GERI-DRYL ALLERGY RELIEF- diphenhydramine hcl tablet, coated RedPharm Drug, Inc.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Diphenhydramine 25mg

Active ingredient (in each tablet)

Diphenhydramine Hydrochloride 25mg

Purpose

Antihistamine

Uses

- temporarily relieves these symptoms due to hay fever and other upper respiratory allergies:
- runny nose
- sneezing
- itching of the nose and throat
- itchy, watery eyes

Warnings

Do not use

- to make a child sleepy
- with any other product containing diphenhydramine, including one used on skin

Ask a doctor before use if you have

- a breathing problem such as chronic bronchitis or emphysema
- trouble urinating due to an enlarged prostate gland
- glaucoma

Ask a doctor or pharmacist before use if you are

• taking sedatives or tranquilizers

When using this product

- excitability may occur, especially in children
- marked drowsiness may occur
- be careful when driving a motor vehicle or operating machinery
- avoid alcoholic drinks
- alcohol, sedatives and tranquilizers may increase drowsiness

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

Keep out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.

Directions

- do not exceed recommended dose
- adults and children 12 years and over, take 1 to 2 tablets every 4-6 hours, as needed; not more than 12 tablets in 24 hours, or as directed by a doctor
- children under 12 years: consult a doctor

Other information

- each tablet contains: calcium 25mg
- store at room temperature 150C 300C (590F 860F)
- for institutional use only

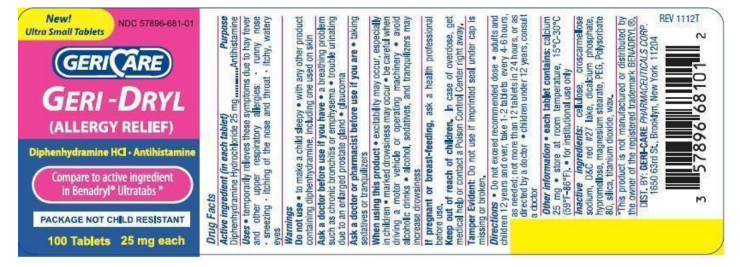
Inactive ingredients

cellulose, croscarmellose sodium, D&C red #27 lake, dicalcium phosphate, hypromellose, magnesium stearate, PEG, polysorbate 80, silica, titanium dioxide, wax.

package label

New! Ultra Small Tablets NDC 57896-681-01

GERICARE GERI-DRYL (ALLERGY RELIEF) Diphenhydramine HCl Antihistamine Compare to active ingredient in Benadryl Ultratabs PACKAGE NOT CHILD RESISTANT 100 Tablets 25 mg each



PACKAGE LABEL.PRINCIPAL DISPLAY PANEL

					1			
	Rx Only		67296-1758-9 TYDRAMINE 25MG 4 Tablets	HCL		7589 9		
99		Lot: 681R04	4X1 E	xp: 04/21		175		
(01)00367296175899 (21)1000000000046 (17)210430 (10)681R04X1	Usual adult dosa Store at controlle Mfg: Ger Broo Dist. by: Redpha	37	3 67296					
GERI-DRY	L ALLERGY	RELIEF						
_	e hcl tablet, coated							
Product Infor	rmation							
Product T ype	Product T ype		Item Code (Sourc	e) NDC:67296-1	.758(NDC:5789	6-681)		
Route of Admini	istration	ORAL						
Active Ingredient/Active Moiety Ingredient Name Basis of Strength DIPHENHYDRAMINE HYDROCHLORIDE (UNII: TC2D6JAD40) (DIPHENHYDRAMINE - UNII:8GTS82S83M) DIPHENHYDRAMINE - HYDROCHLORIDE						Strength 25 mg		
Inactive Ingre	dients							
Inactive Ingredients Ingredient Name								
CARNAUBA WAX (UNII: R12CBM0EIZ)						trength		
SILICON DIO XID	SILICON DIO XIDE (UNII: ETJ7Z6 XBU4)							
CROSCARMELLOSE SODIUM (UNII: M28OL1HH48)								
D&C RED NO. 27 (UNII: 2LRS185U6K)								
HYPROMELLOSES (UNII: 3NXW29V3WO) CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)								
CALCIUM PHO SPHATE, DIBASIC, DIHYDRATE (UNII: 07TSZ97GEP)								
MAGNESIUM STEARATE (UNII: 70097M6I30)								
POLYETHYLENE	POLYETHYLENE GLYCOL 400 (UNII: B697894SGQ)							
POLYSORBATE 80 (UNII: 60ZP39ZG8H)								
TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)								
Product Char	acteristics							
Color				no score				
Shape	CAPSUI	CAPSULE Size 11mm		11mm				
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Flavor	Imprint Code		T;061				
Contains							
Packaging							
# Item Code	Package Description	Marketing Start Date	Marketing End Date				
1 NDC:67296-1758-9	24 in 1 BOTTLE; Type 0: Not a Combination Product	0 1/0 1/20 20					
Marketing Information							
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date				
OTC monograph final	part336	09/01/2011					

Labeler - RedPharm Drug, Inc. (828374897)

Establishment						
Name	Address	ID/FEI	Business Operations			
RedPharm Drug, Inc.		828374897	relabel(67296-1758), repack(67296-1758)			

Revised: 1/2020

RedPharm Drug, Inc.