

KAY RTU FOAMING AB HANDSOAP- chloroxylenol solution
Ecolab Inc.

Drug Facts

Active ingredient

Chloroxylenol 0.5%

Purpose

Antiseptic Handwash

Uses

- for handwashing to decrease bacteria on the skin.

Warnings

- **For external use only**

Do not use

- In eyes

When using this product

- If in eyes, rinse promptly and thoroughly with water
- Discontinue use if irritation and redness develop

Stop use and ask a doctor if

- Skin irritation or redness occurs for more than 72 hours

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- Wet hands to remove soil
- Dispense palmful
- spread to cover hands, rub in well
- air dry, do not rinse or towel dry

Other Information

- for additional information, see Safety Data Sheet (SDS)
- **EMERGENCY HEALTH INFORMATION:** 1 877 231 2615. If located outside the United States and Canada, call collect 952 853 1713 (number is in the US).

Inactive ingredients water (aqua), potassium cocoate, hexylene glycol, sodium sulfate, tetrasodium EDTA, sodium lauryl sulfate, glycerin, caprylyl/capryl glucoside, fragrance, lauryl glucoside, red 40, methylchloroisothiazolinone, yellow 5, methylisothiazolinone

Questions? Call **1-800-529-5458**

Principal display panel and representative label

ECOLAB

Kay RTU Foaming AB Hand Soap

Active Ingredient: Chloroxylenol 0.5%

1200 mL

(40.6 US FL OZ)

1112847

Kay Chemical Company · 8300 Capital Drive

Greensboro, NC 27409-9790 USA

Customer Service: (800) 529-5458

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Directions
■ wash hands to remove soil
■ dispense palmful
■ spread to cover hands, rub in well
■ air dry, do not rinse or towel dry

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This product may be patented: www.ecolab.com/patents
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KAY RTU FOAMING AB HANDSOAP
chloroxylenol solution

Product Information				
Product Type		HUMAN OTC DRUG	Item Code (Source)	NDC:47593-693
Route of Administration		TOPICAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
CHLOROXYLENOL (UNII: 0F32U78V2Q) (CHLOROXYLENOL - UNII:0F32U78V2Q)	CHLOROXYLENOL	5 mg in 1 mL

Inactive Ingredients	
Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
POTASSIUM COCOATE (UNII: F8U72V8ZXP)	
HEXYLENE GLYCOL (UNII: KEH0A3F75J)	
SODIUM SULFATE (UNII: 0YPR65R21J)	
EDETATE SODIUM (UNII: MP1J8420LU)	
SODIUM LAURYL SULFATE (UNII: 368GB5141J)	
GLYCERIN (UNII: PDC6A3C0OX)	
CAPRYLYL/CAPRYL GLUCOSIDE (UNII: E00JL9G9K0)	
LAURYL GLUCOSIDE (UNII: 76LN7P7UCU)	
FD&C RED NO. 40 (UNII: WZB9127XOA)	
METHYLCHLOROISOTHIAZOLINONE (UNII: DEL7T5QRPN)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
METHYLISOTHIAZOLINONE (UNII: 229D0E1QFA)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:47593-693-70	1200 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	12/30/2025	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	505G(a)(3)	12/30/2025	

Labeler - Ecolab Inc. (006154611)