ALBERTSONS PAIN RELIEF PATCH- lidocaine patch Safeway

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.

Albertsons Signature Care Pain Relief Patch with 4% Lidocaine

Active Ingredients

Lidocaine 4% Topical Anesthetic

Purpose

TOPICAL ANESTHETIC

Idications and Usage

Uses: Temporaily relieves minor pains.

Warnings

For external use only.

Do not Use

- more than one patch on your body at a time
- on cut, irritated or swollen skin
- on puncture wounds
- for more than one week without consulting a doctor
- if you are allergic to any active or inactive ingredients
- if pouch is damaged or opened.

When using this product

- use only as directed
- read and follow all directions and warnings on this carton
- do not allow contact with the eyes
- do not use at the same time as other topical analgesics
- do not bandage tightly or apply local heat (such as heating pads) to the area of use

- do not microwave
- dispose of used patch in manner that always keeps product away from children and pets. Used patches still contain the drug product that can produce serious adverse effects if a child or pet chews or ingests this patch.

Stop use and consult a doctor

- condition worsens
- redness is present
- irritation develops
- symptoms persist for more than 7 days or clear up and occur again within a few days
- you experience signs of skin injury, such as pain, swelling or blistering where the product was applied.

If pregnant or breastfeeding, ask a health professional before use.

Keep out of reach of children and pets.

If swallowed, get medical help or contact a Poison Control Center at 1-800-222-222 right away.

Dosage and Administration

Directions Adults and children over 12 years:

- clean and dry affected area
- carefully remove backing from patch starting at a corner
- apply sticky side of patch to affected area
- use one patch for up to 12 hours.
- Discard patch after single use
- Children under 12 years of age: consult a physician.

Other Safety Information

Store in a clean, dry place outside of direct sunlight. Protect from excessive moisture.

Inactive Ingredients

Aluminum Glycinate, Carboxymethylcellulose Sodium, Glycerin, iodopropynyl butylcarbamate, Kaolin, petrolatum, phenoxyethanol, polyacrylic acid, polysorbate 80, Povidone, Propylene Glycol, Sodium Polyacrylate, Tartaric Acid, Titanium Dioxide, Water, 3-(2-ethylhexyloxy) propane-1,2-diol.

Questions

TOLL-FREE

Customer Care Help Line 1-888-423-0139 Mon-Fri 8:30 am-4:30 pm CST

DIST. BY TOPCO ASSOCIATES LLC ELK GROVE VILLAGE, IL 60007 www.topcarebrand.com

Made in China

Albertsons Pain Relief Patch with 4% Lidocaine



Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:21130-960	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
LIDOCAINE (UNII: 98PI200987) (LIDOCAINE - UNII:98PI200987)	LIDOCAINE	4 g in 100 g

Inactive Ingredients	
Ingredient Name	Strength
PETROLATUM (UNII: 4T6H12BN9U)	
CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM (UNII: K6790BS311)	
DIHYDROXYALUMINUM AMINOACETATE (UNII: DO250MG0W6)	
GLYCERIN (UNII: PDC6A3C0OX)	
KAOLIN (UNII: 24H4NWX5CO)	
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)	
POLYACRYLIC ACID (8000 MW) (UNII: 73861X4K5F)	
POLYSORBATE 80 (UNII: 6OZP39ZG8H)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
POVIDONE, UNSPECIFIED (UNII: FZ 989GH94E)	
SODIUM POLYACRYLATE (8000 MW) (UNII: 285CYO341L)	
TARTARIC ACID (UNII: W4888I119H)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
WATER (UNII: 059QF0KO0R)	
IODOPROPYNYL BUTYLCARBAMATE (UNII: 603P14DHEB)	

Product Characteristics			
Color		Score	
Shape	RECTANGLE	Size	
Flavor		Imprint Code	
Contains			

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:21130-960- 06	6 in 1 CARTON	04/01/2022		
1		9 g in 1 PATCH; Type 0: Not a Combination Product			

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part348	04/01/2022	

Labeler - Safeway (009137209)

Establishment				
Name	Address	ID/FEI	Business Operations	
Foshan Aqua Gel Biotech Co.,Ltd.		529128763	manufacture(21130-960)	

Revised: 4/2022 Safeway