DRS PHARMACY TRIPLE ANTIBIOTIC PAIN PLUS ITCH PLUS SCAR- bacitracin zinc, polymyxin b sulfate, neomycin sulfate ointment ointment OL PHARMA TECH, LLC

Active Ingredients

- Bacitracin Zinc 500 units
- Neomycin Sulfate 3.5mg
- Polymyxin B Sulfate 10,000 units
- pramoxine HCL 10mg

Purpose

First aid antibiotic

Uses

First aid to help prevent infection in minor:

- cuts
- scrapes
- burns

Warnings

For external use only

Ask a doctor

Ask a doctor before use if you have

- serious burns
- deep or puncture wounds
- animal bites

Stop use and ask a doctor

Stop use and ask a doctor if

- condition persists or gets worse
- rash or other allergic reaction develops

Do not use

- do not use in the eyes
- do not apply over large areas of the body
- if you are allergic to any of the ingredients
- longer than one week unless directed by your doctor

keep out of the reach of children

If swallowed, get medical help or contact a Poison Control Center right away

Directions

- clean the affected area
- apply a small amount of this product (an amount equal tot the surface area of the tip of a finer) on the area 1 to 3 times daily
- may be covered with a sterile bandage
- store at room temperature

Inactive ingredient

Petrolatum

Questions

www.drspharmacyusa.com

Principal display panel



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Product Type HUMAN OTC DRUG Item Code (Source) NDC:80489-412

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
BACITRACIN ZINC (UNII: 89Y4M234ES) (BACITRACIN - UNII:58H6RW052I)	BACITRACIN ZINC	500 [USP'U] in 1 g		
NEOMYCIN SULFATE (UNII: 057Y626693) (NEOMYCIN - UNII:116QD7X297)	NEOMYCIN	3.5 mg in 1 g		
POLYMYXIN B SULFATE (UNII: 19371312D4) (POLYMYXIN B - UNII: J2VZ 07J96K)	POLYMYXIN B	10000 [USP'U] in 1 g		
PRAMOXINE HYDROCHLORIDE (UNII: 88AYB867L5) (PRAMOXINE - UNII:068X84E056)	PRAMOXINE HYDROCHLORIDE	10 mg in 1 g		

Inactive Ingredients	
Ingredient Name	Strength
PETROLATUM (UNII: 4T6H12BN9U)	

Product Characteristics					
Color	white	Score			
Shape		Size			
Flavor		Imprint Code			
Contains					

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:80489-412- 01	1 in 1 CARTON	02/16/2021		
1		14 g in 1 TUBE; Type 0: Not a Combination Product			
2	NDC:80489-412- 02	1 in 1 CARTON	02/16/2021		
2		28.3 g in 1 TUBE; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M003	02/16/2021		

Labeler - OL PHARMA TECH, LLC (021170377)

Registrant - OL PHARMA TECH, LLC Drs PHARMACY (021170377)

Establishment					
Name	Address	ID/FEI	Business Operations		
OL PHARMA TECH, LLC Drs PHARMACY		021170377	manufacture(80489-412)		

Revised: 4/2024 OL PHARMA TECH, LLC