FUZION AB FOAM- benzalkonium chloride liquid Zep Inc.

66949-111 / 3388 Fuzion AB Foam

Active ingredient

Benzalkonium Chloride 0.13%

Purpose

Antibacterial Hand Wash

Uses

For washing to decrease bacteria on hands.

Warnings

For external use only.

Do not use

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

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.

Stop use and ask a doctor

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

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.
- Press pump to dispense product into hands.
- Massage soap into hands and wrists, emphasizing back of hands, knuckles and

cuticles.

- Rinse hands thoroughly and dry.
- Store at 20 to 25°C (68 to 77°F).
- Dispose in accordance with all applicable federal, state and local regulations.

Inactive Ingredients

Water, Cetrimonium Chloride, Lauryl/Myristyl Amidopropyl Amine Oxide, Glycerin, Di-PPG-2 Myreth-10 Adipate, Tetrasodium Iminodisuccinate, Methylchloroisothiazolinone, Methylisothiazolinone, Fragrance, Glutaral, Citric Acid, Red 4, Yellow 5

Questions or comments?

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



Fuzion AB Foaming Hand Soap

FUZION AB FOAM

benzalkonium chloride liquid

		_		-		_				
•	ro	\sim		c+	0		rm	•	T	n
г	ш	u	u	9				10		

Product Type HUMAN OTC DRUG Item Code (Source) NDC:66949-111

Route of Administration TOPICAL

Active Ingredient/Active Moiety

	Ingredient Name	Basis of Strength	Strength
ı			

BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)

BENZALKONIUM - BENZALKONIUM O.13 g in 100 mL

Inactive Ingredients

Ingredient Name	Strength

METHYLISOTHIAZOLINONE (UNII: 229D0E1QFA)

FD&C RED NO. 4 (UNII: X3W0AM1JLX)

ANHYDROUS CITRIC ACID (UNII: XF417D3PSL)

GLUTARAL (UNII: T3C89M417N)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
CETRIMONIUM CHLORIDE (UNII: UC9PE95IBP)	
DI-PPG-2 MYRETH-10 ADIPATE (UNII: 4IN301M0KJ)	
METHYLCHLOROISOTHIAZOLINONE (UNII: DEL7T5QRPN)	
WATER (UNII: 059QF0KO0R)	
LAUROYL/MYRISTOYL AMIDOPROPYL AMINE OXIDE (UNII: HY906ZW9CY)	
GLYCERIN (UNII: PDC6A3C0OX)	
TETRASODIUM IMINODISUCCINATE (UNII: GYS41J2635)	

F	Packaging						
#	tem Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:66949-111- 16	4800 mL in 1 CASE; Type 0: Not a Combination Product	03/10/2017				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC Monograph Drug	505G(a)(3)	03/10/2017			

Labeler - Zep Inc. (030471374)

Establishment						
Name	Address	ID/FEI	Business Operations			
Zep Inc.		112125310	manufacture(66949-111)			

Revised: 11/2024 Zep Inc.