ALCOHOL FREE FOAMING HAND SANITIZER- benzalkonium chloride soap Betco Corporation, 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.

Alcohol Free Foaming Hand Sanitizer

Alcohol Free Foaming Hand Sanitizer

Active Ingredient

Benzalkonium Chloride 0.13%

Alcohol Free Foaming Hand Sanitizer

Uses

- Hand sanitizer to remove microorganisms on the skin.
- Use this product when soap and water are not available.

Alcohol Free Foaming Hand Sanitizer

Warnings

- For external use only.
- Avoid contact with eyes. If contact occurs, rinse thoroughly with water.
- Discontinue use is irritation or redness develops.
- If irritation persists for more than 72 hours, consult a physician.
- KEEP OUT OF REACH OF CHILDREN.
- If swallowed, get medical help or contact a Poison Control Center right away.

Alcohol Free Foaming Hand Sanitizer

Directions

- Read the entire label before using this product.
- Dispense 2 pumps of product onto palm of hand and rub thoroughly over all surfaces of both hands until dry.

Alcohol Free Foaming Hand Sanitizer

Inactive Ingredients

Deionized IWater, Sodium PCA, PEG/PPG-8/3 Laurate, Dimethicone, PEG-3 Cocamide, Fragrance, Methyl Chloro Isothiazolinone, Methyl Isothiazolinone, D&C Green #5.

Alcohol Free Foaming Hand Sanitizer

Questions or Comments? Phone: (800) 777-9343

MDS information: (800) 891-4965

Alcohol Free Foaming Hand Sanitizer

Purpose

Alcohol Free Foaming Hand Sanitizer

KEEP OUT OF REACH OF CHILDREN

Alcohol Free Foaming Hand Sanitizer



Drug Facts

Active Imgredient

Purpose

Datos del Producto

Desinfectante para las manos destinado a redu

Utilice este producto cuando el jabón y el agua

Evite el contacto con los ojos. En caso de cor

MANTENGA ÉSTE PRODUCTO FUERA DEL.

En caso de ingestión, obtenga asistencia mér texicología de inmediato.

 Lea la etiqueta completa antes de usar es: 💇 Aplique 2º dosis del producto en la palma de superficies de ambas manos hasta que

Deje de usarlo si se desarrolla una imitación c la irritación persiste durante más de 72 ho

Ingrediente Activo

· Sólo para uso externo.

de la piel.

Advertencias

Instrucciones

loruro de benzalconio 0.13%

- Hand sanitizer to reduce microorganisms on the skin.
- . Use this product when soap and water are not available

Warnings

- · For external use only.
- . Avoid contact with eyes. If contact occurs, rinse thoroughly with water
- Discontinue use if irritation or redness develops.
 If irritation persists for more than 72 hours, consult a physician.
- KEEP OUT OF REACH OF CHILDREN.
- If swallowed, get medical help or contact a Poison Control Center right away.

Directions.

- Read the entire label before using this product.
 Dispense 2 pumps of product onto palm of hand and rub thoroughly over all surfaces of both hands until dry.

Imactive Ingredients

Deionized Water, Sodium PCA, PEG/PPG-8/3 Laurate, Dimethicone PEG-3 Cocamide, Fragrance, Magnesium Salts, Methyl Chloro Isothiazolinone, Methyl Isothiazolinone, D&C Green #5

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989-GO BETCO (888-462-3826)

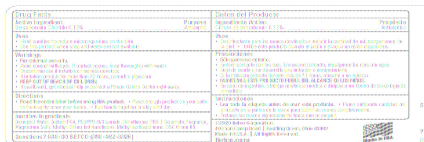
Betca.com





shol Free Foaming Hand Sanitizer

Freshly scented, Alcohol Free Foaming Hand Sanitizer esca perfumada, Espuma sanitizadora para manos sin alcohol









Betco Master - August

Drug Facts Active Ingredient Purpose Inactive Ingredients Deonzed Water, Sodium PCA, PEG/PPG-0/3 Laurate, Dimethicone, PEG-3 Cocamide, Fragran Nation-salam Salts, Methyl Chloro Isothiazolinone, Methyl Isothiazolinone, DAC Green §5.



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Datos del Producto Ingrediente Activo Propósito









Alcohol Free Foaming Hand Sanitizer

Freshly scented, Alcohol Free Foaming Hand Sanitizer Fresca perfumada, Espuma sanitizadora para manos

HAND SANITIZER 752

55 gal. (208 L)

Drug Facts

Active Ingredient

Purpose Antiseptic

Uses

- · Hand sanitizer to reduce microorganisms on the skin.
- . Use this product when soap and water are not available

Warnings

- For external use only.
- Avoid contact with eyes.
- . If contact occurs, rinse thoroughly with water.
- Discontinue use if irritation or redness develops.
- If irritation persists for more than 72 hours, consult a physician.
- KEEP OUT OF REACH OF CHILDREN.
- · If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- · Read the entire label before using this product.
- Dispense 2 pumps of product onto palm of hand and rub thoroughly over all surfaces of both hands until dry.

Inactive Ingredients

Delonized Water, Sodium PCA, PEG/PPG-8/3 Laurate, Dimethicone, PEG-3 Cocamide, Fragrance, Magnesium Salts, Methyl Chloro Isothiazolinone, Methyl Isothiazolinone, D&C Green #5.

Datos del Producto

Ingrediente Activo

Cloruro de benzalconio 0.13%

Usos

- Desinfectante para las manos destinado a red de la piel.
- · Utilice este producto cuando el jabón y el agua

Advertencias

- · Para uso externo únicamente.
- Evite el contacto con los ojos.
- En caso de contacto, enjuáguese los ojos con-
- Deje de usarlo si se desarrolla una irritación o
- Si la irritación persiste durante más de 72 hor
 MANTENER FUERA DEL ALGANCE DE LOS I
- En caso de ingestión, obtenga asistencia médi toxicología de inmediato.

Instrucciones

- Lea toda la etiqueta antes de usar este pr
- Aplique 2 dosis del producto en la palma de la superficies de ambas manos hasta que se seq

SDS No. 752



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Betco Master - 03/201

ALCOHOL FREE FOAMING HAND SANITIZER

benzalkonium chloride soap

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:65601-700
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
BENZALKO NIUM CHLO RIDE (UNII: F5UM2KM3W7) (BENZALKO NIUM - UNII:7N6 JUD5X6 Y)	BENZALKONIUM CHLORIDE	1.3 mg in 1 mL	

Inactive Ingredients	
Ingredient Name	Strength
GRAPEFRUIT OIL (UNII: YR377U58W9)	
BUTYLPHENYL METHYLPROPIONAL (UNII: T7540 GJV69)	
MAGNESIUM NITRATE (UNII: 77CBG3UN78)	
.BETACITRONELLOL, (+/-)- (UNII: 565OK72VNF)	
METHYLCHLORO ISO THIAZO LINO NE (UNII: DEL7T5QRPN)	
CYCLOMETHICONE 4 (UNII: CZ227117JE)	
ACETALDEHYDE (UNII: GO 1N1ZPR3B)	
GERANIOL (UNII: L837108USY)	
HEXAMETHYLINDANO PYRAN (UNII: 14170060AT)	
DIHYDROMYRCENOL (UNII: 46L1B02ND9)	
METHYL DIHYDROJASMONATE (SYNTHETIC) (UNII: 3GW44CIE3Y)	
MYRCENE (UNII: 3M39 CZS25B)	
SODIUM PIDOLATE (UNII: 1V74VH163T)	
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
N-ALKYL DIMETHYL BENZYL AMMONIUM CHLORIDE (C12-C18) (UNII: 9U1Q4T4ZYS)	
LINALOOL, (+/-)- (UNII: D81QY6188E)	
ORANGE OIL (UNII: AKN3KSD11B)	
D&C GREEN NO. 5 (UNII: 8J6RDU8L9X)	
PEG/PPG-15/15 ALLYL ETHER ACETATE (UNII: 8 RP39 FN7AJ)	
HEXYL SALICYLATE (UNII: 8F78EY72YL)	
ALCOHOL (UNII: 3K9958V90M)	
LIME OIL (UNII: UZH29 XGA8 G)	
METHYLISOTHIAZOLINONE (UNII: 229 D0 E1QFA)	
WATER (UNII: 059QF0KO0R)	
DIO XANE (UNII: J8 A3S 10 O7S)	
PEG-6 CO CAMIDE (UNII: YZ6 NLA4O 1E)	
LEMON OIL (UNII: 19 GRO 8 2 4 LL)	

Product Characteristics				
Color	blue	Score		
Shape		Size		

Flavor	Imprint Code	
Contains		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65601-700- 53	50 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	0 1/0 1/20 16	
2	NDC:65601-700- 29	1000 mL in 1 BAG; Type 0: Not a Combination Product	0 1/0 1/20 16	
3	NDC:65601-700- 04	3780 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	0 1/0 1/20 16	
4	NDC:65601-700- 57	550 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	0 1/0 1/20 16	09/11/2020
5	NDC:65601-700- 55	207900 mL in 1 DRUM; Type 0: Not a Combination Product	0 1/0 1/20 16	
6	NDC:65601-700- 03	750 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	09/15/2016	07/15/2020
7	NDC:65601-700- 05	1250 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	09/15/2016	09/11/2020
8	NDC:65601-700- 88	500 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	11/12/2012	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	11/12/2012	

Labeler - Betco Corporation, Ltd. (024492831)

Registrant - Betco corporation, Ltd. (024492831)

Establishment			
Name	Address	ID/FEI	Business Operations
Betco Corporation, Ltd.		024492831	manufacture(65601-700), pack(65601-700), label(65601-700)

Revised: 9/2020 Betco Corporation, Ltd.