# CLEAR ANTI-ITCH- pramoxine hcl, zinc acetate lotion Perrigo Direct, Inc.

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Good Sense 218.002/218AF Clear Anti-Itch Lotion

#### **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 section 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
- before 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, benzoic acid, camphor, citric acid, fragrance, glycerin, hydroxypropyl methylcellulose, Lavandula angustifolia (lavender) oil, polysorbate 40, Rosmarinus officinalis (rosemary) leaf oil, sodium citrate, water

#### **Disclaimer**

\*This product is not manufactured or distributed by Bausch Health US, LLC, distributor of Caladryl® Clear® Lotion.

#### Adverse reaction

Distributed by: Perrigo Direct, Inc.

Peachtree City, GA 30269

www.PerrigoDirect.com

1-888-593-0593

GoodSense is a registed trademark of L. Perrigo Company.

## **Principal Display Panel**

NDC 75981-218-30

**GoodSENSE®** 

Clear Anti-Itch Lotion

External Analgesic/Skin Protectant

Pain and Itch Reliever

Compare to active ingredients of Caladryl® Clear® Lotion\*

6 FL OZ (177 mL)



#### **CLEAR ANTI-ITCH**

pramoxine hcl, zinc acetate lotion

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:75981-218

Route of Administration TOPICAL

### **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>PRAMOXINE HYDROCHLORIDE</b> (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)		
BENZOIC ACID (UNII: 8SKN0B0MIM)		
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)		
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)		
GLYCERIN (UNII: PDC6A3C0OX)		
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)		
LAVENDER OIL (UNII: ZBP1YXW0H8)		
POLYSORBATE 40 (UNII: STI11B5A2X)		
ROSEMARY OIL (UNII: 8LGU7VM393)		
SODIUM CITRATE (UNII: 1Q73Q2JULR)		
WATER (UNII: 059QF0KO0R)		

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:75981- 218-30	177 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	07/24/2008	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M016	07/24/2008	

## Labeler - Perrigo Direct, Inc. (076059836)

## **Registrant -** Consumer Product Partners, LLC (119091520)

Establishment				
Name	Address	ID/FEI	<b>Business Operations</b>	
Consumer Product Partners. LLC		119091514	manufacture(75981-218)	

Revised: 6/2024 Perrigo Direct, Inc.