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**PROCURE Zinc oxide 20%** 

## **Active Ingredient**

Zinc Oxide 20% w/w

## **Purpose**

Skin Protectant

#### Uses

■ helps treat and prevent diaper rash
■ dries the oozing and weeping of
■ poison ivy

■ poison oak ■ poison sumac

## Warnings

For external use only

## When using this product

- do not get into eyes
- do not use over large areas of the body

# Stop use and ask a doctor if

■ the condition worsens ■ symptoms last more than 7 days or clear up and occur again in a few days ■ if you are allergic to any of these ingredients

**Keep out of reach of children.** If swallowed, get medical help or contact a Poison Control Center right away

#### **Directions**

■ for diaper rash  $\blacksquare$  change wet and soiled diapers promptly  $\blacksquare$  cleanse the diaper area and allow to dry  $\blacksquare$  apply ointment liberally with each diaper change, especially at bedtime or anytime when exposure to wet diapers may be prolonged  $\blacksquare$  for poison ivy, oak sumac  $\blacksquare$  apply ointment liberally as often as needed

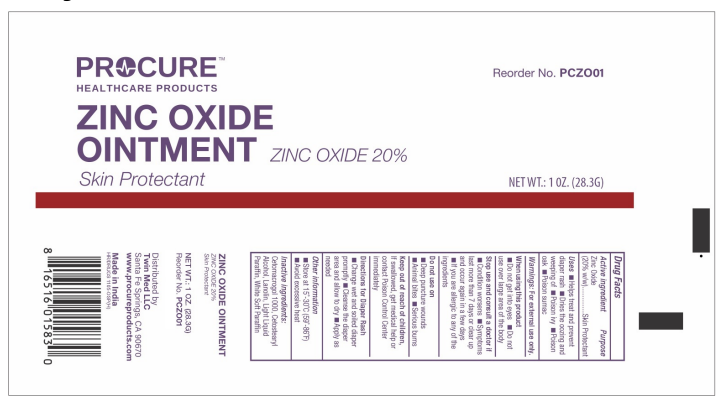
#### Other Information

■ store at 59-86F (15-30C)

## **Inactive Ingredients**

cetostearyl alcohol, cetomacrogol 1000, lanolin, light liquid paraffin, white soft parrafin

## Package Label



## **PROCURE**

zinc oxide ointment

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:55681-304	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety			
Ingredient Name	<b>Basis of Strength</b>	Strength	
ZINC OXIDE (UNII: SOI2LOH54Z) (ZINC OXIDE - UNII:SOI2LOH54Z)	ZINC OXIDE	20 g in 100 g	

Inactive Ingredients		
Ingredient Name	Strength	
LANOLIN (UNII: 7EV65EAW6H)		
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)		
CETETH-20 (UNII: 1835H2IHHX)		
LIGHT MINERAL OIL (UNII: N6K5787QVP)		
PETROLATUM (UNII: 4T6H12BN9U)		

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:55681-304- 03	28.3 g in 1 TUBE; Type 0: Not a Combination Product	05/23/2013			
2	NDC:55681-304- 04	56.6 g in 1 TUBE; Type 0: Not a Combination Product	05/23/2013			
3	NDC:55681-304- 05	425 g in 1 JAR; Type 0: Not a Combination Product	05/23/2013			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M016	05/23/2013		

# **Labeler -** TWIN MED, LLC (009579330)

Revised: 1/2024 TWIN MED, LLC