# ALCOHOL- is opropyl alcohol liquid Dynarex Corporation

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.

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#### Alcohol 99%

Active Ingredient Purpose

Isopropyl Alcohol 99% v/v Antiseptic

#### **PURPOSE**

First aid to help prevent the risk of infection in:

- minor cuts
- scrapes
- burns

#### **WARNINGS**

- For external use only
- Flammable, keep away from fire or flame, heat, spark, electrical.
- Not for use with electrocautinary devices or procedures
- **Ask a doctor before use if you have** deep puncture wounds, animal bites or serious burns.
- **Stop use and ask a doctor if** condition persist or gets worse.

For rubbing and massaging:

- **Caution:** Do not apply to irritated skinor if excessive irritation develops
- Avoid getting into eyes or mucous membranes

#### **INDICATIONS & USAGE**

When using this product:

- do not get in eyes
- do not apply over large areas of the body
- do not use longer than 1 week unless directed by a doctor

#### KEEP OUT OF REACH OF CHILDREN

# Keep out of reach of children.

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

### **DOSAGE & ADMINISTRATION**

#### DIRECTIONS:

- clean the affected area
- apply a small amount of this product on the affected area 1 to 3 times daily
- may be covered with a sterile bandage
- if bandaged let dry first

#### OTHER INFORMATION

- Store at room temperature: 15 deg C to 30 deg C 59 deg F to 86 deg F
- avoid excessive heat
- Does not contain, nor is it intended as a substitute for grain or ethyl alcohol
- Will produce serious gastric disturbances if taken internally

# **INACTIVE INGREDIENT**

**Inactive Ingredient** 

Water

### PRINCIPAL DISPLAY PANEL

DYNAREX 99% ISOPROPYL ALCOHOL

99 IPA.jpg



# **ALCOHOL**

isopropyl alcohol liquid

Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:67777-305		
Route of Administration	TOPICAL				

ı	Active Ingredient/Active Moiety					
ı	Ingredient Name	Basis of Strength	Strength			
	ISOPROPYL ALCOHOL (UNII: ND2M416302) (ISOPROPYL ALCOHOL - UNII:ND2M416302)	ISOPROPYL ALCOHOL	0.99 mL in 1 mL			

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

Packaging						
#	Item Code	Package Description	Marketing	Start Date	Ma	arketing End Date
1 NDC	C:67777-305-01	473 mL in 1 BOTTLE				
Mar	keting Infori	nation				
	keting Infori	nation Application Number or Monog	raph Citation	Marketing Star	t Date	Marketing End Date

# Labeler - Dynarex Corporation (008124539)

# Registrant - Dynarex Corporation (008124539)

Establishment					
Name	Address	ID/FEI	Business Operations		
Blossom Pharmaceuticals		677381470	manufacture(67777-305)		

Revised: 1/2014 Dynarex Corporation