

SOUTH MOON WART REMOVER- glycerin liquid
Shantou South Moon Biotechnology Co., Ltd.

CENTELLA ASIATICA TRITERPENOIDS□HAMAMELIS VIRGINIANA LEAF WATER□ALOE
BARBADENSIS LEAF

1. Clean skin thoroughly and dry first
2. Take an appropriate amount of this product and apply it on the skin where warts grow.
1. Easily remove skin tags, painlessly remove skin warts and moles, and quickly restore smooth skin.
2. The formula is mild and suitable for all types of skin, making it safer to use.
3. Use tea tree oil and saffron plant ingredients to protect and care for the skin.

Please keep out of reach of children. Do not swallow. Please clean your hands before use to ensure the best results from the product. Discontinue use if signs of irritation or rash occur. Store in a cool and dry place.

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AQUA□GLYCERIN

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纸盒

规格尺寸：2.5X2.5X6.5CM



SOUTH MOON WART REMOVER

glycerin liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:84983-013
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CENTELLA ASIATICA TRITERPENOIDS (UNII: 4YS74Q4G4J) (CENTELLA	CENTELLA ASIATICA	0.01 mg

ASIATICA TRITERPENOIDS - UNII:4YS74Q4G4J)	TRITERPENOIDS	in 10 mg
HAMAMELIS VIRGINIANA (WITCH HAZEL) LEAF WATER (UNII: 8FP93ED6H2) (HAMAMELIS VIRGINIANA LEAF WATER - UNII:8FP93ED6H2)	HAMAMELIS VIRGINIANA (WITCH HAZEL) LEAF WATER	0.002 mg in 10 mg
ALOE BARBADENSIS LEAF (UNII: ZY81Z83H0X) (ALOE BARBADENSIS LEAF - UNII:ZY81Z83H0X)	ALOE BARBADENSIS LEAF	0.002 mg in 10 mg

Inactive Ingredients

Ingredient Name	Strength
AQUA (UNII: 059QF0K00R)	9.486 mg in 10 mg
GLYCERIN (UNII: PDC6A3C0OX)	0.5 mg in 10 mg

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:84983-013-01	10 mg in 1 BOTTLE; Type 0: Not a Combination Product	09/17/2025	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M016	09/17/2025	

Labeler - Shantou South Moon Biotechnology Co., Ltd. (457126192)

Registrant - Shantou South Moon Biotechnology Co., Ltd. (457126192)

Establishment

Name	Address	ID/FEI	Business Operations
Shantou South Moon Biotechnology Co., Ltd.		457126192	manufacture(84983-013)

Revised: 11/2025

Shantou South Moon Biotechnology Co., Ltd.