# RAW INSTANT HAND SANITIZER GEL- ethyl alcohol gel Raw Office Inc, The

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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#### Raw Instant Hand Sanitizer

### **Drug Facts**

# **Active Ingredient:**

Ethyl Alcohol 75%

# **Purpose**

Antimicrobial

#### USE:

Suitable for hand, skin and object surface sanitization

# Warnings:

- 1. External use, not oral. Keep out of children's reach.
- 2. Flammable, keep away from fire and flame.
- 3. Use with caution if allergic to alcohol.

**When using this Product** • Avoid contacting face, eyes and broken skin. In case of eye contact, flush with plenty of water and seek medical advice.

**Stop use and ask doctor if •** Irritation or redness occurs.

**Keep out of children's reach.** If swallowed, get medical help or contact poison control center.

*Directions* • Wet hands thoroughly with product and rub into skin until dry. Wet object thoroughly with product until dry. Children under 6 years of age should be supervised by an adult when using.

# Other Information:

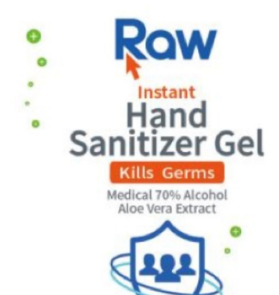
- Store below 43°C (110°F)
- Keep sealed after use

*Inactive Ingredients:* Purified Water, Glycerin, Carbomer

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**MADE IN CHINA** 

# **Packaging**



17 FL OZ (500mL)

# **Drug Facts**

Active Ingredient:

Ethyl Alcohol 65%-75% (v/v) ..

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### Stop use and ask a doctor if

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**Directions** Wet hands thoroughly with product and rub into skin until dry. Wet object surface thoroughly with product until dry. Children under 6 years of age should be supervised by an adult when using.

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Batch No: Production Date: Expiry Date:

Inactive ingredients: Purified Water, Glycerin, Carbomer

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# RAW INSTANT HAND SANITIZER GEL

ethyl alcohol gel

#### **Product Information**

Product Type HUMAN OTC DRUG Item Code (Source) NDC:75353-513

Route of Administration TOPICAL

# **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958 V90M) (ALCOHOL - UNII:3K9958 V90M)	ALCOHOL	70 mL in 100 mL

### **Inactive Ingredients**

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6 A3C0 OX)	
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0 A5MM307FC)	

# **Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:75353-513-10	100 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/15/2020	
2	NDC:75353-513-50	500 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/15/2020	

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part333A	04/15/2020			

# Labeler - Raw Office Inc, The (204127000)

Revised: 4/2020 Raw Office Inc, The