SALICYLIC ACID- medicated plantar wart remover patch Topco Associates LLC

Medicated Plantar Wart Remover

Active ingredient

Salicylic acid 40%

Purpose

Plantar wart remover

Uses

 for the removal of plantar warts on the bottom of the foot. The plantar wart is recognized by its location only on the bottom of the foot, its tenderness, and the interruption of the footprint pattern.

Warnings

For external use only.

Do not use

- on irritated, infected or reddened skin
- on genital warts and warts on the face
- on moles, birthmarks and warts with hair growing from them
- on mucous membranes

Ask a doctor before use if you ahve

- diabetes
- poor blood circulation

Stop use and ask a doctor

if discomfort persists.

Keep out of the reach of children.

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

Directions

- wash affected area, may soak wart in warm water for 5 minutes
- dry area thoroughly
- if necessary, cut medicated patch to fit wart
- apply the adhesive side of patch onto wart
- cover medicated patch with pad to conceal area

repeat procedure every 48 hours as needed (until wart is removed) for up to 12 weeks

Other information

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

Inactive ingredients

acrylic adhesive, acrylic polymer, polyethylene, polyvinyl alcohol

Questions?

Call 1-888-423-0139

Principal Display Panel

TopCare

health

MEDICATED

Plantar

Wart Remover

SALICYLIC ACID 40%

- Cushions, Conceals & Protects While Removing Warts
- Maximum Strength Formula Without the Mess of Liquids

24 MEDICATED PATCHES 24 CUSHIONING PADS



SALICYLIC ACID

medicated plantar wart remover patch

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:36800-378
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	40 mg in 24

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:36800-378- 24	24 in 1 BOX; Type 0: Not a Combination Product	12/06/2017	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M028	12/06/2017		

Labeler - Topco Associates LLC (006935977)

Revised: 2/2024 Topco Associates LLC