# 3 HYDROGEN PEROXIDE- hydrogen peroxide liquid Jell Pharmaceuticals Pvt Ltd.

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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#### **3% Hydrogen Peroxide Spray**

#### **Drug Facts**

#### Active ingredient

Hydrogen Peroxide 3%

#### **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

#### Do not use

- in the eyes or apply over large areas of the body
- longer than 1 week.

### Ask a doctor before use if you have

deep or puncture wounds, animals bites or serious burns

### Stop use and ask a doctor if

- the condition persists or gets worse
- irritation, pain, or redness persists or worsens
- swelling, rash, or fever develops

### Keep out of reach of children.

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

#### **Directions**

- clean the affected area
- spray a small amount of product on the affected area 1 to 3 times a day

- may be covered with a sterile bandage
- if bandaged, let dry first

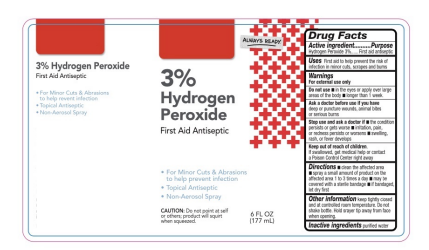
#### Other information

keep tightly closed and at controlled room temperature. Do not shake bottle. Hold srayer tip away from face when opening.

#### Inactive ingredients

purified water

### **Package Labeling:**





### **3 HYDROGEN PEROXIDE**

hydrogen peroxide liquid

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:30400-250

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### **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength	
HYDROGEN PEROXIDE (UNII: BBX060AN9V) (HYDROGEN PEROXIDE - UNII:BBX060AN9V)	HYDROGEN PEROXIDE	30 mg in 1 mL	

### **Inactive Ingredients**

Ingredient Name	Strength

WATER (UNII: 059QF0KO0R)

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:30400-250- 01	177 mL in 1 BOTTLE; Type 0: Not a Combination Product	01/10/2021	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333A	01/10/2021		

## Labeler - Jell Pharmaceuticals Pvt Ltd. (726025211)

# Registrant - Jell Pharmaceuticals Pvt Ltd. (726025211)

Establishment					
Name	Address	ID/FEI	<b>Business Operations</b>		
Jell Pharmaceuticals Pvt Ltd.		726025211	manufacture(30400-250)		

Revised: 5/2022 Jell Pharmaceuticals Pvt Ltd.