#### LORATADINE ALLERGY RELIEF- loratadine tablet Proficient Rx LP

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# **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

NDC 63187-100-10

<sup>†</sup>Compare to the active ingredient of Claritin<sup>®</sup>

**NON-DROWSY**\*

24 HourAllergy Relief

Loratadine Tablets USP, 10 mg

Antihistamine

Indoor & Outdoor Allergies

Relief of:

- Sneezing
- Runny Nose
- Itchy, Watery Eyes
- Itchy Throat or Nose

10 Tablets

When taken as directed. See Drug Facts Panel.

Manufactured by: Ohm Laboratories Inc.

5069178/0908

**Repackaged by: Proficient Rx LP** Thousand Oaks, CA 91320





NDC 63187-100-10

Packaged By: Proficient Rx LP Thousand Oaks, CA 91320

Loratadine (Allergy Relief )10mg #10 Tablets Lot #:00000 SN# MASTER Exp:00/00/00 NDC 63187-100-10

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Loratadine (Allergy Relief )10mg SN# MASTER Tablets #10 Lot #:00000 Exp:00/00/00 NDC 63187-100-10



GTIN: 00363187100100 SN# MASTER Exp. 00/00/00 Lot #:00000

LORATADINE	ALLERO	GY RELIEF							
<b>Product Inform</b>	ation								
Product Type		HUMAN OTC DRUG	Item Code (Source) NDC:63187			NDC:63187-100(N	7-100(NDC:51660-526)		
<b>Route of Administ</b>	ration	ORAL							
Active Ingredier	nt/Active	Majaty							
Active Ingredient/Active Moiety							1-	Charles and the	
					asis of Strengt		Strength		
LORATADINE (UNII: 7AJO3BO7QN) (LORATADINE - UNII:7AJO3BO7QN) LORATADINE					ATADINE		10 mg		
Inactive Ingredi	ents								
		Ingredient Nam	ne				Strength		
STARCH, CORN (UNII: 08232NY3SJ)									
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)									
MAGNESIUM STEARATE (UNII: 70097M6I30)									
Product Charac	teristics								
Color	white (White	to Off White)		Score			no s	core	
Shape	ROUND			Size			6mm	ı	

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Loratadine (Allergy Relief )10mg

#### #10 Tablets

Each tablet contains: Loratadine, USP 10 mg Antihistamine

White to Off White, round, unscored tablet with imprint RX526 on one side and plain on the other side

Product ID: PL010010

Dist. By: Ohm Laboratories Inc. 1385 Livingston Avenue North Brunswick, NJ 08902 Store at 20°-25°C (68°-77°F)

Keep medication out of the reach of children

Flavor		Ir	nprint Code	RX526			
Co	ontains						
Pa	ackaging						
#	ltem Code	Package Description		Marketing Start Date	Marketing End Date		
1	NDC:63187-100- 10	10 in 1 BOTTLE; Type 0: Not a Combination Product	01/	/01/2019			
2	NDC:63187-100- 20	20 in 1 BOTTLE; Type 0: Not a Combination Product	01/	/01/2019			
	NDC:63187-100- 30	30 in 1 BOTTLE; Type 0: Not a Combination Product	on 01/01/2019				
R/	larkating	Information					
Marketing Information							
Marketing Category		Application Number or Monograph Citation		Marketing Start Date	Marketing End Date		
	ANDA ANDA076134			08/28/2003			

Labeler - Proficient Rx LP (079196022)

Establishment						
Name	Address	ID/FEI	Business Operations			
Proficient Rx LP		079196022	REPACK(63187-100), RELABEL(63187-100)			

Revised: 11/2022

Proficient Rx LP