#### AIR AND WATER HAND SANITIZER - CHRISTMAS BERRY- ethyl alcohol gel Pearl World Inc.

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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## AIR&WATER Christmas Berry Hand Sanitizer

#### **Drug Facts**

## Active ingredient

Ethyl Alcohol 70%

## Purpose

Antiseptic

#### Uses

• To decrease bacteria on the skin that could cause disease. • When water, soap and towel are not available. • Recommended for repeated use.

#### Warnings

For external use only: hands.

## Flammable. Keep away from fire or flame.

**When using this product** • Keep out of eyes, ears, or mouth. • In case of eye contact, flush eyes thoroughly with water. • Avoid contact with broken skin. • Do not inhale or ingest.

**Stop use and ask a doctor if** • Redness or irritation develop. • Condition persists for more than 72 hours.

**Keep out of reach of children.**• Children should be supervised by an adult when using this product. • 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. • No rinsing required. • For children under 6, use only under adult supervision. • Not recommended for infants.

**Other information** • Store below 105°F (40°C). • May discolor certain fabrics. • Harmful to wood finishes & plastics.

**Inactive ingredients**: Water (Aqua), Triethanolamine, Carbomer, Aloe Barbadensis Leaf Extract, Fragrance, Glycerin, Propylene Glycol, Tocopheryl Acetate (Vitamin E). Main Contain: FD & C Red 40 (CI 16035), FD & C Yellow 6 (CI 15985), FD & C Violet 2 (CI 60725), FD & C Blue 1 (CI 42090), FD & C Yellow 5 (CI 19140), FD & C Red 33(CI 17200).

## 70% Alcohol

Manufactured for and distributed by Pearl world Inc, NY, NY 10019

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# Packaging





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BATCH: JM129275 MFG: 20200925 EXP: 20220924







# AIR AND WATER HAND SANITIZER - CHRISTMAS BERRY

ethyl alcohol gel

Product Information							
Product T ype	HUMAN OTC DRUG	Item Code (Source) NDC:6993		NDC:69933-403			
Route of Administration	TOPICAL						
Active Ingredient/Active Moiety							
Ingredient Name			Basis of Strengtl	h Strength			
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)			ALCOHOL	70 mL in 100 mL			
Inactive Ingredients							
	Ingredient Name			Strength			

WATER (UNII: 059C	F0KO0R)				
TROLAMINE (UNII:					
CARBOMER HOMO	POLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)				
ALOE VERA LEAF	(UNII: ZY8 1Z8 3H0 X)				
GLYCERIN (UNII: PI	DC6A3C0OX)				
PROPYLENE GLYC	OL (UNII: 6DC9Q167V3)				
.ALPHATOCOPH	EROL ACETATE (UNII: 9E8X80D2L0)				
FD&C RED NO. 40	(UNII: WZB9127XOA)				
FD&C YELLOW NO	<b>D.6</b> (UNII: H77VEI93A8)				
D&C VIOLET NO. 2	2 (UNII: 350 KA7O6 HK)				
FD&C BLUE NO.1	(UNII: H3R47K3TBD)				
FD&C YELLOW NO. 5 (UNII: 1753WB2F1M)					
D&C RED NO.33 (0	JNII: 9DBA0SBB0L)				
Packaging					
# Item Code	Package Description	Marketing Start Date	Marketing End Date		
1 NDC:69933-403- 50	500 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	09/17/2020			

# Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	09/17/2020	

09/17/2020

60 mL in 1 BOTTLE; Type 0: Not a Combination Product

# Labeler - Pearl World Inc. (043130142)

Revised: 9/2020

2 NDC:69933-403-60

Pearl World Inc.