KAZU ANTIBACTERIAL HAND SANITIZING WIPES- benzalkonium chloride cloth N Bridge 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.

benzalkonium chloride

water, disodium EDTA, sodium benzoate, glycerin, polysorbate 20, tocopheryl acetate, phenoxyethanol, citric acid

Antiseptic

keep out of reach of the children

wet hands thoroughly with product and allow to dry without wiping

for children under 6, use only under adult supervision

not recommended for infants

For external use only. Flammable. Keep away from heat or flame.

Do not use

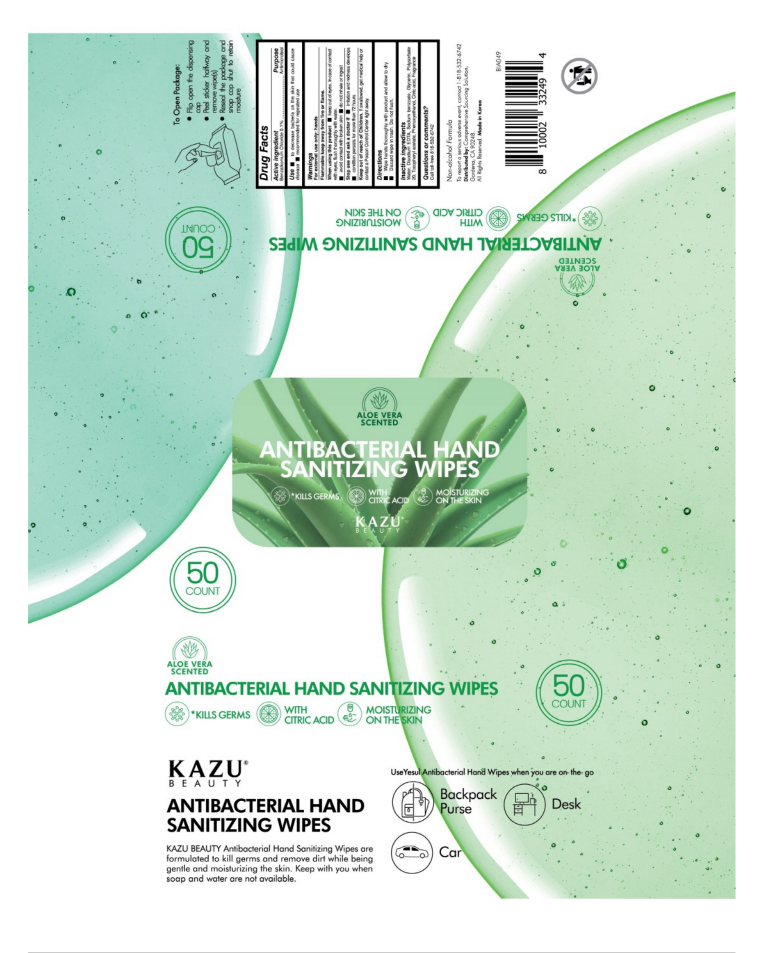
- in children less than 2 months of age
- on open skin wounds

When using this product keep out of eyes, ears, and mouth. In case of contact with eyes, rinse thoroughly with water.

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

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

for external use only



KAZU ANTIBACTERIAL HAND SANITIZING WIPES

benzalkonium chloride cloth

Pro	duct	Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:76966-0017

Route of Administration TOPICAL

Active Ingredient/Active Moiety

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Ingredient Name	Basis of Strength	Strength		
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	0.197 g in 50		

Inactive Ingredients Ingredient Name

Strength

EDETATE DISO DIUM ANHYDRO US (UNII: 8 NLQ36 F6 MM)

SODIUM BENZOATE (UNII: OJ245FE5EU)

PHENO XYETHANOL (UNII: HIE492ZZ3T)

WATER (UNII: 059QF0KO0R)

CITRIC ACID MONOHYDRATE (UNII: 2968 PHW8 QP)

GLYCERIN (UNII: PDC6A3C0OX)

POLYSORBATE 20 (UNII: 7T1F30V5YH)

.ALPHA.-TO COPHEROL ACETATE (UNII: 9E8X80D2L0)

Packaging

# Item Code		Package Description	Marketing Start Date	Marketing End Date
l	1 NDC:76966-0017-1	50 in 1 PACKAGE; Type 0: Not a Combination Product	07/27/2020	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part333A	07/27/2020			

Labeler - N Bridge Inc. (694892752)

Registrant - N Bridge Inc. (694892752)

Establishment

Name	Address	ID/FEI	Business Operations
FIRSTCHAM CO., LTD		689905446	manufacture(76966-0017)

Establishment

Name	Address	ID/FFI	Rusiness Operations			

N Bridge Inc.	694892752	label(76966-0017)

Revised: 7/2020 N Bridge Inc.