DIGITALIS PURPUREA- digitalis 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.

Digitalis purpurea 1M

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(**contains 0.443 mg of the active ingredient per pellet)

Rx Only*

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

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.

lactose, sucrose

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.

1-800-BOIRON-1 (1-800-264-7661), BoironUSA.com Info@boiron.com Distributed by Boiron, Inc. Newtown Square, PA 19073



purea 1 m

DENCE, NOT FDA EVA



Lot: Exp:

3 06960 26623 5

Contains approx. 80 pellets. US Peel for Drugs Facts and instructions for use.

Drug Facts

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

Uses: See symptoms on front panel.

Warnings: Stop use and ask a doctor if symptoms persist for more than 3 days or worsen. If pregnant or breastfeeding, ask a health professional before use. Keep out of reach of children.

Directions: 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.

Other information: ■ Do not use if pellet dispenser seal is broken.

Drug Facts (continued) **Inactive ingredients:** lactose, sucrose



DIGITALIS PURPUREA

digitalis pellet

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0220-1764
Route of Administration	ORAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
DIGITALIS (UNII: F1T8QT9U8B) (DIGITALIS - UNII:F1T8QT9U8B)	DIGITALIS	1 [hp_M] in 1 [hp_M]	

Inactive Ingredients		
Ingredient Name Stro		
LACTOSE, UNSPECIFIED FORM (UNII: J2B2A4N98G)		
SUCROSE (UNII: C151H8M554)		

Product Characteristics			
Color	white	Score	
Shape	ROUND	Size	4mm
Flavor		Imprint Code	

Contains

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
l		NDC:0220-1764- 41	1 [hp_M] in 1 TUBE; Type 0: Not a Combination Product	03/03/1983	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		03/03/1983	

Labeler - Boiron (282560473)

Registrant - Boiron, Inc. (014892269)

Establishment				
Name	Address	ID/FEI	Business Operations	
Boiron		282560473	manufacture(0220-1764)	

Revised: 11/2023 Boiron