#### HAND SANITIZER- benzalkonium chlorida gel Northmed

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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#### kids gel, 1L

### **Active Ingredient**

benzalkonium chloride .2%. Purpose: Antiseptic

#### Purpose

Antiseptic, Hand Sanitizer

#### Use

Hand Sanitizer to help reduce bacteria that potentially can cause disease. For use when soap and water are not available.

### Warnings

For external use only.

#### Do not use

- in children less than 2 months of age
- on open skin wounds

When using this product keep out of eyes, ears, and mouth. In case of contact with eyes, rinse eyes thoroughly with water.

Stop use and ask a doctor if irritation or rash occurs. These may be signs of a serious condition.

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

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#### Directions

Directions. Apply a small amount of gel to the hands and massage. Wait for it to dry. The exposure time is 15-30 seconds. Supervise children under 6 years of age when using this product to avoid swallowing

## Other information

- Store between 41-80F (5-27C).
- Avoid freezing and excessive heat above 40C (104F).

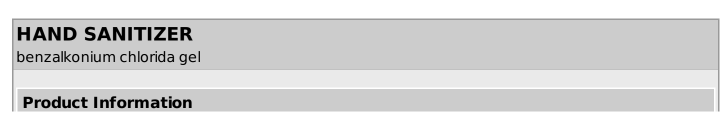
# Inactive ingredients

Water. glycerin, panthenol, aloe vera, hydroxyethyl cellulose, tetrasodium glutamate diacetate, fragrance, chamomile extract, birch leaf extract, elderberry extract.

# Package Label - Principal Display Panel



1L NDC: 78522-102-00



Product Type		HUMAN OTC DRUG Item Code (S		Code (Sou	<b>urce)</b> NDC:78522-102		2-102		
Route of Administration		TOPICAL							
Active Ingredient/Active Moiety									
Ingredient Name						Basis of Strength Streng			
					BENZALKONIUM 2 mg CHLORIDE in 1 L				
Inactive Ingredients									
Ingredient Name							Strength		
CHAMOMILE FLOWER OIL (UNII: 60F80Z61A9)									
HYDROXYETHYL CELLULOSE, UNSPECIFIED (UNII: T4V6TWG28D)									
PANTHENOL (UNII: WV9CM0067Z)									
FRAGRANCE FLORAL ORC0902236 (UNII: R66Z4YW3X0)									
TETRASODIUM GLUTAMATE DIACETATE (UNII: 5EHL50I4MY)									
BIRCH TRITERPENES (UNII: BX09B0RQR0)									
GLYCERIN (UNII: PDC6A3C0OX)									
ALOE VERA LEAF (UNII: ZY81Z83H0X)									
WATER (UNII: 059QF0KO0R)									
Packaging									
# Item Code	Pa	ckage Description		Marketi Da	ng Start Ite		ing End ate		
	. L in 1 BOTTI Product	E; Type 0: Not a Combinat	ion 0	3/16/2020					
Marketing Information									
Marketing Category	Applica	tion Number or Monog Citation	graph		ting Start Date		ting End ate		
OTC monograph not final	part333A			03/16/202	20				

Labeler - Northmed (662588132)

Establishment							
Name	Address	ID/FEI	Business Operations				
Northmed		662588132	manufacture(78522-102) , pack(78522-102) , label(78522-102)				

Revised: 3/2021

Northmed