HUSKY INSTANT HAND SANITIZER- husky instant hand sanitizer solution Canberra Corporation

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.

Husky Instant Hand Sanitizer

For external use only.

Flammable. Keep away from heat or flame.

Do not use in the eyes. In case of contact, immediately flush with water.

Stop use and ask a doctor if irritation or rash appears and lasts.

Keep out of reach of children. If swallowed, contact a physician or poison center.

Ethyl Alcohol 62% v/v......Antibacterial Agent

Keep out of reach of children. If swallowed, contact a physician or poison center.

Water, Glycerin, Propylene Glycol, Isopropyl Mryistate, Aloe Barbadensis Leaf, Carbomer, Tocopheryl Acetate (Vitamin E), Aminomethyl Propanol

63779-056-77 & 63779-056-27:

To decrease bacteria on skin, apply a small amount to palm. Briskly rub, covering hands with product until dry.

Children under 6 years of age should be supervised when using this product.

63779-056-87 & 63779-056-19:

To decrease bacteria on skin, apply a small amount to palm. Briskly rub, covering hands with product until dry.

63779-056-77 & 63779-056-27: For hand cleaning to decrease bacteria on skin that could cause disease. Recommended for repeated use.

63779-056-87: For handwashing to decrease bacteria on skin.

63779-056-19: For hand cleaning to decrease bacteria on skin.

For hand cleaning to decrease bacteria on skin that could cause disease. Recommended for repeated use.

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Drug Facts

Active ingredient

Purpose

Ethyl Alcohol 62% v/v.....Antibacterial Agent

Uses

For hand cleaning to decrease bacteria on skin.

Warnings

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Directions

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Inactive ingredients

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NDC 63779-056-19

LA312J-0513

Net Contents 800 mL

NDC 63779-056-27

Manufactured in the USA for: CANBERRA CORPORATION 3610 N. Holland-Sylvania Rd. Toledo, OH 43615 P 419-841-6616 • F 419-841-7597

63779-056-19.jpg



515

INSTANT HAND SANITIZER

(GEL-TYPE)

Kills 99.9% of common germs that can cause disease in 15 seconds. Formulated to sanitize hands in situations where water is unavailable. Contains Aloe and Vitamin E

Drug Facts

Active ingredient

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Laihase

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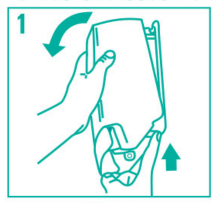
Manufactured in the USA for:

LA453R-0513

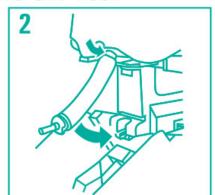
CANBERRA CORPORATION

3610 N. Holland-Sylvania Rd. Toledo, OH 43615

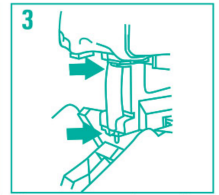
P 419-841-6616 • F 419-841-7597



Open



Load refill and push nozzle back



Secure nozzle and close cover carefully



515

Net Contents 1100 mL NDC 63779-056-77

INSTANT HAND SANITIZER

(GEL-TYPE)

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515

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Manufactured in the USA for:

LA453R-0513

CANBERRA CORPORATION

3610 N. Holland-Sylvania Rd. Toledo, OH 43615

P 419-841-6616 • F 419-841-7597

















Load refill and push nozzle back



Secure nozzle and close cover carefully

63779-056-87.jpg

HUSKY INSTANT HAND SANITIZER

husky instant hand sanitizer solution

Product Information	oduct Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:63779-056	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	62 mL in 100 mL

Inactive Ingredients		
Ingredient Name	Strength	
POLYACRYLIC ACID (250000 MW) (UNII: 9G2MAD7J6W)		
WATER (UNII: 059QF0KO0R)		
GLYCERIN (UNII: PDC6A3C0OX)		
PROPYLENE GLYCOL (UNII: 6 DC9 Q167V3)		
ISOPROPYL MYRISTATE (UNII: 0 RE8 K4LNJS)		
AMINO METHYLPRO PANO L (UNII: LU49 E6626Q)		
ALOE VERA LEAF (UNII: ZY8 1Z8 3H0 X)		
.ALPHATO CO PHERO L ACETATE, DL- (UNII: WR1WPI7EW8)		

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:63779-056-19	237 mL in 1 BOTTLE			
2	NDC:63779-056-87	29 mL in 1 BOTTLE			
3	NDC:63779-056-27	800 mL in 1 BAG			
4	NDC:63779-056-77	1100 mL in 1 BOTTLE, PLASTIC			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333A	05/22/2013		

Labeler - Canberra Corporation (068080621)

Registrant - Kutol Products Company, Inc. (004236139)

Establishment Name Address ID/FEI Business Operations Kutol Products Company, Inc. 004236139 manufacture(63779-056), analysis(63779-056)

Revised: 2/2014 Canberra Corporation