OLOPATADINE HYDROCHLORIDE OPHTHALMIC SOLUTION- olopatadine hydrochloride ophthalmic solution Albertsons Companies

ACTIVE INGREDIENT

Olopatadine (0.2%) (equivalent to olopatadine hydrochloride 0.222%)

PURPOSE

Antihistamine

USE

temporarily relieves itchy eyes due to pollen, ragweed, grass, animal hair and dander

WARNINGS

For external use only

DO NOT USE

- if solution changes color or becomes cloudy
- if you are sensitive to any ingredient in this product
- to treat contact lens related irritation

WHEN USING THIS PRODUCT

- do not touch tip of container to any surface to avoid contamination
- remove contact lenses before use
- wait at least 10 minutes before reinserting contact lenses after use
- do not wear a contact lens if your eye is red

STOP USE AND ASK DOCTOR IF

you experience:

- eye pain
- changes in vision
- increased redness of the eye
- itching worsens or lasts for more than 72 hours

KEEP OUT OF REACH OF CHILDREN

If swallowed, get medical help or contact a Poison Control Center right away.

DIRECTIONS

- adults and children 2 years of age and older:
- put 1 drop in the affected eye(s) once daily, no more than once per day
- if using other ophthalmic products while using this product, wait at least 5 minutes between each product
- replace cap after each use
- children under 2 years of age: consult a doctor

OTHER INFORMATION

- only for use in the eye
- store between 2° to 25°C (36° to 77°F)

INACTIVE INGREDIENTS

Benzalkonium chloride 0.01%, Dibasic sodium phosphate, Edetate disodium, Hydrochloric acid/Sodium hydroxide (adjust pH), Povidone, Sodium chloride, and Water for Injection.

QUESTIONS?

Call 1-888-375-3784

PRINCIPAL DISPLAY PANEL

Olopatadine Hydrochloride Ophthalmic Solution USP, 0.2%

Placeholder Image

OLOPATADINE HYDROCHLORIDE OPHTHALMIC SOLUTION

olopatadine hydrochloride ophthalmic solution

Product Information

Product Type		HUMAN OTC DRUG Item Code (Source)		ource)	NDC:21130-105(NDC:43598-764)		
Route of Admin	nistration	OPHTHALMIC					
Active Ingredient/Active Moiety							
Ingredient Name					Basis of		Strength
-					Streng	Jth	
OLOPATADINE HYDROCHLORIDE (UNII: 2XG66W44KF) (OLOPATADINE - UNII:D27V6190PM)					OLOPATADIN	E	2 mg in 1 mL
Inactive Ingr	adiants						
Ingredient Name							Strength
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)							
SODIUM PHOSPHATE, DIBASIC, ANHYDROUS (UNII: 22ADO53M6F)							
EDETATE DISODIUM (UNII: 7FLD91C86K)							
HYDROCHLORIC ACID (UNII: QTT17582CB)							
POVIDONE K30 (UNII: U725QWY32	2X)					
SODIUM CHLORIDE (UNII: 451W47IQ8X)							
SODIUM HYDROXIDE (UNII: 55X04QC32I)							
WATER (UNII: 059	QF0KO0R)						
Packaging							
# Item Code	P	ackage Descript	ion	Marketing Start Date		Marketing End Date	
1 NDC:21130- 105-02	1 in 1 CARTON			06/25/2021			
1	2.5 mL in 1 BO ⁻ Combination Pr	TTLE, PLASTIC; Type (oduct): Not a				
Marketing	Informat	ion					
Marketing Category	Applica	tion Number or M Citation	onograph	Marketi Da	-	Marl	ceting End Date
ANDA ANDA209				06/25/2021			

Labeler - Albertsons Companies (009137209)

Revised: 3/2021

Albertsons Companies