SAFEHANDS HAND SANITIZER WIPES UNSCENTED- benzalkonium chloride cloth Safehands Distribution Ne, LLC

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.

safeHands Hand Sanitizer Wipes

Active Ingredient

Benzalkonium chloride 0.1%

Purpose

Antibacterial

Uses

 To decrease bacteria on the skin. May be used on the face, arms, and body.

Warnings

For external use only.

Do not use in eyes.

In case of contact, rinse eyes thoroughly with water.

Stop use and ask a doctor if

skin/eye irritation develops.

Keep out of reach of children.

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

Directions

- Pull one sheet from soft pack.
- Clean hands or affected area and discard.
- Close lid after each use to keep wipes fresh.

Inactive Ingredients

Water, Glycerin, Lauryl Glucoside, DMDM Hydantoin, Aloe Barbadensis Leaf Extract, Disodium EDTA, PEG-12 Dimethicone, Panthenol, Chamomilla Recutita (Matricaria) Extract, Iodopropynyl Butylcarbamate, Tocopheryl Acetate, Allantoin

Questions/Comments?

Call toll free 1-866-748-9990

Package Labeling

SafeHands®

- Non-toxic
- Moisturizing

ALCOHOL FREE

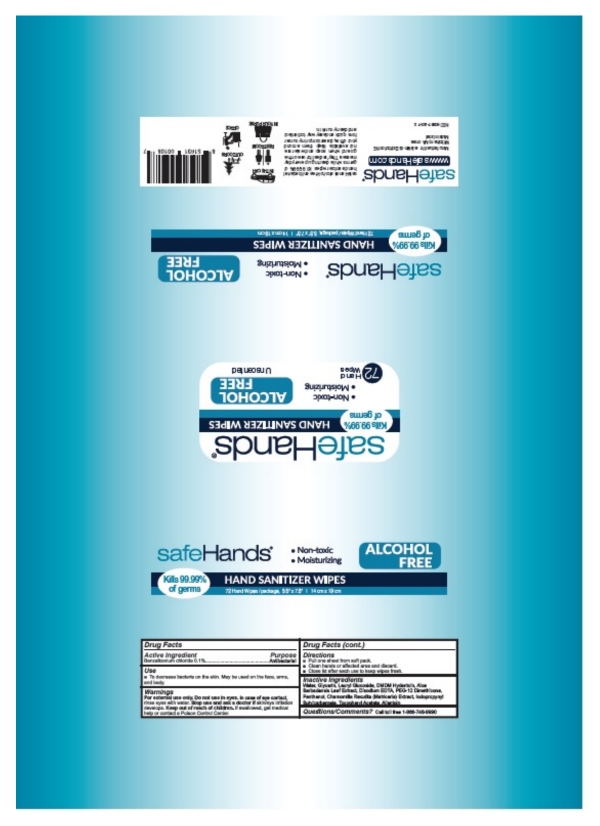
HAND SANITIZEER WIPES

Kills 99.9% of germs

72 Hand Wipes / package, 5.5" x 7.5" | 14 cm x 19 cm

Manufactured for safeHands Distribution NE Wilbraham, MA 01095 Made in Israel

NDC 60867-200-72



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SAFEHANDS HAND SANITIZER WIPES UNSCENTED

benzalkonium chloride cloth

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:60867-200

Active Ingredient/Active Moiety						
Ingredient Name	Basis of Strength	Strength				
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM -	BENZALKONIUM	0.1 g				

Inactive Ingredients				
Ingredient Name	Strength			
WATER (UNII: 059QF0KO0R)				
GLYCERIN (UNII: PDC6A3C0OX)				
LAURYL GLUCOSIDE (UNII: 76LN7P7UCU)				
DMDM HYDANTOIN (UNII: BYR0546TOW)				
ALOE VERA LEAF (UNII: ZY81Z83H0X)				
EDETATE DISODIUM ANHYDROUS (UNII: 8NLQ36F6MM)				
PEG-12 DIMETHICONE (UNII: ZEL54N6W95)				
PANTHENOL (UNII: WV9CM0O67Z)				
MATRICARIA CHAMOMILLA WHOLE (UNII: G0R4UBI2ZZ)				
IODOPROPYNYL BUTYLCARBAMATE (UNII: 603P14DHEB)				
.ALPHATOCOPHEROL ACETATE (UNII: 9E8X80D2L0)				
ALLANTOIN (UNII: 344S277G0Z)				

F	Packaging				
#	tem Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:60867-200- 72	72 in 1 PACKAGE; Type 0: Not a Combination Product	05/12/2021		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333A	05/12/2021		

Labeler - Safehands Distribution Ne, LLC (080026877)

Revised: 11/2021 Safehands Distribution Ne, LLC