

ONOPORDON COMP.- onopordon comp. pellet
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.

Onopordon comp.

Directions: FOR ORAL USE ONLY.

Dissolve pellets under the tongue 3-4 times daily. Ages 12 and older: 10 pellets. Ages 2-11: 5 pellets. Under age 2: Consult a doctor.

Active Ingredients: 100gm contains: 25gm Onopordon (Cotton thistle) 1X, 25gm Primula (Cowslip) 1X, 25gm Hyoscyamus (Henbane) 4X

Inactive Ingredient: Organic sucrose

Prepared using rhythmical processes.

Use: Promotes healthy circulatory support.

KEEP OUT OF REACH OF CHILDREN.

Warnings: Claims based on traditional homeopathic practice, not accepted medical evidence. Not FDA evaluated. Contains sugar. Diabetics and persons intolerant of sucrose (sugar): Consult a doctor before use. Do not use if allergic to any ingredient. 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:



ONOPORDON COMP.

onopordon comp. pellet

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:48951-7090
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ONOPORDUM ACANTHIUM FLOWER (UNII: AP97AUF88E) (ONOPORDUM ACANTHIUM FLOWER - UNII:AP97AUF88E)	ONOPORDUM ACANTHIUM FLOWER	1 [hp_X]
PRIMULA VERIS FLOWER (UNII: W5BET37294) (PRIMULA VERIS FLOWER - UNII:W5BET37294)	PRIMULA VERIS FLOWER	1 [hp_X]
HYOSCYAMUS NIGER LEAF (UNII: 32IT7G8BAW) (HYOSCYAMUS NIGER LEAF - UNII:32IT7G8BAW)	HYOSCYAMUS NIGER LEAF	4 [hp_X]

Inactive Ingredients

Ingredient Name	Strength
SUCROSE (UNII: C151H8M554)	

Product Characteristics

Color	white	Score	no score
Shape	ROUND	Size	3mm
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:48951-7090-2	1350 in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	09/01/2009	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		09/01/2009	

Labeler - Uriel Pharmacy Inc. (043471163)

Establishment

Name	Address	ID/FEI	Business Operations
Uriel Pharmacy Inc.		043471163	manufacture(48951-7090)

Revised: 11/2024

Uriel Pharmacy Inc.