# ANTISEPTIC HAND SANITIZER MINT SCENT- alcohol spray Two's Company, 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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#### ANTISEPTIC HAND SANITIZER SPRAY-MINT SCENT

#### **Drug** Facts

#### Active ingredient

Alcohol 62%

#### Purpose

Antimicrobial

#### Uses

• helps to reduce bacteria on the skin

• recommended for repeated use

#### Warnings

#### For external use only

Flammable: Keep away from heat and flame

#### When using this product

- do not use in or near the eyes. In case of eye contact, rinse eyes thoroughly with water
- do not apply to irritated or broken skin.

#### Stop use and ask a doctor if

- irritation and redness develop
- condition persists for more than 72 hours.

**Keep out of reach of children.** If swallowed, get medical help or contact a Poison Control Center right away.

#### Directions

• lift tab and spray a small amount into the palms of your hands and forearms.

• wet the hands thoroughly with product, rub hands together and allow to dry without wiping • no rinsing required.

• children under 6 years of age should be supervised when using this product • not recommended for infants.

#### Other information

- do not store above 105°F
- may discolor some fabrics or surfaces.

#### **Inactive ingredients**

Fragrance, Glycerin, Propylene Glycol, Water

MINT SCENT CLEANS AND FRESHENS YOUR HANDS AND REDUCES BACTERIA Distributed By

**TWO'S COMPANY INC.** 500 Saw Mill River Rd, Elmsford, NY 10523 / **Made in China** YOU MAY REPORT SERIOUS SIDE EFFECTS TO THE ABOVE ADDRESS

#### Packaging



OUTER PACK LABEL

INNER LABEL





### ANTISEPTIC HAND SANITIZER MINT SCENT

alcohol spray

Product Information							
HUMAN OTC DRUG	Item Code	(Source)	NDC:72762-004				
TOPICAL							
Active Ingredient/Active Moiety							
lient Name		Basis of Strengt	h Strength				
OHOL - UNII:3K9958V90M)		ALCOHOL	0.62 mL in 1 mL				
	TOPICAL	ety lient Name	TOPICAL ety lient Name Basis of Strengt				

Inactive Ingredients				
Ingredient Name	Strength			
GLYCERIN (UNII: PDC6A3C0OX)				
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)				
WATER (UNII: 059QF0KO0R)				

#### Packaging

# Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b> NDC:72762-004- 16	1 in 1 BLISTER PACK	09/01/2018	
<b>1</b> NDC:72762-004- 15	15 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part333E	09/01/2018			
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## Labeler - Two's Company, Inc. (056307960)

Revised: 1/2019

Two's Company, Inc.