JIKIMI GERM KILLER-SPRAY- titanium dioxide, hypochlorous acid liquid MY Corp.,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.

MY Corp - JIKIMI GERM KILLER

titanium dioxide, hypochlorous acid

Coumarin, Linalool, Quercetin, Water

- Do not dilute spray
- Use at strength for maximum effectiveness
- Spray directly, from a distance of about 8 to 10 inches. For equipment or products that are difficult to spray, apply liquid to clean gauze or cotton swab.

keep out of reach of the children

 Wipe with gauze or cotton swab and let air dry. Spray clothes, equipment, and indoor objects for sterilization and deodorization as required.

For external use only.

When using this product keep out of eyes, ears, and mouth. In case of contact with eyes, rinse thoroughly with water.

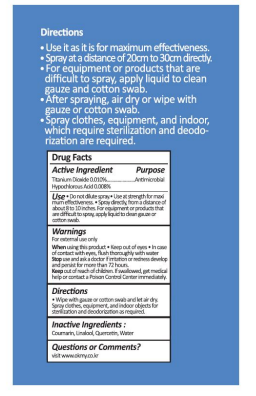
Stop use and ask a doctor if irritation or rash occurs develop and persist for more than 72 hours.

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

for external use only







JIKIMI GERM KILLER-SPRAY

titanium dioxide, hypochlorous acid liquid

Prod	luct	Info	rma	tion
FIUU	uct	11110	шша	uvii

Product Type HUMA	N OTC DRUG Item	em Code (Source)	NDC:71544-0011
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Route of Administration TOPICAL

Active Ingredient/Active Moiety Ingredient Name Basis of Strength TITANIUM DIO XIDE (UNII: 15FIX9 V2JP) (TITANIUM DIO XIDE - UNII:15FIX9 V2JP) HYPOCHLOROUS ACID (UNII: 712K4CDC10) (HYPOCHLOROUS ACID - HYPOCHLOROUS ACID - HYPOCHLOROUS ACID in 60 mL

Inactive Ingredients			
Ingredient Name	Strength		
COUMARIN (UNII: A4VZ22K1WT)			
LINALOOL , (+)- (UNII: F4VNO44C09)			
QUERCETIN (UNII: 9 IKM0 I5T1E)			
WATER (UNII: 059QF0KO0R)			

ı	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:71544-0011- 1	60 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	07/15/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		07/15/2020	

Labeler - MY Corp.,Ltd (688202781)

Registrant - MY Corp.,Ltd (688202781)

Establishment			
Name	Address	ID/FEI	Business Operations
MY Corp.,Ltd		688202781	manufacture(71544-0011), label(71544-0011), pack(71544-0011)

Revised: 7/2020 MY Corp.,Ltd