KREOSOTUM- kreosotum liquid Rxhomeo Private Limited d.b.a. Rxhomeo, Inc

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.

ACTIVE INGREDIENT

KREOSOTUM HPUS 3X and higher

USES

USES: Temporary Relief - Discomfort From Toothache*

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

INDICATIONS

Condition listed above or as directed by the physician

DOSAGE

Adults: Mix 4-6 drops in 1/2 cup of water, take orally three times daily or as directed by a physician. Children (2 years and older): Take half the adult dose.

WARNINGS

This product is to be used for self-limiting conditions

If symptoms do not improve in 4 days, or worsen, discontinue use and seek assistance of health professional

As with any drug, if you are pregnant, or nursing a baby, seek professional advice before taking this product

Keep this and all medication out of reach of children

Do not use if capseal is broken or missing.

Close the cap tightly after use.

INACTIVE INGREDIENTS

Active: As Above; Inactive: ENA 50% v/v and Purified Water q.s.

STORAGE

QUESTIONS OR COMMENTS

www.Rxhomeo.com | 1.888.2796642 | info@rxhomeo.com - Rxhomeo, Inc 2940 Eisenhower St., Suite 100, Carrollton, TX 75007 USA

KREOSOTUM 3X

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USES: Temporary Relief - Discomfort From Toothache*

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NDC 15631-2807 B. No. XXXXXXX MFG XX/XX EXP XX/XX Contents XX ml Drops

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rxhomeo.

Distributed by Rxhomeo, Inc 2940 Eisenhower St., Suite 100, Carrollton, TX 75007 USA

FDA Est. # XXXXXXXXXXX | Made in XXXXXXX | Rxhomeo.com | info@rxhomeo.com

HOMOEOPATHIC MEDICINE



KREOSOTUM kreosotum liquid Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:15631-2807 Route of Administration ORAL

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
WOOD CREOSOTE (UNII: 3JYG22FD73) (WOOD CREOSOTE - UNII:3JYG22FD73)	WOOD CREOSOTE	3 [hp_X] in 1 mL	

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

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:15631- 2807-0	10 mL in 1 BOTTLE; Type 0: Not a Combination Product	05/23/2024	
2	NDC:15631- 2807-1	30 mL in 1 BOTTLE; Type 0: Not a Combination Product	05/23/2024	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		05/23/2024	

Labeler - Rxhomeo Private Limited d.b.a. Rxhomeo, Inc (650833994)

Establishment				
Name	Address	ID/FEI	Business Operations	
Rxhomeo Private Limited d.b.a. Rxhomeo, Inc		650833994	manufacture(15631-2807) , label(15631-2807)	

Revised: 6/2024 Rxhomeo Private Limited d.b.a. Rxhomeo, Inc