MPP INSTANT HAND SANITIZER- isopropyl alcohol gel Biominerales Pharma

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.

MPP INSTANT HAND SANITIZER

DRUG FACTS

Active ingredient

Isopropyl alcohol 70% v/v

Purpose

Antiseptic

Inactive ingredients

Purified Water, Glycerin, Carbomer, Triethanolamine, Hydrogen Peroxide.

Use (s)

Hand sanitizer to help reduce bacteria that potentially can cause disease. For use when water and soap are not available.

Warnings

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

Do not use in children less than 2 months of age or on open skin wounds.

When using this product keep our 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 poison control center right away.

Directions

- 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

Distributed by: MPP Dist. LLC. sales@mppint.com

PRINCIPAL DISPLAY PANEL - 3.785 L Bottle Label

Alcohol Antiseptic 70% Gel MPP INSTANT HAND SANITIZER

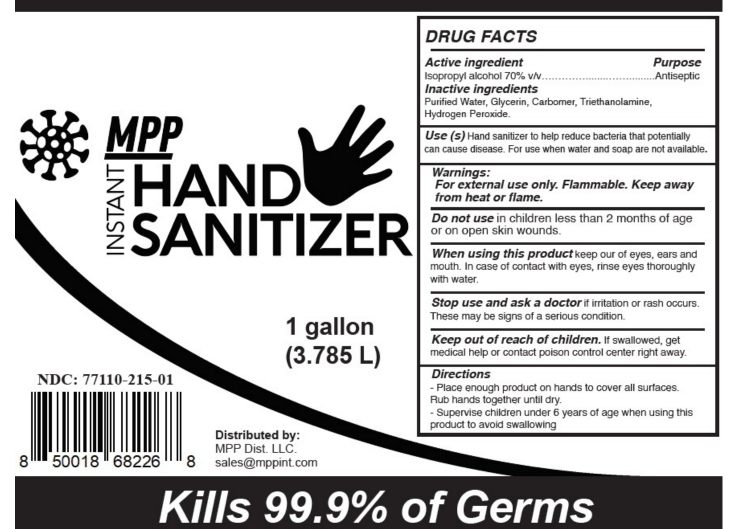
1 gallon (3.785 L)

NDC: 77110-215-01

Distributed by: MPP Dist. LLC. sales@mppint.com

Kills 99.9% of Germs

Alcohol Antiseptic 70% Gel



MPP INSTANT HAND SANITIZER

isopropyl alcohol gel

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:77110-215

Active Ingredient/Active Moiety								
		Ingredient Name		Basis of Stre	ngth	Strength		
ISOPROPYL ALCON UNII:ND2M416302)	HOL (UNII:	ND2M416302) (ISOPROPYL ALCOHOL -		ISOPROPYL ALCOHOL		70 mL in 100 mL		
Inactive Ingredients								
		Ingredient Name				Strength		
WATER (UNII: 059QF0KO0R)								
GLYCERIN (UNII: PDC6A3C0OX)								
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0 A5MM307FC)								
TROLAMINE (UNII: 903K93S3TK)								
HYDRO GEN PERO X	IDE (UNII: I	BBX060AN9V)						
Packaging	IDE (UNII: 1							
Packaging # Item Code		Package Description		ing Start Date	Mark	eting End Dat		
Packaging # Item Code 1 NDC:77110-215-01	3785 mL i	Package Description n 1 BOTTLE; Type 0: Not a Combination Produc	t 07/25/20	20	Mark	eting End Dat		
Backaging Item Code NDC:77110-215-01 NDC:77110-215-06	3785 mL ii 59 mL in 1	Package Description n 1 BOTTLE; Type 0: Not a Combination Produc BOTTLE; Type 0: Not a Combination Product	t 07/25/20 07/25/20	20 20	Mark	eting End Dat		
Participation Item Code 1 NDC:77110-215-01 2 NDC:77110-215-06 3 NDC:77110-215-04	3785 mL ii 59 mL in 1 236 mL in	Package Description n 1 BOTTLE; Type 0: Not a Combination Produc BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product	t 07/25/20 07/25/20 07/25/20	20 20 20	Mark	eting End Dat		
 NDC:77110-215-01 NDC:77110-215-06 NDC:77110-215-04 NDC:77110-215-03 	3785 mL i 59 mL in 1 236 mL in 472 mL in	Package Description n 1 BOTTLE; Type 0: Not a Combination Produc BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product	 07/25/20 07/25/20 07/25/20 07/25/20 	20 20 20 20 20	Mark	eting End Dat		
Partial State Item Code IDC:77110-215-01 NDC:77110-215-04 NDC:77110-215-04 NDC:77110-215-04 NDC:77110-215-04	3785 mL i 59 mL in 1 236 mL in 472 mL in	Package Description n 1 BOTTLE; Type 0: Not a Combination Produc BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product	t 07/25/20 07/25/20 07/25/20	20 20 20 20 20	Mark	eting End Dat		
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Partial State Item Code IDC:77110-215-01 NDC:77110-215-04 NDC:77110-215-04 NDC:77110-215-04 NDC:77110-215-04	3785 mL i 59 mL in 1 236 mL in 472 mL in	Package Description n 1 BOTTLE; Type 0: Not a Combination Produc BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product	 07/25/20 07/25/20 07/25/20 07/25/20 	20 20 20 20 20	Mark	eting End Dat		
Packaging Item Code 1 NDC:77110-215-01 2 NDC:77110-215-06 3 NDC:77110-215-04 4 NDC:77110-215-02 5 NDC:77110-215-02	3785 mL i 59 mL in 1 236 mL in 472 mL in 944 mL in	Package Description n 1 BOTTLE; Type 0: Not a Combination Product BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product 1 BOTTLE; Type 0: Not a Combination Product	 07/25/20 07/25/20 07/25/20 07/25/20 	20 20 20 20 20	Mark	eting End Dat		
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Labeler - Biominerales Pharma (117489663)

Establishment							
Name	Address	ID/FEI	Business Operations				
Biominerales Pharma		117489663	MANUFACTURE(77110-215)				

Revised: 7/2020

Biominerales Pharma