ETHYL ALCOHOL- ethyl alcohol gel Marine Essence Biosciences Corporation of USA

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.

Ethyl Alcohol Hand Sanitizer

Ethanol alcohol 80% v/v

.....Antiseptic

Hand sanitizer to help reduce bacteria that potentially can cause disease.

glycerin, hydrogen peroxide, carbomer 940, triethanolamine, purified water USP

Topical - Administration to a particular spot on the outer surface of the body. The E2B term TRANSMAMMARY is a subset of the term TOPICAL

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

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

In children less than 2 months of age

On open skin wounds

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.

Place enough product on hands to cover all surfaces. Rub hands together until dry.

Supervise children under 6 years of age when using this product to avoid swallowing.

Other Information

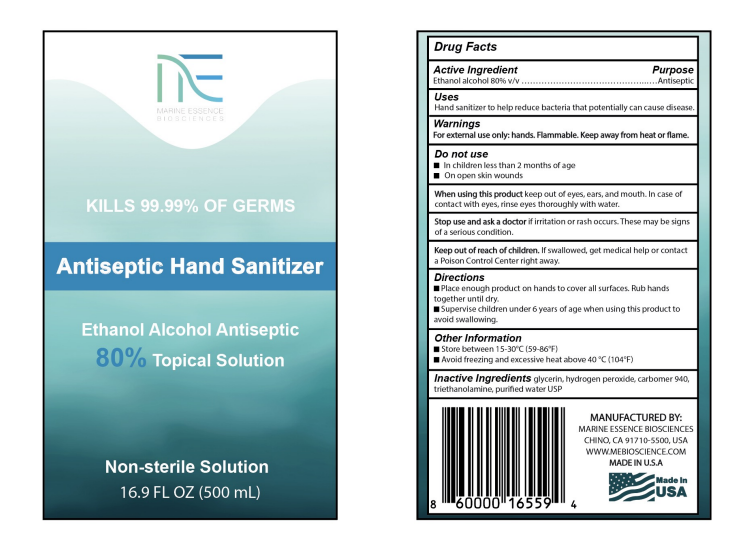
Store between 15-30C (59-86F)

Avoid freezing and excessive heat above 40C (104F)

Gel form

Warnings for external use only-hands

To help reduce bacteria on the hands that can potentially cause disease.



ETHYL ALCOHOL

ethyl alcohol gel

Product Information						
Product T ype	HUMAN OTC DRUG Iter			NI	NDC:73764-002	
Route of Administration	TOPICAL					
A .1 T 31 ./A .1 3.F	• .					
Active Ingredient/Active Mo	iety					
Ingredient Name			Basis of Strength		Strength	
ALCOHOL (UNII: 3K9958V90M) (AL	ALCOHOL		80 mL in 100 mL			
Inactive Ingradiants						
Inactive Ingredients						
Ingredient Name				Strength		
HYDROGEN PEROXIDE (UNII: BBX)	60AN9V)					
	60AN9V)					
HYDRO GEN PERO XIDE (UNII: BBX) WATER (UNII: 059QF0KO0R) GLYCERIN (UNII: PDC6A3C0OX)	60 AN9 V)					

Packaging										
# Item Code	Package Description				ting Date	Marketing End Date				
1 NDC:73764- 002-01		1 BOTTLE, PUMP; Type 9: Other Type of Part 3 Con /Device/Biological Product)	03/23/2020							
2 NDC:73764- 002-02	236 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product				0					
3 NDC:73764- 002-03	177 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product				0					
4 NDC:73764- 002-04	473 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product				03/23/2020					
5 NDC:73764- 002-05	250 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product				06/01/2020					
Marketing Information										
Marketing C	ategory	Application Number or Monograph Citation	n Marketing S	tart Date	Marke	ting End Date				
OTC monograph		part333A	03/23/2020		At	Ling Lind D				

Labeler - Marine Essence Biosciences Corporation of USA (116587566)

Establishment								
Name	Address	ID/FEI	Business Operations					
Marine Essence Biosciences Corporation of USA		116587566	manufacture(73764-002), pack(73764-002), label(73764- 002)					

Revised: 6/2020

Marine Essence Biosciences Corporation of USA