

Merck Animal Health

EQUINE PRODUCTS

Reference Guide



We are Driven by Passion for Science and Animal Health

Merck Animal Health is not just a company. It's where the science of healthy animals meets the commitment of horse health professionals. We work every day to bring you innovative products and trusted support. We are building on a rich history of providing animal health solutions. However, to help solve the challenges equine veterinarians face, we're not resting on our history alone. Merck Animal Health continues to make significant investments in research and development each year.



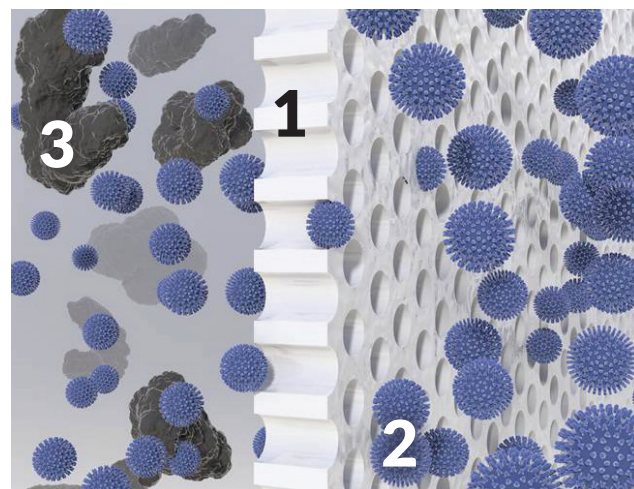
“The science behind our protection builds off a rich history of innovation but doesn't stop there. Our researchers are behind the scenes looking for tomorrow's solutions. We're listening to what horse owners are concerned with and anticipating tools veterinarians will need. I'm proud of our portfolio today and even more excited about how it will look in the future.”

Wendy E. Vaala,
V.M.D., Dipl. ACVIM
Merck Animal Health

Innovation Backed by Science You Can Trust

Antigen Purification System™

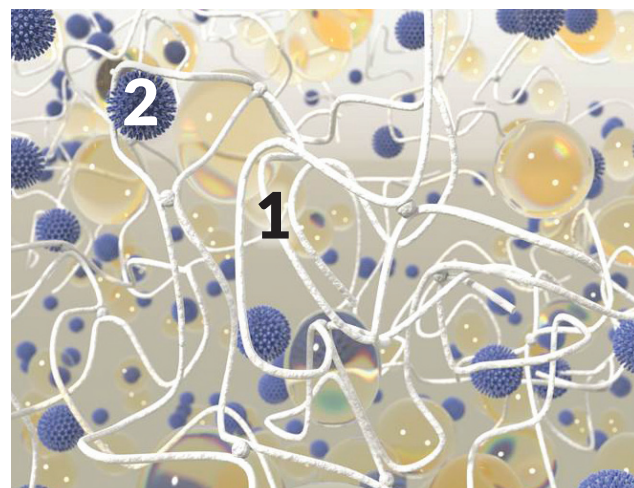
A vaccine can never be too safe. Our technology, known as the Antigen Purification System (APS), has been utilized for more than 20 years to help remove extraneous protein and cellular debris. Using this method of filtration purification allows concentration of antigen while minimizing the presence of extraneous protein and cellular debris that can contribute to vaccine-associated adverse events. By purifying the vaccines with the APS, we reduce the debris that can cause undesirable injection site reactions in the horse.



The microfilter (1) technology in the Merck APS helps purify the vaccine antigen (2) by filtering out unwanted extraneous proteins (3) that may be involved with injection site reactions.

Exclusive Havlogen® Adjuvant

Our killed vaccines are highly efficacious, in part, because of our exclusive Havlogen adjuvant. Havlogen is an emulsive, lipid-based, carbopol polymer cross-linking adjuvant. Havlogen stimulates the immune system to produce high, long-lasting levels of protection through the slow release and gradual absorption of antigen. Due to the composition of Havlogen, the vaccine maintains suspension without separation and settling in the vial—resulting in consistency and potency in every dose. By combining our APS system and Havlogen adjuvant, we are able to produce a line of killed virus vaccines that are highly efficacious and have an exceptional safety profile—shown to be 98% reaction-free in field safety trials¹.



Havlogen is a proprietary adjuvant that is comprised of a lipid-based, carbopol polymer cross-linking suspension (1) that, when combined with the antigen (2), enhances antigen presentation through the slow release and gradual absorption of the antigen.

“The only thing a vaccine should provide is protection. That’s why Merck uses state-of-the-art technology in all its products to minimize risk of reactions and provide consistency in each and every dose.”

D. Craig Barnett, DVM
Merck Animal Health Equine
Professional Services



¹ Data on file. Merck Animal Health.

Vaccines



PRESTIGE® 5 + WNV ENCEPHALOMYELITIS - RHINOPNEUMONITIS - INFLUENZA - WEST NILE VIRUS VACCINE

EASTERN & WESTERN, KILLED VIRUS,
KILLED FLAVIVIRUS CHIMERA
TETANUS TOXOID

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against Eastern and Western encephalomyelitis viruses, EIV, EHV-1, EHV-4, tetanus, and West Nile virus. Duration of immunity has been shown to be at least 6 months for EIV. Duration of immunity for Eastern and Western encephalomyelitis viruses, EHV-1, EHV-4, tetanus, and West Nile virus has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against virus shedding of EIV, EHV-1 and EHV-4, and encephalitis and viremia caused by West Nile virus.

1 x 10 mL, 10 x 1 mL



PRESTIGE® 5 ENCEPHALOMYELITIS - RHINOPNEUMONITIS - INFLUENZA VACCINE

EASTERN AND WESTERN, KILLED VIRUS
TETANUS TOXOID

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against Eastern and Western encephalomyelitis viruses, EIV, EHV-1, EHV-4 and tetanus. Duration of immunity has been shown at six months for EIV. Duration of immunity for Eastern and Western Encephalomyelitis Viruses, EHV-1, EHV-4 and tetanus has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against virus shedding of EIV and EHV-1.

1 x 10 mL, 10 x 1 mL



PRESTIGE® 4 ENCEPHALOMYELITIS - INFLUENZA VACCINE

EASTERN AND WESTERN, KILLED VIRUS
TETANUS TOXOID

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against Eastern and Western encephalomyelitis viruses, EIV and tetanus. Duration of immunity has been shown at six months for EIV. Duration of immunity has not been established for Eastern and Western Encephalomyelitis and Tetanus. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against virus shedding of EIV.

1 x 10 mL, 10 x 1 mL



PRESTIGE® 3 + WNV ENCEPHALOMYELITIS - WEST NILE VIRUS VACCINE

EASTERN & WESTERN, KILLED VIRUS, KILLED FLAVIVIRUS
CHIMERA
TETANUS TOXOID

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against Eastern and Western encephalomyelitis viruses, tetanus and West Nile Virus. Duration of immunity has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against encephalitis and viremia caused by West Nile Virus.

1 x 10 mL, 10 x 1 mL



PRESTIGE® 3 ENCEPHALOMYELITIS VACCINE

EASTERN AND WESTERN, KILLED VIRUS
TETANUS TOXOID

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against Eastern and Western encephalomyelitis viruses and tetanus. Duration of immunity has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov.

1 x 10 mL, 10 x 1 mL



PRESTIGE® 2 EQUINE RHINOPNEUMONITIS - INFLUENZA VACCINE

KILLED VIRUS

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against EIV, EHV-1 and EHV-4. Duration of immunity has been shown at six months for EIV. Duration of immunity for EHV-1 and EHV-4 has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against virus shedding of EIV and EHV-1.

1 x 10 mL, 10 x 1 mL



PRESTIGE® EHV 1&4 EQUINE RHINOPNEUMONITIS VACCINE

KILLED VIRUS

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against EHV-1 and EHV-4. Duration of immunity has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against virus shedding of EHV-1 and EHV-4.

1 x 10 mL



PRESTIGE® WNV WEST NILE VIRUS VACCINE

KILLED FLAVIVIRUS CHIMERA

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against West Nile Virus. Duration of immunity has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product has been shown to be effective against encephalitis and viremia caused by West Nile Virus.

1 x 10 mL, 10 x 1 mL



PRESTIGE® PRODIGY® EQUINE RHINOPNEUMONITIS VACCINE

KILLED VIRUS

This product has been shown to be effective for the vaccination of healthy horses six months of age or older against abortion and respiratory disease caused by EHV-1. Duration of immunity has not been established. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov.

1 x 20 mL, 10 x 2 mL



PRESTIGE® Tetanus TETANUS TOXOID

This product has been shown to be effective for the vaccination of healthy horses, cattle, swine and sheep six months of age or older against tetanus.

10 x 1 mL



PRESTIGE® EQUIRAB® RABIES VACCINE

KILLED VIRUS

This product has been shown to be effective for the vaccination of healthy horses 4 months of age or older against rabies virus. Duration of immunity for rabies virus is at least 14 months. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov.

1 x 10 mL, 10 x 1 mL



FLU AVERT® I.N. EQUINE INFLUENZA VACCINE

MODIFIED LIVE VIRUS - FOR INTRANASAL USE ONLY

This product has been shown to be effective for the vaccination of healthy horses 11 months of age or older against disease caused by EIV. Duration of immunity has been shown to be at least 6 months. For more information regarding safety and efficacy data, go to productdata.aphis.usda.gov. This product contains influenza A/Equine 2/Kentucky/91 (H3N8). Efficacy was demonstrated against A/Equine 2/Kentucky/91 (H3N8) and the duration of immunity was demonstrated against A/Equine/Kentucky/99 (H3N8). This product has been shown to be effective against virus shedding of EIV.

10 x 1 mL

Vaccine Chart

Vaccine	Tetanus	WNV	Rabies	EEE/WEE	Influenza	EHV 1&4	EHV-1 Abortion & Respiratory
PRESTIGE® 5 + WNV	●	●		●	●	●	
PRESTIGE® 5	●			●	●	●	
PRESTIGE® 4	●			●	●		
PRESTIGE® 3 + WNV	●	●		●			
PRESTIGE® 3	●			●			
PRESTIGE® 2					●	●	
PRESTIGE® EHV 1&4						●	
PRESTIGE® WNV		●					
PRESTIGE® Tetanus	●						
PRESTIGE® PRODIGY®							●
PRESTIGE® EQUIRAB®			●				
FLU AVERT® I.N.					●		

Pharmaceuticals



BANAMINE® Paste/Injectable (flunixin meglumine)

BANAMINE® (flunixin meglumine) Paste and BANAMINE® (flunixin meglumine) Injectable Solution are recommended for the alleviation of inflammation and pain associated with musculoskeletal disorders in the horse. BANAMINE® Injectable is also recommended for the alleviation of visceral pain associated with colic in the horse.

100 mL vial, 250 mL vial
30 g syringe

IMPORTANT SAFETY INFORMATION

BANAMINE® Paste: Not for use in horses intended for food. There are no known contraindications to this drug when used as directed. Do not use in horses showing hypersensitivity to flunixin meglumine. The effect of BANAMINE® Paste on pregnancy has not been determined. Concomitant use of BANAMINE® Paste with other anti-inflammatory drugs such as NSAIDs and corticosteroids should be avoided or closely monitored. BANAMINE® Injectable Solution: Not for use in horses intended for food. There are no known contraindications to this drug when used as directed. Intra-arterial injection should be avoided to avoid adverse reactions. Do not use in horses showing hypersensitivity to flunixin meglumine. The effect of BANAMINE® Injectable Solution on pregnancy has not been determined. Concomitant use of BANAMINE® Injectable Solution with other anti-inflammatory drugs such as NSAIDs and corticosteroids should be avoided or closely monitored. In horses, rare instances of anaphylactic-like reactions some of which have been fatal have been reported, primarily following intravenous use.



PANACUR® (fenbendazole) Paste 10%

PANACUR® (fenbendazole) Paste 10% is indicated for the control of large strongyles (*Strongylus edentates*, *S. equinus*, *S. vulgaris*), encysted early third stage (hypobiotic), late third stage and fourth stage cyathostome larvae, small strongyles, pinworms (*Oxyuris equi*), ascarids (*Parascaris equorum*) and arteritis caused by fourth stage larvae of *Strongylus vulgaris* in horses.

PANACUR® (fenbendazole) Paste 10% is approved for use concomitantly with an approved form of trichlorfon. Trichlorfon is approved for the treatment of stomach bots (*Gastrophilus spp.*) in horses.

12 x 25 g syringes

IMPORTANT SAFETY INFORMATION

Consult your veterinarian for assistance in the diagnosis, treatment and control of parasitism. When using PANACUR® (fenbendazole) Paste 10% concomitantly with trichlorfon, refer to the manufacturer's labels for use and cautions for trichlorfon. Do not use in horses intended for human consumption.



PANACUR® (fenbendazole) PowerPac

PANACUR® (fenbendazole) PowerPac is indicated for the control of large strongyles (*Strongylus edentates*, *S. equinus*, *S. vulgaris*), encysted early third stage (hypobiotic), late third stage and fourth stage cyathostome larvae, small strongyles, pinworms (*Oxyuris equi*), ascarids (*Parascaris equorum*) and arteritis caused by fourth stage larvae of *Strongylus vulgaris* in horses.

PANACUR® (fenbendazole) Paste 10% is approved for use concomitantly with an approved form of trichlorfon. Trichlorfon is approved for the treatment of stomach bots (*Gastrophilus spp.*) in horses.

5 x 57 g syringes

IMPORTANT SAFETY INFORMATION

Consult your veterinarian for assistance in the diagnosis, treatment and control of parasitism. When using PANACUR® (fenbendazole) Paste 10% concomitantly with trichlorfon, refer to the manufacturer's labels for use and cautions for trichlorfon. Do not use in horses intended for human consumption.

PROTAZIL®

(1.56% diclazuril) Antiprotozoal Pellets

PROTAZIL® (1.56% diclazuril) Antiprotozoal Pellets are indicated for the treatment of equine protozoal myeloencephalitis (EPM) caused by *Sarcocystis neurona* in horses.

2.4 lb. pail

IMPORTANT SAFETY INFORMATION

Use of PROTAZIL® (1.56% diclazuril) Antiprotozoal Pellets is contraindicated in horses with known hypersensitivity to diclazuril. Safe use in horses used for breeding purposes, during pregnancy, or in lactating mares has not been evaluated. The safety of PROTAZIL® (1.56% diclazuril) Antiprotozoal Pellets with concomitant therapies in horses has not been evaluated. For use in horses only. Do not use in horses intended for human consumption. Not for human use. Keep out of reach of children. For complete safety information, please read label.



DOLOREX® (butorphanol tartrate injection)

DOLOREX® (butorphanol tartrate injection) is indicated for the relief of pain associated with colic in adult horses and yearlings. Clinical studies in the horse have shown that butorphanol tartrate injection alleviates abdominal pain associated with torsion, impaction, intussusception, spasmodic and tympanic colic and postpartum pain.

10 mg/mL 50 mL vial

IMPORTANT SAFETY INFORMATION

FOR USE IN HORSES ONLY. NOT FOR USE IN HORSES INTENDED FOR FOOD. DOLOREX®, a potent analgesic, should be used with caution with other sedative or analgesic drugs as these are likely to produce additive effects. There are no well controlled studies using butorphanol in breeding horses, weanlings, and foals. Therefore, the drug should not be used in these groups. In clinical trials in horses, the most commonly observed side effect was slight ataxia which lasted 3 to 10 minutes. Marked ataxia was reported in 1.5% of the 327 horses treated. Mild sedation was reported in 9% of the horses.



E-SE® (selenium, vitamin E)

E-SE® Injection is recommended for the control of the following clinical signs when associated with myositis (Selenium-Tocopherol Deficiency) syndrome: rapid respiration, profuse sweating, muscle spasms and stiffness, elevated SGOT.

100 mL vial

IMPORTANT SAFETY INFORMATION

Intravenous administration if elected should be by slow injection. Selenium is toxic if administered in excess. Anaphylactoid reactions, some of which have been fatal, have been reported in horses. Medications have been reported to cause major adverse reactions in horses should be avoided when E-SE® is administered. Not to be used in horses intended for food. For complete product information, please see package insert.



REGU-MATE® (altrenogest) Solution 0.22%

REGU-MATE® (altrenogest) Solution 0.22% is indicated to suppress estrus in mares. Suppression of estrus allows for a predictable occurrence of estrus following drug withdrawal. This facilitates the attainment of regular cyclicity during the transition from winter anestrus to the physiological breeding season. Suppression of estrus will also facilitate management of prolonged estrus conditions. Suppression of estrus may be used to facilitate scheduled breeding during the physiological breeding season.

1,000 mL bottle

IMPORTANT SAFETY INFORMATION

REGU-MATE® (altrenogest) Solution 0.22% is contraindicated for use in mares having a previous or current history of uterine inflammation. Pregnant women or women who suspect they are pregnant should not handle REGU-MATE®.



SALIX® (furosemide injection)

SALIX® is an effective diuretic possessing a wide therapeutic range. Pharmacologically it promotes the rapid removal of abnormally retained extracellular fluids. The rationale for the efficacious use of diuretic therapy is determined by the clinical pathology producing the edema. SALIX® is indicated for the treatment of edema, (pulmonary congestion, ascites) associated with cardiac insufficiency and acute noninflammatory tissue edema.

The continued use of heart stimulants, such as digitalis or its glycosides is indicated in cases of edema involving cardiac insufficiency.

50 mL vial

IMPORTANT SAFETY INFORMATION

SALIX® (furosemide injection) is a highly effective diuretic-saluretic which if given in excessive amounts may result in dehydration and electrolyte imbalance, enhancing the risk of circulatory collapse, thrombosis, and embolism. The animal should be observed for early signs of fluid depletion with electrolyte imbalance, and corrective measures should be administered. See package insert for full information regarding contraindications, warnings, and precautions. SALIX® (furosemide injection) may lower serum calcium levels and cause tetany in rare cases of animals having an existing hypocalcemic tendency. Milk taken from dairy cattle during treatment and for 48 hours (four milkings) after the last treatment must not be used for food. Cattle must not be slaughtered for food within 48 hours following last treatment. Do not use in horses intended for human consumption.

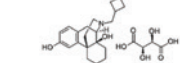
To learn more about Merck Animal Health equine products please contact your equine sales representative or call 1-800-521-5767

DOLOREX®

Intervet/Merck Animal Health
(butorphanol tartrate injection)

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

DESCRIPTION DOLOREX (butorphanol tartrate injection) is a totally synthetic, centrally acting, narcotic agonist-antagonist analgesic with potent antitussive activity. It is a member of the phenanthrene series. The chemical name is Morphinan-3, 14-diol, 17-(cyclobutylmethyl)-, (-)-, (S- (R*, R*)):- 2, 3- dihydroxybutanedioate (1:1) (salt). It is a white crystalline, water soluble substance having a molecular weight of 477.55; its molecular formula is C₂₁H₂₉NO₂•C₄H₆O₆



Each mL of DOLOREX contains 10 mg butorphanol base (as butorphanol tartrate), 3.3 mg citric acid, 6.4 mg sodium citrate, 4.7 mg sodium chloride, and 0.1 mg benzethonium chloride, q.s. with water for injection.

COMPARATIVE PHARMACOLOGY In animals, butorphanol has been demonstrated to be 4 to 30 times more potent than morphine and pentazocine (Talwin®-V) respectively! In humans, butorphanol has been shown to have 5 to 7 times the analgesic activity of morphine and 20 times that of pentazocine.^{2,3} Butorphanol has 15 to 20 times the oral antitussive activity of codeine or dextromethorphan in dogs and guinea pigs.⁴

As an antagonist, butorphanol is approximately equivalent to nalorphine and 30 times more potent than pentazocine.¹

Cardiopulmonary depressant effects are minimal after treatment with butorphanol as demonstrated in dogs⁵, humans^{6,7} and horses.⁸ Unlike classical narcotic agonist analgesics which are associated with decreases in blood pressure, reduction in heart rate, and concomitant release of histamine, butorphanol does not cause histamine release! Furthermore, the cardiopulmonary effects of butorphanol are not distinctly dosage related but rather reach a ceiling effect beyond which further dosage increases result in relatively lesser effects.

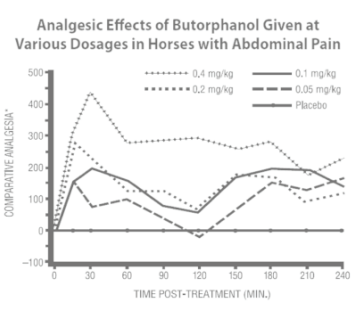
Reproduction studies performed in mice and rabbits revealed no evidence of impaired fertility or harm to the fetus due to butorphanol tartrate. In the female rat, parenteral administration was associated with increased nervousness and decreased care for newborn, resulting in a decreased survival rate of the newborn. This nervousness was seen only in the rat species.

EQUINE PHARMACOLOGY

Following intravenous injection in horses, butorphanol is largely eliminated from the blood within 3 to 4 hours. The drug is extensively metabolized in the liver and excreted in the urine.

In ponies, butorphanol given intramuscularly at a dosage of 0.22 mg/kg was shown to alleviate experimentally induced visceral pain for about 4 hours.⁹

In horses, intravenous dosages of butorphanol ranging from 0.05 to 0.4 mg/kg were shown to be effective in alleviating visceral and superficial pain for at least 4 hours.



A definite dosage-response relationship was detected in that butorphanol dosage of 0.1 mg/kg was more effective than 0.05 mg/kg, but not different from 0.2 mg/kg, in alleviating deep abdominal pain.

ACUTE EQUINE STUDIES

Rapid intravenous administration of butorphanol at a dosage of 2.0 mg/kg (20 times the recommended dosage) to a previously unmedicated horse resulted in a brief episode of inability to stand, muscle fasciculation, a convulsive seizure of 6 seconds duration, and recovery within 3 minutes. The same dosage administered after 10 successive daily 1.0 mg/kg dosages of butorphanol resulted only in transient

sedative effects. During the 10-day course of administration at 1.0 mg/kg (10 times the recommended use level) in 2 horses, the only detectable drug effects were transient behavioral changes typical of narcotic agonist activity. These included muscle fasciculation about the head and neck, dysphoria, lateral nystagmus, ataxia, and salivation. Repeated administration of butorphanol at 1.0 mg/kg (10 times the recommended dosage) every 4 hours for 48 hours caused constipation in one of two horses.

SUBACUTE EQUINE STUDIES

Horses were found to tolerate butorphanol given intravenously at dosages of 0.1, 0.3, and 0.5 mg/kg every 4 hours for 48 hours followed by once daily injections for a total of 21 days. The only detectable drug effects were slight transient ataxia observed occasionally in the high dosage group. No clinical, laboratory, or gross or histopathologic evidence of any butorphanol-related toxicity was encountered in the horses.

INDICATIONS DOLOREX (butorphanol tartrate injection) is indicated for the relief of pain associated with colic in adult horses and yearlings. Clinical studies in the horse have shown that butorphanol tartrate injection alleviates abdominal pain associated with torsion, impaction, intussusception, spasmodic and tympanic colic, and postpartum pain.

WARNING FOR USE IN HORSES ONLY. NOT FOR USE IN HORSES INTENDED FOR HUMAN CONSUMPTION.

CAUTION DOLOREX, a potent analgesic, should be used with caution with other sedative or analgesic drugs as these are likely to produce additive effects.

There are no well controlled studies using butorphanol in breeding horses, weanlings, and foals. Therefore, the drug should not be used in these groups.

ADVERSE REACTIONS In clinical trials in horses, the most commonly observed side effect was slight ataxia which lasted 3 to 10 minutes. Marked ataxia was reported in 1.5% of the 327 horses treated. Mild sedation was reported in 9% of the horses.

DOUSAGE The recommended dosage in the horse is 0.1 mg butorphanol per kilogram of body weight (0.05 mg/lb) by intravenous injection. This is equivalent to 5 mL DOLOREX for each 1000 lb body weight. The dose may be repeated within 3 to 4 hours, but treatment should not exceed 48 hours. Preclinical model studies and clinical field trials in horses demonstrate that the analgesic effects of butorphanol are seen within 15 minutes following injection and persist for about 4 hours.

HOW SUPPLIED DOLOREX is supplied in 50 mL vials (Order Code No. 017070). Store at or below 25°C (77°F)..

REFERENCES

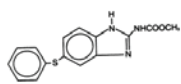
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Intervet Inc d/b/a Merck Animal Health
Madison, NJ 07940 Rev. 08/20
By: INTERVET INTERNATIONAL GmbH,
Unterschleissheim - Germany

PANACUR®

Intervet/Merck Animal Health
(benzbendazole)
25 gram
Paste 10% (100 mg/g) Equine Dewormer

DESCRIPTION: PANACUR® (fenbendazole) Paste 10% contains the active anthelmintic, fenbendazole. The chemical name of fenbendazole is methyl 5-(phenylthio)-2-benzimidazole carbamate. The chemical structure is:



Each gram of PANACUR® (fenbendazole) Paste 10% contains 100 mg of fenbendazole and is flavored with artificial apple-cinnamon liquid.

ACTIONS: The antiparasitic action of PANACUR® (fenbendazole) Paste 10% is believed to be due to the inhibition of energy metabolism in the parasite.

INDICATIONS: PANACUR® (fenbendazole) Paste 10% is indicated for the control of large strongyles (*Strongylus edentatus*, *S. equinus*, *S. vulgaris*), encysted early third stage (hyobiotic), late third stage and fourth stage cyathostome larvae, small strongyles, pinworms (*Oxyuris equi*), ascarids (*Parascaris equorum*), and arteritis caused by fourth stage larvae of *Strongylus vulgaris* in horses. PANACUR® (fenbendazole) Paste 10% is approved for use concomitantly with an approved form of trichlorfon. Trichlorfon is approved for the treatmentof stomach bots (*Gasterophilus spp.*) in horses. Refer to the manufacturer’s label for directions foruse and cautions for trichlorfon.

PRECAUTIONS: Side effects associated with PANACUR® (fenbendazole) Paste 10% could not be established in well-controlled safety studies in horses with single doses as high as 454 mg/ lb (1,000 mg/kg) and 15 consecutive daily doses of 22.7 mg/lb (50 mg/kg). Particularly with higher doses, the lethal action of fenbendazole may cause the release of antigens by the dying parasites. This phenomenon may result in either a local or systemic hypersensitive reaction. As with any drug, these reactions should be treated symptomatically. PANACUR® (fenbendazole) Paste 10% has been evaluated for safety in pregnant mares during all stages of gestation with doses as high as 11.4 mg/lb (25 mg/kg) and in stallions with doses as high as 11.4 mg/lb (25 mg/kg). No adverse effects on reproductivity were detected. The recommended dose for control of 4th stage larvae of Strongylus vulgaris, 4.6 mg/lb (10 mg/kg) daily for 5 consecutive days, has not been evaluated for safety in stallions or pregnant mares.

Internal Parasites: Regular deworming at intervals of six to eight weeks may be required due to the possibility of reinfection.

Migrating Tissue Parasites: In the case of 4th stage larvae of *Strongylus vulgaris*, treatment and retreatment should be based on the life cycle andthe epidemiology. Treatment should be initiated in the spring and repeated in the fall after a six month interval.

Optimum Deworming Program for control of *S. vulgaris*: Optimum reduction of *S. vulgaris* infections is achieved by reducing the infectivity of the pastures. When horses are running on pasture, in temperate North America, maximum pasture infectivity occurs in October-December. If horses are removed from those pastures in January, pasture infectivity will decline to zero by July 1. Egg production of *S. vulgaris* is minimal from January through April, peaking in August and declining to minimal values in December.

Recommended Deworming Program:
** December 1, February 1, **April 1**, June 1, August 1, **October 1**.

The two treatments that are in bold type are the recommended periods when the 5 day

treatment regimen for the control of the migrating larvae of *S.vulgaris* should be performed.
**For other areas in the world, retreatment periods for the migrating larvae of *S. vulgaris* may be different; consult with your veterinarian.

CAUTIONS: Keep this and all medications out of the reach of children.

When using PANACUR® (fenbendazole) Paste 10% concomitantly with trichlorfon, refer to the manufacturer’s labels for use and cautions for trichlorfon.

WARNING: Do not use in horses intended for human consumption

DOSAGE: PANACUR® (fenbendazole) Paste 10% is administered orally at a rate of 2.3 mg/lb (5 mg/kg) for the control of large strongyles, small strongyles, and pinworms. One syringe will deworm a 1,100 lb horse. For foals and weanlings (less than 18 months of age) where ascarids are a common problem, the recommended dose is 4.6 mg/lb (10 mg/kg); one syringe will deworm a 550 lb horse. For control of encysted early third stage (hypobiotic), late third stage and fourth stage cyathostome larvae, and fourth stage larvae of *Strongylus vulgaris*, the recommended dose is 4.6 mg/lb (10 mg/kg) daily for 5 consecutive days; administer one syringe for each 550 lbs body weight per day.

SEE PRECAUTIONS FOR RETREATMENT RECOMMENDATIONS.

DIRECTIONS FOR USE:

- Determine the weight of the horse.
- Remove syringe tip.
- Turn the dial ring until the edge of the ring nearest the tip lines up with zero.
- Depress plunger to advance paste to tip.
- Now set the dial ring at the graduation nearest the weight of the horse (do not underdose).
- Horse’s mouth must be free of food.
- Insert nozzle of syringe through the interdental space and deposit the paste on the back of the tongue by depressing the plunger.

HOW SUPPLIED: PANACUR® (fenbendazole) Paste 10.% Equine Dewormer is supplied in 25 g syringes. **Store at or below 25°C (77°F).**

CONSULT YOUR VETERINARIAN FOR ASSISTANCE IN THE DIAGNOSIS, TREATMENT AND CONTROL OF PARASITISM.

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147109 R2
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