CLEAR ANTI-ITCH- pramoxine hcl, zinc acetate lotion Target Cop

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Up & Up 218.001-218AE

Active ingredients

Pramoxine HCI 1%

Zinc acetate 0.1%

Purpose

External analgesic

Skin protectant

Uses

- •for the temporary relief of pain and itching associated with minor skin irritations and rashes due to poison ivy, poison oak, or poison sumac
- •dries the oozing and weeping of poison: •ivy •oak •sumac

Warnings

For external use only

When using this product

• do not get into 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 right away.

Directions

- shake well
- befor applying wash affected area of skin

adults and children 2 years of age and older - apply to affected area not more than 3 to 4 times daily

children under 2 years - ask a doctor

Other information

store at room temperature (59°-77° F)

Inactive ingredients

alcohol, camphor, citric acid, diazolidinyl urea, fragrance, glycerin, hydroxypropyl methylcellulose, methylparaben, oil of lavender, oil of rosemary, polysorbate 40, propylene glycol, propylparaben, purified water, sodium citrate

Questions

Call 1-800-910-6874

This product is not manufactured or distributed by Johnson & Johnson Consumer Products Company, Distributor of Caladryl Clear Lotion*

Dist. by Target Corp., Mpls., MN 55403 Made in U.S.A. with U.S. and foreign components 2015 Target Brands, Inc. Shop Target.com

principal display panel

NDC 11673-218-30

Compare to active ingredints in Caladryl

clear anti-tich lotion

external analgesic/skin protectant

drying action plus itch relief

up & up

6 FL OZ (177 mL)



CLEAR ANTI-ITCH

pramoxine hcl, zinc acetate lotion

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:11673-218
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
PRAMOXINE HYDROCHLORIDE (UNII: 88AYB867L5) (PRAMOXINE - UNII:068X84E056)	PRAMOXINE HYDROCHLORIDE	10 mg in 1 mL	
ZINC ACETATE (UNII: FM5526K07A) (ZINC CATION - UNII:13S1S8SF37)	ZINC ACETATE	1 mg in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
ALCOHOL (UNII: 3K9958V90M)			
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)			
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)			
DIAZOLIDINYL UREA (UNII: H5RIZ 3MPW4)			
GLYCERIN (UNII: PDC6A3C0OX)			
HYPROMELLOSES (UNII: 3NXW29V3WO)			
METHYLPARABEN (UNII: A2I8C7HI9T)			
LAVENDER OIL (UNII: ZBP1YXW0H8)			

ROSEMARY OIL (UNII: 8LGU7VM393)	
POLYSORBATE 40 (UNII: STI11B5A2X)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
PROPYLPARABEN (UNII: Z8IX2SC10H)	
WATER (UNII: 059QF0KO0R)	
SODIUM CITRATE (UNII: 1Q73Q2JULR)	

ı	Packaging			
	# Item Code	# Item Code Package Description		Marketing End Date
	1 NDC:11673- 218-30	177 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	03/31/1996	

Marketing Information			
Marketing Application Number or Monograph Marketing Start Mar Category Citation Date		Marketing End Date	
OTC monograph not final	part347	03/31/1996	
IIIIdi			

Labeler - Target Cop (006961700)

Registrant - Vi-Jon, LLC (790752542)

Establishment			
Name	Address	ID/FEI	Business Operations
Vi-Jon, LLC		790752542	manufacture(11673-218)

Establishment			
Name	Address	ID/FEI	Business Operations
Vi-Jon, LLC		088520668	manufacture(11673-218)

Revised: 12/2022 Target Cop