# ALCOHOL WIPE- alcohol cloth YAHON ENTERPRISE CO.,LTD

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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Yahon 001-06

## **Active Ingredient(s)**

Ethyl Alcohol 75% (alc/vol)

### **Purpose**

**Antiseptic** 

#### Use

Sanitizing wipes to help reduce bacteria on the skin or surfaces

# **Warnings**

Flammable, keep away from heat, spark, electrical fire or flame.

Avoid contact with eyes. If in eyes, remove contact lenses (if any), rinse slowly with water or get medical help.

Stop use and ask a doctor if you have deep or puncture wounds, animal bites serious burns

Irritation and redness develop or condition persists 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**

Open cap, remove seal and pull wipes out.

Discard after single use and do not flush

Keep cap and seal tightly closed between uses to prevent from drying out.

#### Other information

Do not apply internally. Store in a cool dry place below 30°C

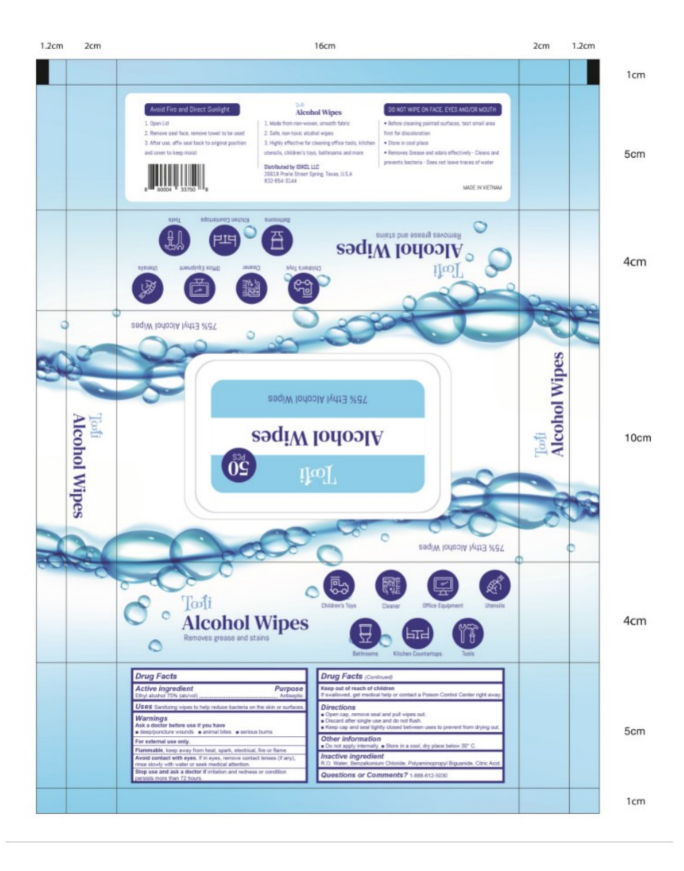
# **Inactive ingredients**

Water, Benzalkonium Chloride, Polyaminopropyl Biguanide, Citric Acid

# Package Label - Principal Display Panel







# ALCOHOL WIPE alcohol cloth Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:79618-001

Route of Administration TOP
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Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	75 mL in 100		

Inactive Ingredients		
Ingredient Name	Strength	
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)		
CITRIC ACID ACETATE (UNII: DSO12WL7AU)		
WATER (UNII: 059QF0KO0R)		
POLYAMINOPROPYL BIGUANIDE (UNII: DT9D8Z79ET)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:79618-001- 06	50 in 1 PACKET; Type 0: Not a Combination Product	07/21/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	07/21/2020	

# Labeler - YAHON ENTERPRISE CO.,LTD (555347945)

Establishment				
Name	Address	ID/FEI	<b>Business Operations</b>	
YAHON ENTERPRISE CO.,LTD		555347945	manufacture(79618-001)	

Revised: 12/2022 YAHON ENTERPRISE CO.,LTD