HARRIS TEETER ALL DAY ALLERGY D- cetirizine hcl, pseudoephedrine hcl tablet, extended release

Harris Teeter, LLC

Harris Teeter, LLC All Day Allergy D Drug Facts

Active ingredients (in each extended release tablet)

Cetirizine HCl 5 mg

Pseudoephedrine HCl 120 mg

Purpose

Antihistamine Nasal Decongestant

Uses

- temporarily relieves these symptoms due to hay fever or other upper respiratory allergies:
- runny nose
- sneezing
- itchy, watery eyes
- itching of the nose or throat
- nasal congestion
- reduces swelling of nasal passages
- temporarily relieves sinus congestion and pressure
- temporarily restores freer breathing through the nose

Warnings

Do not use

- if you have ever had an allergic reaction to this product or any of its ingredients or to an antihistamine containing hydroxyzine.
- if you are now taking a prescription monoamine oxidase inhibitor (MAOI) (certain drugs for depression, psychiatric, or emotional conditions, or Parkinson's disease), or for 2 weeks after stopping the MAOI drug. If you do not know if your prescription drug contains an MAOI, ask a doctor or pharmacist before taking this product.

Ask a doctor before use if you have

- heart disease
- thyroid disease
- diabetes
- glaucoma
- high blood pressure
- trouble urinating due to an enlarged prostate gland

• liver or kidney disease. Your doctor should determine if you need a different dose.

Ask a doctor or pharmacist before use if you are

taking tranquilizers or sedatives.

When using this product

- do not use more than directed
- drowsiness may occur
- avoid alcoholic drinks
- alcohol, sedatives, and tranquilizers may increase drowsiness
- be careful when driving a motor vehicle or operating machinery

Stop use and ask a doctor if

- an allergic reaction to this product occurs. Seek medical help right away.
- you get nervous, dizzy, or sleepless
- symptoms do not improve within 7 days or are accompanied by fever

If pregnant or breast-feeding:

- if breast-feeding: not recommended
- if pregnant: 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 break or chew tablet; swallow tablet whole

adults and children 12 years and over	take 1 tablet every 12 hours; do not take more than 2 tablets in 24
	hours.
adults 65 years and over	ask a doctor
children under 12 years of age	ask a doctor
consumers with liver or kidney	ask a doctor
disease	

Other information

• store between 20° to 25°C (68° to 77°F)

Inactive ingredients

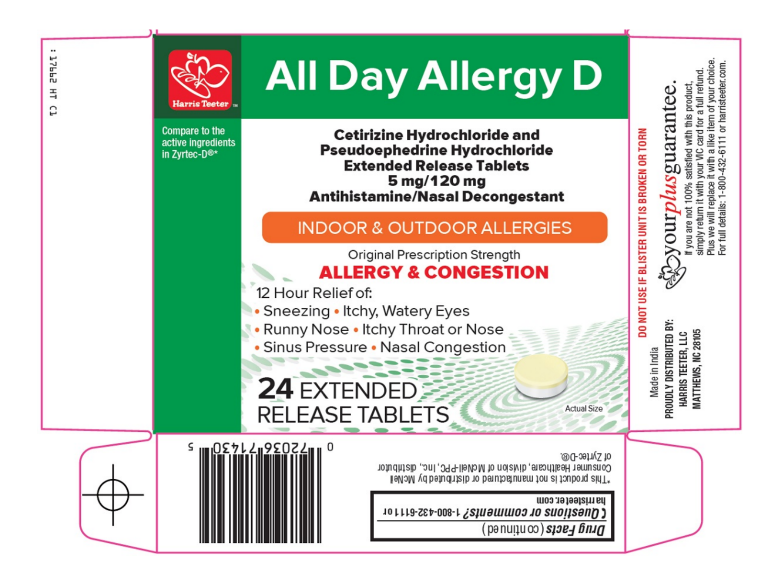
colloidal silicon dioxide, hydroxypropyl cellulose, hypromellose, lactose monohydrate, magnesium stearate, microcrystalline cellulose, talc, yellow iron oxide

Questions or comments?

Principal Display Panel

All Day Allergy D Compare to the active ingredients in Zyrtec-D® Cetirizine Hydrochloride and Pseudoephedrine Hydrochloride Extended Release Tablets 5 mg/120 mg Antihistamine/Nasal Decongestant INDOOR & OUTDOOR ALLERGIES Original Prescription Strength ALLERGY & CONGESTION 12 Hour Relief of: Sneezing – Itchy, Watery Eyes Runny Nose – Itchy Throat or Nose Sinus Pressure – Nasal Congestion 24 EXTENDED RELEASE TABLETS

Actual Size



$\overline{+}$	Drug Facts Active ingredients Purpose (in each extended release tablet) Cetirizine HCI 5 mg	 reduces swelling of r temporarily relieves s 	espiratory allergies: neezing itchy, watery eyes or throat nasal congestion	
	Drug Facts (continued)	Drug Facts (con	tinued)	
	<i>Warnings</i> Do not use ■ if you have ever had an allergic reaction to this	∎you get nervous, diz	zy, or sleepless prove within 7 days or are	
	product or any of its ingredients or to an antihistamine containing hydroxyzine. If you are now taking a prescription monoamine oxidase inhibitor (MAOI) (certain drugs for depression, psychiatric, or emotional conditions, or Parkinson's disease), or for 2 weeks after stopping the	Keep out of reach of ch		
	MAOI drug. If you do not know if your prescription drug contains an MAOI, ask a doctor or pharmacist before taking this product.	Directions ■ do not break or che	w tablet; swallow tablet whole	
	Ask a doctor before use if you have ■ heart disease ■ thyroid disease ■ diabetes ■ glaucoma ■ high blood pressure	adults and children 12 years and over	take 1 tablet every 12 hours; do not take more than 2 tablets in 24 hours.	
	 trouble urinating due to an enlarged prostate gland liver or kidney disease. Your doctor should determine if you need a different dose. 	adults 65 years and o <i>v</i> er	aska doctor	
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	Stop use and ask a doctor if ■ an allergic reaction to this product occurs. Seek medical help right away.		mon ohy drate, magnes ium stearate,	

HARRIS TEETER ALL DAY ALLERGY D

cetirizine hcl, pseudoephedrine hcl tablet, extended release

Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:69256-176	
Route of Administration	ORAL				
Active Ingradiant/Active	Moioty				
Active Ingredient/Active	wolety				
	Ingredient Name		Basis of St	rength	Strength
CETIRIZINE HYDRO CHLO RIDE (UNII: 640047KTOA) (CETIRIZINE - UNII: YO7261ME24)				0	
	E (UNII: 640047KTOA) (CETIRIZINE	-	CETIRIZINE HYDRO	OCHLORIDE	5 mg
UNII:YO7261ME24)	E (UNII: 640047KTOA) (CETIRIZINE ILORIDE (UNII: 6V9V2RYJ8N) (PSE		CETIRIZINE HYDRO PSEUDOEPHEDRIN HYDROCHLORIDE		U

		Ingredient Name				Strength	
SILICON DIO X	IDE (UN	II: ETJ7Z6 XBU4)					
HYDRO XYPRO	PYL CE	LLULOSE (1600000 WAMW) (UNII: RFW2ET671P)					
HYPRO MELLO	SES (UI	NII: 3NXW29V3WO)					
LACTOSE MO	NOHYD	RATE (UNII: EWQ57Q8I5X)					
MAGNESIUM S	TEARA	Г Е (UNII: 70097M6I30)					
MICROCRYST	ALLINE	CELLULOSE (UNII: OP1R32D61U)					
TALC (UNII: 7S	EV7J4R1	U)					
FERRIC OXIDE	YELLO	W (UNII: EX438O2MRT)					
Product Cha	aracter	ristics					
Color	WHITE (one side white one side light yellow)			Score		no score	
Shape	hape ROUND			Size		2mm	
-							
Flavor				Imprint Code	5	029;5;120	
Flavor				Imprint Code	5	029;5;120	
Flavor				Imprint Code	5	029;5;120	
Flavor				Imprint Code	5	029;5;120	
Flavor Contains				Imprint Code	5	029;5;120	
Flavor Contains Packaging	de	Package Description	Mark				
Flavor Contains Packaging # Item Co		U	Mark 07/24/	eting Start Date			
Flavor Contains Packaging # Item Co 1 NDC:69256-1	76-62 2	U		eting Start Date			
Flavor Contains Packaging # Item Co 1 NDC:69256-1	76-62 2	4 in 1 CARTON		eting Start Date			
Flavor Contains Packaging # Item Co 1 NDC:69256-1	76-62 2	4 in 1 CARTON		eting Start Date			
Flavor Contains Packaging # Item Co 1 NDC:69256-1 1	.76-62 2	4 in 1 CARTON in 1 BLISTER PACK; Type 0: Not a Combination Product		eting Start Date			
Flavor Contains Packaging # Item Co 1 NDC:69256-1 1 SMarketing	.76-62 2 1 g Info	4 in 1 CARTON in 1 BLISTER PACK; Type 0: Not a Combination Product rmation	07/24/	eting Start Date 2015	Marke	eting End Dat	
Flavor Contains Packaging	.76-62 2 1 g Info	4 in 1 CARTON in 1 BLISTER PACK; Type 0: Not a Combination Product rmation	07/24/	eting Start Date	Marke		

Labeler - Harris Teeter, LLC (048463103)

Revised: 12/2019

Harris Teeter, LLC