

**BIOPURE HAND SANITIZER SPRAY-COASTAL GLOW- hand sanitizer
spray liquid
Quest USA Corp**

BIOPURE HAND SANITIZER SPRAY-Coastal Glow

Active Ingredient(s)

Ethyl Alcohol 70% v/v

Purpose

Antiseptic

Use

- Hand sanitizer to decrease bacteria on the skin.
- Recommended for repeated use.

Warnings

For external use only.

Flammable, keep away from fire or flame.

Do not use

Stop use and ask doctor

if irritation and rash occurs. These may be signs of serious condition.

When using this product

keep out of eyes, ears and mouth. In case of contact with eyes, rinse eyes thoroughly with water. Avoid contact with broken skin.

Stop use and ask doctor if irritation and rash occurs. These may be signs of serious condition.

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

Directions

Directions

- Spray enough product in your palm to thoroughly cover your hands.
- Rub hands together briskly until dry. Recommended for repeated use.
- Children under 6 years of age should be supervised by an adult when using this product.

Other information

- Protect the product in this container from excessive heat and direct sun.
- Store below 104°F (40°C).
- May discolor certain fabrics.

Inactive ingredients

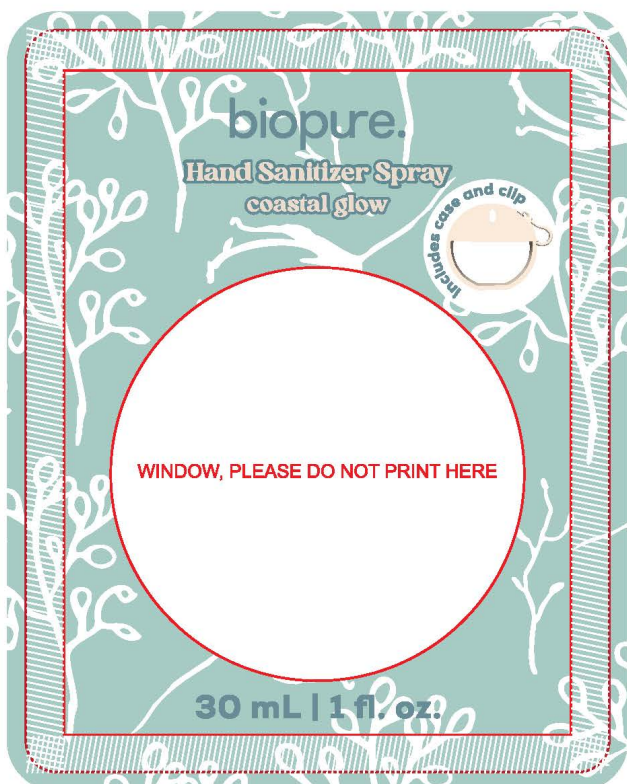
Water, Glycerin, Fragrance, Aloe Barbadensis Leaf Juice, Carbomer, Aminoethyl Propanol, Tocoheryl Acetate

Questions

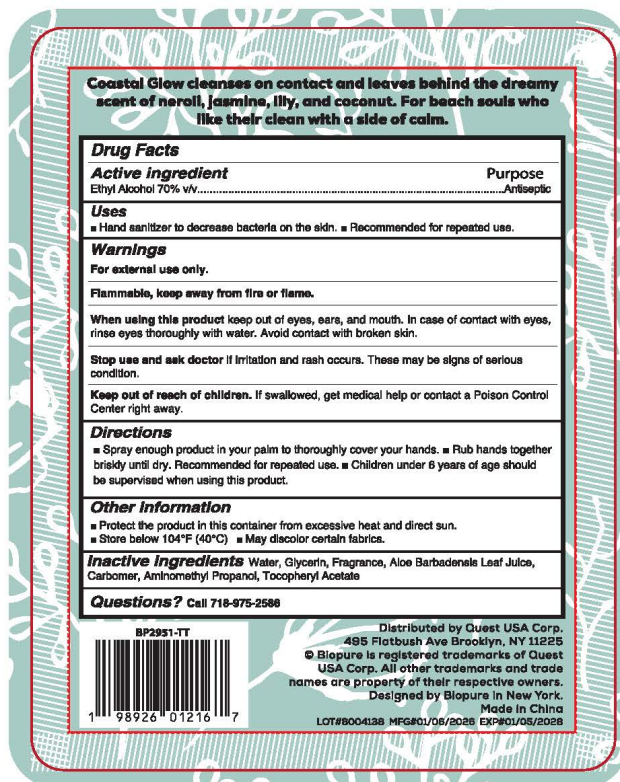
Questions? call 718-975-2586

Package Label - Alcohol Hand Sanitizer Spray-Coastal Glow

FRONT



BACK



FRONT



BACK

INSIDE



OUTSIDE



BOTTLE, CASE AND CARABINER PANTONE 9202 C



PANTONE®
2212 U

BIOPURE HAND SANITIZER SPRAY-COASTAL GLOW

hand sanitizer spray liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:78691-037
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	16 g in 30 mL

Inactive Ingredients

Ingredient Name	Strength
AMINOMETHYL PROPANOL (UNII: LU49E6626Q)	
WATER (UNII: 059QF0KO0R)	
ALOE BARBADENSIS LEAF (UNII: ZY81Z83H0X)	
GLYCERIN (UNII: PDC6A3C00X)	
CARBOMER (UNII: 0A5MM307FC)	
ALPHA-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:78691-037-00	30 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	01/15/2026	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M003	01/15/2026	

Labeler - Quest USA Corp (079869689)

Revised: 1/2026

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