

STING RELIEF PAD- benzocaine, isopropyl alcohol swab

Acme United 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.

First Aid Only Sting Relief Pad

Drug Facts

Active Ingredients

Benzocaine, 6%

Isopropyl alcohol 60% w/v

Purpose

Topical Analgesic

Antiseptic

Use

For the temporary relief of pain and itching associated with minor burns, 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

Keep out of the 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 or older, apply to affected area not more than 3 to 4 times daily
- Children under 2 years: consult physician

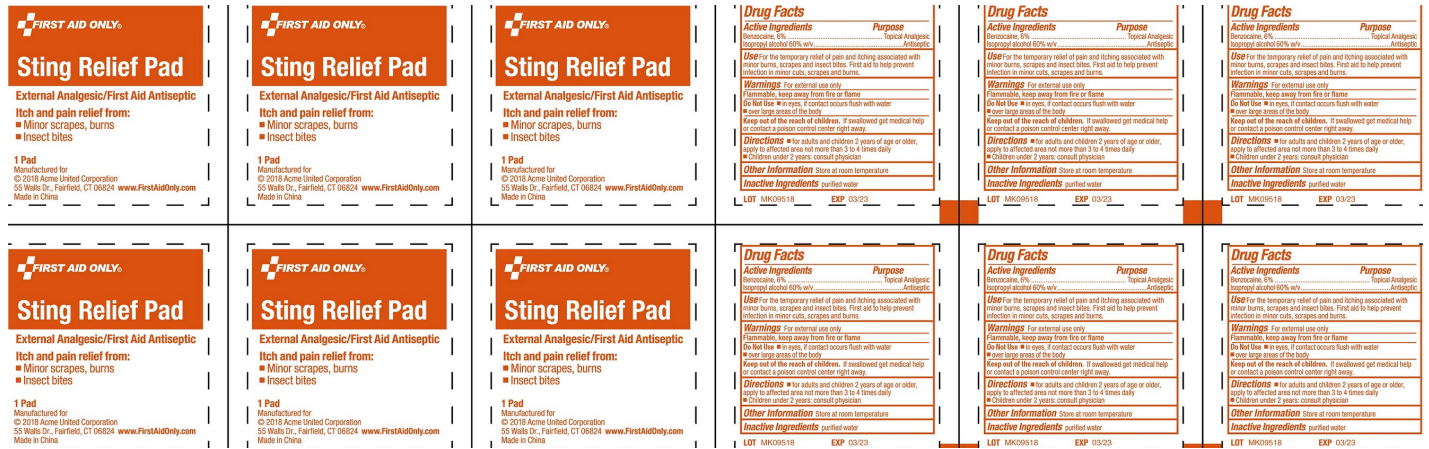
Other Information

Store at room temperature

Inactive Ingredients

purified water

Package Labeling:



STING RELIEF PAD

benzocaine, isopropyl alcohol swab

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:0924-5202(NDC:59050-059)
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZOCAINE (UNII: U3RSY48JW5) (BENZOCAINE - UNII:U3RSY48JW5)	BENZOCAINE	60 mg in 1 g
ISOPROPYL ALCOHOL (UNII: ND2M416302) (ISOPROPYL ALCOHOL - UNII:ND2M416302)	ISOPROPYL ALCOHOL	600 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0K00R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0924-5202-01	0.34 g in 1 POUCH; Type 0: Not a Combination Product	05/09/2018	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part348	05/09/2018	

Labeler - Acme United Corporation (001180207)

Registrant - Acme United Corporation (001180207)

Establishment

Name	Address	ID/FEI	Business Operations
Acme United Corporation		045924339	relabel(0924-5202) , repack(0924-5202)

Establishment

Name	Address	ID/FEI	Business Operations
Acme United Corporation		080119599	repack(0924-5202) , relabel(0924-5202)

Revised: 5/2018

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