# JOJO SIWA HAND SANITIZER- benzalkonium chloride liquid Ashtel Studios, 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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### Jojo Siwa Hand Sanitizer

**Drug Facts** 

#### **Active Ingredient**

Benzalkonium Chloride 0.1%

#### **Purpose**

Antiseptic

**Use** To help reduce bacteria and germs on the skin.

**WARNING** Flammable. Keep away from fire or flame. For external use only • Stop use and ask a doctor if irritation or redness develops and persists.

• **Keep out of reach of children**. • In case of accidental digestion, seek professional assistance or contact a Poison Control Center immediately.

**Directions** • Place enough product in palm to cover hands and rub hands together briskly until dry.

- Children under 6, use only under adult supervision.
- Not recommended for infants.

**Other Information** • Do not store above 100° F (38° C). • May discolor some fabrics. • Harmful to wood finishes and plastics.

**Inactive Ingredients** • Aqua (Water), Hydroxyethylcellulose, Phenoxyethanol, Disodium EDTA, Polysorbate 20, Parfum (Fragrance).

#### Smart Care ®

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#### QUESTIONS OR COMMENTS?

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#### **Packaging**





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**Smart Care®** 

### JOJO SIWA HAND SANITIZER

benzalkonium chloride liquid

#### **Product Information**

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70108-022
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Route of Administration TOPICAL

## Active Ingredient/Active Moiety

- 1	, , , , , , , , , , , , , , , , , , ,		
	Ingredient Name	Basis of Strength	Strength
	BENZALKO NIUM CHLO RIDE (UNII: F5UM2KM3W7) (BENZALKONIUM -	BENZALKONIUM	0.1 g
1	UNII:7N6JUD5X6Y)	CHLORIDE	in 100 mL

## **Inactive Ingredients**

indeave ingredients		
Ingredient Name	Strength	
WATER (UNII: 059QF0KO0R)		
HYDROXYETHYL CELLULOSE (4000 MPA.S AT 1%) (UNII: ZYD53NBL45)		
PHENO XYETHANO L (UNII: HIE492ZZ3T)		
EDETATE DISO DIUM ANHYDRO US (UNII: 8 NLQ36 F6 MM)		
POLYSORBATE 20 (UNII: 7T1F30V5YH)		

#### Packaging

ı		88			
ı	#	Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
ı	1	NDC:70108-022-01	53 mL in 1 BOTTLE; Type 0: Not a Combination Product	11/0 1/20 19	

## **Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	11/0 1/20 19	

## Labeler - Ashtel Studios, Inc. (148689180)

Revised: 10/2019 Ashtel Studios, Inc.