HEALTHY ACCENTS ALLERGY RELIEF- loratadine tablet DZA Brands LLC

DZA Brands, LLC Allergy Relief Drug Facts

Active ingredient (in each tablet)

Loratadine 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. (1-800-222-1222)

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 or kidney disease	ask a doctor

Other information

- do not use if printed foil under cap is broken or missing (bottle only)
- store between 20° to 25°C (68° to 77°F)

Inactive ingredients

lactose monohydrate, magnesium stearate, povidone, pregelatinized starch

Questions or comments?

1-866-322-2439

Principal Display Panel

Compare to Claritin® Tablets active ingredient

non-drowsy*

allergy relief

24 hour relief of:

sneezing

runny nose

itchy, watery eyes

itchy throat or nose

loratadine tablets, 10 mg – antihistamine

indoor & outdoor allergies

original prescription strength

NO GLUTEN

actual size

30 tablets

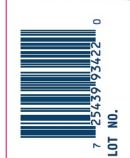
^{*}When taken as directed. See drug facts panel.

and code from package. For product questions or concerns, contects 1-866-322-2439.
Please include UPC number SCAR BOROUGH, ME 04074 SALISBURY, NC 28147 DISTRIBUTED BY: DZA BRANDS, LLC

Questions or comments? 1-866-322-2439

actose monohydrate, magnesium stearate, povidone, pregelatinized starch Inactive ingredients

Drug Facts (continued)



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

If pregnant or breast-reading, ask a health professional before use. Keep out of reach of children. In case of overdose, get medical help or contact a Polson Control Center light away (1-800-222-1222)

Stop use and ask a doctor if an allergic reaction to this product occurs. Seek medical help Drug Facts (continued)

allergy relief

30 tablets for 30 days of relief ontains 1 bothe with 30 tablets

indoor & outdoor allergies

antihistamine

loratadine tablets,10mg

cause drowsiness.

When using this product do not take more than directed. Taking more than directed may Ask a doctor before use if you have liver or kidney disease. Your doctor should determine if you need a different dose.

Do not use if you have ever had an allergic reaction to this product or any of its ingredients *SbuinneW*

Uses temporarily relieves these symptoms due to hay fever or other upper respliatory allergles: \blacksquare runny nose \blacksquare tichy, watery eyes \blacksquare sneezing \blacksquare itching of the nose or throat

Antihistamine

Loratadine 10 mg Active ingredient (in each tablet)

Drug Facts

NDC 55316-612-65 Compare to Claritin® Tablets active ingredient

allergy relief

loratadine tablets, 10mg • antihistamine indoor & outdoor allergies

original prescription strength





*When taken as directed. See drug facts panel.

30 tablets

itchy, watery eyesitchy throat or nose

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24 hour relief of:

sneezingrunny nose



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QUESTIONS OF COMMENTS? 1-866-322-2439

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Directions adults and children 6 years and o <i>v</i> er	nsrit enorm on y lisb feblet f en let in 24 hours

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this product occurs. Seek medical help nght away. Stop use and ask a doctor it an allengic reaction to

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disease. Your doctor should determine if you need a Ask a doctor before use if you have liver or kidney

this product or any of its ingredients Do not nae il you have ever had an allergic reaction to *SbujujeM*

Ifching of the nose or throat

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Active ingredient

Drug Facts

allergy relief

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loratadine tablets, 10mg • antihistamine

*When taken as directed. See drug facts panel.



NDC 55316-612-46 Compare to Claritin® Tablets active ingredient

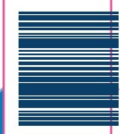
allergy relief

loratadine tablets, 10mg • antihistamine

indoor & outdoor allergies original prescription strength

24 hour relief of:

- gluten free sneezing
- runny nose
- itchy, watery eyes
- itchy throat or nose





loratadine tablets, IOmg • **antihistamine**

*When taken as directed. See drug facts panel.



10 tablets

HEALTHY ACCENTS ALLERGY RELIEF

loratadine tablet

Product	Information
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Product TypeHUMAN OTC DRUGItem Code (Source)NDC:55316-612

Route of Administration ORAL

Active Ingredient/Active Moiety

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

Inactive Ingredients

Ingredient Name	Strength	
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)		
MAGNESIUM STEARATE (UNII: 70097M6I30)		
PO VIDO NE, UNSPECIFIED (UNII: FZ989GH94E)		

Product Characteristics

Color	WHITE	Score	no score
Shape	OVAL	Size	8 mm
Flavor		Imprint Code	L612
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:55316-612-46	10 in 1 CARTON	12/15/2007		
1		1 in 1 BLISTER PACK; Type 0: Not a Combination Product			
2	NDC:55316-612-65	1 in 1 CARTON	07/07/2008		
2		30 in 1 BOTTLE; Type 0: Not a Combination Product			
3	NDC:55316-612-75	1 in 1 CARTON	08/28/2009		
3		90 in 1 BOTTLE; Type 0: Not a Combination Product			
4	NDC:55316-612-87	1 in 1 CARTON	02/18/2008		
4		300 in 1 BOTTLE; Type 0: Not a Combination Product			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA076301	12/15/2007	

Labeler - DZA Brands LLC (090322194)

Revised: 12/2019 DZA Brands LLC