

SALICYLIC ACID- medicated callus removers patch
LEADER/ Cardinal Health 110, Inc.

Leader Medicated Callus Removers

Active ingredient

Salicylic acid 40%

Purpose

Callus remover

Use

- for the removal of hard calluses
- relieves pain by removing calluses

Warnings

For external use only.

Do not use

- if you are a diabetic
- if you have poor blood circulation
- on irritated skin or any area that is infected or reddened

If discomfort persists see your doctor or podiatrist

Keep out of reach of children.

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

Directions

- wash affected area and dry thoroughly
- if necessary, cut medicated patch to fit callus
- apply adhesive side down of medicated patch onto callus
- cover medicated patch with pad
- after 48 hours, remove medicated patch
- repeat procedure every 48 hours as needed for up to 14 days (until callus is removed)
- may soak callus in warm water for 5 minutes to assist in removal
- continued wearing of pad (without patch) will help prevent recurrence of calluses

Other information

store between 15°C to 30°C (59°F to 86°F)

Inactive ingredients

acrylic adhesive, acrylic polymer, polyethylene, polyvinyl alcohol

Questions?

call 1-866-964-0939

Principal Display Panel

LEADER

Medicated

Callus

Removers

with Salicylic Acid 40%

- Effective Callus Removal Treatment
- Protect Against Pressure and Pain
- **4 MEDICATED PATCHES/ 6 PROTECTIVE PADS**



SALICYLIC ACID

medicated callus removers patch

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70000-0333
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	400 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
POLYVINYL ALCOHOL (UNII: 532B59J990)	
VINYL ACETATE (UNII: L9MK238N77)	
HIGH DENSITY POLYETHYLENE (UNII: UG00KM4WR7)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:70000-0333-1	4 in 1 CARTON	02/21/2018	
1		1 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 Drug	M030	02/21/2018	

Labeler - LEADER/ Cardinal Health 110, Inc. (063997360)

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LEADER/ Cardinal Health 110, Inc.