

LIQUID PRO HAND SANITIZER- benzalkonium chloride gel
Saerom International, 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.

Liquid Pro Hand Sanitizer Gel

Drug Facts

Active ingredient

Benzalkonium Chloride 0.12%

Purpose

Antiseptic

Uses

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

Warnings

For external use only

When using this product

keep out of eyes. In case of contact, flush eyes with water.

Stop use and consult a doctor if

irritation or redness develops.

Keep out of reach of children.

If swallowed, get medical assistance or contact a Poison Control Center immediately.

Directions

- Rub thoroughly over all surfaces of both hands.
- Rub hands together briskly until dry.

Inactive ingredients

aloe barbadensis leaf extract, fragrance, hydroxyethylcellulose, propylene glycol, water

Package Labeling:75774-000-01

Liquid Pro™ Hand Sanitizer with moisturizing aloe is formulated without parabens * sulfates * phthalates * dyes * harsh, skin drying, alcohol. Recommended for repeated use.

LIQUID PRO™

Hand Sanitizer Gel with Aloe

For Sensitive Skin
 Anti-Microbial Protection
 Kills Germs
 Long Lasting
 No Need to Rinse
 Alcohol Free

1 Gallon (3.79L)

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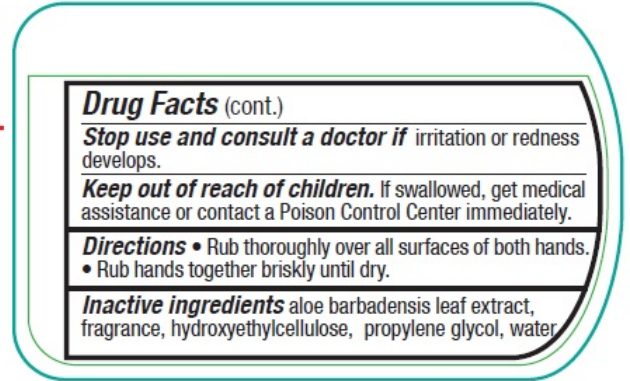
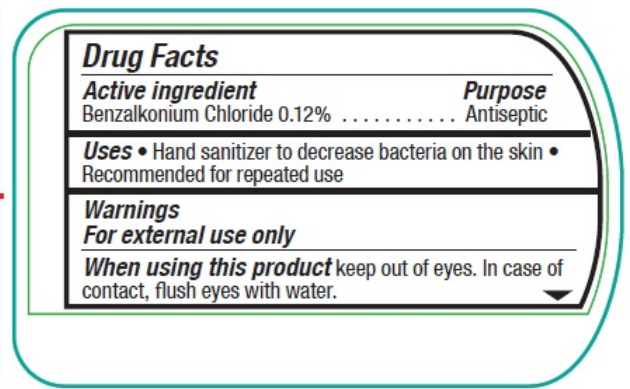
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Dist. by Saerom International
 70 Railroad Avenue, Ridgefield Park, NJ 07660
 201-931-0033 Made in USA



Package Labeling:75774-000-02



LIQUID PRO HAND SANITIZER

benzalkonium chloride gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:75774-000
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	1.2 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
HYDROXYETHYL CELLULOSE, UNSPECIFIED (UNII: T4V6TWG28D)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
WATER (UNII: 059QF0KO0R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:75774-000-01	3790 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/13/2020	
2	NDC:75774-000-02	60 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/13/2020	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	04/13/2020	

Labeler - Saerom International, Inc. (033235947)

Revised: 4/2020

Saerom International, Inc.