ALCOHOL WIPES- alcohol cloth Vara Home Usa Llc

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.

Alcohol Wipes

Active Ingredient(s)

Alcohol 75% v/v. Purpose: Antiseptic

Purpose

Antiseptic, Alcohol wipes

Use

- For hand santitizing to decrease bacteria on the skin.
- Apply topically to the skin to help prevent cross contamination.
- Not recommended for repeated use.
- Dries in seconds.

Warnings

Flammable. Keep away from fire or flame.

For external use only.

Do not use

in or contact the eyes.

Stop use and ask a doctor

if too much skin irritation and sensitivity develops or increases

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

Directions

- Gently pull back resealable label, remove and use wipe as required.
- Reseal back after use to avoid evaporation of alcohol.

Other information

- Store at room temperature 15-30 (59-86°F).
- Lot No. Manufacture date and expiration date can be found on package.

Inactive ingredients

water

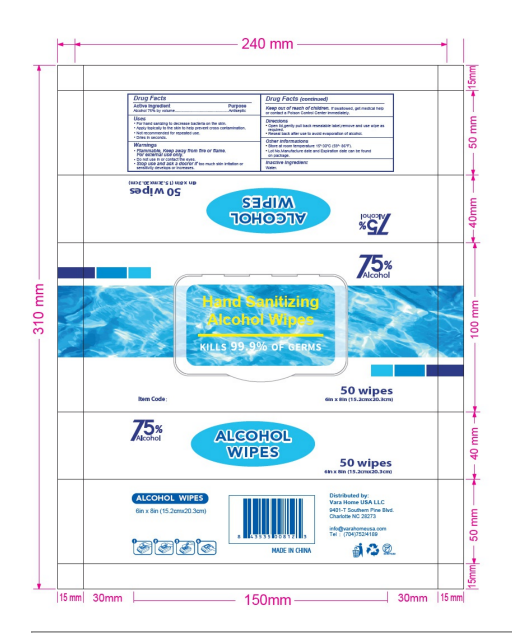
Package Label - Principal Display Panel

1 wipe NDC: 79060-001-01



10 wipes NDC: 79060-001-02

120mm





We recinnend not to completely remove this label. Close the label and lid to preserve the alcohol content.

ALCOHOL WIPES

alcohol cloth

Dи	Λd	nct	Info	rma	tion

Product Type HUMAN OTC DRUG Item Code (Source) NDC:79060-001(NDC:75162-1001)

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	0.75 mL in 1 g

Inactive Ingredients

	Ingredient Name	Strength
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WATER (UNII: 059QF0KO0R)

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:79060-001-01	1 in 1 PACKAGE	06/12/2020		
1		1.35 g in 1 PATCH; Type 0: Not a Combination Product			
2	NDC:79060-001-02	10 in 1 PACKAGE	06/12/2020		
2	1.35 g in 1 PATCH; Type 0: Not a Combination Product				
3	NDC:79060-001-03	50 in 1 PACKAGE	06/12/2020		
3		1.35 g in 1 PATCH; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333A	06/12/2020		

Labeler - Vara Home Usa Llc (802033196)

Registrant - Shandong Mainclean Medical Products Co., Ltd. (418445935)

Establishment					
Name	Address	ID/FEI	Business Operations		
Shandong Mainclean Medical Products Co., Ltd.		418445935	manufacture(79060-001), relabel(79060-001)		

Revised: 6/2020 Vara Home Usa Llc