# PULLIO HAND SANITIZING WIPES LAVENDER- benzalkonium chloride cloth YES SALES

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.

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# **Drug Facts**

#### ACTIVE INGREDIENT

Benzalkonium Chloride 0.13%

#### INACTIVE INGREDIENTS

Water, Glycerin, Sodium Benzoate, Ethylhexylglycerin, Polysorbate 20, Lavandula Angustifolia (Lavender) Extract, Butylene Glycol, 1,2-Hexanediol, Disodium EDTA, Citric Acid, Tocopheryl Acetate, Fragrance

#### **PURPOSE**

Antiseptic

#### WARNINGS

## For external use only

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# Stop use and ask a doctor if

- hypersensitivity symptoms such as erythema, itching and dermatitis happen
- skin irritation happens

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#### Do not use

- in combination with soap or antibacterial cleansing agents
- the product for a long time in the same area as swelling, inflammation or sickness may occur due to absorption through the skin

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#### When using this product

- aviod using repeatedly in the same area, skin irritation may occur
- avoid getting into the eyes (if contact occurs, wash well with clean water)
- if following abnormal symptoms persist, discontinue use :

Irritation around the eyes, ears, mucous membranes, including the mouth, under the skin irritation and rashes

## KEEP OUT OF REACH OF CHILDREN

#### Keep out of reach of children

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

#### Uses

■ Instant hand antiseptic to decrease bacteria on the skin

## **Directions**

- Clean with wipes and let dry
- Do not flush

## Other information

- Read the directions and warnings before use
- Avoid freezing and excessive heat above 40C (104F)

# PACKAGE LABEL.PRINCIPAL DISPLAY PANEL



# PULLIO HAND SANITIZING WIPES LAVENDER benzalkonium chloride cloth Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:80618-020 Route of Administration TOPICAL

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>BENZALKONIUM CHLORIDE</b> (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	0.13 in 100	

Inactive Ingredients		
Ingredient Name	Strength	
WATER (UNII: 059QF0KO0R)		
GLYCERIN (UNII: PDC6A3C0OX)		
SODIUM BENZOATE (UNII: OJ245FE5EU)		
Ethylhexylglycerin (UNII: 147D247K3P)		
Polysorbate 20 (UNII: 7T1F30V5YH)		
LAVANDULA ANGUSTIFOLIA SUBSP. ANGUSTIFOLIA FLOWERING TOP (UNII: 9 YT4B71U8P)		
Butylene Glycol (UNII: 3XUS85K0RA)		
1,2-Hexanediol (UNII: TR046 Y3K1G)		
EDETATE DISODIUM ANHYDROUS (UNII: 8 NLQ36 F6 MM)		
CITRIC ACID MO NO HYDRATE (UNII: 2968 PHW8 QP)		
.ALPHATO COPHERO L ACETATE (UNII: 9E8 X80 D2L0)		

l	Packaging				
ı	#	Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
l	1	NDC:80618-020-01	60 in 1 PACKET; Type 0: Not a Combination Product	09/01/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	09/01/2020	

# Labeler - YES SALES (080733755)

# **Registrant -** YES SALES (080733755)

Establishment				
Name	Address	ID/FEI	Business Operations	
NAICO		694725335	manufacture(80618-020)	

Revised: 9/2020 YES SALES