ANTI-BACTERIAL HAND EUCALYPTUS SPEARMINT- alcohol gel Bath & Body Works, Inc.

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.

DRUG FACTS

ACTIVE INGREDIENT

Alcohol 71%

PURPOSE

Antiseptic

USE

Decrease bacteria on hands.

WARNINGS

For external use only.

When using this product keep out of eyes. Stop use and ask a doctor if irritation or redness develops.

FLAMMABLE

Keep away from flame or high heat.

KEEP OUT OF REACH OF CHILDREN

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

DIRECTIONS

• Rub a dime sized drop into hands.

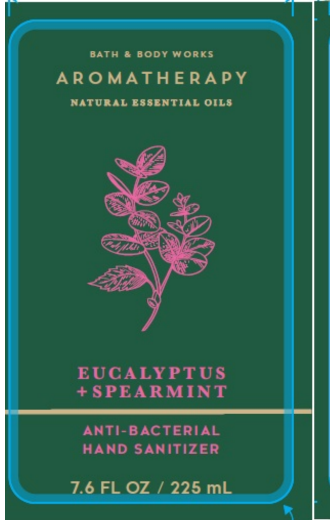
INACTIVE INGREDIENTS

Water (Aqua, Eau), Fragrance (Parfum), Carbomer, Lactose, Aminomethyl Propanol, Isopropyl Myristate, Cellulose, Propylene Glycol, Hydroxyethyl Urea, Tocopheryl Acetate, Wheat Amino Acids, Aloe Barbadensis Leaf Juice, Butyrospermum Parkii (Shea) Butter Extract, Hydroxypropyl Methylcellulose, Retinyl Palmitate, Ultramarines (CI 77007), Ext. Violet 2 (CI 60730), Yellow 5 (CI 19140).

COMPANY INFORMATION

Bath & Body Works, Distr. Reynoldsburg, Ohio 43068 1-800-395-1001

PRODUCT PACKAGING





ANTI-BACTERIAL HAND EUCALYPTUS SPEARMINT

alcohol gel

Droduct	Information
Product	Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:62670-5912

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name		Basis of Strength	Strength
I	ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	68 mL in 100 mL

Inactive 1	lngredients
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Ingredient Name	Strength
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WATER (UNII: 059QF0KO0R)

F	Packaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:62670-5912- 0	29 mL in 1 BOTTLE; Type 0: Not a Combination Product	11/11/2020	
2	NDC:62670-5912- 1	73 mL in 1 BOTTLE; Type 0: Not a Combination Product	11/11/2020	
3	NDC:62670-5912-3	225 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	11/11/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	11/11/2020	

Labeler - Bath & Body Works, Inc. (878952845)

Revised: 11/2020 Bath & Body Works, Inc.