NEO-TERRAMYCIN 100/100D- neomycin-oxytetracycline powder Phibro Animal Health

Neo-Terramycin® 100/100D (neomycin-oxytetracycline)
TYPE A MEDICATED ARTICLE

(Antibiotic)

Active Drug Ingredients:

CAUTION:

CAUTION: Federal law restricts medicated feed containing this veterinary feed directive (VFD) drug to use by or on the order of a licensed veterinarian.

CAUTION: For use in manufacturing medicated animal feeds only.

CAUTION: Certain components of animal feeds, including medicated premixes, possess properties that may be a potential health hazard or a source of personal discomfort to certain individuals who are exposed to them. Human exposure should, therefore, be minimized by observing the general industry standards for occupational health and safety.

Precautions such as the following should be considered: dust masks or respirators and protective clothing should be worn; dust-arresting equipment and adequate ventilation should be utilized; personal hygiene should be observed; wash before eating or leaving a work site; be alert for signs of allergic reactions—seek prompt medical treatment if such reactions are suspected.

STORE IN A DRY, COOL PLACE

STORE AT OR BELOW 25°C (77°F), EXCURSIONS PERMITTED UP TO 40°C (104°F)

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Phibro Animal Health Corporation, Teaneck, NJ 07666

SEE BACK PANEL FOR COMPLETE MIXING DIRECTIONS
USE DIRECTIONS AND WARNINGS
Approved by FDA under NADA # 094-975
8853000
101-9076-08
Net Weight 50 lb (22.7 kg)
FOR USE IN DRY FEEDS ONLY. NOT FOR USE IN LIQUID FEED SUPPLEMENTS.

MIXING AND USE DIRECTIONS

Mix Neo-Terramycin 100/100D with nonmedicated milk replacer to provide the following drug concentrations:

Indications for Use	Oxytetracycline and Neomycin Amount	lb. of Neo-Terramycin 100/100D per ton ¹			
CALVES (milk replacer)					
For calves (up to 250 lb) for treatment of bacterial enteritis caused by <i>E. coli</i> susceptible to oxytetracycline; treatment and control of colibacillosis (bacterial enteritis) caused by <i>E. coli</i> susceptible to neomycin	10 mg/lb of body weight daily Feed continuously for 7-14 days Treatment should continue 24 to 48 hours beyond remission of disease symptoms.	20 ²			
CALVES, BEEF CATTLE, AND	CALVES, BEEF CATTLE, AND NONLACTATING DAIRY CATTLE				
Treatment of bacterial enteritis caused by <i>E. coli</i> and bacterial pneumonia (shipping fever complex) caused by <i>Pasteurella multocida</i> susceptible to oxytetracycline; treatment and control of colibacillosis (bacterial enteritis) caused by <i>E. coli</i> susceptible to neomycin	10 mg/lb of body weight daily Feed continuously for 7-14 days Treatment should continue 24 to 48 hours beyond remission of disease symptoms.	20 ²			

WARNING: A withdrawal period has not been established in preruminating calves; do not use in calves to be processed for veal. At the 10 mg/lb level, withdraw 5 days before slaughter.

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

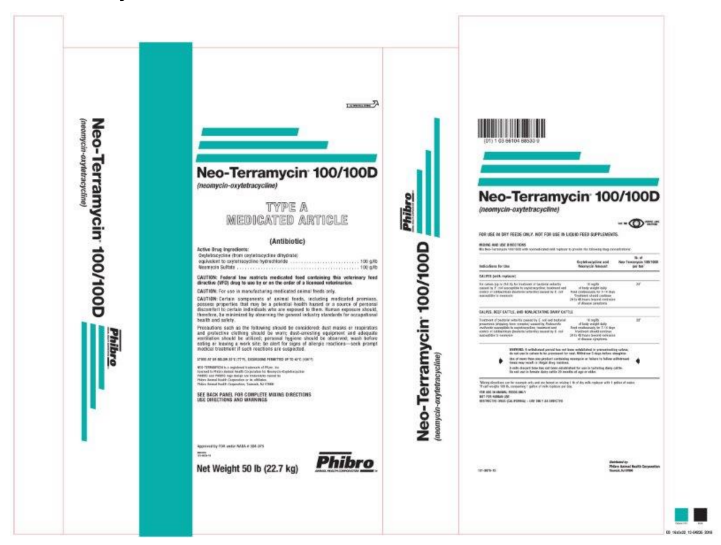
A milk discard time has not been established for use in lactating dairy cattle. Do not use in female dairy cattle 20 months of age or older.

 1 Mixing directions are for example only and are based on mixing 1 lb of dry milk replacer with 1 gallon of water.

 2 If calf weighs 100 lb, consuming 1 gallon of milk replacer per day.

FOR USE IN ANIMAL FEEDS ONLY NOT FOR HUMAN USE RESTRICTED DRUG (CALIFORNIA) - USE ONLY AS DIRECTED

Neo-Terramycin 100/100D



NEO-TERRAMYCIN 100/100D neomycin-oxytetracycline powder **Product Information** VFD TYPE A MEDICATED ARTICLE ANIMAL **Item Code** NDC:66104-**Product Type DRUG** 8853 (Source) **Route of Administration ORAL Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength OXYTETRACYCLINE HYDROCHLORIDE** (UNII: 4U7K4N52ZM) **OXYTETRACYCLINE** 100 g (OXYTETRACYCLINE ANHYDROUS - UNII:SLF0D9077S) **HYDROCHLORIDE** in 0.45 kg

NEOMYCIN SULFATE

100 g in 0.45 kg

Inactive Ingredients	
Ingredient Name	Strength
MINERAL OIL (UNII: T5L8T28FGP)	
SODIUM ALUMINOSILICATE (UNII: 058TS43PSM)	
SUCROSE (UNII: C151H8M554)	

P	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:66104-8853-0	22.7 kg in 1 BAG		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
NADA	NADA094975	07/08/2009		

Labeler - Phibro Animal Health (006989008)

Registrant - Phibro Animal Health (006989008)

Establishment			
Name	Address	ID/FEI	Business Operations
Sichuan Long March Pharmaceutical Co. Ltd.		544881741	api manufacture

Revised: 8/2023 Phibro Animal Health