LICENSED HAND SANITIZER CRAYOLA 01- alcohol spray Shenzhen Lantern Scicence Co.,Ltd.

88112 Licensed Hand Sanitizer Spray Crayola

Drug Facts

Active Ingredient

Active ingredient Purpose Ethyl Alcohol 70% v/v Antiseptic

Uses

Decreases bacieria on skin

Warning

For external use only

Do not use

If you are allergic to any of the ingredients

When using this product

Do not get into eyes. If contact occurs, rinse thoroughly with water. Do not use in or near eyes, In case of eye contact, flush eyes thoroughly with water. Discontinue if skin becomes irritated and ask for a doctor.

Stop use and ask a doctor

If irritation or rash develops and continues for more than 72 hours.

keep out of reach of children

If swallowed, get medical help or contact a Poison Control Center immediately.

Directions

Adults and children 6 years and older Spray onto hands and rub together unil dry. Recommended for repeat use. Children under 6 years of age should be supervised when using this product.

Inactive ingredients

Water, Fragrance, Glycerin, ACRYLATES/C10-30 ALKYL ACRYLATE CROSSPOLYMER, Aminomethyl Propanol, Tocopheryl Acetate, Aloe Barbadensis Gel, Denatonium Benzoate.

other Information

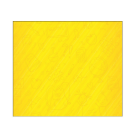
Storebetween15-30C(59-86F) Avoid freezing and excessive heat above 40C (104F)

Spray 3-4 times each time. Used in the skin area of the hands.

packing

packing







LICENSED HAND SANITIZER CRAYOLA 01

alcohol spray

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:54860-434
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	70 mL in 100 mL

Inactive Ingredients			
Ingredient Name	Strength		
AMINOMETHYLPROPANOL (UNII: LU49E6626Q)			
ALOE VERA LEAF (UNII: ZY81Z83H0X)			
GLYCERIN (UNII: PDC6A3C0OX)			
WATER (UNII: 059QF0KO0R)			

ACRYLATES/C10-30 ALKYL ACRYLATE CROSSPOLYMER (60000 MPA.S) (UNII: 8Z5ZAL5H3V)			
.ALPHATOCOPHEROL ACETATE (UNII: 9E8X80D2L0)			
DENATONIUM BENZOATE (UNII: 4YK5Z54AT2)			

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:54860- 434-01	3 in 1 BOX	02/14/2025		
1		28 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M003	02/14/2025		

Labeler - Shenzhen Lantern Scicence Co.,Ltd. (421222423)

Establishment				
Name	Address	ID/FEI	Business Operations	
Shenzhen Lantern Science Co.,Ltd.		421222423	manufacture(54860-434)	

Revised: 2/2025 Shenzhen Lantern Scicence Co.,Ltd.