BROMPTON AND LANGLEY LAVENDER VANILLA HAND SANITIZER LAVENDER VANILLA- alcohol denat gel

Cita International Limited

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.

Active Ingredient

Alcohol 62%

□**Purpose:**□ Antiseptic

 \Box Use

To decrease bacteria on the skin that could cause disease.

Keep out of reach of children.

Warnings

- Keep away from flame or fire
- for external use only-hands.
- keep out of eyes.
- stop use and ask a Doctor if irritation or redness persists for more than 72 hours
- do not inhale or ingest. if swallowed, get medical help or contact a poison control center right away.

Directions

- spray a thumb size amount into hands.
- rub hands together gently making sure to cover all areas in between fingers
- let hands air dry

Inactive Ingredients

water (aqua/eau), glycerin, propylen glycol, Isopropyl alcohol, phenoxythenol, acrylates,

lauryl glycol ether, carbomer, Vitamin E, fragrance (parfum).

Brompton and Langley Lavender Vanilla Scentend Hand Sanitizer 15mL 7/11/2016 image027.png



alcohol denat gel

Product Information

Product TypeHUMAN OTC DRUGItem Code (Source)NDC:70805-006

Route of Administration TOPICAL

Active Ingredient/Active Moiety

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	Ingredient Name	Basis of Strength	Strength	
	ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	0.26 g in 26 g	

Inactive Ingredients			
Ingredient Name	Strength		
CARBOMER INTERPOLYMER TYPE A (55000 CPS) (UNII: 59TL3WG5CO)			
WATER (UNII: 059QF0KO0R)			
GLYCERIN (UNII: PDC6A3C0OX)			
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)			
ISOPROPYL ALCOHOL (UNII: ND2M416302)			
PHENOXYETHANOL (UNII: HIE492ZZ3T)			
METHO XY PEG-40 (UNII: 6AXS45P1QU)			
ISOPROPYL MYRISTATE (UNII: 0 RE8 K4LNJS)			
.ALPHATO COPHEROL ACETATE (UNII: 9E8X80D2L0)			

Product Characteristics			
Color		Score	
Shape		Size	
Flavor	VANILLA (Lavender Vanilla)	Imprint Code	
Contains			

l	Packaging			
	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:70805-006- 01	15 g in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	07/13/2016	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333E	06/03/2016		

Labeler - Cita International Limited (665926739)

Establishment					
Name	Address	ID/FEI	Business Operations		
Cita International Limited		665926739	manufacture(70805-006)		

Revised: 7/2016 Cita International Limited