#### CUPRUM OXYDATUM NIGRUM- cupric oxide pellet Boiron

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.

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#### Cuprum oxydatum nigrum 200CK

Cuprum oxydatum nigrum 200CK HPUS

Active ingredient\*\*: See product name on front panel (\*\*contains 0.443 mg of the active ingredient per pellet).

Uses: See symptoms on front panel.

Relieves nervousness in children \*

Stop use and ask a doctor if symptoms persist for more than 3 days or worsen.

If pregnant or breast-feeding, ask a health professional before use.

### Keep out of reach of children.

Adults and children: At the onset of symptoms, dissolve 5 pellets under the tongue 3 times a day until symptoms are relieved or as directed by a doctor.

lactose, sucrose

## BoironUSA.com Info@boiron.com

**1-800-BOIRON-1** (1-800-264-7661) Distributed by Boiron, Inc. Newtown Square, PA 19073

Do not use if pellet dispenser seal is broken.

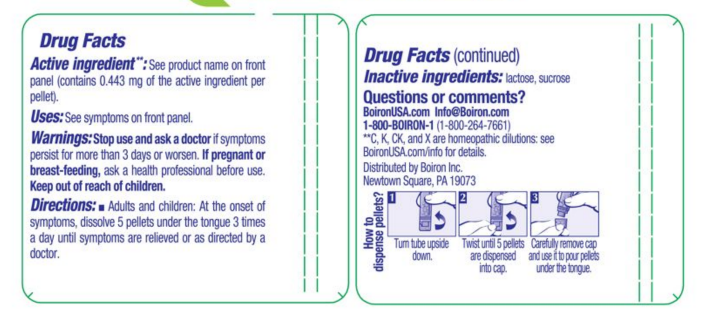
Contains approx.80 pellets.

How to dispense pellets? Turn tube upside down. Twist until 5 pellets are dispensed into cap. Carefully remove the cap and use it to pour pellets under the tongue.

\*CLAIMS BASED ON TRADITIONAL HOMEOPATHIC PRACTICE NOT ACCEPTED MEDICAL EVIDENCE. NOT FDA EVALUATED.

\*\*C,K,CK, and X are homeopathic dilutions: see BoironUSA.com/info for details.

Cuprum oxydatu Relieves nervou	sness in childre	en* BOIR	O.N <sup>®</sup>	NOT ACCEPTED MEDICAL EVIDE
Made in Frank HPUS NDC 0220-10		OPATHIC ME	DICINE	
Do not use if pellet				nigrum 200 ck
	LOT	EXP		DA EV/
3	06961	62041	8	RACT
Peel for D	Contains appro rugs Facts and	x. 80 pellets. d instructions f	US for use.	



CUPRUM OXYDATUN cupric oxide pellet	4 NIGRUM					
Product Information						
Product Type	HUMAN OTC DRUG Item		ode (Source)	NDC:0220-1620		
Route of Administration	ORAL					
Active Ingredient/Active Moiety						
Ingredient Name			Basis of Strength	Strength		
CUPRIC OXIDE (UNII: V1XJQ704R4) (CUPRIC CATION - UNII:8CBV67279L)			CUPRIC CATION	200 [kp_C] in 200 [kp_C]		

In	active Ingre	edients					
			Ingredie	ent Name	3		Strength
LA	CTOSE, UNSPE	CIFIED FORM	<b>1</b> (UNII: J2B2A4	N98G)			
su	CROSE (UNII: C	151H8M554)					
Pr	oduct Char	acteristics	5				
Color		white	Se	Score			
Shape		ROUND	Si	ze	4mm		
Flavor			In	Imprint Code			
Co	ntains						
Pa	ackaging						
#	ltem Code	P	Package Description		I	Marketing Start Date	Marketing End Date
	NDC:0220- 1620-41	200 [kp_C] in Product	1 TUBE; Type	0: Not a Co	ombination	01/01/2024	
Μ	arketing	Informa	tion				
	Marketing Category	Applic	cation Numb Cita	per or Mo ition	nograph	Marketing Start Date	Marketing End Date
	approved					01/01/2024	

# Labeler - Boiron (282560473)

Registrant - Boiron, Inc. (014892269)

Establishment						
Name	Address	ID/FEI	<b>Business Operations</b>			
Boiron		282560473	manufacture(0220-1620)			

Revised: 4/2024

Boiron