WET WIPE- biopure antibacterial hand wipe fresh scent 75 ct cloth Osike Cosmetics 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.

79866-012

WET WIPE

Active Ingredient(s)

Benzalkonium Chloride 0.13%Antimicroblal

Purpose

Antimicroblal

Use

- Hand sanitizer to help reduce bacteria
- •For use when soap and water are not available.

Warnings

For external use only.

Do not use

- in children less than 2 months old.
- on open skin wounds

keep out of eyes, ears, and mouth. In case of contact with eyes, rinse eyes 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.

Directions

- Apply to hands, allow to air dry without wiping.
- ●Children under 6 years of age should be supervised when using this product

Other information

- ●Store in a cool, dry place, between 15°C-30°C□59°F-86°F□
- ●Avoid freezing and excessive heat above 40°C 104°F

Inactive ingredients

Aloe Barbadensis Extract, Alpha-Tocopherol, Fragrance, Glycerin, Phenoxyethanol, Propylene Glycol, Purified Water

Package Label - Principal Display Panel

79866-012-01 75



WET WIPE

biopure antibacterial hand wipe fresh scent 75 ct cloth

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:79866-012	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	0.13 in 100		

Inactive Ingredients	
Ingredient Name	Strength
GLYCERIN (UNII: PDC6A3C0OX)	

ALOE VERA LEAF (UNII: ZY81Z83H0X)	
WATER (UNII: 059QF0KO0R)	
.ALPHATOCOPHEROL, DL- (UNII: 7QWA1RIO01)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	

l	Packaging				
	# Item Code Package Description		Marketing Start Date	Marketing End Date	
		NDC:79866-012- 01	75 in 1 DRUM; Type 0: Not a Combination Product	12/07/2020	

Marketing Information			
Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
part333A	12/07/2020		
	Application Number or Monograph Citation	Application Number or Monograph Marketing Start Citation Date	

Labeler - Osike Cosmetics Co., Ltd. (415803943)

Establishment				
Name	Address	ID/FEI	Business Operations	
Osike Cosmetics Co., Ltd.		415803943	manufacture(79866-012)	

Revised: 1/2022 Osike Cosmetics Co., Ltd.