1. Lees P, Higgins AJ. Flunixin inhibits prostaglandin E, production in equine inflammation. Res Vet Sci. 1984;
- 2. Oldensvik K. Pharmacokinetics of flunixin and its effect on prostaglandin F₂₀ alpha metabolite concentrations after oral and intravenous administration in heifers. J Vet Pharmacol Ther. 1995; 18:254-

Distributed by: Intervet Inc. d/b/a Merck Animal Health Madison, NJ 07940

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MATRIX®

(ALTRENOGEST

Oral Solution, 2.2 mg altrenogest per ml (0.22%)

FOR USE IN SWINE ONLY

- Before using this drug, read package insert for complete product Information
 Always use tile MATRIX® Dosing Device to administer this product.

It is a violation of Federal law to use this drug product other than as directed in the labeling or as directed by vour veterinarian

WARNINGS

Withdrawal Periods



Animals intended for human consumption must not be slaughtered within 21 days of the last treatment with this drug product.

User Safety Warnings

Not for use in humans. Keep out of reach of children. Skin contact must be avoided as MATRIX® is readily absorbed through unbroken skin, and exposure may result in serious side effects to both men and women. Always wear vinyl, neoprene, or nitrile protective gloves when handling this product or when in contact with equipment or surfaces contaminated by this product. This product can penetrate latex or other types of porous gloves. Latex gloves are not protective. Read all labeling prior to use.

PREGNANT WOMEN OR WOMEN WHO MAY BE PREGNANT SHOULD NOT HANDLE MATRIX® enogest). WOMEN OF CHILDBEARING AGE SHOULD EXERCISE EXTREME CAUTION WHEN

Accidental absorption, such as absorption through the skin, could lead to a disruption of the menstrual cycle equipment or surfaces that come in contact with MATRIX® should be adequately cleaned and nated to prevent human exposure.

Always use the MATRIX® Dosing Device to administer this product. The MATRIX® bottle is designed only tor use with the MATRIX® Dosing Device. Use without the device increases the risk of human exposure.

People who should not handle MATRIX®:*

- Women who are or may be pregnant.
- 2. Anyone with blood clots or clotting disorders, or with a history of these events.
 3. Anyone with a history of heart disease or stroke.
- 4. Women with known or suspected breast cancer.
- 5. People with known or suspected estrogen-dependent cancer. Women with vaginal bleeding of unknown cause.
- 7. People with tumors which developed during the use of oral contraceptives or other
- estrogen-containing products. 8. Anyone with liver dysfunction or disease.
- *Based on known effects of long-term progestin use in humans.

Accidental exposure

MATRIX® is readily absorbed from contact with the skin. In addition, this oil-based product can penetrate latex or other types of porous gloves. Always wear vinyl, neoprene, or nitrile protective gloves when handling this product. Latex gloves are not protective. If MATRIX® gets inside gloves by damage or spilling, the covered skin may absorb more of the drug. Side effects after a single exposure may be possible; however, continued daily exposure has the potential for more serious side effects.

In case of accidental exposure Skin Exposure and/or clothing contamination: Wash skin immediately with soap and water, and launder clothing with detergent

Eye Exposure: Immediately flush with plenty of water for 15 minutes. Get medical attention. If wearing tact lenses, flush eyes immediately with water before removing lenses. If Swallowed: On or induce vomiting. Seek medical attention immediately. MATRIX® contains an oil.

Vomiting should be supervised by a physician because of possible pulmonary damage via aspiration of the oil

base. If possible, bring the labeling to the physician. Reported HUMAN effects from exposure

Side effects have been reported in women and men following accidental exposure to altrenogest products,

- including MATRIX®, either through handling of the product or contact with contaminated surfaces.
- Reproductive side effects reported in women included abnormal or absent menstrual cycles.
 Reproductive side effects reported in men included decreased libido.
- Other side effects reported in women and men included headaches, fever, abdominal pain, nausea, diarrhea, vomiting, and rashes.

Storage, Handling, and Disposal

Store MATRIX® solution bottle and Dosing Device when loaded with solution for continued use at or below room temperature, 77°F (25°C). Close tightly. Refer to the MATRIX® Dosing Device label for equipment **cleaning instructions.** Place empty drug containers, waste from rinsing tile Dosing Device, protective gloves, or other articles that contact this product in a leak-resistant container tor disposal in accordance with applicable Federal, state, and local regulations.

Questions? Comments?

- To report side effects, contact Merck at 1-800-211-3573, or www.merck-animal-health-usa.com
- To obtain product information, including safety data sheet (SDS), call 1-800-441-8272
- For additional information about reporting side effects for animal drugs, contact FDA at 1-888-FDA-VETS or online at: www.fda.gov/reportanimalae



Net Contents: 6 x 1.000 ml (33.8 fl. oz.) bottles

Manufactured for: Intervet Inc. (d/b/a Merck Animal Health), Madison, NJ 07940 Made in France

Approved by FDA under NADA # 141-222



Rev. 8/2022

PRODUCT

Approved by FDA under NADA # 141-063

Nuflor®-S

Injectable Solution 300 mg/mL

For intramuscular use in swine except for nursing piglets and swine of reproductive age intended for breeding.

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

DESCRIPTION: Nuflor®-S Injectable Solution is a sterile solution of the synthetic, broad-spectrum antibiotic florfenicol, Each milliliter of sterile Nuflor®-S Injectable Solution contains 300 mg of florfenicol, 250 mg N-methyl-2-pyrrolidone (NMP), 150 mg propylene glycol and polyethylene glycol a.s.

INDICATIONS: Nuflor®-S Injectable Solution is indicated for treatment of swine respiratory disease associated with Actinobacillus pleuropneumoniae. Pasteurella multocida, Salmonella Choleraesuis, Streptococcus suis, Bordetella bronchiseptica, and Glaesserella (Haemophilus) parasuis in swine except for nursing piglets and swine of reproductive age intended for

DOSAGE AND ADMINISTRATION: Nuflor®-S Injectable Solution should be administered by intramuscular injection to swine at a dose rate of 15 mg/kg (1 mL/45 lb) body weight. A second dose should be administered 48 hours later. The injection should be given only in the neck musculature. If a positive response is not noted within 24 hours after the second injection, the diagnosis should be re-evaluated, and/or an alternative treatment may be considered. Administered dose volume should not exceed 10 mL per

Nuflor®-S	DOSAGE GUIDE FOR SWINE
ANIMAL	IM Nuflor®-S DOSAGE
WEIGHT (lbs)	(1 mL/ 45 lb Body Weight)
	(mL)
22	0.5
45	1
90	2
135	3
180	4
225	5
270	6

WARNINGS: NOT FOR HUMAN USE. KEEP OUT OF REACH OF CHILDREN. This product contains materials that can be irritating to skin and eyes. Avoid direct contact with skin, eyes and clothing. In case of accidental eye exposure, flush with water for 15 minutes. In case of accidental skin exposure, wash with soap and water. Remove contaminated clothing Consult a physician if irritation persists. Accidental injection of this product may cause local irritation Consult a physician immediately. Reproductive and developmental toxicities have been reported in laboratory animals following high, repeated exposures to NMP. Pregnant women should wear gloves and exercise caution or avoid handling this product. The Safety Data Sheet (SDS) contains more detailed occupational safety information

For customer service, adverse effects reporting and/or a copy of the SDS, call 1-800-211-3573. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS or online at www.fda.gov/reportanimalae

PRECAUTIONS: Not for use in animals intended for breeding purposes. The effects of florfenicol on porcine reproductive performance, pregnancy and lactation have not been determined.

Intramuscular injection in swine may result in local tissue reaction which could persist up to 21 days post-dosing. This may result in trim loss of edible tissue at slaughte

RESIDUE WARNINGS: Swine intended for human consumption must not be slaughtered within 11 days of the last intramuscular treatment.

ADVERSE REACTIONS: Perianal inflammation, rectal eversion, rectal prolapse and diarrhea may occur transiently following treatment. Decreased feed and water consumption may occur if the labeled dosage

CLINICAL PHARMACOLOGY: The pharmacokinetic disposition of florfenicol was evaluated in 20 pigs following a single IM injection of Nuflor®-S at the labeled dose of 15 mg/kg BW. The mean ± standard deviation maximum plasma concentration (C_{max}) and the time to reach C_{max} (T_{max}) of florfenicol were 3.42 \pm 0.82 $\mu g/mL$ and 4.70 ± 2.15 hours, respectively. The mean ± standard deviation area under the drug concentration-time curve between times 0 and the last quantifiable concentration (AUC_{0.100}) and the terminal half-life (T_{1/2}) of florfenicol were $70.34 \pm 23.78 \, \mu g^*$ hours/mL and $11.21 \pm 3.73 \, hours$,

MICROBIOLOGY: Florfenicol is a synthetic, broad-spectrum antibiotic active against many Gram-negative and Gram-positive bacteria isolated from domestic animals. It acts by binding to the 50S ribosomal subunit and inhibiting bacterial protein synthesis. Florfenicol is generally considered a

bacteriostatic drug, but exhibits bactericidal activity against certain bacterial species.

In vitro activity of florfenicol has been demonstrated against commonly isolated pathogens associated with swine respiratory disease. Isolates tested were obtained from pre-treatment lung samples from representative non-enrolled pigs at each study site and post-treatment lung samples from pigs in the florfenicol-treated and saline-treated groups that died or were euthanized during the study, or were classified as treatment failures at the end of the study. The minimum inhibitory concentrations (MICs) of florfenicol for swine respiratory pathogens from clinical studies were determined using dilution methods. These susceptibility test methods were adequately controlled with the inclusion and acceptable performance of appropriate reference strains. The results are presented in Table 1.

Table 1. Florfenicol minimum inhibitory concentration (MIC) values* for indicated target pathogens isolated from a multi-site field study evaluating swine respiratory disease in the U.S. in 2001.

Indicated pathogens	Number of Isolates	MIC ₅₀ ** (μg/mL)	MIC ₉₀ ** (μg/mL)	MIC Range (µg/mL)
Actinobacillus pleuropneumoniae	100	0.25	0.5	0.25-1
Pasteurella multocida	107	0.5	0.5	0.25-0.5
Bordetella bronchiseptica	49	2	2	0.5-4
Glaesserella parasuis	36	0.5	0.5	≤0.12-1.0
Streptococcus suis	62	2	2	1-2
Salmonella Choleraesuis	36	4	4	2-4

* The correlation between in vitro susceptibility data and the clinical effectiveness of florfenicol is unknown.

**The lowest MIC to encompass 50% and 90% of the most susceptible isolates, respectively

EFFECTIVENESS: In a multi-site natural infection field study, a total of 620 growing pigs with clinical signs of SRD (rectal temperature of ≥ 104.5°F, and a depression score (on a scale of 0 [absent] to 3 [severe]) of ≥ 2, and a dyspnea score (on a scale of 0 [absent] to 3 [severe]) of ≥ 2) were treated with either florfenicol (15 mg/kg BW IM given on Days 0 and 2) or an equivalent volume of saline. reatment success (rectal temperature of < 104°F, and a depression score of 0 or 1, and a dyspnea score of 0 or 1) was evaluated on Day 6. The treatment success rate was statistically significantly different (p < 0.0001) and higher in the florfenicol-treated group (72%) than in the saline-treated control group (33.1%).

361060 R1

ANIMAL SAFETY: A safety study was conducted in 40 healthy crossbred growing pigs. Pigs were administered florfenicol by IM injection in the neck at 1X, 3X, or 5X the labeled dose (15, 45, or 75 mg/kg BW, respectively) for 3X the labeled duration of treatment (6 injections at 48-hour intervals), or 10X the labeled dose (150 mg/kg BW) administered as two injections 48 hours apart. Test article-related diarrhea (moderate), anal swelling/erythema (mild to moderate), and injection site swelling (mild to moderate) were seen in all florfenicol-treated groups after dosing, most frequently in the 3X and 5X groups. Although these findings were considered clinically relevant, the incidence and severity in the 1X group was considered within acceptable limits. Test article-related decreases in feed and water consumption and an associated decrease in body weight were seen in the 3X and 5X groups. Test article-related changes in some serum chemistry parameters and decreased numbers of white blood cells were seen in the 3X, 5X, and/or 10X groups; the changes were generally minimal and not considered clinically significant. Most changes in drug-related, in-life parameters did not become apparent until after dosing was extended beyond the labeled duration of two injections, 48 hours apart.

Injection site irritation was evaluated in a safety study using 20 healthy crossbred growing pigs administered florfenicol at 15 mg/kg BW IM in the neck as two injections 48 hours apart. Mild injection site swelling was seen in up to approximately 32% of the pigs by 4 days post-injection and was resolved by 16 days post-injection. Gross and histopathologic evaluation showed that injection site discoloration and inflammation was present at 7 and 14 days post-injection, and absent at 21, 28, and 42 days post-injection.

STORAGE CONDITIONS: Store between 2-30°C (36-86°F). Do not store above 30°C (86°F). Protect from light when not in use. Use within 30 days of first puncture and puncture a maximum of 30 times. If more than 30 punctures are anticipated, the use of multidosing equipment is recommended. When using a draw-off spike or needle with hore diameter larger than 18 gauge, discard any product remaining in the vial immediately after use.

HOW SUPPLIED: Nuflor®-S (florfenicol) Injectable Solution is packaged in 100 mL glass sterile multiple dose vials.

Approved by FDA under NADA # 141-063

Florfenicol (active ingred.) made in China. Formulated in Germany. Copyright © 2021, Intervet Inc., a subsidiary of Merck & Co. Inc. Madison, NJ 07940 All rights reserved. Rev. 09/2021



398967 R2 INU240 03

P.G. 600[®]

(serum gonadotropin and chorionic gonadotropin for injection)

Approved by FDA under NADA # 140-856

FOR ANIMAL USE ONLY

DESCRIPTION

Gilts normally reach puberty (begin experiencing normal estrous cycles and exhibiting regular estrus or heat) at any time between six and eight months of age, although some gilts will not have exhibited their first estrus at ten months of age. Age at first estrus is influenced by several factors including breed type, season of the year, environmental conditions, and management practice (Hurtgen, 1986).

Sows normally exhibit estrus three to seven days after weaning their litters; however, some otherwise healthy sows may not exhibit estrus for 30 days or more after weaning (Dial and Britt, 1986). The causes of delayed return to estrus in healthy sows are poorly understood, but probably include season of the year (so-called seasonal anestrus; Hurtgen, 1979), adverse environmental conditions, such as high ambient temperatures (Love, 1978), and the number of previous litters, because the condition is more prevalent after the first litter than after later litters (Hurtgen, 1986).

P.G. 600 (serum gonadotropin and chorionic gonadotropin for injection) is a combination of serum gonadotropin (Pregnant Mare Serum Gonadotropin or PMSG) and chorionic gonadotropin (Human Chorionic Gonadotropin or HCG) for use in prepuberal gilts (gilts that have not yet exhibited their first estrus) and in sows at weaning. It is supplied in freeze-dried form with sterile diluent for reconstitution.

In gilts and sows, the action of serum gonadotropin is similar to the action of Follicle-Stimulating Hormone (FSH), which is produced by the animals' anterior pituitary gland. It stimulates the follicles of the ovaries to produce mature ova (eggs), and it promotes the outward signs of estrus (heat).

The action of chorionic gonadotropin in gilts and sows is similar to the action of Luteinizing Hormone (LH), which is also produced by the animals' anterior pituitary gland. It causes the release of mature ova from the follicles of the ovaries (ovulation), and it promotes the formation of corpora lutea, which are necessary for the maintenance of pregnancy once the animals have become pregnant.

The combination of serum gonadotropin and chorionic gonadotropin in P.G. 600 induces fertile estrus in most prepuberal gilts and weaned sows three to seven days after administration (Schilling and Cerne, 1972; Britt et al., 1986; Bates et al., 1991). The animals may then be mated or, in the case of gilts, mating may be delayed until the second estrus after treatment.

NOTE: P.G. 600 IS INTENDED AS A MANAGEMENT TOOL TO IMPROVE REPRODUCTIVE EFFICIENCY IN SWINE NOTE: F.G. 50013 INTERIORS A AMANAGEMENT TOUL OF WITHOUT REPRODUCT, ESTRUS DETECTION AND OTHER ASPECTS OF REPRODUCTIVE MANAGEMENT MUST BE ADEQUATE. IF YOU ARE IN DOUBT ABOUT THE ADEQUACY OF YOUR BREEDING PROGRAM, CONSULT YOUR VETERINARIAN.

PG. 600 is available

FIVE DOSE VIALS (order Code No. PG-720-5) - One vial containing white freeze-dried powder, and one vial containing sterile diluent. When reconstituted, the five dose vial (25 mL) of P.G. 600 contains

SERUM GONADOTROPIN (PMSG) CHRIONIC GONADOTROPIN (HCG) 1000 II I (equivalent to 1000 USP Units chorionic gonadotropin)

INDICATIONS FOR USE

PREPUBERAL GILTS: P.G. 600 is indicated for induction of fertile estrus (heat) in healthy prepuberal (non-cycling) gilts over five and one-half months of age and weighing at least 85 kg (187 lb.).

SOWS AT WEANING: P.G. 600 is indicated for induction of estrus in healthy weaned sows experiencing delayed

CALITIONS

Treatment will not induce estrus in gilts that have already reached puberty (begun to cycle). Gilts that are less than five and one-half months of age or that weigh less than 85 kg (187 lb.) may not be mature enough to continue normal estrus cycles or maintain a normal pregnancy to full term after treatment.

Treatment will not induce estrus in sows that are returning to estrus normally three to seven days after weaning. Delayed return to estrus is most prevalent after the first litter; the effectiveness of P.G. 600 has not been established after later litters. Delayed return to estrus often occurs during periods of adverse environmental conditions, and sows mated under such conditions may farrow smaller than normal litters

DOSAGE AND ADMINISTRATION

One dose (5 mL) of reconstituted P.G. 600, containing 400 IU serum gonadotropin (PMSG) and 200 IU chorionic gonadotropin (HCG), should be injected into the gilt or sow's neck behind the ear.

Prepuberal gilts should be injected when they are selected for addition to the breeding herd. Sows should be injected at weaning during periods of delayed return to estrus.

DIRECTIONS FOR USE

FIVE DOSE VIAL: Using a sterile syringe and a sterile 0.90 x 38 mm (20 G x 1½") hypodermic needle, transfer approximately 5 mL of the sterile diluent into the vial of freeze-dried powder. Shake gently to dissolve the powder. Transfer the dissolved product back into the vial of diluent and shake gently to mix. Inject one dose (5 mL) of the reconstituted solution into the gilt's or sow's neck subcutaneously behind the ear.

STORAGE PRECAUTIONS

Store at 36-46°F (2-8°C).

Once reconstituted, P.G. 600 should be used immediately. Unused solution should be disposed of properly and not

Spent hypodermic needles and syringes generated as a result of the use of this product must be disposed of properly in accordance with all applicable Federal, State and local regulations.

Bates, R.O., B.N. Day, J.H. Britt, L.K. Clark and M.A. Brauer (1991). Reproductive performance of sows treated with a combination of Pregnant Mare's Serum Gonadotropin and Human Chorionic Gonadotropin at weaning in the summer. Journal of Animal Science 69:894.

Britt, J.H., B.N. Day, S.K. Webel and M.A. Brauer (1989). Induction of fertile estrus in prepuberal gilts by treatment with a combination of Pregnant Mare's Serum Gonadotropin and Human Chorionic Gonadotropin. Journal of Animal Science 67:1148.

Dial, G.D., and J.H. Britt (1986). The clinical endocrinology of reproduction in the pig. In: D.A. Morrow (ed.). Current Therapy in Theriogenology 2. W.B. Sanders Company, Philadelphia. p. 905.

Hurtgen, J.P. (1979). Seasonal breeding patterns in female swine. Ph.D. Dissertation, University of Minnesota. Hurtgen, J.P. (1986). Noninfectious infertility in swine. In: D.A. Morrow (ed.) Current Therapy in Theriogenology 2.

W.B. Sanders Company, Philadelphia. p. 962. Love, R.J. (1978). Definition of a seasonal infertility problem in pigs. Veterinary Record 103:443.

Schilling, E., and F. Cerne (1972). Induction and synchronization of oestrus in prepubertal gilts and anoestrus sows

by a PMS/HCG-compound. Veterinary Record 91:471.

Manufactured for: Intervet Inc., a subsidiary of Merck & Co. Inc. Madison, NJ, 07940 USA

INTERVET INTERNATIONAL GmbH UNTERSCHLEISSHEIM, GERMANY

Rev. 10/2022

SAFE-GUARD® (fenbendazole) DEWORMER

MUST BE MIXED BEFORE FEEDING ACCORDING TO DIRECTIONS

AND PERMITTED CLAIMS.

FOR USE IN MANUFACTURED FEEDS ONLY.

ACTIVE DRUG INGREDIENT: Fenbendazole 200 grams per kilogram (90.7 grams per pound).

INERT INGREDIENTS: Roughage Products or Roughage Products and Calcium Carbonate: and Mineral Oil.

FOR THE TREATMENT AND CONTROL OF:

Lunaworms: Adult *Metastrongylus apri*, adult *Metastrongylus pudendotectus*, Gastrointestinal worms: Adult and larvae (L3, L4 stages, liver, lung, intestinal forms) large roundworms (Ascaris suum), Adult nodular worms (Oesophagostomum dentatum, O. quadrispinulatum), Adult small stomach

worms (*Hyostrongylus rubidus*), Adult and larvae (L2, L3, L4 stages - intestinal mucosal forms) whipworms (*Trichuris suis*); and **Kidney worms:** Adult and larvae Stephanurus dentatus.

DRUG FEEDING RATE:

9 mg fenbendazole per kg body weight (4.08 mg fenbendazole per pound) to be fed as the sole ration over a period of 3 to 12 days.

MIXING DIRECTIONS:

Thoroughly mix SAFE-GUARD® 20% Type A medicated article with non-medicated swine feed according to the table below to obtain the proper concentration in the Type B medicated feed. The following table gives examples of how some Type B medicated feed concentrations can be prepared:

Swine Type B Medicated Feed Instructions				
Pounds of Type A Medicated	Resulting Fenbendazole			
Article to Add per Ton of Feed	Concentration in Type B			
to Make a Type B Medicated	Medicated Feed [grams/ton			
Feed	(grams/pound)]			
11.03	1,000 (0.5)			
195.59	17,740 (8.9)			

Thoroughly mix SAFE-GUARD® 20% Type A medicated article with non-medicated swine feed according to the table below to obtain the proper concentration in the complete Type C medicated feed. Prepare an intermediate pre-blend of the Type A medicated article prior to mixing in a complete feed. Thoroughly mix the required amount of Type A medicated article in a convenient quantity of feed ingredients (a dilution of one part Type A medicated article and nine parts grain carrier is suggested), then thoroughly mix this pre-blend with the rest of the feed ingredients to ensure complete and uniform distribution of the Type A medicated article.

The following table gives examples of how some complete Type C medicated feeds can be prepared:

Swine Type C Medicated Feed Instructions					
Pounds of Type A Medicated Article to Add per Ton to Make a Type C Medicated Feed	Resulting Fenbendazole Concentration in Type C Medicated Feed [grams/ton (grams/pound)]				
0.11	10 (0.005)				
3.31	300 (0.15)				

FEEDING DIRECTIONS:

Feed as the sole ration for three (3) to twelve (12) consecutive days. No prior withdrawal of feed or water necessary.

Type C medicated swine feeds containing SAFE-GUARD® 20% can be fed nelleted or as a meal

WARNING: KEEP THIS AND ALL DRUGS OUT OF THE REACH OF CHILDREN. NOT FOR USE IN HUMANS. The Safety Data Sheet (SDS) contains more detailed occupational safety information. For customer service, adverse effects reporting, and/or a copy of the SDS, call 1-800-211-3573. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDAVETS, or http://www.fda.gov/reportanimalae.

OTHER WARNINGS: Parasite resistance may develop to any dewormer. All dewormers require accurate dosing for best results. Following the use of any dewormer, effectiveness of treatment should be monitored. A decrease of effectiveness over time may indicate the development of resistance to the dewormer administered. The parasite management plan should be adjusted accordingly based on regular monitoring.



Withdrawal Periods: Swine must not be slaughtered for human consumption within 4 days following last treatment with this drug

CONSULT YOUR VETERINARIAN FOR ASSISTANCE IN THE DIAGNOSIS, TREATMENT, AND CONTROL OF PARASITISM. STORE AT OR BELOW 25°C (77°F). EXCURSIONS UP TO 40°C (104°F) ARE PERMITTED.

SAFE-GUARD® AQUASOL

SUSPENSION CONCENTRATE, ANTIPARASITIC

APPROVED BY FDA UNDER NADA # 141-449

200 MG FENBENDAZOLE/ML

For oral administration via drinking water

DESCRIPTION: Safe-Guard® AquaSol is a suspension concentrate containing fenbendazole, an antiparasitic. Each mL of Safe-Guard® AquaSol contains 200 mg of fenbendazole.

Chickens: Safe-Guard® AguaSol is indicated for the treatment and control of adult *Ascaridia galli* in broiler chickens and replacement chickens and for the treatment and control of adult A. galli and Heterakis gallinarum in breeding chickens and laying hens.

Swine: Safe-Guard® AquaSol is indicated for swine, except for nursing piglets, for the treatment and control of: **Lungworms:** Adult *Metastrongylus apri*, Adult *Metastrongylus pudendotectus*, **Gastrointestinal** worms: Adult and larvae (L3, L4 stages, liver, lung, intestinal forms) large roundworms (Ascaris suum); Adult nodular worms (*Desophagostomum dentatum*, *O. quadrispinulatum*); Adult small stomach worms (*Hyostrongylus rubidus*); Adult and larvae (L2, L3, L4 stages - intestinal mucosal forms) whipworms (*Trichuris* suis), and Kidney worms: Adult and larvae Stephanurus dentatus.

DOSAGE AND ADMINISTRATION:

Chickens: Safe-Guard® AquaSol must be administered orally to chickens via the drinking water at a daily dose of 1 mg/kg BW (0.454 mg/lb) for 5 consecutive days.

Swine: Safe-Guard® AquaSol must be administered orally to swine via the drinking water at a daily dose of 2.2 mg/kg BW (1 mg/lb) for 3 consecutive days.

Consult your veterinarian for assistance in the diagnosis, treatment, and control of parasitism.

GENERAL MIXING DIRECTIONS:

Dose calculation

The daily dose of 1 mg fenbendazole per kg BW (0.454 mg/lb) is equivalent to 0.005 mL Safe-Guard® AquaSol per kg BW (0.00227 mL/lb). The required daily volume of product is calculated from the total estimated body weight [kg] of the entire group of chickens to be treated. Please use the following formula:

Total estimated body weight [kg] of the chickens to be treated x 0.005 mL = mL Safe-Guard® AquaSol/day

Examples

Total body weight of birds to be treated	Volume of Safe-Guard® AquaSol per day.	Volume of Safe-Guard® AquaSol (for 5 days)
5,000 kg (11,000 lb)	25 mL	5 x 25 mL = 125 mL
10,000 kg (22,000 lb)	50 mL	5 x 50 mL = 250 mL
80,000 kg (176,000 lb)	400 mL	5 x 400 mL = 2,000 mL
320,000 kg (704,000 lb)	1,600 mL	5 x 1,600 mL = 8,000 mL

Follow the instructions in the order described below to prepare the medicated water. The medicated water must be prepared daily prior to each administration

Dase calculation.

The daily dose is 2.2mg fenbendazole per kg BW (1 mg/lb) which is equivalent to 0.011mL Safe-Guard®

AquaSol per kg BW (0.0045 mL/lb). The required daily volume of product is calculated from the total estimated body weight [kg] of the entire group of pigs to be treated. Please use the following formula:

Total estimated body weight [kg] of the pigs to be treated x 0.011mL = mL Safe-Guard® AguaSol/day

Examples

Total body weight of pigs to be treated	Volume of Safe-Guard® AquaSol per day.	Volume of Safe-Guard® AquaSol (for 3 days)
10,000kg (22,000 lb)	110 mL	3 x 110mL = 330 mL
80,000kg (176,000 lb)	880 mL	3 x 880mL = 2640 mL
320,000 kg (704,000 lb)	3520 mL	3 x 3520mL = 10,560 mL

Follow the instructions in the order described below to prepare the medicated water. The medicated water must be prepared daily prior to each administration

Prepare a 1 to 1 dilution (pre-dilution) of Safe-Guard AquaSol in water: 1) Calculate the volume of Safe-Guard® AquaSol to be administered dails

2) Select a measuring device capable of accurately measuring a volume of at least twice the calculated Safe-Guard® AguaSol daily volume

[Note: If the total volume of the 1 to 1 dilution needed exceeds the volume of the largest available measuring device, divide the total volume into two or more smaller batches of 1 to 1 dilution, prepared following the steps below. Safe-Guard® AquaSol should always be measured by adding it to a measuring device that already contains an equivalent volume of water.

3) Pour a volume of water equal to the calculated volume of product needed into the measuring device.

Shake the product well before mixing.

5) Fill up the measuring device containing the water with the calculated volume of the product to obtain the

[Note: If more than the required amount of the product is accidently poured into the measuring device, discard the entire contents and repeat the process from Step 3 above.] 6) Add the 1 to 1 dilution of Safe-Guard® AquaSol in water to the water supply system as described below. Be careful to avoid any accidental spill or loss of 1 to 1 dilution which may inadvertently result in less than the

7) Rinse the container used to prepare the 1 to 1 dilution of Safe-Guard® AquaSol with additional water, and add the rinse water to the medicated water tank or the stock suspension tank of the dosing pump.

For use with a medication tank:

Add the entire 1 to 1 dilution of Safe-Guard® AquaSol in water to the medication tank containing the volume of drinking water usually consumed by the animals in 3 to 24 hours. Stir the medicated water in the medication tank until the medicated water is visibly homogeneous. The medicated water should appear hazy. No further stirring during administration is necessary

Add the entire 1 to 1 dilution of Safe-Guard® AquaSol in water to the water in the stock suspension tank of the dosing pump. The volume of water in the stock suspension container has to be calculated taking as a basis the present injection rate of the dosing pump and the volume of drinking water usually consumed by the animals over a period of 3 to 24 hours. Stir until the content in the stock suspension tank is visibly homogeneous. The medicated water should appear hazy.

At concentrations of up to 5 mL/L stock suspension (1 g fenbendazole/L) no stirring is required. At concentrations from 5 mL up to 75 mL of product /L stock suspension (1,000 mg to 15,000 mg fenbendazole/L) and within up to 8 hours during the treatment administration period no stirring of the stock suspension is required. If the administration period exceeds 8 hours, but being no longer than 24 hours, the stock suspension container needs to be equipped with a stirring device.

During treatment, all animals must have sole and unrestricted access to the medicated water. After complete consumption of the medicated water, the animals should have access to non-medicated drinking water ad libitum. Ensure that the total amount of medicated water offered is consumed.

USER SAFETY WARNINGS: Not for use in humans. Keep out of reach of children. Protective gloves should be used and care should be taken when handling the product to avoid skin and eye exposure and accidental ingestion. Accidental exposure may result in skin and eye irritation. Accidental ingestion may cause gastrointestinal disturbances and hypersensitivity reactions in humans. For customer service, adverse effects reporting, and/or a copy of the SDS, call 1-800-211-3573. For additional information about adverse drug experience reporting for animal drugs, contact FDA at 1-888-FDA-VETS, or http://www.fda.gov/ AnimalVeterinary/SafetvHealth

RESIDUE WARNING: Chickens: No withdrawal period is required when used according to labeling. Swine: Swine intended for human consumption must not be slaughtered within 2 days from the last

OTHER WARNINGS: Parasite resistance may develop to any dewormer. All dewormers require accurate dosing for best results. Following the use of any dewormer, effectiveness of treatment should be monitored. A decrease of effectiveness over time may indicate the development of resistance to the dewormer administered. The parasite management plan should be adjusted accordingly based on regular monitoring.

Chickens: Six pivotal dose confirmation studies and five field effectiveness studies were conducted to evaluate the effectiveness of Safe-Guard® AquaSol oral suspension against adult A. galli in broiler chickens and replacement chickens and against A. galli and H. gallinarum in breeding chickens and laying hens. Safe-Guard® AquaSol was administered orally in drinking water at 1 mg fenbendazole/kg body weight/day for 5 consecutive days. The chickens were necropsied 7 to 8 days after the last treatment, and adult worms in the intestines and ceca of the chickens in the control and treated groups were counted to determine percent

Three dose confirmation studies were conducted in European Union (EU), using 105 Rhode Island Red breed hens (2 years old) for each study. In all three studies, the efficacy against *A. galli* (97.9%, 97.3%, and 93.9%) and H. gallinarum (99.8%, 96.9%, and 97.3%) was greater than 90%. A fourth dose confirmation study was conducted in the United States (US) using 264 Rhode Island Red breed hens (12 months old). In the study, the efficacy against adult *A. galli* and *H. gallinarum* was 98.7% and 99.2%, respectively. A fifth dose confirmation study was conducted in the US using 176 Cobb breed broiler chickens (4 to 5 weeks old). In the study, the efficacy against adult *A. galli* was 99.4%. A sixth dose confirmation study was conducted in the US using 176 Ross breed broiler chickens (4 to 5 weeks old). In the study, the efficacy against adult A. galli was 100%

A field effectiveness study was conducted in the EU in a flock of 13,244 Hv-Line layer breed replacement chickens (13 weeks old). Fifteen chickens were necropsied before treatment initiation, and 15 chickens were necropsied seven days after treatment for worm counts. The efficacy against adult A. galli was 90.2%. A second field effectiveness study was conducted in the US using 550 Ross breed broiler chickens (4 to 5 weeks old). The efficacy against adult *A. galli* was 100%. A third field effectiveness study was conducted in the US using 550 White Leghorn breed replacement chickens (14 weeks old). The efficacy against adult A. galli and H. gallinarum was 100% and 88.9%, respectively. A fourth field effectiveness study was conducted in the US using 550 Cobb breed breeder hens (63 weeks old). The efficacy against adult A. galli and H. gallinarum was 97.6% and 95.3%, respectively. A fifth effectiveness study was conducted in the US using 550 Cobb breed broiler chickens (4 to 5 weeks old). The efficacy against adult A. galli was 100%.

The pivotal dose confirmation studies and field effectiveness studies demonstrated substantial evidence of effectiveness of Safe-Guard® AquaSol at the dose of 1 mg fenbendazole/kg body weight/day for 5 consecutive days against adult A. galli in broiler chickens and replacement chickens and against adult A. galli and *H. gallinarum* in breeding chickens and laying hens.

Swine: A multi-site, masked, negative-controlled dose confirmation field study was conducted to provide substantial evidence of the effectiveness of Fenbendazole (FBZ) Suspension (20% w/v) administered orally in drinking water to pigs for three consecutive days to provide a dose of 2.2 mg FBZ/kg body weight daily against the dose-limiting worm Trichuris suis (T. suis). A common protocol was implemented in two different geographical locations and with two different investigators.

Weaned, growing-finishing pigs of approximately 6 weeks of age were used in the study. Each study site selected pigs from one source herd verified to be free of *T. suis* infection. Barrow and gilt breeds representative of U.S. commercial production were used. Housing, management, and husbandry procedures were typical of commercial production practice. A complete feed, adequate to meet the nutritional needs of the study animals, was offered to the animals in self-feeders throughout the study. The feed did not contain antibiotics, anthelmintics, or any other medication.

Fifty-six days prior to treatment administration, all suitable study candidates were orally dosed with approximately 4000 embryonated T. suis eggs. A natural field isolate of T. suis collected in April 2010 from a sow located on a commercial farrow to wean operation located in the U.S. was used. Individual fecal samples were obtained from each candidate animal 46, 47, and 48 days after *T. suis* inoculation and analyzed for the presence of *T. suis* eggs. Animals with at least two fecal examinations positive for *T. suis* eggs were

In each study, 24 healthy pigs were randomized to two treatment groups (FBZ treated and non-medicated) by first blocking by weight in blocks of 4 pigs each and within each weight block, fecal egg count (FEC) in blocks of 2 pigs. The two pigs with the two lowest FEC counts within a weight block were randomized one per treatment group and the two pigs with the highest FEC counts within a weight block were randomized one per treatment group. The two animals assigned to the same treatment group within the same weight block were then assigned to the same pen. Six pens of 2 pigs each were used per treatment group.

Non-medicated water consumption of the pigs in each treatment pen was measured prior to treatment administration to estimate the amount of water required for dosing on each day of the treatment period. The amount of FBZ Suspension administered in drinking water to the study pigs was calculated from pretreatment body weights. Medicated water was prepared on each treatment day by diluting FBZ Suspension in drinking water to provide a daily dose of 2.2 mg FBZ/kg body weight to the FBZ treated group. The control group received non-medicated drinking water.

Only two pigs at the Minnesota (MN) site that were treated with FBZ had abnormal post-treatment observations ("loose stools"). These two pigs had exhibited abnormal fecal consistency prior to treatment with FBZ. There were no abnormal observations made at the California (CA) site on pigs after FBZ administration. There were no abnormal post-treatment observations attributed to administration of FBZ at either study site. The study animals were euthanized after either 8 or 9 days following the last FBZ administration for retrieval of the large intestinal tract. Adult *T.suis* worms attached to the tract and in the contents of the tract were

Adequacy of infection was demonstrated at both study sites by having more than 6 non-medicated pigs (11 of the 12 non-medicated animals in MN and 9 of the 12 non-medicated animals in CA) with adult *T. suis* worm counts of 100 or more per animal.

There was a significant treatment effect in T.suis worm counts between medicated and non-medicated treatment groups at each study site (p=0.0006 in MN and p=0.0003 in CA). The percent reduction in T.suis worm counts in the FBZ medicated animals was greater than 90% (98.5% in MN and 98.6% in CA) compared to the non-medicated animals using transformed data (geometric means).

Palatability: A pivotal palatability study was conducted to evaluate the palatability of 20% Fenbendazole Suspension in pigs through voluntary consumption of medicated water when offered for approximately 5 hours a day over 3 consecutive days at a dose of 2.2 mg fenbendazole/kg body weight (BW) per day (label dose). The average percent of medicated water consumed was 98.18%, thus the study demonstrated that 20% Fenbendazole Suspension has acceptable palatability.

ANIMAL SAFETY:

Chickens: Two margin of safety studies (growing broiler chickens and laying hens at peak egg production) and one reproductive safety study (broiler breeder chickens) were conducted. These studies supported the safety of Safe-Guard® AguaSol in broiler chickens, replacement chickens, laving hens and breeding chickens. when administered in drinking water at 1 mg fenbendazole/kg body weight/day for 5 consecutive days.

The margin of safety in broiler chickens was conducted in 480 broiler chickens. Safe-Guard® AquaSol was administered orally as medicated drinking water to three groups of 120 chickens (60 male and 60 female in each group) at 1, 3, and 5 mg fenbendazole/kg body weight/day (1, 3, and 5 times the recommended label dose) for 15 consecutive days (3 times the recommended duration). Another group of 120 chickens (60 male and 60 female) was provided non-medicated drinking water and used as a control group. In all chickens, feed and water intake, body weights, clinical health, and mortality were recorded. Hematology and clinical chemistry parameters were evaluated in 24 chickens from each group. At the end of the treatment phase, gross necropsies were performed on 48 chickens from each group, and organs weights were evaluated. Histopathologic examinations were performed on 48 chickens each from the control and 5 mg fenbendazole/kg body weight groups. No clinically significant effects related to the administration of Safe Guard® AquaSol were observed for any of the parameters evaluated.

The margin of safety in laying hens was conducted in 144 laying hens. Safe-Guard® AquaSol was administered orally as medicated drinking water to three groups of 36 hens at 1, 3, and 5 times the recommended label dose (1, 3, and 5 mg fenbendazole/kg body weight/day) for 15 consecutive days (3 times the recommended duration). Another group of 36 hens was provided non-medicated drinking water and used as a control group. In all hens, feed and water intake, body weights, clinical health, mortality, egg production, and egg quality parameters (including egg shell thickness and strength, egg weight, and Haugh unit) were evaluated. Hematology and clinical chemistry parameters were evaluated in 12 hens from each group. At the end of the treatment phase, gross necropsies were performed on 12 hens from each group, and organs weights were evaluated. Histopathologic examinations were performed on 12 hens each from the control and 5 mg fenbendazole/kg body weight groups. No clinically significant effects related to the administration of Safe-Guard® AquaSol were observed for any of the parameters evaluated.

The reproductive safety in broiler breeding chickens was conducted in 220 broiler breeder chickens. Safe-Guard® AquaSol was administered orally as medicated drinking water to a group of 110 breeding chickens (10 male and 100 female) at 3 mg fenbendazole/kg body weight/day (3 times the recommended label dose) for 21 consecutive days (4 times the recommended duration). Another group of 110 breeding chickens (10 male and 100 female) were provided non-medicated drinking water and used as a control group. The parameters evaluated in the study included feed and water intake, body weights, clinical health, egg production and weight, fertility, hatchability, and 14-day old chick viability. Necropsy of unhatched eggs was performed to record the percentage of dead embryos and dead and culled hatchlings. At the end of the treatment phase, 30 breeding chickens (10 male and 20 female) from each group were necropsied; and gross pathology and weights of testes and female reproductive tracts were evaluated. Histopathologic evaluations were performed on the gross lesions collected during the necropsy. No clinically significant effects related to the administration of Safe-Guard® AquaSol were observed for any of the parameters evaluated.

Swine: Animal safety was established using a combination of swine pharmacokinetic, physiologic, and pharmacologic data that provided a basis for bridging the safety data of Safe-Guard® Type A medicated article (NADA 131-675) to Safe-Guard® AquaSol oral suspension in swine.

STORAGE INFORMATION: Store at room temperature 30°C (86°F). Once opened, do not store the container above 25°C (77°F). Do not freeze, Use within 6 months after opening. Use the medicated water within 24

HOW SUPPLIED: 1 Liter and 1 Gallon (3,785 mL) HDPE plastic containers

For Patent Information: http://www.merck.com/product/patent/home.html. **Use Only as Directed**

Copyright© 2021 Intervet Inc., a subsidiary of Merck & Co., Inc. Madison, NJ 07940 Fenbendazole (active ingred.) made in China. Formulated in France.

Distributed by: Intervet Inc. 2 Giralda Farms Madison, NJ 07940

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WHAT DRIVES YOU. DRIVES US.





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