LORATADINE ALLERGY RELIEF- loratadine tablet OHM LABORATORIES INC.

Drug Facts

ACTIVE INGREDIENT(S)

Loratadine USP, 10 mg

PURPOSE

Antihistamine

USE(S)

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

- runny nose
- itchy, watery eyes
- sneezing
- 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.

Ask a doctor before use if you have

Liver 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 if

An allergic reaction to this product occurs. Seek medical help right away.

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

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

OTHER INFORMATION

- store between 20 and 25° C (68 and 77° F)
- protect from excessive moisture
- TAMPER EVIDENT: DO NOT USE IF IMPRINTED SEAL IS BROKEN OR MISSING FROM BOTTLE.

INACTIVE INGREDIENTS

Corn starch, lactose monohydrate, magnesium stearate, pregelatinized starch

QUESTIONS?

Call 1-800-406-7984

PRINCIPAL DISPLAY PANEL - 10 mg Tablet Bottle Label

Original Prescription Strength NDC 51660-526-53

NON-DROWSY*

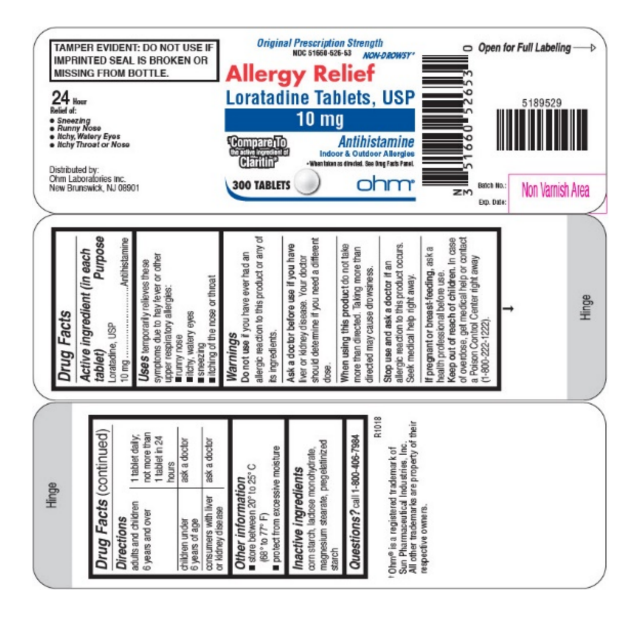
Allergy Relief Loratadine Tablets, USP 10 mg

[†]Compare To the active ingredient of Claritin[®]

Antihistamine Indoor & Outdoor Allergies * When taken as directed. See Drug Facts Panel.

300 TABLETS

ohm®



LORATADINE ALLERGY RELIEF

oratadine tablet					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:51660-526	
Route of Administration	ORAL				
Active Ingredient/Active N	Лоiety				
	Ingredient Name		Basis of S	trength	Strength
LORATADINE (UNII: 7AJO3BO7QN) (LORATADINE - UNII:7AJO3BO7QN) LORATADINE			2	10 mg	
Inactive Ingredients					
Ingredient Name			St	Strength	
	8				0
STARCH, CORN (UNII: 08232NY3					U
STARCH, CORN (UNII: 08232NY3 LACTOSE MONOHYDRATE (UN	SJ)				U

P	Product Characteristics						
С	olor	white (White to Off White)	Score		no score		
s	hape	ROUND	Size		6 mm		
F	lavor		mprint Co	ode	RX526		
С	ontains						
_							
P	ackaging						
#	Item Code	Package Description	Market	ting Start Date	Marketing l	End Date	
1	NDC:51660-526-01	100 in 1 BOTTLE; Type 0: Not a Combination Product	11/0 1/20	17			
2	NDC:51660-526- 05	500 in 1 BOTTLE; Type 0: Not a Combination Product	11/01/20	17			
3	NDC:51660-526-53	300 in 1 BOTTLE; Type 0: Not a Combination Product	11/0 1/20	17			
4	NDC:51660-526- 30	30 in 1 BOTTLE; Type 0: Not a Combination Product	02/01/20) 19			
5	NDC:51660-526-31	30 in 1 BLISTER PACK; Type 0: Not a Combination Produ	ct 11/01/20	17			
Marketing Information							
ľ	Marketing Categor	y Application Number or Monograph Citation	Marketi	ng Start Date	Marketing E	and Date	
A	NDA	ANDA076134	11/0 1/20 17	7			

Labeler - OHM LABORATORIES INC. (184769029)

Registrant - Ranbaxy Pharmaceuticals Inc. (937890044)

Establishment

Name	Address	ID/FEI	Business Operations
Ohm Laboratories Inc.		184769029	MANUFACTURE(51660-526)

Revised: 3/2019

OHM LABORATORIES INC.