# GOOD NEIGHBOR PHARMACY ITCH RELIEF- diphenhydramine hcl, zinc acetate spray AMERISOURCEBERGEN DRUG CORPORATION

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.

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### Good Neibor Pharmacy Extra Strength Itch Relief Spray

### **Active Ingredient**

Diphenhydramine hydrochloride 2% Topical analgesic

Zinc acetate 0.1% Skin protectant

#### Uses

for

temporary relieves pain and itching due to:

insect bites

minor burns

sunburn

minor skin irritations

minor cuts

scrapes

rashes due to poison ivy, poison oak, and poison sumac

dries the oozing and weeping of poison:

ivy

oak

sumac

#### **Directions**

do not use more often than directed

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

children under 2 years: ask a doctor

#### 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 doctor Before use

on chicken pox on measles

### When using this product

do not get into eyes

### Stop Use and ask physician

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.

#### Warning

for external use only

#### **Purposes**

Itch relief

### **Inactive Ingredients**

Purified Water, Glycerin, SD Alcohol, Povidone, Povidonee (K-30), Trolamine

#### Prinicpal dispaly panel



Diphenhydramine hydrochloride 2%

Topical analgesic

Zinc acetate 0.1% Skin protectant

Relieves Itch and Pain associated with insects bites & rashes due to posinon ivy Oak& sumac

#### GOOD NEIGHBOR PHARMACY ITCH RELIEF

diphenhydramine hcl, zinc acetate spray

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:46122-572	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ZINC ACETATE (UNII: FM5526K07A) (ZINC CATION - UNII:13S1S8SF37)	ZINC ACETATE	0.1 g in 100 mL	
<b>DIPHENHYDRAMINE HYDRO CHLO RIDE</b> (UNII: TC2D6JAD40) (DIPHENHYDRAMINE - UNII:8GTS82S83M)	DIPHENHYDRAMINE HYDROCHLORIDE	2 g in 100 mL	

Inactive Ingredients			
Ingredient Name	Strength		
WATER (UNII: 059QF0KO0R)			
GLYCERIN (UNII: PDC6A3C0OX)			
ALCOHOL (UNII: 3K9958V90M)			
PO VIDO NE K30 (UNII: U725QWY32X)			
TROLAMINE (UNII: 903K93S3TK)			

ı	Packaging			
	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:46122-572- 46	59 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	04/14/2014	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part348	0 4/14/20 14	

# Labeler - AMERISOURCEBERGEN DRUG CORPORATION (007914906)

## **Registrant -** Weeks & LEO, Inc. (005290028)

Establishment				
Name	Address	ID/FEI	Business Operations	
Weeks & Leo		005290028	manufacture(46122-572)	

Revised: 4/2019 AMERISOURCEBERGEN DRUG CORPORATION