### CLEAN CARE PLUS HAND SANITIZER- alcohol gel AT Bio Pharm 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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### **Drug Facts**

ethanol

Water, Glycerin, butylene glycol, carbomer, triethanolamine, rosmarinus officinalis (rosemary) leaf extract, paeonia suffruticosa bark extract, aloe barbadensis leaf extract, calendula officianalis flower extract

Antiseptic

instant healthcare personnel hand antiseptic to reduce bacteria that potentially can cause disease

instant hand antiseptic to decrease bacteria on the skin recommended for repeated use

hand sanitizer to help reduce bacteria on the skin

KEEP OUT OF REACH OF THE CHILDREN

Apply to clean, dry hands. Apply sufficient amount to thoroughly wet all surfaces of hands and fingers. Rub onto hands until dry.

Supervise children in the use of this product.

Dispense appropriate amount on your palm and thoroughly spread on both hands and rub into the skin until dry

For external use only.

Flammable, keep away from fire or flame.

When using this product keep out of eyes. If contact with eyes occurs, rinse promptly and thoroughly with water.

Stop use and ask a doctor if significant irritation or sensitization develops.

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

for external use only



## **CLEAN CARE PLUS HAND SANITIZER**

alcohol gel

Product Information									
Product Type			HUMAN OTC DRUG	Item Code (Source)		NDC:73932-0006			
Route of Administration			TOPICAL						
Active Ingredient/Active Moiety									
Ingredient Name Basis of Stren							Strength		
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M) ALCOHOL							70 g in 100 mL		
Inactive Ingredients									
Ingredient Name							Strength		
GLYCE	<b>RIN</b> (UNII: PDO	C6A3C0OX)							
TROLA									
WATER	(UNII: 059QF								
BUTYLENE GLYCOL (UNII: 3XUS85K0RA)									
Packaging									
# It	em Code		Package Description	Ν	larketing Start Dat	e Ma	arketing End Date		
1 NDC:	73932-0006-1	500 mL in 1 BO	TTLE; Type 0: Not a Combinatio	on Product 05	5/26/2020				
Marketing Information									

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	05/26/2020	

# Labeler - AT Bio Pharm Co Ltd (695742996)

# Registrant - AT Bio Pharm Co Ltd (695742996)

Establishment									
Name	Address	ID/FEI	<b>Business Operations</b>						
AT Bio Pharm Co Ltd		695742996	manufacture(73932-0006)						

Revised: 5/2020

AT Bio Pharm Co Ltd