

EQUISUL-SDT- sulfadiazine and trimethoprim suspension
Aurora Pharmaceutical, Inc.

EQUISUL-SDT®
(Sulfadiazine/Trimethoprim)
Oral Suspension

For use in horses only.

Approved by FDA under NADA # 141-360

CAUTION

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

DESCRIPTION

EQUISUL-SDT is a broad-spectrum antimicrobial from the potentiated sulfonamide class of chemotherapeutic agents. These two drugs block different sequential steps in the biosynthesis of nucleic acids. Sulfadiazine inhibits bacterial synthesis of dihydrofolic acid by competing with para-aminobenzoic acid. Trimethoprim blocks the production of tetrahydrofolic acid from dihydrofolic acid by reversibly inhibiting dihydrofolate reductase. The effect of the dual action is to reduce the minimum inhibitory concentration of each agent (synergism) and to convert a bacteriostatic action to a bactericidal action. Sulfadiazine is the non-proprietary name for 4-amino-N-2-pyrimidinylbenzenesulfonamide. Trimethoprim is the non-proprietary name for 5-[(3,4,5-trimethoxyphenyl) methyl]-2,4-pyrimidinediamine.

Figure 1. Structure of sulfadiazine

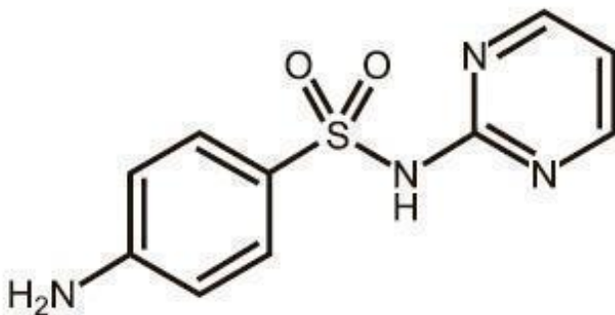
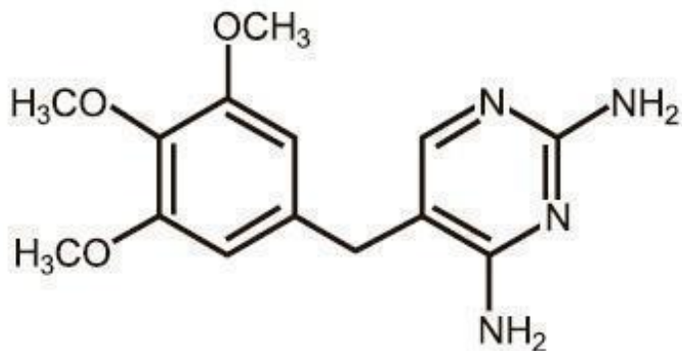


Figure 2. Structure of trimethoprim



Each mL of EQUISUL-SDT contains 400 mg combined active ingredients (333 mg sulfadiazine and 67 mg trimethoprim) in an aqueous suspension.

INDICATION

EQUISUL-SDT is indicated for the treatment of lower respiratory tract infections in horses caused by susceptible strains of *Streptococcus equi* subsp. *zooepidemicus*.

DOSAGE AND ADMINISTRATION

Shake well before use.

Administer EQUISUL-SDT orally at the dosage of 24 mg combined active ingredients per kilogram body weight (10.9 mg/lb) twice daily for 10 days. EQUISUL-SDT can be administered by volume at 2.7 mL per 45.4 kg (2.7 mL/100 lb) body weight.

For 280 mL and 560 mL bottles with draw-off cap:

Remove cap. Peel off white foil backed bottle seal and replace cap. Peel off outer cap seal exposing (hole) opening. Push an oral tip syringe into the cap opening. Invert and draw out appropriate volume of EQUISUL-SDT solution. (Note: Do not remove syringe while the bottle is inverted as possible spillage may result.) Detach syringe and administer orally at the dosage of 24 mg combined active ingredients per kilogram body weight (10.9 mg/lb) twice daily for 10 days. EQUISUL-SDT can be administered by volume at 2.7 mL per 45.4 kg (2.7 mL/100 lb) body weight.

CONTRAINDICATIONS

EQUISUL-SDT is contraindicated in horses with a known allergy to sulfadiazine, sulfonamide class antimicrobials, or trimethoprim.

WARNING

Do not use in horses intended for human consumption.

HUMAN WARNINGS

Not for use in humans. For use in animals only. Keep this and all drugs out of

the reach of children. Consult a physician in the case of accidental human exposure.

Antimicrobial drugs, including sulfonamides, can cause mild to severe allergic reactions in some individuals. Avoid direct contact of the product with the skin, eyes, mouth, and clothing. Persons with a known sensitivity to sulfonamides or trimethoprim should avoid exposure to this product. If an allergic reaction occurs (e.g., skin rash, hives, difficulty breathing, facial swelling) seek medical attention.

PRECAUTIONS

Prescribing antibacterial drugs in the absence of a proven or strongly suspected bacterial infection is unlikely to provide benefit to treated animals and may increase the risk of development of drug-resistant animal pathogens.

The administration of antimicrobials, including sulfadiazine and trimethoprim, to horses under conditions of stress may be associated with acute diarrhea that can be fatal. If acute diarrhea or persistent changes in fecal consistency are observed, additional doses of EQUISUL-SDT should not be administered and appropriate therapy should be initiated.

The safe use of EQUISUL-SDT has not been evaluated in breeding, pregnant, or lactating horses. Potentiated sulfonamides should only be used in pregnant or lactating mares when the benefits to the mare justify the risks to the fetus. Use of potentiated sulfonamides during pregnancy has been associated with an increased risk of congenital abnormalities that may be related to folate deficiency. In humans, sulfonamides pass through the placenta, are excreted in milk, and may cause hyperbilirubinemia-induced neurotoxicity in nursing neonates.

Decreased hematopoietic activity and blood dyscrasias have been associated with the use of elevated doses and/or prolonged administration of potentiated sulfonamides. EQUISUL-SDT should be discontinued if prolonged clotting times, or decreased platelet, white blood cell or red blood cell counts are observed.

Sulfonamides should be used with caution in horses with impaired hepatic function. Although rare, sulfonamide use has been associated with fulminant hepatic necrosis in humans.

Neurologic abnormalities have been reported in several species following administration of potentiated sulfonamides. In horses, potentiated sulfonamides have been associated with gait alterations and behavior changes that resolved after discontinuation of the drug.

The safe use of EQUISUL-SDT has not been evaluated in horses less than 1 year of age.

ADVERSE REACTIONS

Adverse reactions reported during a field study of 270 horses of various breeds, ranging from 1 to 25 years of age, which had been treated with either EQUISUL-SDT (n = 182) or with a saline control (n = 88) are summarized in Table 1. At least one episode of loose stool of varying severity was observed in 69 of 182 (38%) of the EQUISUL-SDT-treated horses, and 29 of 88 (33%) saline control horses. Of those animals experiencing loose stool, 2 of 182 (1.1%) of the EQUISUL-SDT-treated horses and 0 of 88 (0%)

placebo-treated horses were removed from the study due to diarrhea (defined as at least one episode of watery stool). Both cases of diarrhea in this study were self-limiting and resolved without treatment within 5–10 days after discontinuation of EQUISUL-SDT.

Table 1. Number of Horses with Adverse Reactions During the Field Study with EQUISUL-SDT

Adverse Reactions	Equisul-SDT (n=182)	Saline control (n=88)
Loose stool (including diarrhea)	69 (38%)	29 (33%)
Colic	3 (1.6%)	2 (2.2%)
Diarrhea	2 (1.1%)	0 (0%)

To report suspected adverse events, for technical assistance or to obtain a copy of the SDS, contact Aurora Pharmaceutical, Inc. at 1-888-215-1256 or www.aurorapharmaceutical.com. 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.

CLINICAL PHARMACOLOGY

Following oral administration, EQUISUL-SDT is rapidly absorbed and widely distributed throughout body tissues. Sulfadiazine levels are usually highest in the kidney, while the tissue concentration in other tissues is only slightly lower than plasma concentrations. Concentrations of trimethoprim are usually higher in the lungs, kidney, and liver than in the blood. Sulfadiazine and trimethoprim are both eliminated primarily by renal excretion, both by glomerular filtration and tubular secretion. Urine concentrations of both sulfadiazine and trimethoprim are several-fold higher than blood concentrations.¹ Sulfadiazine and trimethoprim are 20% and 35% bound to plasma protein, respectively. Administration of sulfadiazine and trimethoprim with food has no apparent effect on absorption of sulfadiazine but the absorption of trimethoprim is decreased.

Based on a study in fed horses, trimethoprim concentrations following repeat oral administration of 24 mg/kg EQUISUL-SDT to 6 horses reached peak concentration in 0.5 to 12.0 hours. The median plasma elimination half-life was 3 hours, with a range of 2.31 to 4.96 hours. Peak sulfadiazine concentrations were reached within 1.0 to 12.0 hours in the same study. The median plasma elimination half-life for sulfadiazine was approximately 7.80 hours, with a range of 6.78 to 10.39 hours. Only minor accumulation of both drugs was observed following repeat oral administration of EQUISUL-SDT and both drugs reached steady state by day 3. Sulfadiazine and trimethoprim key steady state parameters associated with administration in 6 fed horses over a period of 7 days are found in Table 2.

Table 2. Median (Range) of sulfadiazine and trimethoprim pharmacokinetics parameters following repeat dosing of 24 mg/kg bid EQUISUL-SDT for 7 days to six horses in fed condition

Drug	Sulfadiazine	Trimethoprim
Tmax (hr)	4.75 (1.0012.00)	8.50 (0.5012.00)
Cmax (µg/mL)	17.63 (10.1031.15)	0.78 (0.601.14)
AUC 012 (last dose) (hr*µg/mL)	159.35 (73.90282.54)	5.47 (3.3110.91)
T 1/2 (hr)	7.80 (6.7810.39)	3.00 (2.314.96)

1 Kahn CM, Line S, eds. The Merck Veterinary Manual. 10th Ed. Merck & Co. 2010.

MICROBIOLOGY

EQUISUL-SDT is the combination of the sulfonamide sulfadiazine and trimethoprim. These two drugs block sequential steps in nucleic acids biosynthesis. Sulfadiazine inhibits bacterial synthesis of dihydrofolic acid by competing with para-aminobenzoic acid. Trimethoprim blocks the production of tetrahydrofolic acid from dihydrofolic acid by reversibly inhibiting dihydrofolate reductase. The two drugs act synergistically, reducing the minimum inhibitory concentration of each, while enhancing the bacteriostatic action of each separately to a bactericidal action when combined.

EQUISUL-SDT administered as a combined sulfadiazine-trimethoprim dose of 24 mg/kg body weight twice daily for 7 days provided concentrations of sulfadiazine and trimethoprim with T>MIC90 (%T) values of 100% and 98% respectively. The minimum inhibitory concentration (MIC) values for EQUISUL-SDT against indicated pathogens isolated from lower respiratory tract infections in horses enrolled in a 2010–2011 effectiveness field study are presented in Table 3. All MICs were determined in accordance with the Clinical and Laboratory Standards Institute (CLSI) Approved Standard M31-A3 using a broth microdilution system and 3% lysed horse blood.

Table 3. Trimethoprim/sulfadiazine minimum inhibitory concentration (MIC) values* of isolates recovered from horses with lower respiratory infection caused by *Streptococcus equi* subsp. *zooepidemicus* treated with EQUISUL-SDT in the U.S. (2010-2011)

Treatment Outcome	Success	Failure
Number of Isolates	65 [†]	46
Time of Sample Collection	Pre-Treatment	Pre-Treatment
MIC 50 [‡] (µg/mL)	0.25/4.75	0.25/4.75
MIC 90 [‡] (µg/mL)	0.25/4.75	0.25/4.75
MIC Range (µg/mL)	0.12/2.4 to 0.5/9.5	0.12/2.4 to 0.5/9.5

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

† One isolate of *S. equi* subsp. *zooepidemicus* was not tested.

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

EFFECTIVENESS

A negative control, randomized, masked, field study evaluated the effectiveness of EQUISUL-SDT administered at 24 mg/kg body weight, orally, twice daily for 10 days for the treatment of lower respiratory tract infections in horses caused by *Streptococcus equi* subsp. *zooepidemicus*. In this study, a total of 182 horses were treated with EQUISUL-SDT, and 88 horses were treated with saline. One hundred seventy-three horses (112 EQUISUL-SDT and 61 saline) were included in the statistical analysis. Therapeutic success was characterized by absence of fever and no worsening of clinical signs at Day 5 and Day 10, and significant clinical improvement or resolution of clinical signs of lower respiratory tract infection by Day 17. The observed success rates are 58.9% (66/112) and 14.8% (9/61) for the EQUISUL-SDT and saline-treated groups, respectively.

Table 4 summarizes the statistical analysis results on the overall success rate.

Table 4. Overall Clinical Effectiveness Results

	Equisul-SDT	Saline	P-value*
Least Square Means	61%	13.1%	0.0123

* P-value and estimated success rates are based on back-transformed mean estimates from the statistical analysis.

ANIMAL SAFETY

In a target animal safety study, EQUISUL-SDT was administered orally to 32 healthy adult horses at 0 (0×), 24 (1×), 72 (3×), or 120 (5×) mg/kg twice daily for 30 days. Loose stool was the most common abnormal observation. Observations of loose stool (pellets with liquid or unformed/cowpile stool) occurred more often in horses treated with EQUISUL-SDT with the incidence of loose stool increasing in a dose related manner. All incidents of loose stool were self-limiting and resolved without treatment.

Horses in all EQUISUL-SDT groups demonstrated statistically significantly higher mean serum creatinine concentrations, and those in the 3× and 5× groups demonstrated statistically significantly higher mean serum albumin concentrations. Statistically higher mean neutrophil counts and mean serum gamma glutamyl transferase (GGT) activity were seen in the 1× and 5× groups. Individual animal creatinine, GGT, and albumin concentrations remained within the reference range. Individual animal elevations in absolute neutrophil counts ranged up to $7.09 \times 10^3/\text{mCL}$ (reference range: $1.96\text{-}5.31 \times 10^3/\text{mCL}$).

Based upon blood concentrations obtained during the study, it was noted that the sulfadiazine and trimethoprim plasma concentrations did not increase in proportion to dose. For sulfadiazine, a 3× and 5× dose resulted in an average exposure of 2.0× and 2.6× the concentrations observed following a 1× dose. For trimethoprim, the corresponding values were 2.5× and 3.5× as compared to the 1× dose. Furthermore, marked intersubject variability, particularly with sulfadiazine, resulted in substantial

overlap of individual subject blood levels across the three dosing groups.

STORAGE CONDITIONS

Store upright at 59°– 86° F (15°– 30° C). Brief periods up to 104° F (40° C) are permitted. Protect from freezing.

HOW SUPPLIED

EQUISUL-SDT is available in the following package sizes:

135 mL

280 mL

560 mL

900 mL

MANUFACTURED IN THE USA

MANUFACTURED BY:

Aurora Pharmaceutical, Inc.

NORTHFIELD, MN 55057

1-888-215-1256

Patent 6,800,631 6,410,543 6,211,185

Note: See insert for additional information, precautions and side effects

PRINCIPAL DISPLAY PANEL - 900 mL Bottle Label

NDC 51072-020-00

EQUISUL-SDT®

(Sulfadiazine/Trimethoprim)

Antimicrobial

Oral Suspension

400 mg/mL

Approved by FDA under NADA # 141-360

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

For use in horses only.

900 mL

REORDER No: 28001

IN 50-1274 02/2021

aurora pharmaceutical®

NOTE: See insert for additional information, precautions and side effects.

NDC 51072-020-00

EQUISUL-SDT[®]

(Sulfadiazine/Trimethoprim)

Antimicrobial
Oral Suspension
400 mg/mL

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

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900 mL

INDICATION: For the treatment of lower respiratory tract infections in horses caused by susceptible strains of *Streptococcus equi* subsp. *zooepidemicus*.

ACTIVE INGREDIENTS: Each mL contains 400 mg combined active ingredients (333 mg sulfadiazine and 67 mg trimethoprim).

DOSAGE AND ADMINISTRATION: **Shake well before use.** Administer EQUISUL-SDT orally at the dosage of 24 mg combined active ingredients per kilogram body weight (10.9 mg/lb) twice daily for 10 days. EQUISUL-SDT can be administered by volume at 2.7 mL per 45.4 kg (2.7 mL/100 lb) body weight.

WARNING: Do not use in horses intended for human consumption.

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MANUFACTURED IN THE USA

REORDER NO: 28001

MANUFACTURED BY:
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1-888-215-1256

IN 50-1274 02/2021

Patent 6,800,631 6,410,543
6,211,185

3 51072 02000 8

OPEN

OBSERVE LABEL DIRECTIONS

TAKE TIME



PRINCIPAL DISPLAY PANEL - 280 mL Bottle Label with draw-off cap

NDC 51072-020-03

EQUISUL-SDT[®]

(Sulfadiazine/Trimethoprim)

Antimicrobial

Oral Suspension

400 mg/mL

Approved by FDA under NADA # 141-360

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

For use in horses only.

280 mL

REORDER No: 28003

IN 50-1346 01/2021

aurora pharmaceutical[®]

Discard 60 days after removing bottle seal.

Date of bottle seal removal: _____

NOTE: See insert for additional information, precautions and side effects.

MANUFACTURED IN THE USA

REORDER NO: 28003

MANUFACTURED BY:
Aurora Pharmaceutical, Inc.
NORTHFIELD, MN 55057
1-888-215-1256

IN 50-1346 01/2021

Patent 6,800,631 6,410,543 6,211,185



NDC 51072-020-03

EQUISUL-SDT

(Sulfadiazine/Trimethoprim)

Antimicrobial

Oral Suspension

400 mg/mL

Approved by FDA under NADA # 141-360

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

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280 mL



INDICATION: For the treatment of lower respiratory tract infections in horses caused by susceptible strains of *Streptococcus equi* subsp. *zooepidemicus*.

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WARNING: Do not use in horses intended for human consumption.

STORAGE: Store upright at 59°–86° F (15°–30° C). Brief periods up to 104° F (40° C) are permitted. Protect from freezing.

OPEN

OBSERVE LABEL DIRECTIONS



TAKE TIME

EQUISUL-SDT

sulfadiazine and trimethoprim suspension

Product Information

Product Type	PRESCRIPTION ANIMAL DRUG	Item Code (Source)	NDC:51072-020
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SULFADIAZINE (UNII: 0N7609K889) (SULFADIAZINE - UNII:0N7609K889)	SULFADIAZINE	333 mg in 1 mL
TRIMETHOPRIM (UNII: AN164J8Y0X) (TRIMETHOPRIM - UNII:AN164J8Y0X)	TRIMETHOPRIM	67 mg in 1 mL

Product Characteristics

Color	WHITE	Score	
Shape		Size	
Flavor	APPLE	Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:51072-020-00	900 mL in 1 BOTTLE		
2	NDC:51072-020-01	135 mL in 1 BOTTLE		
3	NDC:51072-020-02	560 mL in 1 BOTTLE		
4	NDC:51072-020-03	280 mL in 1 BOTTLE		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NADA	NADA141360	08/05/2013	

Labeler - Aurora Pharmaceutical, Inc. (832848639)

Establishment

Name	Address	ID/FEI	Business Operations
Aurora Pharmaceutical, Inc.		832848639	MANUFACTURE

Revised: 6/2025

Aurora Pharmaceutical, Inc.