## I.C. IVY BLOCK- zinc acetate lotion R & R Lotion, 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.

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### I.C.<sup>®</sup> Ivy Block

**Drug Facts** 

## **Active Ingredients**

Zinc Acetate 0.1%

### Purpose

Skin Protectant

#### Uses

Poison Ivy, Oak, Sumac Protectant

## Warnings

For external use only - When using this product

- do not get into eyes
- If contact occurs, rinse eyes thoroughly with water
- 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 before use

- Apply 15 minutes before exposure
- Apply generously to dry, exposed skin
- Allow to dry
- Apply every 4 hours for continued protection or sooner if needed
- After possible exposure with skin irritants, remove using IC Ivy Cleanser
- Children under 2 year ask a doctor.

## **Other Information**

Protect from excessive heat (48°C/120°F)

### **Inactive Ingredients**

Di Water, Cyclopentasiloxane (and) Quaternium - 90 Bentonite (and) Trifluoromethyl C1-4 Alkyl Dimethicone (and) Propylene Carbonate, Bentonite, Dimethicone, Polyacrylamide (and) c13-14 Isoparaffin (and) Laureth-7, Glycerin, Magnesium Aluminum Silicate, Phenoxyethanol (and) Ethylhexlglycerin, PVP, Aloe Barbadensis Leaf Juice, Disodium EDTA, Hamamelis Virginiana (Witch Hazel) Water (and) Phenoxyethanol, Citric Acid

### **Questions or Comments?**

1-888-860-7424

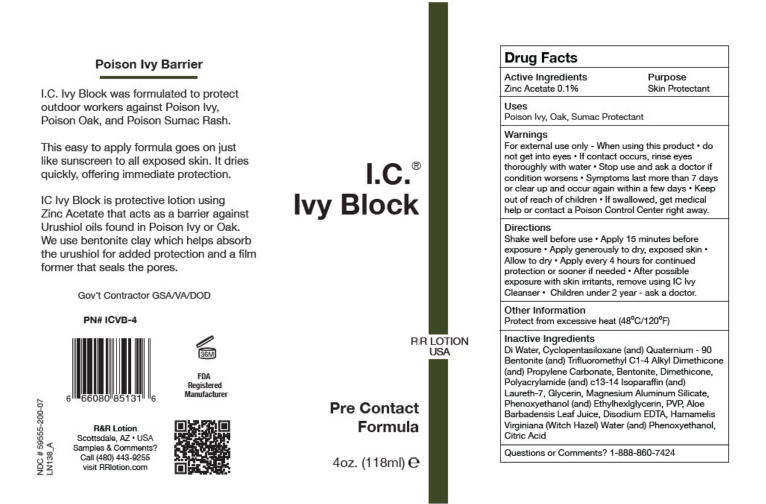
## **PRINCIPAL DISPLAY PANEL - 118 ml Bottle Label**

I.C.<sup>®</sup> Ivy Block

R&R LOTION USA

Pre Contact Formula

4oz. (118ml)  $\mathrm{e}$ 



zinc acetate lotion						
Product Information						
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:5	IDC:59555-200	
Route of Administration	TOPICAL					
Active Ingredient/Active	Moiety					
Ingredient Name Basis of Strengt			igth	Strength		
ZINC ACETATE (UNII: FM5526K07A) (ZINC CATION - UNII:13S1S8SF37) ZINC ACETATE				1 mg in 1 ml		
Inactive Ingredients					Ctwo worth	
Inactive Ingredients	Ingredient Name				Strength	
-	•				Strength	
CYCLOMETHICONE 5 (UNII: 0TH QUATERNIUM-90 (UNII: BE5VYX4	T5PCIOR)				Strength	
	T5PCIOR) IVP8)				strengtn	

PROPYLENE CARBONATE (UNII: 8D08K3S51E)	
POLYACRYLAMIDE (10000 MW) (UNII: E2KR9C9V2I)	
C13-14 ISOPARAFFIN (UNII: E4F12ROE70)	
LAURETH-7 (UNII: Z95S6G8201)	
GLYCERIN (UNII: PDC6A3C0OX)	
MAGNESIUM ALUMINUM SILICATE (UNII: 6M3P64V0NC)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)	
POVIDONE, UNSPECIFIED (UNII: FZ989GH94E)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
EDETATE DISODIUM (UNII: 7FLD91C86K)	
HAMAMELIS VIRGINIANA TOP (UNII: UDA30A2JJY)	
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)	

## **Product Characteristics**

Color	WHITE	Score
Shape		Size
Flavor		Imprint Code
Contains		

# Packaging

#	ltem Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:59555- 200-03	50 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	09/21/2021			
2	NDC:59555- 200-04	59 mL in 1 BOTTLE, DISPENSING; Type 0: Not a Combination Product	09/21/2021			
3	NDC:59555- 200-07	118 mL in 1 BOTTLE, DISPENSING; Type 0: Not a Combination Product	09/21/2021			
4	NDC:59555- 200-08	236 mL in 1 BOTTLE, DISPENSING; Type 0: Not a Combination Product	09/21/2021			
5	NDC:59555- 200-10	946 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	09/21/2021			
6	NDC:59555- 200-11	3785 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	09/21/2021			
7	NDC:59555- 200-01	5 mL in 1 PACKET; Type 0: Not a Combination Product	09/21/2021			
8	NDC:59555- 200-16	4 mL in 1 PACKET; Type 0: Not a Combination Product	09/21/2021			
Marketing Information						
	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		

09/21/2021

OTC monograph final part347

Revised: 9/2021