

**FIRST AID ONLY STING RELIEF PAD- benzocaine, isopropyl alcohol liquid**  
**Acme United Corporation**

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**First Aid Only Sting Relief Pad**

***Drug Facts***

***Active ingredients***

Benzocaine 6.0%

Isopropyl Alcohol 60% w/v

**Purpose**

Topical Anesthetic

Antiseptic

For the temporary relief of pain and itching associated with minor scrapes and insect bites.

First aid to help prevent infection in minor cuts, scrapes, and burns.

**Warnings**

For external use only

Flammable, keep away from fire or flame

***Do Not Use***

- **in eyes, if contact occurs flush with water ■over large areas of the body**

Consult a doctor ■if condition worsens ■if symptoms last for more than seven days or clear up and occur again ■for deep puncture wounds

**Keep out of reach of children.**

If swallowed, get medical help or contact a Poison Control center right away.

***Directions***

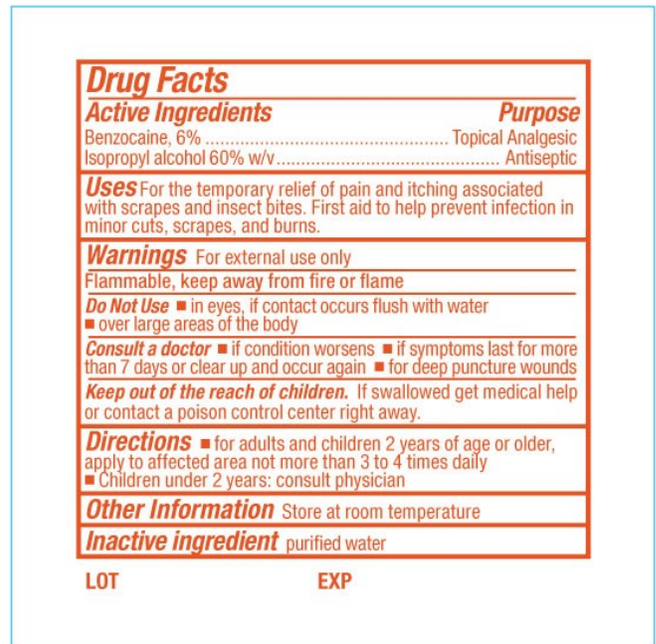
■ For adults and children 2 years of age and older, apply to affected area not more than 3 to 4 times daily

■Children under 2 years: consult physician

**Inactive ingredient**

Purified Water

**Other information** store at room temperature



pouch label

<b>FIRST AID ONLY STING RELIEF PAD</b>				
benzocaine, isopropyl alcohol liquid				
<b>Product Information</b>				
<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:0924-5204(NDC:59050-414)	
<b>Route of Administration</b>	TOPICAL			
<b>Active Ingredient/Active Moiety</b>				
Ingredient Name		Basis of Strength	Strength	
BENZOCAINE (UNII: U3RSY48JW5) (BENZOCAINE - UNII:U3RSY48JW5)		BENZOCAINE	60 mg in 1 mL	
ISOPROPYL ALCOHOL (UNII: ND2M416302) (ISOPROPYL ALCOHOL - UNII:ND2M416302)		ISOPROPYL ALCOHOL	600 mg in 1 mL	
<b>Inactive Ingredients</b>				
Ingredient Name			Strength	
WATER (UNII: 059QF0KO0R)				
<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC 0924			

1	NDC:0924-5204-05	100 in 1 BOX	02/25/2022	
1		0.4 mL in 1 POUCH; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)		
2	NDC:0924-5204-04	50 in 1 BOX	02/25/2022	
2		0.4 mL in 1 POUCH; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)		
3	NDC:0924-5204-03	25 in 1 BOX	02/25/2022	
3		0.4 mL in 1 POUCH; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)		
4	NDC:0924-5204-02	10 in 1 BOX	02/25/2022	
4		0.4 mL in 1 POUCH; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)		
5	NDC:0924-5204-01	0.4 mL in 1 POUCH; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)	02/25/2022	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M003	02/25/2022	

**Labeler** - Acme United Corporation (001180207)

## Establishment

Name	Address	ID/FEI	Business Operations
Acme United Corporation		045924339	relabel(0924-5204) , repack(0924-5204)

## Establishment

Name	Address	ID/FEI	Business Operations
Acme United Corporation		080119599	relabel(0924-5204) , repack(0924-5204)

## Establishment

Name	Address	ID/FEI	Business Operations
Acme United Corporation		117825595	manufacture(0924-5204)

Revised: 11/2024

Acme United Corporation