OSTEODORON AM- osteodoron am powder Uriel Pharmacy 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.

Osteodoron AM

Directions: FOR ORAL USE ONLY.

Take in the morning. Use in combination with Osteodoron PM Powder. Ages 12 and older: 1/8 teaspoon. Ages 2-11: 1/16 teaspoon. Under age 2: Consult a doctor.

Active Ingredients: Cucurbita (Squash) 3X, Apatite (Nat. calcium fluorophosphate) 6X,

Fluorite (Calcium fluoride) 6X, Quartz (Rock crystal) 6X

Inactive Ingredient: Lactose

Prepared using rhythmical processes.

Uses: For healthy bone and teeth development.

KEEP OUT OF REACH OF CHILDREN.

Warnings: Claims based on traditional homeopathic practice, not accepted medical evidence. Not FDA evaluated. Do not use if allergic to any ingredient. Contains lactose. Consult a doctor before use for serious conditions or if conditions worsen or persist. If pregnant or nursing, consult a doctor before use. Do not use if safety seal is broken or missing.

Questions? Call 866.642.2858 Uriel, East Troy, WI 53120 shopuriel.com Lot:





Osteodoron AM

Homeopathic Powder net wt. 1.7 oz (50g)



OSTEODORON AM

osteodoron am powder

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:48951-7190	
Route of Administration	ORAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
CUCURBITA PEPO FLOWER (UNII: 413MGP37HQ) (CUCURBITA PEPO FLOWER - UNII:413MGP37HQ)	CUCURBITA PEPO FLOWER	3 [hp_X] in 1 g	
CALCIUM FLUORIDE (UNII: O3B55K4YKI) (FLUORIDE ION - UNII:Q80VPU4080)	CALCIUM FLUORIDE	6 [hp_X] in 1 g	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4) (SILICON DIOXIDE - UNII:ETJ7Z6XBU4)	SILICON DIOXIDE	6 [hp_X] in 1 g	
FLUORAPATITE (UNII: M4CM1H238J) (FLUORAPATITE - UNII:M4CM1H238J)	FLUORAPATITE	6 [hp_X] in 1 g	

Inactive Ingredients			
Ingredient Name	Strength		
LACTOSE, UNSPECIFIED FORM (UNII: J2B2A4N98G)			

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:48951- 7190-4	50 g in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	09/01/2009	

Marketing Information			
Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
	09/01/2009		
4	• •	Citation Date	

Labeler - Uriel Pharmacy Inc. (043471163)

Establishment				
Name	Address	ID/FEI	Business Operations	
Uriel Pharmacy Inc.		043471163	manufacture(48951-7190)	

Revised: 12/2024 Uriel Pharmacy Inc.