ISOPROPYL RUBBING ALCOHOL 50%- isopropyl alcohol liquid MY IMPORTS LLC

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

AMERICAN REDCROSS ISOPROPYL RUBBING ALCOHOL 50%

Active ingredient (by volume)

Isopropyl alcohol (50% concentrate)

Purpose

topical antimicrobial

Uses

- to decrease germs in minor cuts and scrapes
- helps relieve minor muscular aches due to exertion

Warnings

For external use only

- flammable, keep away from fire and flame
- will produce serious gastric disturbances if taken internally

Ask a doctor before use if you have deep puncture wounds or serious burns

When using this product

- do not get into eyes or mucous membranes
- use only in a well-ventilated area

Stop use and ask a doctor if condition persists or gets worse

Keep out of reach of children.

In case of an accidental ingestion, contact a Poison Control Center immediately

Directions

- apply to skin directly of with clean gauze, cotton or swab
- for rubbing apply liberally and rub with hands

Other information

- does not contain, nor is intended as a substitute for grain or ethyl alcohol
- keep bottle tightly closed

Inactive ingredient

Water

PRINCIPAL DISPLAY PANEL

ISOPROPYL RUBBING ALCOHOL 50%

TOPICAL ANTIMICROBIAL

12 FL.OZ (354 mL)



ISOPROPYL RUBBING ALCOHOL 50%

isopropyl alcohol liquid

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:51628-4232	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ISOPROPYL ALCOHOL (UNII: ND2M416302) (ISOPROPYL ALCOHOL - UNII:ND2M416302)	ISOPROPYL ALCOHOL	50 mL in 100 mL	

Inactive Ingredients				
	Ingredient Name	Strength		
WATER (UNII: 059QF0KO0R)				

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:51628- 4232-1	355 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/18/2020	
2	NDC:51628- 4232-2	473 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/18/2020	

final	part333A		33, 23, 2323		
OTC monograph i	ot		06/18/2020		
Marketing Category	Application	n Number or Monogra Citation	ph Marketing Date		arketing End Date
Marketing Information					
3 NDC:51628- 4232-3	946 mL in 1 BOTTLE Combination Produc	E, PLASTIC; Type 0: Not a ct	06/18/2020		

Labeler - MY IMPORTS LLC (195767988)

Registrant - Jell Pharmaceuticals Pvt. Ltd. (726025211)

Establishment				
Name	Address	ID/FEI	Business Operations	
Jell Pharmaceuticals Pvt. Ltd.		726025211	manufacture(51628-4232)	

Revised: 2/2022 MY IMPORTS LLC