ALL DAY ALLERGY- cetirizine hydrochloride tablet, film coated Rite Aid Corporation

Rite Aid Corporation All Day Allergy Drug Facts

Active ingredient (in each tablet)

Cetirizine HCl 10 mg

Purpose

Antihistamine

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

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.

Ask a doctor before use if you have

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

- 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.

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. (1-800-222-1222)

Directions

years and over	one 10 mg tablet once daily; do not take more than one 10 mg tablet in 24 hours. A 5 mg product may be appropriate for less severe symptoms.
adults 65 years and over	ask a doctor
children under 6 years of	ask a doctor
age	
consumers with liver or kidney disease	ask a doctor

Other information

- store between 20-25°C (68-77°F)
- do not use if printed foil under cap is broken or missing

Inactive ingredients

corn starch, FD&C blue no. 1 aluminum lake, hypromellose, lactose monohydrate, magnesium stearate, polydextrose, polyethylene glycol, povidone, titanium dioxide, triacetin

Questions or comments?

1-800-719-9260

Principal Display Panel

Compare to the active ingredient in Zyrtec[®]

24 HOUR

ORIGINAL PRESCRIPTION STRENGTH

ALL DAY ALLERGY

CETIRIZINE HYDROCHLORIDE TABLETS, 10 mg

ANTIHISTAMINE

ACTUAL SIZE INDOOR & OUTDOOR ALLERGIES 24 HOUR RELIEF OF Sneezing • Runny nose • Itchy, watery eyes Itchy throat or nose 90 TABLETS



ALL DAY ALLERGY

cetirizine hydrochloride tablet, film coated

Product Infor	mation								
Product Type		HUMAN OTC D	RUG	ltem Code	(Source)	NDC:1182	2-1002		
Route of Admin	istration	ORAL					2.11022-1002		
Route of Autim		OTHE							
Active Ingred	ient/Active	Moiety							
	Ingre	edient Name	•		Basis of	Strength	Strengt		
CETIRIZINE HYDROCHLORIDE (UNII: 640047KTOA) (CETIRIZ UNII:YO7261ME24))A) (CETIRIZI	NE -	CETIRIZ INE 10 HYDROCHLORIDE		10 mg		
Inactive Ingredients									
STARCH CORN /	INIII: กรวรวทกวง	Ingredier	nt Name			S	trength		
STARCH, CORN (U FD&C BLUE NO. 1									
HYPROMELLOSE,			3WO)						
LACTOSE MONOH			,						
MAGNESIUM STE									
POLYDEXTROSE (
POLYETHYLENE G	GLYCOL, UNSP	ECIFIED (UNII: 3	3WJQ0SDW1A	()					
POVIDONE, UNSP									
TITANIUM DIOXID		V2JP)							
TRIACETIN (UNII: >	(HX3C3X673)								
Product Char	acteristics								
Color						no score			
Shape			Size			10mm			
Flavor			Imprint Code			4H2			
Contains									
Packaging									
# Item Code	Pa	ackage Desci	ription	Ma	arketing Start Date		ting End ate		
1 NDC:11822- 1002-4	1 in 1 CARTON			09/14	4/2021				
1	120 in 1 BOTTLE; Type 0: Not a Combination Product			on					
2 NDC:11822- 1002-2	1 in 1 CARTON				4/2021				
2	30 in 1 BOTTLE; Type 0: Not a Combination Product								
3 NDC:11822- 1002-1	14 in 1 CARTON				7/2021				
3 NDC:11822-	1 in 1 BLISTER PACK; Type 0: Not a Combination Product								
4 1002-3	1 in 1 CARTON 60 in 1 BOTTLE; Type 0: Not a Combination				7/2021				
A	OU IN T BOITL	.⊑; Type U: Not a	Combinatio	n					

4		Product							
5	NDC:11822- 1002-8	1 in 1 CARTON	03/23/2022						
5		90 in 1 BOTTLE; Type 0: Not a Combination Product							
6	NDC:11822- 1002-7	1 in 1 CARTON	10/19/2022						
6		300 in 1 BOTTLE; Type 0: Not a Combination Product							
Marketing Information									
	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date					
٨N	IDA	ANDA078336	09/14/2021						

Labeler - Rite Aid Corporation (014578892)

Revised: 3/2023

Rite Aid Corporation