



TIM-DOR-LAT- timolol - dorzolamide - latanoprost pf solution/ drops
ImprimisRx NJ

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Store at 20° to 25° C (68° to 77° F)

Bottle Label

Sterile 5ml Bottle		
Timolol-Dorzolamide-Latanoprost P-F		70261-521-05
(0.5/2/0.005)%		
Ophthalmic Drops		
Compounded for a licensed professional or patient use by		
		
ImprimisRx NJ 1705 Route 46 West, Suite 6A Ledgewood, NJ 07852 (844) 446-6979		
Each mL contains: Timolol 5mg, Dorzolamide 20mg, Latanoprost 0.05mg, Sodium Chloride 3.8mg, Sodium citrate 1.5mg, Pluronic L64 0.001ml, Polysorbate 80 NF 0.0005 ml and Sterile Water q.s. Sodium Hydroxide may have been used to adjust pH.		
Store under refrigeration 2-8°C (36-46°F). This medicine was compounded for you at the direction of your prescriber. Protect from light. Rx Only - Not for resale		
Lot#		
Use By:		
Date Compounded: mm/dd/yyyy hh:mm xx		

TIM-DOR-LAT

timolol - dorzolamide - latanoprost pf solution/ drops

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:70261-521
Route of Administration	OPHTHALMIC		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
DORZOLAMIDE HYDROCHLORIDE (UNII: QZO5366EW7) (DORZOLAMIDE - UNII:9JDX055TW1)	DORZOLAMIDE	20 mg in 1 mL
TIMOLOL MALEATE (UNII: P8Y54F701R) (TIMOLOL ANHYDROUS - UNII:5JKY92S7BR)	TIMOLOL ANHYDROUS	5 mg in 1 mL
LATANOPROST (UNII: 6Z5B6HVF6O) (LATANOPROST - UNII:6Z5B6HVF6O)	LATANOPROST	0.05 mg in 1 mL

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:70261-521-05	5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product	01/01/2018	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		01/01/2018	

Labeler - ImprimisRx NJ (931390178)

Revised: 5/2018

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