UNIVERSAL 70% RUBBING ALCOHOL WITH WINTERGREEN AND GLYCERINisopropyl alcohol liquid Universal Distribution Center LLC

70% Isopropyl Rubbing Alcohol with Wintergreen & Glycerin

Drug Facts

Active ingredients (by volume):

Isopropyl alcohol (70% conc.)

Purpose

first aid antiseptic

Uses

• first aid to help prevent the risk of infection in minor cuts, scrapes and burns

Warnings

For external use only; flammable, keep away from fire or flame, heat, spark, electrical

Ask a doctor before use if you have

deep punctured wounds, animal bites or serious burns

When using this product

- do not get into eyes
- do not apply over large areas of the body
- do not use longer than one week unless directed by a doctor

Stop using this product if

• condition persists or gets worse

Keep this and all drugs out of the reach of children.

In case of accidental ingestion, seek professional assistance or contact a Poison control center (1-800-222-1222) immediately.

Directions

- clean effected area
- apply small amount of this product on the area 1-3 times daily
- May be covered with a sterile bandage
- If bandaged, let dry first

Other information

- store at room temperature
- does not contain, nor is intended as a substitute for grain or ethyl alcohol will produce serious gastric disturbances if taken internally.

Inactive Ingredients

Purified Water, Methyl Salicylate, Glycerin, FD&C Green 3, FD&C Yellow 5

For Rubbing & Massaging

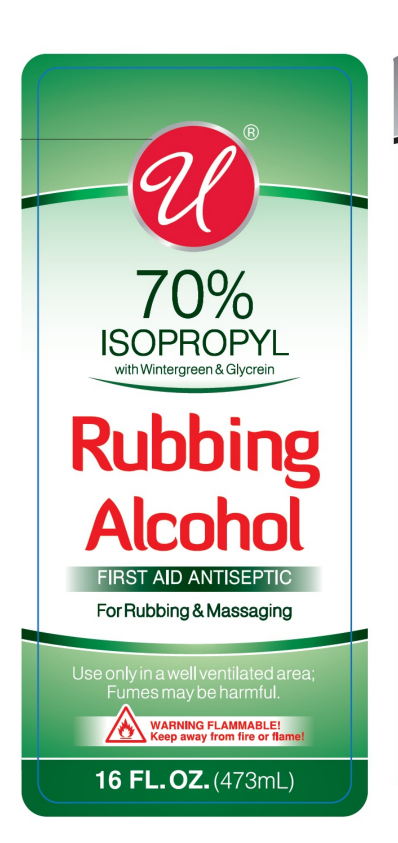
Use only in a well ventilated area; Fumes may be harmful.

TAMPER EVIDENT: DO NOT USE IF THE UNDER CAP PRINTED SAFETY FOIL IS BROKEN OR MISSING.

Made in Turkey

Mfd for and Distributed By:
Universal Distribution Center LLC
330 Applegarth Road,
Monroe Township, NJ 08831
www.universaldc.com

Packaging



Isopropyl rubbing alcohol 70% by volume

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UNIVERSAL 70% RUBBING ALCOHOL WITH WINTERGREEN AND GLYCERIN

isopropyl alcohol liquid

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:52000-429

TOPICAL Route of Administration

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
, (, (ISOPROPYL ALCOHOL	70 mL in 100 mL			

Inactive Ingredients				
Ingredient Name	Strength			
WATER (UNII: 059QF0KO0R)				
METHYL SALICYLATE (UNII: LAV5U5022Y)				
GLYCERIN (UNII: PDC6A3C0OX)				
FD&C GREEN NO. 3 (UNII: 3P3ONR6O1S)				
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)				

Ш	P	ackaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:52000- 429-16	473 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/17/2025	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M003	06/17/2025		

Labeler - Universal Distribution Center LLC (019180459)

Revised: 6/2025 Universal Distribution Center LLC