

DERMAMED- skin protectant ointment
DermaRite Industries, LLC

DermaMed

Active ingredient

Aluminum Hydroxide Gel 2%

Purpose

Skin protectant

Uses

Dries the oozing and weeping of poison: ivy, oak, or sumac, or other skin irritations.

Warnings

For external use only.

Avoid contact with eyes. In case of contact, flush thoroughly with water.

Warnings

Stop and ask a doctor if

- condition worsens
- symptoms last more than 7 days or clear up and occur again within a few days

Warnings

Keep out of reach of children.

In case of accidental ingestion contact a Physician or Poison Control Center right away.

Directions

Apply liberally to the affected area as needed or as directed by a physician.

Other information

- Store at room temperature (59°- 86°F)
- You may report a serious adverse event to DermaRite Industries, PO Box 7209, North Bergen, NJ 07047.

Inactive ingredients

Petrolatum, Lanolin, Water, Propylene Glycol, Stearyl Alcohol, Calcium Carbonate, Magnesium Hydroxide, Zinc Chloride, DMDM Hydantoin, Methylparaben, Propylparaben, Sodium Laureth Sulfate, Lanolin Alcohol, Cholecalciferol, Zea Mays (Corn) Oil, Retinyl Palmitate, Citric Acid, Sodium Chloride

Questions?

Call 800-337-6296 Mon - Fri 9AM - 5PM EST.

DermaMed Package Label and Principal Display Panel

NDC 61924-214-04



SKIN PROTECTANT

pH balanced

Soothes and promotes healing

DermaRite

REORDER #00214

Net Wt. 106 g (3.75 oz.)

Drug Facts

Active ingredient	Purpose
Aluminum Hydroxide Gel 2%	Skin protectant

Uses
 ■ Dries the oozing and weeping of poison: ivy, oak, or sumac, or other skin irritations.

Warnings
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Stop use and ask a doctor if ■ condition worsens ■ symptoms last more than 7 days or clear up and occur again within a few days.
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Questions? Call 1-800-337-6296 Mon - Fri 9AM - 5PM EST.

Patient Name

Room #



7 14196 21404 1

DermaRite Industries LLC • 7777 West Side Ave. North Bergen, NJ 07047 • www.dermarite.com

MADE IN THE USA 100832




DermaMed™
OINTMENT

SKIN PROTECTANT

DermaRite®

REORDER #00214

CONTENTS: 24 TUBES - 106g (3.75 oz.) Each



1 0 7 1 4 1 9 6 2 1 4 0 4 8

DermaRite Industries LLC
7777 West Side Avenue
North Bergen, NJ 07047
www.dermarite.com

**MADE
IN THE
USA**
100832

DERMAMED

skin protectant ointment

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:61924-214	
Route of Administration	TOPICAL			
Active Ingredient/Active Moiety				
Ingredient Name		Basis of Strength	Strength	
ALUMINUM HYDROXIDE (UNII: 5QB0T2IUN0) (ALUMINUM HYDROXIDE - UNII:5QB0T2IUN0)		ALUMINUM HYDROXIDE	2 g in 100 g	
Inactive Ingredients				
Ingredient Name			Strength	
CHOLECALCIFEROL (UNII: 1C6V77QF41)				
LANOLIN (UNII: 7EV65EAW6H)				
LANOLIN ALCOHOLS (UNII: 884C3FA9HE)				
SODIUM LAURETH SULFATE (UNII: BPV390UAP0)				
MAGNESIUM HYDROXIDE (UNII: NBZ3QY004S)				
PROPYLENE GLYCOL 1-(2-METHYLBUTYRATE) (UNII: 9Q5W5G6461)				
VITAMIN A PALMITATE (UNII: 1D1K0N0VVC)				
DMDM HYDANTOIN (UNII: BYR0546TOW)				
ANHYDROUS CITRIC ACID (UNII: XF417D3PSL)				
CORN OIL (UNII: 8470G57WFM)				
METHYLPARABEN (UNII: A2I8C7HI9T)				
SODIUM CHLORIDE (UNII: 451W47IQ8X)				
ZINC CHLORIDE (UNII: 86Q357L16B)				
PROPYLPARABEN (UNII: Z8IX2SC1OH)				
STEARYL ALCOHOL (UNII: 2KR89I4H1Y)				
PETROLATUM (UNII: 4T6H12BN9U)				
CALCIUM CARBONATE (UNII: H0G9379FGK)				
WATER (UNII: 059QF0KO0R)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:61924-214-04	106 g in 1 TUBE; Type 0: Not a Combination Product	10/18/2016	
Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M016	10/18/2016		

Labeler - DermaRite Industries, LLC (883925562)

Registrant - DermaRite Industries, LLC (883925562)

Establishment

Name	Address	ID/FEI	Business Operations
DermaRite Industries, LLC		883925562	manufacture(61924-214)

Revised: 12/2024

DermaRite Industries, LLC