# BENADRYL EXTRA STRENGTH ITCH STOPPING- diphenhydramine hydrochloride gel Johnson & Johnson Consumer Inc.

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# Benadryl® EXTRA STRENGTH ITCH STOPPING GEL

# **Drug Facts**

## **Active ingredient**

Diphenhydramine HCl 2%

## **Purpose**

**Topical Analgesic** 

#### Use

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

# **Warnings**

For external use only.

## Do not use

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

#### 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

- condition worsens
- symptoms persist for 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**

- 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

## **Inactive ingredients**

SD alcohol 38-B, camphor, citric acid, diazolidinyl urea, glycerin, hydroxypropyl methylcellulose, methylparaben, propylene glycol, propylparaben, purified water, sodium citrate

## Questions or comments?

call **1-877-717-2824** (toll free) or **215-273-8755** (collect) www.benadryl.com

Distributed by:

JOHNSON & JOHNSON CONSUMER INC.

Skillman, NJ 08558

## PRINCIPAL DISPLAY PANEL - 103 mL Tube Label

**Benadryl®** 

®

**Cooling relief for** 

most outdoor itches

**EXTRA STRENGTH** 

For Skin Use Only

**ITCH STOPPING** 

GEL

**Diphenhydramine HCI 2% /** 

**Topical Analgesic** 

Poison Minor Insect Ivy, Mosquito Cuts Sunburn

# 3.5 FL OZ (103 mL)



# BENADRYL EXTRA STRENGTH ITCH STOPPING

diphenhydramine hydrochloride gel

<b>Product Information</b>	oduct Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69968-0242		
Route of Administration	TOPICAL				

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
DIPHENHYDRAMINE HYDROCHLORIDE (UNII: TC2D6JAD40) (DIPHENHYDRAMINE - UNII:8GTS82S83M)	DIPHENHYDRAMINE HYDROCHLORIDE	20 mg in 1 mL		

Inactive Ingredients		
Ingredient Name	Strength	
ALCOHOL (UNII: 3K9958V90M)		
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)		
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)		
DIAZOLIDINYL UREA (UNII: H5RIZ3MPW4)		
GLYCERIN (UNII: PDC6A3C0OX)		
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)		
METHYLPARABEN (UNII: A2I8C7HI9T)		
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)		
PROPYLPARABEN (UNII: Z8IX2SC10H)		
WATER (UNII: 059QF0KO0R)		
SODIUM CITRATE, UNSPECIFIED FORM (UNII: 1Q73Q2JULR)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69968- 0242-3	103 mL in 1 TUBE; Type 0: Not a Combination Product	10/01/2013	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	10/01/2013	

Labeler - Johnson & Johnson Consumer Inc. (118772437)

Revised: 1/2024 Johnson & Johnson Consumer Inc.