GYUN E ZERO STERILIZATION TISSUE- alcohol liquid Sunmedical

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.

Active Ingredients

Active ingredients: ISOPROPYL ALCOHOL 70.0%(w/w)

INACTIVE INGREDIENT

Inactive ingredients:

Water, EDTA-2Na, Sodium Benzoate, Methyl Propanediol, Caprylyl Glycol, Ethylhexylglycerin, Octyldodeceth-16

Purpose

Purpose: SANITIZER

Warnings

Warnings:

First Aid

- 1. Wash with clean water when it gets into your eyes, and consult your doctor if you have any problems.
- 2. Take emergency measures to eat or swallow contents and Consult a doctor immediately

Precautions for Use

- 1.Please follow the indicated usage and standard usage.
- 2.Be careful not to let the contents touch your eyes.
- 3.Do not inhale or drink.
- 4. Ventilate enough when using in an enclosed space.

KEEP OUT OF REACH OF CHILDREN

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Uses

Uses:

- It is a sterilized tissue that can be used for various purposes in hospitals, kitchen utensils, and students' hygiene.
- Isopropyl alcohol is a typical raw material for disinfectants.

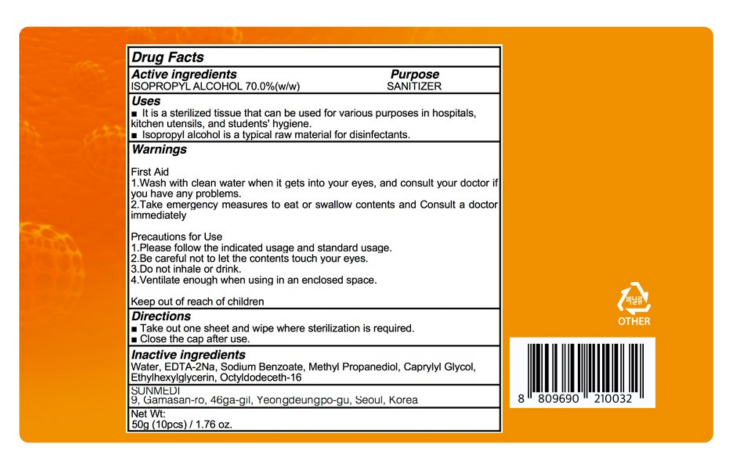
Directions

Directions:

- Take out one sheet and wipe where sterilization is required.
- Close the cap after use.

PACKAGE LABEL - Gyun-e zero sterilization tissue (10pcs, 50g)







각종 유해 세균 살균 소독

Drug Facts

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SUNMELDI 9, Gamasan-ro, 46ga-gil, Yeongdeungpo-gu, Seoul, Korea

Net Wt: 580g(75pcs) / 20.46 oz



GYUN E ZERO STERILIZATION TISSUE

alcohol liquid

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:75578-020
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength

ISOPROPYL ALCOHOL (UNII: ND2M416302) (ISOPROPYL ALCOHOL - UNII:ND2M416302) | ISOPROPYL ALCOHOL | 70 g in 100 g

Inactive Ingredients			
Ingredient Name	Strength		
Water (UNII: 059QF0KO0R)			
Sodium Benzoate (UNII: OJ245FE5EU)			
MethylPropanediol (UNII: N8F53B3R4R)			
CAPRYLYL GLYCOL (UNII: 00 YIU5438 U)			
Ethylhexylglycerin (UNII: 147D247K3P)			
Octyldodeceth-16 (UNII: RJV574G0DE)			

1	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:75578-020-01	50 g in 1 CONTAINER; Type 0: Not a Combination Product	04/01/2020		
2	NDC:75578-020- 02	580 g in 1 CONTAINER; Type 0: Not a Combination Product	04/01/2020		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	04/01/2020	

Labeler - Sunmedical (688954380)

Registrant - Sunmedical (688954380)

Establishment			
Name	Address	ID/FEI	Business Operations
Sunmedical		688954380	manufacture(75578-020)

Revised: 4/2020 Sunmedical