ZINC OXIDE- zinc oxide ointment ointment Dynarex Corporation

1190, 1191, 1192 Zinc Oxide Ointment 20%

Active Ingredient

Zinc Oxide 20%

Purpose

Skin Protectant

Use(s)

- Helps treat chafed skin associated with diaper rash
- Dries the oozing and weeping of poison ivy, oak, and sumac

Warnings

For External Use Only

When using this product

Avoid contact with eyes.

Stop use and ask a doctor if

• Condition worsens • Symptoms last more than 7 days or clear up and occur again within a few days

Keep out of reach of children

If swallowed, get medical help or contact a Poison Control Center (1-800-222-1222) right away.

Directions

- For diaper rash: Change wet and soiled diapers promptly, cleanse the diaper area, and allow to dry. Apply ointment liberally as often as necessary with each diaper change, especially at bedtime or anytime when exposure to wet diapers may be prolonged.
- For poison ivy, oak, and sumac: Apply as needed.

Other Information

• Store at room temperature between 15º-30ºC (59º-86ºF)

- Avoid excessive heat
- Tamper evident. Do not use if seal is damaged.

Inactive Ingredients

Cetomacrogol 1000. Cetostearyl Alcohol Light Liquid Paraffin, White Soft Paraffin

Questions?

1-888-Dynarex Monday - Friday, 9AM - 5PM EST.

Label



Label 1192

Label



Label



Label 1191

ZINC OXIDE

CETETH-20 (UNII: 1835H2IHHX)

CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)

ZINC OXIDE					
zinc oxide ointment ointment					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code	(Source)	NDC	::67777-223
Route of Administration	TOPICAL				
Active Ingredient/Active	Moiety				
Ingred	ient Name		Basis of Stren	gth	Strength
ZINC OXIDE (UNII: SOI2LOH54Z) (ZINC OXIDE - UNII:SOI2LOH	54Z)	ZINC OXIDE		200 mg in 1 g
Inactive Ingredients					
	Ingredient Name				Strength
WHITE PETROLATUM (UNII: B6E5	W8RQJ4)				
LIGHT MINERAL OIL (UNII: N6K57	87QVP)				

Product Characteristics				
Color		Score		
Shape	FREEFORM	Size		
Flavor		Imprint Code		
Contains				

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:67777-223- 02	72 in 1 CASE	02/29/2024		
1	NDC:67777-223- 11	1 in 1 BOX			
1	NDC:67777-223- 01	28.4 g in 1 TUBE; Type 0: Not a Combination Product			
2	NDC:67777-223- 04	72 in 1 CASE	02/29/2024		
2	NDC:67777-223- 13	1 in 1 BOX			
2	NDC:67777-223- 03	56.7 g in 1 TUBE; Type 0: Not a Combination Product			
3	NDC:67777-223- 06	12 in 1 CASE	02/29/2024		
3	NDC:67777-223- 05	425 g in 1 JAR; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M016	02/29/2024		

Labeler - Dynarex Corporation (008124539)

Revised: 2/2024 Dynarex Corporation