FORMICA RUFA- formica rufa pellet 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

FORMICA RUFA HPUS 1X and higher

USES

Arthrtitis, Gout

INDICATIONS

Condition listed above or as directed by the physician

DOSAGE

Adults- Take 4 or 6 Pellets by mouth, three times daily or as suggested by physician. Children 2 years and older- take 1/2 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

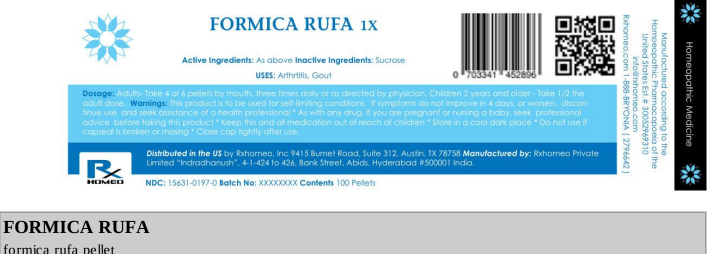
Sucrose

STORAGE

Store in a cool dark place

QUESTIONS OR COMMENTS

www.Rxhomeo.com | 1.888.2796642 | info@rxhomeo.com Rxhomeo, Inc 9415 Burnet Road, Suite 312, Austin, TX 78758



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P	roduct Informa	tion									
Product T ype			HUMAN OTC DRUG	Item Code (Source))	NDC:15631-0197				
R	oute of Administra	ation	ORAL								
A	Active Ingredient/Active Moiety										
							trength	Strength			
FORMICA RUFA (UNII: 55H0 W83JO5) (FORMICA RUFA - UNII:55H0 W83JO5)						FORMICA RUFA 1 [hp_X]		1 [hp_X]			
Inactive Ingredients											
Ingredient Name							Strength				
SUCROSE (UNII: C151H8 M554)											
Packaging											
#	Item Code		Package Description		Marketing	Start Date	Marketii	ng End Date			
1	NDC:15631-0197-0	100 in 1 PACKA	GE; Type 0: Not a Combination Pro		0 1/0 1/20 18			-			
2	NDC:15631-0197-1	200 in 1 PACKA	GE; Type 0: Not a Combination Pro	duct	0 1/0 1/20 18						
3	NDC:15631-0197-2	400 in 1 PACKA	GE; Type 0: Not a Combination Pro	duct	0 1/0 1/20 18						
4	NDC:15631-0197-3	750 in 1 PACKA	GE; Type 0: Not a Combination Pro	duct	0 1/0 1/20 18						
5			SE, Type 0. Not a Combination 110								
	NDC:15631-0197-4		AGE; Type 0: Not a Combination Pr		0 1/0 1/20 18						
6		2500 in 1 PACKA		oduct	0 1/0 1/20 18 0 1/0 1/20 18						
6		2500 in 1 PACKA	AGE; Type 0: Not a Combination Pr	oduct							
6		2500 in 1 PACKA	AGE; Type 0: Not a Combination Pr	oduct							
		2500 in 1 PACKA 12500 in 1 PACK	AGE; Type 0: Not a Combination Pr	oduct							
N	NDC:15631-0197-5	2500 in 1 PACKA 12500 in 1 PACK	AGE; Type 0: Not a Combination Pr	oduct Product	0 1/0 1/20 18	s Start Date	Marketi	ng End Date			
N	NDC:15631-0197-5	2500 in 1 PACKA 12500 in 1 PACK Cormation ry Applicat	AGE; Type 0: Not a Combination Pr AGE; Type 0: Not a Combination F	roduct Product	0 1/0 1/20 18	g Start Date	Marketi	ng End Date			

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

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Establishment									
Name	Address	ID/FEI	Business Operations						
Rxhomeo Private Limited d.b.a. Rxhomeo, Inc		650833994	manufacture(15631-0197), label(15631-0197)						

Revised: 3/2020

Rxhomeo Private Limited d.b.a. Rxhomeo, Inc