# ZEP FUZION FS ANTIMICROBIAL FOAMING HAND CLEANER- benzalkonium chloride liquid Zep Inc.

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### 66949-132 / 0996 Fuzion FS Antimicrobial Foaming

#### **□**Active ingredient

Benzalkonium Chloride 0.13%

#### **Purpose**

Antiseptic Hand Wash

#### Uses

- Hand washing to decrease bacteria on the skin.
- For use in food processing facilities.

#### Warnings

For external use only.

#### Do not use

Do not use in the eyes; if in eyes, rinse promptly and thoroughly with water.

# Stop use and ask doctor

**Stop use and ask doctor if** skin irritation or redness persists for more than 72 hours.

# When using this product

- Do not swallow.
- If swallowed, do not induce vomiting and if individual is conscious, give large quantities of water to drink and consult a physician immediately.

# Keep out of reach of children and pets

**Keep out of reach of children and pets.** Children must be supervised in use of this product.

# **□** Directions

• Wet hands with water.

- Place hands under dispenser and apply liquid soap.
- Massage soap into hands and writsts, emphasizing back of hands, knuckles and cuticles.
- Rise hands thoroughly and dry.

#### **| | | Other information**

- Store at 20 to 25°C (68 to 77°F).
- Do not freeze.
- Dispose in accordance with all applicable federal, state and local regulations.

### **□Inactive ingredients**

Water, Cocamidopropyl Hydroxysultaine, Lauramine Oxide, Didecyldimonium Chloride, PEG-6 Cocamide, Phenoxyethanol, Iodopropynyl Butylcarbamate, Methylisothiazolinone, Hexanediol

#### **||Questions or comments?**

Call 1-800-I-BUY-ZEP (1-800-428-9937)



#### ZEP FUZION FS ANTIMICROBIAL FOAMING HAND CLEANER

benzalkonium chloride liquid

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:66949-132	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety			
Ingredient Name	<b>Basis of Strength</b>	Strength	
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZ ALKONIUM CHLORIDE	0.13 g in 100 mL	

#### **Inactive Ingredients**

Ingredient Name	Strength	
HEXANEDIOL (UNII: ZIA319275I)		
WATER (UNII: 059QF0KO0R)		
COCAMIDOPROPYL HYDROXYSULTAINE (UNII: 62V75NI93W)		
LAURAMINE OXIDE (UNII: 4F6FC4MI8W)		
PEG-6 COCAMIDE (UNII: YZ6NLA4O1E)		
DIDECYLDIMONIUM CHLORIDE (UNII: JXN40O9Y9B)		
PHENOXYETHANOL (UNII: HIE492ZZ3T)		
IODOPROPYNYL BUTYLCARBAMATE (UNII: 603P14DHEB)		
METHYLISOTHIAZOLINONE (UNII: 229D0E1QFA)		

Packaging				
# Ite	m Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:	66949-132-	4800 mL in 1 CASE; Type 0: Not a Combination Product	04/12/2006	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	505G(a)(3)	04/12/2006		

# **Labeler -** Zep Inc. (030471374)

Establishment				
Name	Address	ID/FEI	Business Operations	
Zep Inc.		112125310	manufacture(66949-132)	

Revised: 10/2023 Zep Inc.