### PROCURE MEDICATED- aluminum hydroxide gel 2% ointment Twin Med LLC

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### **Active Ingredient**

Aluminum Hydroxide Gel 2%

### **Purpose**

Skin Protectant

#### Uses

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• Dries the oozing and weeping of poison ivy, oak and sumac

### **Directions**

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- Apply as needed
- Children under 6 months of age: ask a doctor

## **Inactive Ingredients**

Calcium Carbonate, Cholecalciferol, Citric Acid, Ethylhexylglycerin, Glyceryl Monostearate, Glyceryl Stearate/ PEG-100 Stearate, Lanolin, Magnesium Hydroxide, Petrolatum, Phenoxyethanol, Propylene Glycol, Retinyl Palmitate, Sodium Chloride, Sodium Laureth Sulfate, Stearyl Alcohol, Water, Zea Mays Corn Oil, Zinc Chloride

### **Warnings**

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For external use only

When using this product

When using this product do not get into the eyes

# Stop use and ask a doctor if

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

## Keep out of reach of children

**Keep out of reach of children.** In case of accidental ingestion contact a physician or Poison Control Center right away.



# **Drug Facts**

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Purpose

Item No. PCMG35

LICOC - Drive the series and was

..Skin Protectant

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#### Directions

- Apply as needed
- Children under 6 months of age: ask a doctor

#### Other Information

■ Store between 15°-30°C (59°-86°F)

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#### Questions or Comments:

Call 1-877-894-6633 8 am - 6 pm EST M-F

Patient Name:



Medicated
OINTMENT

with Aluminum
Hydroxide Gel



Manufactured for:

Twin Med LLC.

11333 Greenstone Ave Santa Fe Springs, CA <u>90670</u>

ProCureProducts.com

Made in India

Poison ivy, oak and sumac drying ointment



NDC: 55681-029-04

NET WT 3.75 OZ (106g)



### PROCURE MEDICATED

aluminum hydroxide gel 2% ointment

# **Product Information**

Product Type HUMAN OTC DRUG Item Code (Source) NDC:55681-029

**Route of Administration** TOPICAL

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
<b>ALUMINUM HYDROXIDE</b> (UNII: 5QB0T2IUN0) (ALUMINUM HYDROXIDE - UNII:5QB0T2IUN0)	ALUMINUM HYDROXIDE	2 g in 100 g

Inactive Ingredients	
Ingredient Name	Strength
ZEA MAYS (CORN) OIL (UNII: 8470G57WFM)	
GLYCERYL MONOSTEARATE (UNII: 2300U9XXE4)	
LANOLIN (UNII: 7EV65EAW6H)	
MAGNESIUM HYDROXIDE (UNII: NBZ3QY004S)	
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)	
CALCIUM CARBONATE (UNII: H0G9379FGK)	
CHOLECALCIFEROL (UNII: 1C6V77QF41)	
CITRIC ACID (UNII: 2968PHW8QP)	
GLYCERYL STEARATE/PEG-100 STEARATE (UNII: RD25J5V947)	
PETROLATUM (UNII: 4T6H12BN9U)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
RETINYL PALMITATE (UNII: 1D1K0N0VVC)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SODIUM LAURETH SULFATE (UNII: BPV390UAP0)	
STEARYL ALCOHOL (UNII: 2KR89I4H1Y)	
WATER (UNII: 059QF0KO0R)	
ZINC CHLORIDE (UNII: 86Q357L16B)	

Product Characteristics			
Color	white (White to off-white)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

F	Packaging				
#	tem Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:55681-029- 04	106 g in 1 TUBE; Type 0: Not a Combination Product	07/14/2025		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M016	07/14/2025	

# Labeler - Twin Med LLC (009579330)

Revised: 9/2025 Twin Med LLC