#### PROTECT HAND SANITIZER- alcohol solution Boya Biotechnology Company 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.

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## Active Ingredient(s)

Alcohol 75% v/v. Purpose: Antiseptic

## Purpose

Antiseptic

## Use

For hand washing to decrease bacteria on the skin

## Warnings

For external use only. Flammable. Keep away from fire or flame

#### Do not use

Do not use in the eyes. In case of contatc, rinse eyes thoroughly with water.

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.

Stop use and ask a doctor if redness or irritation develops and persist 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

- Wash hands thoroughly with product and allow to dry without wiping.
- Supervise children in the use of this product to avoid swallowing.

## Other information

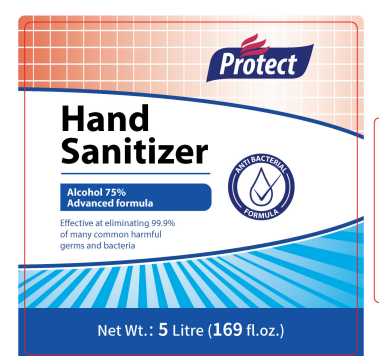
- Store between 15-30C (59-86F)
- May discolor fabrics

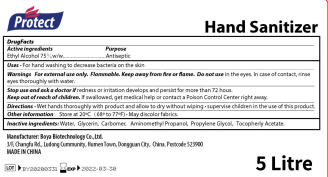
## **Inactive ingredients**

Water, Glycerin, Carbomer, Aminomethyl Propanol, Propylene Glycol, Tocopherol Acetate













	Instant Hand Sanitizer		
Protect	Drug Facts         Manufacturer           Active ingredients         Purpose           Ethyl Alcohol 75 % w/wAntiseptic		
WATERLESS	Uses · For hand washing to decrease bacteria on the skin Warnings For external use only. Flammable. Keep away from fire or flame. Do not use in the eyes. In case		
ANTI-BACTERIAL HAND SANITIZER GEL	of contact, rinse eyes thoroughly with water. Stop use and ask a doctor if redness or irritation develops and persist for more than 72 hous. Keep out of reach of children. If swallowed, get medical		
A STEPACTER	Drug Facts       Purpose         Active ingredients       Purpose         Ethyl Alcohol 75 % w/wAntiseptic       Uses · For hand washing to decrease bacteria on the skin         Warnings For external use only. Flammable. Keep       away from fire or flame. Do not use in the eyes. In case         of contact, rinse eyes thoroughly with water.       Stop use and ask a doctor if redness or irritation         develops and persist for more than 72 hous.       Keep out of reach of children. If swallowed, get medical         help or contact a Poison Control Center right away.       Directions · Wet hands thoroughly with product and         allow to dry without wiping · supervise children in the       use of this product.         Other information:       Store at 20°C (68° to 77°F) · May         discolor fabrics.       Inactive ingredients: Water, Glycerin, Carbomer,         Aminomethyl Propanol, Propylene Glycol, Tocopherly       Acetate.		
FORMULA	Other information:       Store at 20°C (68° to 77°F) • May         discolor fabrics.		
Effective at eliminating 99.9% of many common harmful germs and bacteria	Inactive ingredients: Water, Glycerin, Carbomer, Aminomethyl Propanol, Propylene Glycol, Tocopherly Acetate.		
40ml <b>e</b> 1.35 fl.oz	LOT BY20200331 SEXP 2022-03-30		



		<b>DrugFacts</b>	
	<b>Protect</b>	Active ingredients	Purpose
		Ethyl Alcohol 75%w/v Uses · For hand washing	
		bacteria on the skin	<u> </u>
Uand		Warnings For extern	
Hand		Flammable. Keep awa flame. Do not use in t	
	TIBACTER	contact, rinse eyes the	oroughly with water.
Sanitiz		Stop use and ask a do irritation develops and	
		than 72 hous.	
Alashal 750/		Keep out of reach of c	
Alcohol 75% Advanice formula		swallowed, get medic Poison Control Center	al help or contact a
Advance formula	ORMMUL	Directions • Wet hand	s thoroughly with
Effective at eliminating 99.9	%	product and allow to a	ary without wiping •
of many common harmful germs and bacteria		supervise children in t	
gernis and bacteria		product. Other information :	Store at 20°C (68°
		to 77°F) • May discolor	
		Inactive ingredients:	Water,Glycerin,
		Carbomer, Aminomet ene Glycol, Tocopherly	hyl Propanol,Propyl-
		Manufacturer: Boya B 3/F, Changfu Rd., Ludong C	
Net W	/t.: 100ml	Dongguan City, Chin MADE IN	a. Postcode 523900
	NITIZER		
lcohol solution	NITIZER		
lcohol solution Product Information	NITIZER HUMAN OTC DRUG	Item Code (Source)	NDC:73885-001
lcohol solution Product Information Product Type		Item Code (Source)	NDC:73885-001
lcohol solution Product Information Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:73885-001
lcohol solution Product Information Product Type Route of Administration Active Ingredient/Active	HUMAN OTC DRUG TOPICAL Moiety		
lcohol solution <b>Product Information</b> Product Type Route of Administration Active Ingredient/Active In	HUMAN OTC DRUG TOPICAL Moiety agredient Name	Basis of Stren	gth Strength
lcohol solution <b>Product Information</b> Product Type Route of Administration Active Ingredient/Active In	HUMAN OTC DRUG TOPICAL Moiety		
	HUMAN OTC DRUG TOPICAL Moiety agredient Name	Basis of Stren	gth Strength
llcohol solution Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M)	HUMAN OTC DRUG TOPICAL Moiety gredient Name (ALCOHOL - UNII:3K9958V90M)	Basis of Stren	gth Strength 75 mL in 100 mL
llcohol solution Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M) Inactive Ingredients	HUMAN OTC DRUG TOPICAL MOiety gredient Name (ALCOHOL - UNII:3K9958V90M)	Basis of Stren	gth Strength
lcohol solution Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M) Inactive Ingredients GLYCERIN (UNII: PDC6A3C0OX	HUMAN OTC DRUG TOPICAL MOiety agredient Name (ALCOHOL - UNII:3K9958V90M) Ingredient Name	Basis of Stren	gth Strength 75 mL in 100 mL
lcohol solution Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M) Inactive Ingredients GLYCERIN (UNII: PDC6 A3C00X CARBOMER 934 (UNII: Z135WT9	HUMAN OTC DRUG TOPICAL MOiety agredient Name (ALCOHOL - UNII:3K9958V90M) Ingredient Name	Basis of Stren	gth Strength 75 mL in 100 mL
lcohol solution Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M) Inactive Ingredients GLYCERIN (UNII: PDC6A3C0OX CARBOMER 934 (UNII: Z135WT9 WATER (UNII: 059QF0K00R)	HUMAN OTC DRUG TOPICAL Moiety agredient Name (ALCOHOL - UNII:3K9958V90M) Ingredient Name	Basis of Stren	gth Strength 75 mL in 100 mL
lcohol solution Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M) Inactive Ingredients GLYCERIN (UNII: PDC6A3C00X CARBOMER 934 (UNII: Z135WTS WATER (UNII: 059QF0K00R) PROPYLENE GLYCOL (UNII: 61	нимам отс DRUG торісаL Moiety agredient Name (ALC HOL - UNII:3K9958V90M) липетанти Name ) 2028) 2028	Basis of Stren	gth Strength 75 mL in 100 mL
NICOHOL SOLUTION Product Information Product Type Route of Administration Active Ingredient/Active In ALCOHOL (UNII: 3K9958V90M)	HUMAN OTC DRUG TOPICAL MOiE TOPICAL MOIE	Basis of Stren	gth Strength 75 mL in 100 mL

P	Packaging						
#	Item Code		Package Description	Marketing Start Date	Marketing End Date		
1	NDC:73885-001-01	100	mL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
2	NDC:73885-001-02	40 m	nL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
3	NDC:73885-001-03	59 m	nL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
4	NDC:73885-001-04	250	mL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
5	NDC:73885-001-05	500	mL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
6	NDC:73885-001- 06	1000	0 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
7	NDC:73885-001-07	5000	0 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/26/2020			
Marketing Information							
Marketing Category		ry	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final		raph not final part333A		04/26/2020			

Labeler - Boya Biotechnology Company Limited (403701144)

# Establishment

Name	Address	ID/FEI	<b>Business Operations</b>
Boya Biotechnology Company Limited		403701144	manufacture(73885-001)

Revised: 4/2020

Boya Biotechnology Company Limited