LORATADINE- loratadine tablet Select Brand

Drug Facts

ACTIVE INGREDIENT (IN EACH TABLET)

Loratadine USP, 10 mg

PURPOSE

Antihistamine

USES

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

ladilite and children 6 vears 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 or 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 BLISTER UNITS ARE TORN, BROKEN OR SHOW ANY SIGNS OF TAMPERING.

INACTIVE INGREDIENTS

Corn starch, lactose monohydrate, magnesium stearate, pregelatinized starch

QUESTIONS?

Call 1-800-406-7984

PRINCIPAL DISPLAY PANEL *Compare to the active ingredient of Claritin[®]

NDC 15127-715-30

select $brand_{\mathbb{R}}$

NON-DROWSY

Allergy Relief

Loratadine Tablets USP, 10 mg

Antihis tamine

INDOOR & OUTDOOR ALLERGIES

RELIEF OF:

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

24 Hour Allergy Relief

30 Tablets

When taken as directed. See Drug Facts Panel.

Distributed by: SELECT BRAND DISTRIBUTORS

5069402/1008



LORATADINE loratadine tablet

	duct Informat	ion							
Product T ype			HUMAN OTC DRUG	Ite m	Item Code (Source)		NDC:151	27-715	
Route of Administration			ORAL						
Acti	ve Ingredient	/Active Moi	ety						
Ing			redient Name		Basis of Strength		Strength		
LORATADINE (UNII: 7AJO3BO7QN) (I			LORATADINE - UNII:7AJO3BO7QN)			LORATADINE		10 mg	
Inac	tive Ingredie	nts							
			Ingredient Nan	ne			5	Strength	
	RCH, CORN (UNII:	,							
	FOSE MONOHYD								
	NESIUM STEARA		7M6I30) UNII: 08232NY3SJ)						
Pro	duct Characte	ristics							
Colo		white (White to	Off-White)		Score		no s	no score	
Shape ROUND				Size		6 mm			
Flavor				Imprint Code		RX5	RX526		
Cont	ains								
Pac	kaging								
	Item Code	Pack	age Description	Marketi	ing Start Da	te Ma	Marketing End Date		
#	C:15127-715-10	10 in 1 BL	ISTER PACK						
	C:15127-715-30	30 in 1 BL	ISTER PACK						
1 NE		60 in 1 BC	DTTLE						
1 NE 2 NE	C:15127-715-60								
2 NE3 NE	0C:15127-715-60 0C:15127-715-01	100 in 1 B	OTTLE						
 1 NE 2 NE 3 NE 		100 in 1 B	OTTLE						
 1 NE 2 NE 3 NE 4 NE 	c:15127-715-01 rketing Info	ormation	OTTLE						
1 NE 2 NE 3 NE 4 NE	C:15127-715-01	ormation	OTTLE on Number or Monogr	raph Citation	Marketin	ng Start Date	Marketi	ng End Date	

Labeler - Select Brand (043562370)

Registrant - Ranbaxy Pharmaceuticals Inc. (937890044)

Establishment									
Name	Address	ID/FEI	Business Operations						
Ohm Laboratories Inc.		051565745	manufacture(15127-715)						

Revised: 10/2012