RHUS TOX 6X/12X- rhus toxicodendron liquid MedCara Pharmaceuticals, LLC

Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.

RHUS TOX 6x/12x

ACTIVE INGREDIENTS equal amounts of: Rhus Toxicodendron 6X, 12X HPUS*

USES: Temporarily relieves irritation from red, swollen, itchy skin.**

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DIRECTIONS: Adults and children above 12 years: 3 droppers full 1 time per week for 3 weeks, then 1 time per month thereafter, hold solution in mouth for 30 seconds then swallow. Can be taken at any time of year but ideal to begin administering during Poison Ivy dormant phase (winter) often starting in February in the Midwest.

KEEP OUT OF THE REACH OF CHILDREN. In case of overdose (or accidental ingestion) get medical help or contact a Poison Control Center right away.

WARNINGS:

- Consult a physician for use in children under 12 years of age.
- IF PREGNANT OR BREAST-FEEDING, ask a health care professional before use.
- **KEEP OUT OF THE REACH OF CHILDREN.** In case of overdose (or accidental ingestion) get medical help or contact a Poison Control Center right away.
- Do not use if **TAMPER EVIDENT** seal is broken or missing.

INACTIVE INGREDIENTS: Ethyl Alcohol USP, Purified Water.

QUESTIONS & COMMENTS?

MedCara / Conrad, IA 50621 / 855-409-5496

www.outdoorjoes.com

*The letters "HPUS" indicate the components in the products are officially monographed in the Homeopathic Pharmacopeia of the United States.

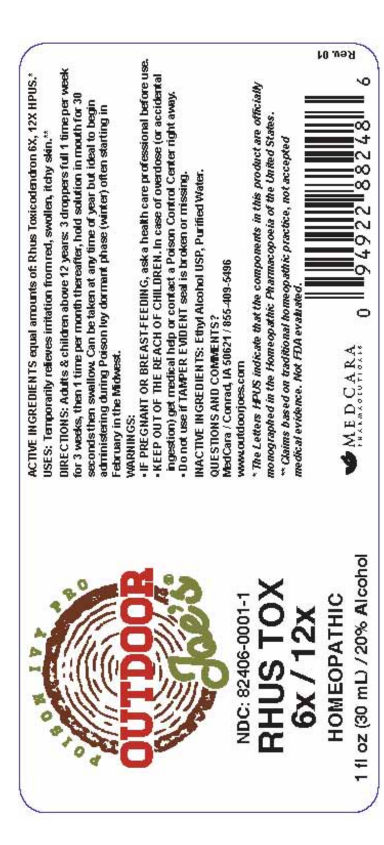
**Claims based on traditional homeopathic practice, not accepted medical evidence. Not FDA evaluated.

NDC: 82406-0001-1

RHUS TOX 6X / 12X

Homeopathic

1 fl oz (30 mL) / 20% Alcohol



RHUS TOX 6X/12X

rhus toxicodendron liquid

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:82406-0001	
Route of Administration	ORAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
TOXICODENDRON PUBESCENS LEAF (UNII: 610182RP7A) (TOXICODENDRON PUBESCENS LEAF - LINII: 610182RP7A)	TOXICODENDRON PUBESCENS LEAF	12 [hp_X]	

Inactive Ingredients			
Ingredient Name	Strength		
ALCOHOL (UNII: 3K9958V90M)			
WATER (UNII: 059QF0KO0R)			

l	Packaging				
	#	# Item Code Package Description		Marketing Start Date	Marketing End Date
	1	NDC:82406- 0001-1	30 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product	02/17/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		02/17/2022	

Labeler - MedCara Pharmaceuticals, LLC (966701166)

Establishment				
Name	Address	ID/FEI	Business Operations	
OHM Pharma, Inc		030572478	manufacture(82406-0001)	

Revised: 3/2022 MedCara Pharmaceuticals, LLC