

ANTI ITCH TOPICAL ANALGESIC- diphenhydramine hydrochloride, zinc acetate cream
Chain Drug Marketing Associations Inc

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.

Quality Choice Extra Strength Itch Stopping Cream 1 oz. 94731 ZDP 2020

Active Ingredients Purpose

Diphenhydramine HCl 2%.....Topical analgesic

Zinc acetate, 0.1%..... Skin protectant

Uses

- temporarily relieves pain and itching associated with:
- insect bites
- minor burns
- minor skin irritations
- sunburn
- minor cuts
- scrapes
- rashes due to poison ivy, oak, and sumac
- dries the oozing and weeping of poison:
- ivy
- oak
- sumac

Warnings

For external use only

Do not use

- with any other product containing diphenhydramine, even one taken by mouth
- on large areas of the body

Ask a doctor before use

- on chicken pox
- on measles

When using this product □avoid contact with the eyes

Stop use and ask a doctor if

- condition worsens or does not improve within 7 days
- symptoms persist for more than 7 days or clear up and occur again within a few days

If pregnant or breast-feeding, ask a health professional before use.

Keep out of reach of children

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

Directions

- do not use more than directed
- 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 of age: ask a doctor

Other information

- store at room temperature 20-25°C (68-77°F)
- protect from excessive heat (40°C/104°F), humidity, and light

inactive ingredients

cetostearyl alcohol, glyceryl stearate/peg-100 stearate, methylparaben, propylene glycol, propylparaben, purified water

DISTRIBUTED BY:

C.D.M.A. INC.

43157 W. 9 Mile Road

Novi, MI 48375

www.qualitychoice.com

Made in China



NDC 63868-708-28

*Compare to the Active Ingredients in Benadryl® Extra Strength Itch Stopping Cream

Extra Strength Itch Stopping Cream

Relief from most outdoor itches

Diphenhydramine HCl 2% (Topical Analgesic)
Zinc Acetate 0.1% (Skin Protectant)

Temporary Relief of Itching & Pain from Minor Skin Irritations & Rashes due to Insect Bites, Poison Ivy, Oak & Sumac

Extra Strength Itch Stopping Cream

Relief from most outdoor itches

Diphenhydramine HCl 2% (Topical Analgesic)
Zinc Acetate 0.1% (Skin Protectant)



NET WT. 1 OZ (28 g)

Extra Strength Itch Stopping Cream



Distributed by C.D.M.A., Inc.®
43157 W 9 Mile Rd
Novi, MI 48375
www.qualitychoice.com
Questions: 800-935-2362

MADE IN CHINA

94731-ZDP-BX-R1
OM-MD-ZDP124-R1

LOT & EXP.

This product is not manufactured or distributed by Johnson & Johnson Consumer Inc., distributor of Benadryl® Extra Strength Itch Stopping Cream.

Drug Facts (continued)

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

Directions

- do not use more than directed
- adults and children 2 years of age and older: apply to affected areas not more than 3 to 4 times daily
- children under 2 years of age: ask a doctor

Other information

- store at room temperature 20-25°C (68-77°F)
- protect from excessive heat (40°C/104°F), humidity, and light

Inactive ingredients Cetearyl alcohol, glyceryl stearate, PEG-100 stearate, methylparaben, propylene glycol, propylparaben, purified water.

Drug Facts

Active ingredients Diphenhydramine HCl 2%, Topical analgesic
Zinc acetate 0.1%, Skin protectant

Purposes

Uses

- temporarily relieves pain and itching associated with:
 - insect bites ■ minor burns ■ minor skin irritations
 - sunburn ■ minor cuts ■ scrapes
 - rashes due to poison ivy, oak and sumac
- dress the sores and weeping of poison:
 - ivy ■ oak ■ sumac

Warnings

For external use only

Do not use

- with any other product containing diphenhydramine, even one taken by mouth ■ on large areas of the body
- on chicken pox ■ on measles

Ask a doctor before use

- on chicken pox ■ on measles

When using this product, avoid contact with eyes.

Stop use and ask a doctor if

- symptoms persist for more than 7 days or clear up and occur again within a few days

KEEP OUTER CAP FOR COMPLETE INFORMATION.

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Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:63868-708
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Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
DIPHENHYDRAMINE HYDROCHLORIDE (UNII: TC2D6JAD40) (DIPHENHYDRAMINE - UNII:8GTS82S83M)	DIPHENHYDRAMINE HYDROCHLORIDE	2 g in 100 g
ZINC ACETATE (UNII: FM5526K07A) (ZINC CATION - UNII:13S1S8SF37)	ZINC ACETATE	0.1 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0K00R)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)	
GLYCERYL STEARATE/PEG-100 STEARATE (UNII: RD25J5V947)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:63868-708-28	1 in 1 CARTON	04/08/2020	
1		28 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 not final	part348	04/08/2020	

Labeler - Chain Drug Marketing Associations Inc (011920774)

Revised: 7/2020

Chain Drug Marketing Associations Inc