LORATADINE- loratadine tablet Advanced Rx Pharmacy of Tennessee, LLC

Loratadine 10mg Tablets

Drug Facts

Active Ingredient (in each tablet)

Loratadine, 10 mg USP

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 useif you ever have had an allergic reaction to this product or any of its ingredients.

Ask a doctor before use if you haveliver or kidney disease. Your doctor should determine if you need a different dose.

When using this product do not take more than directed. Taking more than directed may cause drowsiness.

Stop use and ask a doctor ifan allergic reaction to this product occurs. Seek medical help right away.

If pregnanct 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

adults and children 6 years of age and	1 tablet daily; not more than 1 tablet in 24
over	hours
children under 6 years of age	ask a doctor
consumers with liver or kidney disease	ask a doctor

Other Information

- Tamper-evident: do not use if foil seal under cap, printed with "SEALED for YOUR PROTECTION" is missing, open or broken.
- Store between 20° 25°C (68°-77°F)

Inactive ingredientscorn starch, lactose monohydrate, magnesium stearate, povidone

Questions or comments?

Call 1-800-874-7464 Monday to Friday 9 AM to 5 PM EST

How Supplied/ Storage and Handling

Loratidine is supplied as:

Bottles of 30 Tablets NDC: 80425-0352-01

store at 20-25°C (68-77°F) (see USP Controlled Room Temperature)

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL



LORATADINE

loratadine tablet

l	Product	Information	

Product Type HUMAN OTC DRUG Item Code (Source) NDC:80425-0352(NDC:16571-822)

Route of Administration ORAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
LORATADINE (UNII: 7AJO3BO7QN) (LORATADINE - UNII:7AJO3BO7QN)	LORATADINE	10 mg

Ingredient Name	Strength
ingredient Name	Stite

LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
POVIDONE (UNII: FZ 989GH94E)	
STARCH, CORN (UNII: O8232NY3SJ)	

Product Characteristics				
Color	white (white to off-white)	Score	no score	
Shape	ROUND	Size	6mm	
Flavor		Imprint Code	10;p	
Contains				

F	Packaging			
#	tem Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:80425- 0352-1	30 in 1 BOTTLE; Type 0: Not a Combination Product	07/31/2023	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA214684	07/31/2023	

Labeler - Advanced Rx Pharmacy of Tennessee, LLC (117023142)

Establishment				
Name	Address	ID/FEI	Business Operations	
Advanced Rx Pharmacy of Tennessee, LLC		117023142	repack(80425-0352)	

Revised: 7/2023 Advanced Rx Pharmacy of Tennessee, LLC