LEVOCETIRIZINE DIHYDROCHLORIDE- levocetirizine dihydrochloride tablet, coated

Target Corporation

Drug Facts

Active ingredient (in each tablet)

Levocetirizine dihydrochloride USP, 5 mg

Purpose

Antihistamine

Uses

temporarily relieves these symptoms due to hay fever or other respiratory allergies:

- runny nose
- sneezing
- itchy, watery eyes
- itching of the nose or throat

Warnings

Do not use

- if you have kidney disease
- if you have ever had an allergic reaction to this product or any of its ingredients or to an antihistamine containing cetirizine

Ask a doctor before use if you have

• ever had trouble urinating or emptying your bladder

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 doctor if

- you have trouble urinating or emptying your bladder
- an allergic reaction to this product occurs. Seek medical help right away.

If pregnant or breast-feeding:

• if breast-feding: 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

adults 65 years of age and older	ask a doctor
adults and children 12-64 years of age	 take 1 tablet (5 mg) once daily in the evening do not take more than 1 tablet (5 mg) in 24 hours 1/2 tablet (2.5 mg) once daily in the evening may be appropriate for less severe symptoms
children 6-11 years of age	 take 1/2 tablet (2.5 mg) once daily in the evening do not take more than 1/2 tablet (2.5 mg) in 24 hours
children under 6 years of age	do not use
consumers with kidney disease	• do not use

Other information

- store between 20° and 25°C (68° and 77°F)
- (Bottle only)- Safety Sealed: do not use if carton was opened or if printed foil inner seal on bottle is torn or missing
- (Blister only)- Safety Sealed: Do not use if seal is broken or if individual blister unit is open or torn

Inactive ingredients

colloidal silicon dioxide, hypromellose, lactose monohydrate, magnesium stearate, microcrystalline cellulose, polyethylene glycol, titanium dioxide

Questions or comments?

Call 1-888-375-3784

Carton Label

Carton label



Bottle Label

Bottle label

original NDC 11673-847-35 scription strength tablets USP, **5 mg** antihistamine

Active ingredient (in each tablet) Purpose Levocetirizine dihydrochloride USP, 5 mg...... Antihistamine Uses temporarily relieves these symptoms due to hay fever or other respiratory allergies: ■ runny nose ■ sneezing ■ itchy, watery eyes ■ itching of the nose or throat Warnings Do not use ≡ if you have kidney disease ≡ if you have ever had an allergic reaction to this product or any of its ingredients or to an antihistamine containing cetirizine Ask a doctor before use if you have ■ ever had trouble urinating or emptying your bladder 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 wou have trouble urinating or emptying your bladder ■ 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

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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 adults 65 years of age and older ■ ask a doctor adults and children 12-64 years of age

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LEVOCETIRIZINE DIHYDROCHLORIDE

levocetirizine dihydrochloride tablet, coated

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:11673-847(NDC:43598-735)

Route of Administration ORAL

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength levocetirizine dihydrochloride (UNII: SOD6A38AGA) (levocetirizine levocetirizine 5 mg UNII:6U5EA9RT2O) dihydrochloride

Inactive Ingredients		
Ingredient Name	Strength	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)		
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)		
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		
HYPROMELLOSES (UNII: 3NXW29V3WO)		
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)		

POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)

Product Characteristics						
Color	white	Score	2 pieces			
Shape	OVAL	Size	9mm			
Flavor		Imprint Code	L			
Contains						

Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:11673-847- 80	1 in 1 CARTON	12/31/2018			
1		80 in 1 BOTTLE; Type 0: Not a Combination Product				
2	NDC:11673-847- 16	2 in 1 CARTON	12/01/2020			
2		80 in 1 BOTTLE; Type 0: Not a Combination Product				
3	NDC:11673-847- 35	1 in 1 CARTON	12/31/2018			
3		35 in 1 BOTTLE; Type 0: Not a Combination Product				
4	NDC:11673-847- 79	2 in 1 BLISTER PACK	12/31/2018			
4		5 in 1 BLISTER PACK; Type 0: Not a Combination Product				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
ANDA	ANDA210375	12/31/2018			

Labeler - Target Corporation (006961700)

Revised: 8/2020 Target Corporation