

**NEOMYCIN - neomycin sulfate liquid**  
**Durvet, Inc.**

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**NEOMYCIN ORAL SOLUTION**

**FOR ANIMAL USE ONLY**

Indicated for the treatment and control of colibacillosis (bacterial enteritis) caused by *Escherichia coli* susceptible to neomycin sulfate in cattle (excluding veal calves), swine, sheep and goats.

Restricted Drug Use Only As Directed (California)  
For Oral Use in Animals Only

**Dosage and Administration**

Administer to cattle (excluding veal calves), swine, sheep and goats at a dose of 10 mg neomycin sulfate per pound of body weight in divided doses for a maximum of 14 days.

**Dosage Schedule for treatment of colibacillosis**

<u>Pounds of Body Weight</u>	<u>Amount of Neomycin Solution Per Day in Divided Doses</u>
25 lb	1.2 mL (1/4 tsp*)
50 lb	2.5 mL (1/2 tsp*)
100 lb	5 mL (1 tsp*)
300 lb	15 mL (1 tbsp*)
591.5 lb	29.5 mL (1 fl oz)

\* Teaspoon (tsp) / Tablespoon (tbsp) is equal to U.S. Standard Measure

Neomycin Solution may be given undiluted or diluted with water.

**Herd Treatment:** Each bottle will treat 9,600 pounds of body weight. Therefore, estimate the total number of pounds body weight of the animals to be treated and administer on (1) fluid ounce of each 600 pounds. The product should be added to the amount of drinking water to be consumed in 12-24 hours. Provide medicated water as the sole source of water each day until consumed, followed by non-medicated water as required. Fresh medicated water should be prepared each day.

**Individual Animal Treatment:** To provide 10 mg neomycin sulfate per pound of body weight, mix one (1) teaspoon in water or milk for each 100 pounds of body weight. Administer daily either as a drench in divided doses or in the drinking water to be consumed in 12-24 hours.

**PRECAUTIONS**

To administer the stated dosage, the concentration of neomycin required in medicated water must be adjusted to compensate for variation in age and weight of animal, the nature and severity of disease signs, and environmental temperature and humidity, each of which affects water consumption.

**KEEP OUT OF REACH OF CHILDREN**

**Contains per mL: neomycin sulfate 200 mg equivalent to 140 mg neomycin**

If symptoms persist after using this preparation for 2 or 3 days, consult a veterinarian. If symptoms such as fever, depression, or going off feed develop, oral neomycin is not indicated as the sole treatment since systemic levels of neomycin are not obtained due to low absorption from the gastrointestinal tract.

## Important

Treatment should continue 24 to 48 hours beyond remission of disease symptoms, but not to exceed a total of 14 consecutive days. Animals not drinking or eating should be treated individually by drench.

## Residue Warnings

Not for human use. Keep out of reach of children. Discontinue treatment prior to slaughter by at least the number of days listed below for appropriate species:

Cattle.....	1 day
Sheep.....	2 days
Swine and Goats.....	3 days

A withdrawal period has not been established for this product in pre-ruminating calves. Do not use in calves to be processed for veal.

A milk discard period has not been established for this product in lactating dairy cattle. Do not use in female dairy cattle 20 months of age or older.

**Use of more than one product containing Neomycin or failure to follow withdrawal times may result in illegal drug residues.**

Store at controlled room temperature 20° to 25°C (68 to 77°F) (see USP).

**TAKE TIME OBSERVE LABEL DIRECTIONS**

**NEOMYCIN LIQUID**

**ANTIBACTERIAL**

**FOR ORAL USE ONLY**

**ANADA 200-379, Approved by FDA**

NDC 30798-340-31

**FOR ANIMAL USE ONLY**

Indicated for the treatment and control of colibacillosis (bacterial enteritis) caused by *Escherichia coli* susceptible to neomycin sulfate in cattle (excluding veal calves), swine, sheep and goats.

Restricted Drug-Use Only As Directed (California)  
For Oral Use in Animals only

**Dosage and Administration:** Administer to cattle (excluding veal calves), swine, sheep and goats at a dose of 10 mg neomycin sulfate per pound of body weight in divided doses for a maximum of 14 days.

**Dosage Schedule for Treatment of Colibacillosis:**

Pounds of Body Weight	Amount of Neomycin Oral Solution Per Day in Divided Doses
25 lb	1.2 mL (1/4 tsp*)
50 lb	2.5 mL (1/2 tsp*)
100 lb	5 mL (1 tsp*)
300 lb	15 mL (1 tbsp*)
591.5 lb	29.5 mL (1 fl oz)

Neomycin Solution may be given undiluted or diluted with water.

**Here Treatment:** Each bottle will treat 9,454 pounds body weight. Therefore, estimate the total number of pounds body weight of the animals to be treated and administer one (1) fluid ounce for each 591.5 pounds. The product should be added to the amount of drinking water to be consumed in 12-24 hours. Provide medicated water as the sole source of water each day until consumed, followed by non-medicated water as required. Fresh medicated water should be prepared each day.

**Individual Animal Treatment:** To provide 10 mg neomycin sulfate per pound of body weight, mix 5 mL (1 tsp) in water or milk for each 100 pounds body weight. Administer daily either as a drench in divided doses or in the drinking water to be consumed in 12-24 hours.

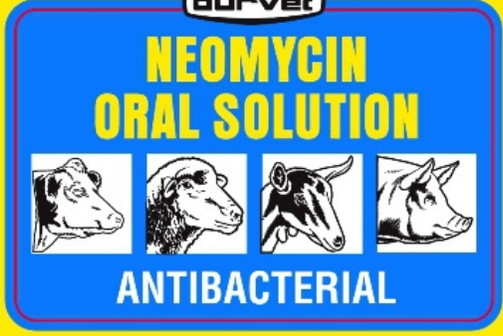
**PRECAUTION:** To administer the stated dosage, the concentration of neomycin required in medicated water must be adjusted to compensate for variation in age and weight of animal, the nature and severity of disease signs, and environmental temperature and humidity, each of which affects water consumption.

N-1050-05 Rev. 9-07

Manufactured for  
**DJURVET, INC.**  
Blue Springs, Missouri 64014  
www.durvvet.com



**TAKE TIME  
OBSERVE LABEL  
DIRECTIONS**



**NEOMYCIN  
ORAL SOLUTION**

**ANTIBACTERIAL**

**KEEP OUT OF REACH OF CHILDREN**

Contains per mL: neomycin sulfate 200 mg equivalent to 140 mg neomycin

If symptoms persist after using this preparation for 2 or 3 days, consult a veterinarian. If symptoms such as fever, depression, or going off feed develop, oral neomycin is not indicated as the sole treatment since systemic levels of neomycin are not obtained due to low absorption from the gastrointestinal tract.

**Important:** Treatment should continue 24 to 48 hours beyond remission of disease symptoms, but not to exceed a total of 14 consecutive days. Animals not drinking or eating should be treated individually by drench.

**Residue Warnings:** Not for human use. Keep out of reach of children. Discontinue treatment prior to slaughter by at least the number of days listed below for appropriate species:

Cattle .....	1 day
Sheep .....	2 days
Swine and Goat .....	3 days

A withdrawal period has not been established for this product in pre-ruminating calves. Do not use in calves to be processed for veal.

A milk discard period has not been established for this product in lactating dairy cattle. Do not use in female dairy cattle 20 months of age or older.

**Use of more than one product containing Neomycin or failure to follow withdrawal times may result in illegal drug residues.**

Store at controlled room temperature 20° to 25° C (68 to 77° F) (see USP).

\* Teaspoon (tsp) ; Tablespoon (tbsp) is equal to U.S. Standard Measure.

**ANADA#:** 200-379 Approved by F.D.A.

Lot No.                      Exp. Date



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**NEOMYCIN**

neomycin liquid

**Product Information**

<b>Product Type</b>	OTC ANIMAL DRUG LABEL	<b>Item Code (Source)</b>	NDC:30798-340
<b>Route of Administration</b>	ORAL	<b>DEA Schedule</b>	

**Active Ingredient/Active Moiety**

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
NEOMYCIN SULFATE (Neomycin)	Neomycin	140 mg in 1 mL

**Packaging**

<b>#</b>	<b>Item Code</b>	<b>Package Description</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:30798-340-31	473 mL in 1 BOTTLE		
2	NDC:30798-340-35	3823 mL in 1 JUG		

**Marketing Information**

<b>Marketing Category</b>	<b>Application Number or Monograph Citation</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
ANADA	ANADA200379	07/31/2007	

**Labeler** - Durvet, Inc. (056387798)

Revised: 1/2013

Durvet, Inc.