NEKVNRO WART REMOVER PATCH- wart remover patch patch Shenzhen Xingqi Technology Co., Ltd.

ACTIVE INGREDIENT

TALC 1%

PURPOSE

wart removal

USES

- 1, Clean and wipe the skin around the wart.
- 2, Remove the wart removal tablet, Apply it to the wart, Cover it completely
- 3, 1 capsule can last for 8-12hours, Recommended 2-3 times a day

warning

- 1, external use, Avoid contact with eyes and mouth.
- 2,If irritation or abnormality occurs, stop using and wash with water
- 3, Please keep this product out of the reach of children

Dosage and administration

For external use only.

Do not use

- 1, external use, Avoid contact with eyes and mouth.
- 2, If irritation or abnormality occurs, stop using and wash with water
- 3, Please keep this product out of the reach of children

When using section

- 1, external use, Avoid contact with eyes and mouth.
- 2, If irritation or abnormality occurs, stop using and wash with water
- 3, Please keep this product out of the reach of children

STOP USE

- 1, external use, Avoid contact with eyes and mouth.
- 2, If irritation or abnormality occurs, stop using and wash with water
- 3, Please keep this product out of the reach of children

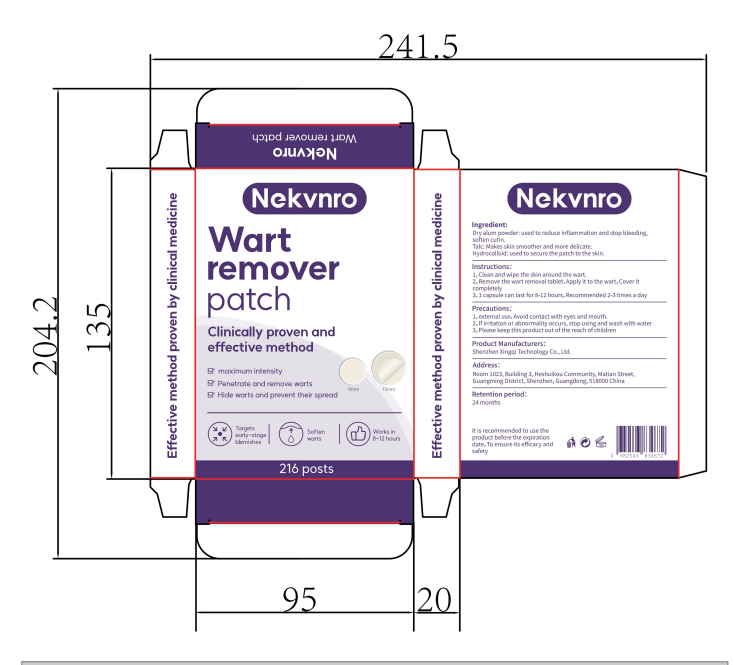
KEEP OUT OF REACH OF CHILDREN

Please keep this product out of the reach of children

INACTIVE INGREDIENT

POTASSIUM ALUM, SODIUM ALGINATE

PRINCIPAL DISPLAY PANEL



NEKVNRO WART REMOVER PATCH wart remover patch patch Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:84613-052

Route of Administration

TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
TALC (UNII: 7SEV7J4R1U) (TALC - UNII:7SEV7J4R1U)	TALC	1 g in 100 g

Inactive Ingredients

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Ingredient Name	Strength			
POTASSIUM ALUM (UNII: 1L24V9R23S)	1 g in 100 g			
SODIUM ALGINATE (UNII: C269C4G2ZQ)				

Packaging

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	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:84613-052-	30 g in 1 BOX; Type 0: Not a Combination Product	08/08/2024	

Marketing Information

Marketing information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M028	08/08/2024		

Labeler - Shenzhen Xingqi Technology Co., Ltd. (974493783)

Establishment				
Name	Address	ID/FEI	Business Operations	
Shenzhen Xingqi Technology Co., Ltd.		974493783	manufacture(84613-052)	

Revised: 8/2024 Shenzhen Xingqi Technology Co., Ltd.