AFTER BITE WIPE INSECT STING RELIEF- benzocaine, alcohol swab Adventure Ready Brands

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.

After Bite Wipe Insect Sting Relief

Active Ingredients

Benzocaine, 6%

SD alcohol, 60%

Purpose

Topical Analgesic Antiseptic

Use

- temporarily relieves pain and itching due to minor stings and insect bites
- to help reduce bacteria on the skin

Warnings

For external use only.

Flammable, keep away from fire or flame.

Do not use

- over large areas of the body
- in the eyes
- on broken skin or deep puncture wounds

Stop use and ask a doctor if

condition worsens or if 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

- adults and children 2 year of age and older: apply to affected area not more than 3 to 4 times daily.
- children under 2 years of age: consult a doctor

Other information

Store at room temperature

Inactive Ingredient

Purified water

Drug Facts (continued)

Warnings

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Inactive ingredient Purified Water

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LOT:



Adventure Ready Brands 944 Industrial Park Rd. Littleton, NH 03561

Drug Facts

Active ingredients

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AFTER BITE WIPE INSECT STING RELIEF

benzocaine, alcohol swab

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:90107-3623

TOPICAL **Route of Administration**

Active Ingredient/Active Moiety

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I	Ingredient Name	Basis of Strength	Strength		
I	BENZOCAINE (UNII: U3RSY48JW5) (BENZOCAINE - UNII:U3RSY48JW5)	BENZOCAINE	6 mg in 100 mg		
I	ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	60 mg in 100 mg		

Inactive Ingredients

Ingredient Name	Strength
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WATER (UNII: 059QF0KO0R)

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;	t Item Code	Package Description	Marketing Start Date	Marketing End Date
		1000 mg in 1 PACKAGE; Type 0: Not a Combination Product	12/0 1/20 20	
	NDC:90107-3623-	100 in 1 BOX	12/0 1/20 20	

2 1	000 mg in 1 PACKET; Type 0: Not a Combination Produc	t			
Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not fina	l part333E	12/01/2020			

Labeler - Adventure Ready Brands (064437304)

Revised: 12/2020 Adventure Ready Brands