

SIGNATRY ANTIMICROBIAL FOAM- chloroxylenol liquid
GOJO Industries, Inc.

SIGNATRY Antimicrobial Foam Soap

Active ingredient

Chloroxylenol 0.3%

Purpose

Antimicrobial

Uses

- Handwash to help decrease bacteria on the skin
- Recommended for repeated use

Warnings

For external use only

When using this product do not use in or near the eyes. In case of contact, rinse eyes thoroughly with water.

Stop use and ask a doctor if irritation or rash appears and lasts

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- Wet hands
- Apply a small amount of product and work into a lather
- Rinse well and dry hands completely

Inactive ingredients

Water (Aqua), Alcohol, Ammonium Laureth Sulfate, Ammonium Lauryl Sulfate, Propylene Glycol, Ammonium Xylenesulfonate, Cocamide MEA, Glycerin, Isopropyl Alcohol, Lactic Acid, Retinyl Palmitate, Simmondsia Chinensis (Jojoba) Seed Oil, Tetrasodium EDTA, Tocopheryl Acetate, Zea Mays (Corn) Oil, Ammonium Sulfate, Fragrance (Parfum), Methylchlorisothiazolinone, Methylisothiazolinone, Red 4 (CI 14700), Yellow 6 (CI 15985)



S6162-04



Made in U.S.A. for,
 Hecho en los E.E.U.U.
 Distributed by,
 Distribuido por:
Signatry, Inc.
 Akron, OH 44309
1-800-321-9647
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**ANTIMICROBIAL
 FOAM SOAP**
**ESPUMA DE JABÓN
 ANTIMICROBIANA**

1.25L (42 FL OZ)

6162-640-SIG-F

Drug Facts

Active Ingredient	Purpose
Chloroxylenol 0.3%.....	Antimicrobial

Uses

- Handwash to help decrease bacteria on the skin
- Recommended for repeated use

Warnings
For external use only

When using this product do not use in or near the eyes. In case of contact, rinse eyes thoroughly with water.

Stop use and ask a doctor if irritation or rash appears and lasts

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- Wet hands
- Apply product and thoroughly cover hands with lather
- Rinse well and dry hands completely

Drug Facts (continued)

Directions

- Wet hands
- Apply product and thoroughly cover hands with lather
- Rinse well and dry hands completely

Inactive Ingredients Water (Aqua), Alcohol, Ammonium Laureth Sulfate, Ammonium Lauryl Sulfate, Propylene Glycol, Ammonium Xylenesulfonate, Cocamide MEA, Glycerin, Isopropyl Alcohol, Lactic Acid, Retinyl Palmitate, Simmondsia Chinensis (Jojoba) Seed Oil, Tetrasodium EDTA, Tocopheryl Acetate, Zea Mays (Corn) Oil, Ammonium Sulfate, Fragrance (Parfum), Methylchloroisothiazolinone, Methylisothiazolinone, Red 4 (CI 14700), Yellow 6 (CI 15985)

5362-644-ES-E

SIGNATRY ANTIMICROBIAL FOAM			
chloroxylenol liquid			
Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:21749-581
Route of Administration	TOPICAL		
Active Ingredient/Active Moiety			
Ingredient Name		Basis of Strength	Strength

CHLOROXYLENOL (UNII: 0F32U78V2Q) (CHLOROXYLENOL - UNII:0F32U78V2Q)	CHLOROXYLENOL	0.003 mg in 1 mL
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Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
ALCOHOL (UNII: 3K9958V90M)	
AMMONIUM LAURETH-2 SULFATE (UNII: 698O4Z48G6)	
AMMONIUM LAURYL SULFATE (UNII: Q7AO2R1M0B)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
AMMONIUM XYLENESULFONATE (UNII: 4FZY6L6XCM)	
COCO MONOETHANOLAMIDE (UNII: C80684146D)	
GLYCERIN (UNII: PDC6A3C0OX)	
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
LACTIC ACID (UNII: 33X04XA5AT)	
VITAMIN A PALMITATE (UNII: 1D1K0N0VVC)	
JOJOBA OIL (UNII: 724GKU717M)	
EDETATE SODIUM (UNII: MP1J8420LU)	
.ALPHA.-TOCOPHEROL ACETATE, D- (UNII: A7E6112E4N)	
CORN OIL (UNII: 8470G57WFM)	
AMMONIUM SULFATE (UNII: SU46BAM238)	
METHYLCHLOROISOTHIAZOLINONE (UNII: DEL7T5QRPN)	
METHYLISOTHIAZOLINONE (UNII: 229D0E1QFA)	
FD&C RED NO. 4 (UNII: X3W0AM1JLX)	
FD&C YELLOW NO. 6 (UNII: H77VEI93A8)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:21749-581-89	1200 mL in 1 PACKAGE; Type 0: Not a Combination Product	09/09/2004	
2	NDC:21749-581-90	1250 mL in 1 PACKAGE; Type 0: Not a Combination Product	09/09/2004	
3	NDC:21749-581-43	1500 mL in 1 PACKAGE; Type 0: Not a Combination Product	09/09/2004	11/26/2024
4	NDC:21749-581-67	2000 mL in 1 PACKAGE; Type 0: Not a Combination Product	09/09/2004	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	505G(a)(3)	09/09/2004	

Labeler - GOJO Industries, Inc. (004162038)

Establishment

Name	Address	ID/FEI	Business Operations
GOJO Industries, Inc.		036424534	manufacture(21749-581)

Establishment

Name	Address	ID/FEI	Business Operations
GOJO Industries, Inc.		088312414	label(21749-581) , manufacture(21749-581)

Revised: 11/2024

GOJO Industries, Inc.