# EMOJI WATERMELON HAND SANITIZER- benzalkonium chloride gel Townley 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 Ingredient**

Benzalkonium Chloride 0.1%

□**Purpose:**□ Antibacterial

 $\Box$ *Use* 

To decrease bacteria on the skin that could cause disease.

Keep out of reach of children.

#### □ Warnings

- For external use only-hands.
- **Keep out of eyes.** Avoid contact with broken skin.
- **Stop use and ask a doctor if** irritation or redness develops.
- **Do not inhale or ingest.** If swallowed, get medical help or contact a poison control center right away.

#### Directions

- Rub a dime sized drop into hands.
- For children under 6 use under adult supervision.

#### **Inactive Ingredients**

water (aqua/eau), glycerin, coceth-7, PPG-1-PEG-9 lauryl glycol ether, carbomer, hydrogenated castor oil, fragrance (parfum).

**May Contain** 

Red 40 (CI 16035), Red 33 (CI 17200), Blue 1 (CI 42090), Yellow 5 (CI 19140).

Hand Sanitizer

## **EJ0223WA**

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DIE LINE—DO NOT PRINT



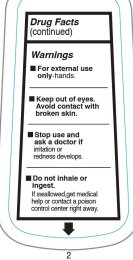
CLEAR ADHESIVE STOCK DO NOT PRINT



CONFIRM WEIGHTS + FLAVORS + WARNING White Base X2









benzalkonium chloride gel

#### **Product Information**

Product TypeHUMAN OTC DRUGItem Code (Source)NDC:58737-197

Route of Administration TOPICAL

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

Ingredient Name	Basis of Strength	Strength
(,	BENZALKONIUM CHLORIDE	0.059 g in 59 mL

Inactive Ingredients	
Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6A3C0OX)	
COCETH-7 CARBOXYLIC ACID (UNII: 35KO064932)	
CARBOMER HOMOPOLYMER TYPE C (ALLYL PENTAERYTHRITOL CROSSLINKED) (UNII: 4Q93RCW27E)	
PEG-40 CASTOR OIL (UNII: 4ERD2076EF)	
PPG-1-PEG-9 LAURYL GLYCOL ETHER (UNII: 5R8J43K25L)	
SO DIUM HYDRO XIDE (UNII: 55X04QC32I)	

Pa	Packaging			
#	Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:58737-197-01	59 mL in 1 BOTTLE; Type 0: Not a Combination Product	0 1/18/20 17	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333E	0 1/18/20 17		

### Labeler - Townley Inc. (016956158)

Establishment				
Name	Address	ID/FEI	<b>Business Operations</b>	
Shenzhen Lantern Science Co., Ltd.		421222423	manufacture(58737-197)	

Revised: 1/2017 Townley Inc.