PHARMACYS PRESCRIPTION HAND SANITIZER- alcohol spray American Consumer Products Corp

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.

Pharmacys Prescription Hand Sanitizer

Active Ingredients Active ingredients - Ethyl Alcohol 80%

Purpose Purpose - Antiseptic

Warnings Warnings - For external use only. Do not ingest or swallow. Flammable. Keep away from fire or flame. Do not apply around eyes. Do not use in ears & mouth.

Indications & Usage

When using this product, avoid contact with eyes. In case of contact, flush eyes with water.

Stop Use

Stop use and ask a doctor if redness or irritation develops and persists for more than 72 hours.

Keep out of reach of children

Keep out of reach of children. Do not use on children less than 2 months of age. Supervise use in children under 6 years of age to prevent accidental swallowing. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

Directions - Apply directly onto hands, rub into skin until dry. Apply directly onto surfaces: knobs, handles, steering wheels, light swithches, grocery carts, countertops, etc and let air dry.

Other Information

Other information - store at 20° C (68° to 77° F).

-may discolor fabrics.

Inactive Ingredients

INACTIVE INGREDIENTS: Purified Water (Aqua), Glycerin, Hydrogen Peroxide

Distributed By: Distributed By: American Consumer Products Corp Vernon, CA 90058

Pharmacys Prescription Hand Sanitizer



PHARMACYS PRESCRIPTION HAND SANITIZER

alcohol spray

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:72197-023

Active Ingredient/Active Moiety					
	Ingredient Name Basis of Strengt			Basis of Strength	Strength
A	ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M) ALCOHOL			ALCOHOL	80 mL in 100 mL
Inactive Ingredients					
	Ingredient Name				Strength
WATER (UNII: 059QF0KO0R)					
GLYCERIN (UNII: PDC6A3C0OX)					
HYDROGEN PEROXIDE (UNII: BBX060AN9V)					
Packaging					
	ackaging				
#	Item Code		Package Description	Marketing Start Date	Marketing End Date
# 1		3785. Pro du	41 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination	_	U
	Item Code NDC:72197-023-		41 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination	Date	U
1	Item Code NDC:72197-023-	Pro du	41 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination ct	Date	U
1	Item Code NDC:72197-023- 99	Produ 1for	41 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination ct	Date	U
1	Item Code NDC:72197-023- 99 /Iarketing In	Produ 1fori gory	41 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination ct nation Application Number or Monograph Citation	Date 04/17/2020	Date

Labeler - American Consumer Products Corp (081101181)

Revised: 4/2020

American Consumer Products Corp