

**LAMWAVE ANTIMICROBIAL HAND SANITIZERS GEL- bacteriostatic gel gel**  
**Guangzhou Youmei Biotechnology Co., Ltd.**

*Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, [click here](#).*

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Store below 110°F(43°C). May discolor certain fabrics or surfaces.

ethanol

pure water

Carbomer

trimethylpentylenediol / adipic acid / glycerine crosslinked aggregate

triethylamine

Disinfection

Sterilization

Put enough product on your palm to cover hands and rub hands together briskly until dry.

allantoin

Fermentation products of bacillus

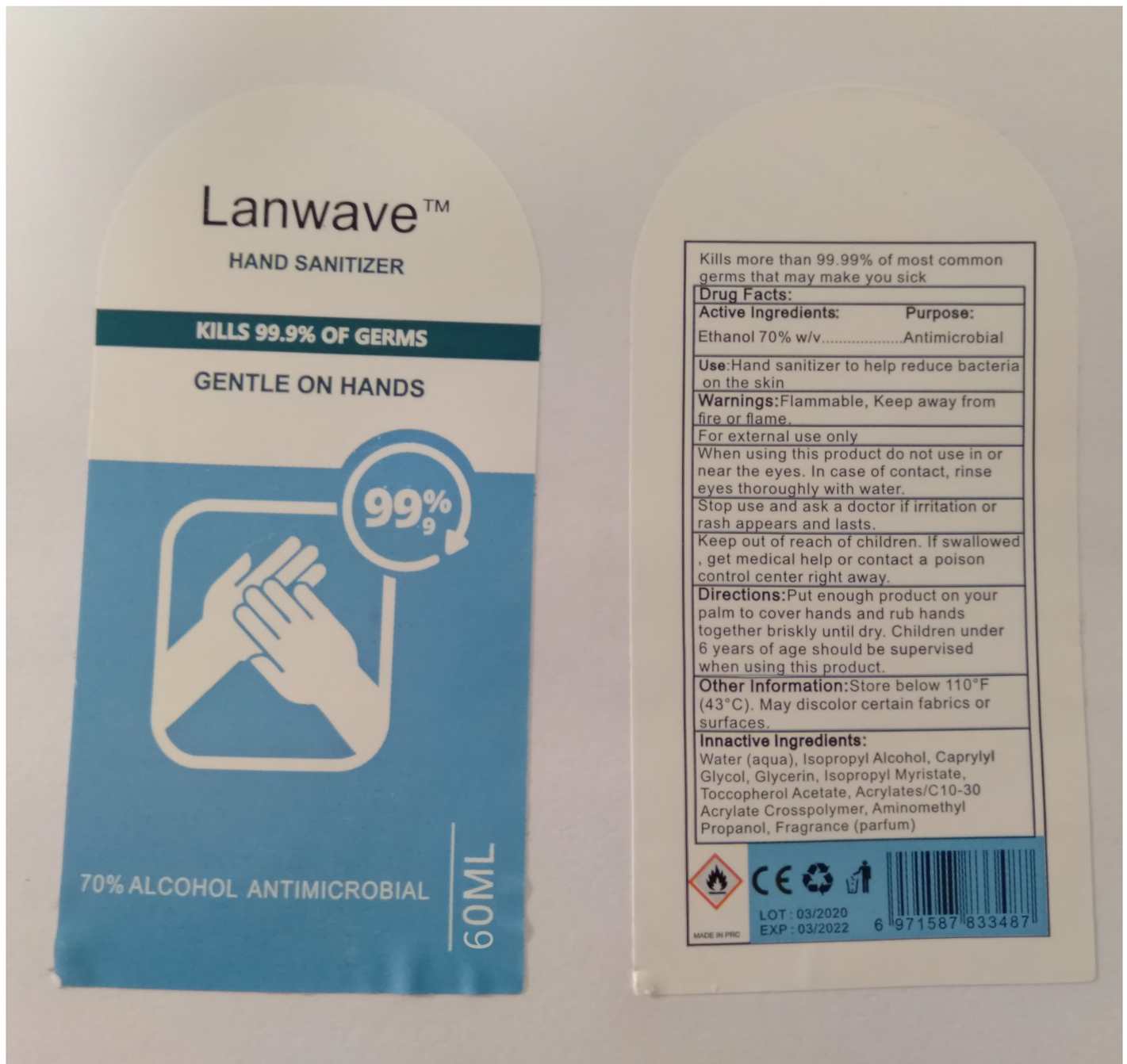
keep out of children

1.Flammable, Keep away from fire or flame

2.For external use only When using this product do not use in or near the eyes. In case of contact, rinse eyes thoroughly with water.

3.Stop use and ask a doctor if irritation or rash appears and lasts.

4.Keep out of reach of children. If swallowed get medical help or contact a poison control center right away.



**LAMWAVE ANTIMICROBIAL HAND SANITIZERS GEL**  
 bacteriostatic gel gel

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:41878-003
Route of Administration	EXTRACORPOREAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
BACILLUS AMYLOLIQUEFACIENS (UNII: 1M5Y26M37U) (BACILLUS AMYLOLIQUEFACIENS - UNII:1M5Y26M37U)	BACILLUS AMYLOLIQUEFACIENS	0.15 mg in 300 mL
ALLANTOIN (UNII: 344S277C07) (ALLANTOIN - UNII:344S277C07)	ALLANTOIN	0.15 mg

ALLANTOIN (UNII: 344327700Z) (ALLANTOIN - UNII:344327700Z)

ALLANTOIN

in 300 mL

**Inactive Ingredients**

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
ALCOHOL (UNII: 3K9958V90M)	
TRIMETHYL-1,3-PENTANEDIOL (UNII: WT1X081P0L)	
TRIETHYLAMINE (UNII: VOU728O6AY)	
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)	

**Product Characteristics**

Color		Score	
Shape	BULLET	Size	
Flavor		Imprint Code	
Contains			

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:41878-003-01	300 mL in 1 BOTTLE; Type 0: Not a Combination Product	04/16/2020	

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		04/16/2020	

**Labeler** - Guangzhou Youmei Biotechnology Co., Ltd. (418782063)**Establishment**

Name	Address	ID/FEI	Business Operations
Guangzhou Youmei Biotechnology Co., Ltd.		418782063	manufacture(41878-003)

Revised: 4/2020

Guangzhou Youmei Biotechnology Co., Ltd.