

**ANTI BACTERIAL HAND SANITIZER- ethyl alcohol gel**  
**Dolgencorp Inc.**

*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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**Active Ingredients :**

Ethyl Alcohol 62%

**Purpose:**

Antiseptic

**Warnings:** For external use only.

Flammable. Keep away from fire or flame.

Do not apply around eyes.

Do not use in ears or mouth.

When using this product, avoid contact with eyes. In case of contact, flush eyes with water.

Stop use or ask for a doctor if redness or irritation develops and persists for more than 72 hours.

Keep out of reach of children. Children must be supervised in use of this product.

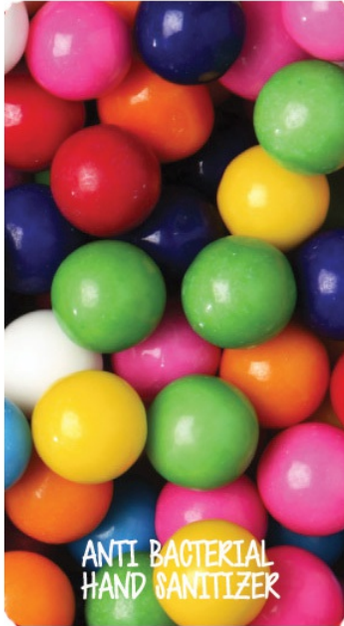
**Directions :**

Pump as needed into your palms and thoroughly spread on both hands. Rub into skin until dry.

**Inactive ingredients:** Water, Glycerin, Propylene Glycol, Carbomer, Triethanolamine, Aloe Barbadenis Leaf Juice

**Other information:** Do not store in temperature over 118 F.

**Uses:** Hand sanitizer to help decrease bacteria on the skin when water, soap & towel are not available. Recommended for repeated use.



**Drug Facts**

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1 fl oz / 29mL  
MADE IN CHINA

## ANTI BACTERIAL HAND SANITIZER

ethyl alcohol gel

### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:559 10-988
<b>Route of Administration</b>	TOPICAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	62 mL in 100 mL

## Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
GLYCERIN (UNII: PDC6A3C0OX)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
CARBOMER INTERPOLYMER TYPE A (55000 CPS) (UNII: 59TL3WG5CO)	
TROLAMINE (UNII: 9O3K93S3TK)	

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:55910-988-20	29 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	06/08/2015	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	06/08/2015	

**Labeler** - Dolgencorp Inc. (068331990)

Revised: 6/2015

Dolgencorp Inc.