SALICYLIC ACID- medicated callus removers patch AmerisourceBergen Drug Corporation

GNP Medicated Callus Removers

Active ingredient

Salicylic acid 40%

Purpose

Callus remover

Uses

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

Warnings

For external use only.

Do not use

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

Stop use and ask a doctor if

discomfort persists

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

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 1-888-423-0139

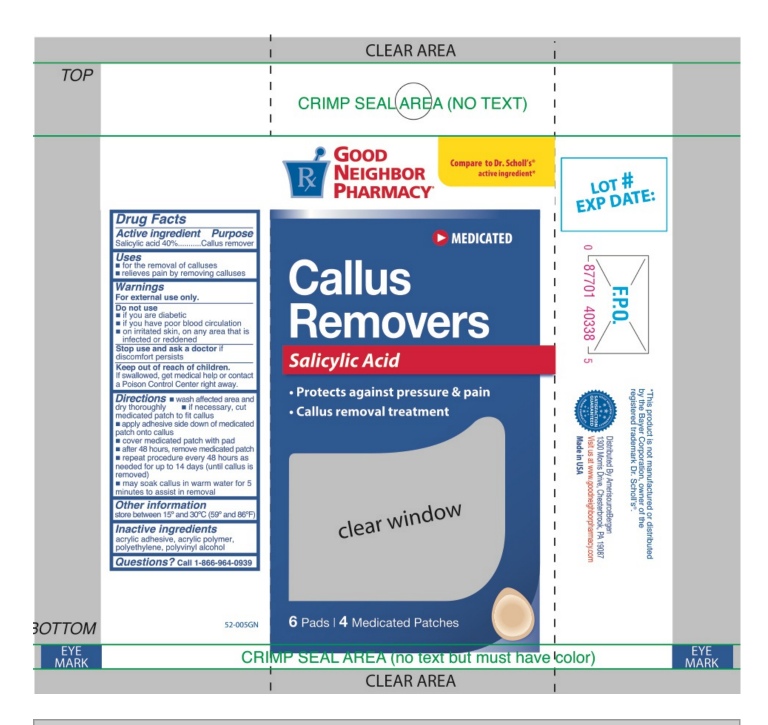
Principal Display Panel Good Neighbor Pharmacy

Callus Removers

Salicylic Acid

- Callus removal treatment
- Protect against pressure and pain

4 Medicated Patches/6 Pads



SALICYLIC ACID

Product Information

medicated callus removers patch

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:46122-524

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: 0414PZ4LPZ) (SALICYLIC ACID - UNII:0414PZ4LPZ)	SALICYLIC ACID	40 ma in 4

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

ı	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:46122-524- 13	4 in 1 PACKAGE; Type 0: Not a Combination Product	12/21/2017	

Marketing In	larketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC Monograph Drug	M030	12/21/2017			

Labeler - AmerisourceBergen Drug Corporation (007914906)

Revised: 1/2024 AmerisourceBergen Drug Corporation