

XIMONTH SUNFLOWER XANTHIUM SEED OIL- glycerin oil
Guangdong Ouhoe Biotechnology Co., Ltd.

ACTIVE INGREDIENTS

XANTHIUM SIBIRICUM ROOT, HELIANTHUS ANNUUS (SUNFLOWER) SEED OIL, ANGELICA
DAHURICA ROOT EXTRACT

purpose

Clear the nasal cavity and relieve nasal discomfort; remove foreign objects blocking the nasal cavity and make the nasal cavity clean and unobstructed; solve acute and chronic rhinitis.

use

Apply an appropriate amount of this product to the affected area and wait for it to be absorbed.

WARNINGS

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.

Do not use

Discontinue use if signs of irritation or rash occur.

Stop use

Discontinue use if signs of irritation or rash occur.

Please keep out of reach of children. Do not swallow.

Store in a cool and dry place.

GLYCERIN, LANOLIN

纸盒（多页）
规格尺寸：长5X宽2.4X高7.8X1X5CM



XIMONTH SUNFLOWER XANTHIUM SEED OIL

glycerin oil

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:84744-052
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
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XANTHIUM SIBIRICUM ROOT (UNII: 8555ORT735) (XANTHIUM SIBIRICUM ROOT - UNII:8555ORT735)	XANTHIUM SIBIRICUM ROOT	3 mg in 10 mg
SUNFLOWER OIL (UNII: 3W1JG795YI) (SUNFLOWER OIL - UNII:3W1JG795YI)	SUNFLOWER OIL	1 mg in 10 mg
ANGELICA DAHURICA ROOT (UNII: 1V63N2S972) (ANGELICA DAHURICA ROOT - UNII:1V63N2S972)	ANGELICA DAHURICA ROOT	1.5 mg in 10 mg

Inactive Ingredients	
Ingredient Name	Strength
LANOLIN (UNII: 7EV65EAW6H)	3 mg in 10 mg
GLYCERIN (UNII: PDC6A3C0OX)	1.5 mg in 10 mg

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:84744-052-01	10 mg in 1 BOX; Type 0: Not a Combination Product	10/22/2024	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M016	10/22/2024	

Labeler - Guangdong Ouhoe Biotechnology Co., Ltd. (843651433)

Registrant - Guangdong Ouhoe Biotechnology Co., Ltd. (843651433)

Establishment			
Name	Address	ID/FEI	Business Operations
Guangdong Ouhoe Biotechnology Co., Ltd.		843651433	manufacture(84744-052)

Revised: 10/2024

Guangdong Ouhoe Biotechnology Co., Ltd.