KUNDAL PURE AND SAFE SANITIZING TISSUE (BENZALKONIUM CHLORIDE)benzalkonium chloride liquid THESKINFACTORY Co., Ltd.

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

Aloe Barbadensis Leaf Extract Camellia Sinensis Leaf Extract Chamomilla Recutita (Matricaria) Extract Phenoxyethanol Cetylpyridinium Chloride Caprylyl Glycol Cocamidopropyl PG-Dimonium Chloride Phosphate Citric Acid

hand sanitizing tissue to help reduce bacteria on the skin

keep out of reach of the children

pull ouy wipes and sanitize hands when necessary

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



KUNDAL PURE AND SAFE SANITIZING TISSUE (BENZALKONIUM CHLORIDE)

benzalkonium chloride liquid

Product Informa	tion							
Product T ype		HUMAN OTC DRUG Item Code (Source)		NDC:74773	NDC:74773-0027			
Route of Administra	Administration TOPICAL							
Active Ingredien	t/Active Moi	ety						
Ingredient Name					Basis of St	trength	Strengtl	
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6 JUD5X6 Y)					BENZALKONIUM CHLORIDE		0.1 g in 100 g	
Inactive Ingredie	ents							
Ingredient Name						S	Strength	
SODIUM BENZOATE (UNII: OJ245FE5EU)								
ALOE VERA LEAF (UNII: ZY8 1Z8 3H0 X)								
GREEN TEA LEAF (UNII: W2ZU1RY8B0)								
WATER (UNII: 059QF0K00R)								
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)								
MATRICARIA RECUT								
SODIUM CITRATE (U	NII: 1Q73Q2JUL	R)						
POLYSORBATE 20 (UNII: 7T1F30V5Y	H)						
Packaging								
# Item Code		Package Description		Marketi	ng Start Date	Marketi	ng End Dat	
1 NDC:74773-0027-1	120 g in 1 BOTT	LE; Type 0: Not a Combinat	ion Product	09/09/20	20			
Marketing Inf	ormation							
Marketing Catego	ry Applicat	ion Number or Monogra	ph Citation	Marke	ting Start Date	Market	ing End Dat	
OTC monograph not fi	nal part333A			09/09/20)20			

Labeler - THESKINFACTORY Co., Ltd. (694804099)

Registrant - THESKINFACTORY Co., Ltd. (694804099)

Establishment										
Name	Address	ID/FEI	Business Operations							
C&TECH CORPORATION		688204698	manufacture(74773-0027)							

Establishment

Name	Address	ID/FEI	Business Operations
THESKINFACTORY Co., Ltd.		694804099	label(74773-0027)