# MANNITOL 20% - mannitol injection Neogen Corporation

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

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NeogenVet Mannitol Injection 20%

## **INDICATIONS:**

Mannitol Injection 20% is indicated for use as an osmotic diuretic in canine species. Mannitol is essentially inert metabolically. When given parenterally, it is freely filtered at the glomerulus which produces osmotic diuresis as more than 90% of the mannitol injected escapes reabsorption.

## Each 100 mL Contains:

Mannitol USP......20 g
Water for Injection......q.s.

This solution contains 1098 m0smols/Liter

# **Dosage and Administration:**

The usual canine dosage administered intravenously is 1.5 - 2.0 g per Kg body weight given over a 30 minute period. This is approximately 3.4-4.5 mL/lb of body weight.

## Note:

Crystals of mannitol may form in a 20% saturated solution of mannitol. Dissolve the crystals by warming in hot water or autoclaving for 15 minutes. Cool to body temperature before administering. This is a single dose vial that contains no preservatives. Use entire contents when first opened.

# Store at temperatures between 15° and 30°C (59°-86°F)

RMS 92-384

Made in the USA

Item No. 09061

NDC 59051-8061-5

Mannitol Injection 20%

Sterile Solution

NeogenVet

Net Contentes: 100 mL

Lot No.

Exp. Date:

Neogen

Manufactured by: Nova-Tech, Grand Island, NE 68801

Manufactured for: Neogen Corporation, Lexington, KY 40511

859-254-1221

animalsafety.neogen.com

L566-0518

#### FOR ANIMAL USE ONLY

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

# **IKEEP OUT OF REACH OF CHILDREN**

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# **MANNITOL 20%**

mannitol injection

# **Product Information**

Product TypePRESCRIPTION ANIMAL DRUGItem Code (Source)NDC:59051-8061Route of AdministrationINTRAVENOUS

# **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
MANNITOL (UNII: 3OWL53L36A) (MANNITOL - UNII:3OWL53L36A)	MANNITOL	20 g in 100 mL

# **Inactive Ingredients**

Ingredient Name	Strength

WATER (UNII: 059QF0KO0R)

Packaging							
#	Item Code	Package Description	Marke	ting Start Date	Marketing End Date		
1	NDC:59051-8061-5	100 mL in 1 VIAL, SINGLE-USE					
Marketing Information							
Marketing Information							
M	arketing Category	Application Number or Monograph	Citation	Marketing Start Dat	Marketing End Date		
un	approved drug other			06/20/2011			

# Labeler - Neogen Corporation (042125879)

# Registrant - Nova-Tech, Inc (196078976)

Establishment							
Name	Address	ID/FEI	Business Operations				
Nova-Tech, Inc		196078976	manufacture				

Revised: 11/2019 Neogen Corporation