## CYCLOSPORINE/CHONDROITIN SULFATE PF- cyclosporine/chondroitin sulfate pf emulsion Imprimis Rx 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.

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Store at 20° to 25° C (68° to 77° F)

Sterile 5.5ml Bottle

## Cyclosporine 0.1% Chondroitin Sulfate

Preservative Free Ophthalmic Emulsion

Compounded for a licensed professional or patient use by



1705 Route 46 West, Suite 6A Ledgewood, NJ 07852 (973)328-8756

#### NDC 70261-514-05

Each mL contains: Cyclosporine USP 1mg
Inactive ingredients: Chondroitin Sulfate USP,
Glycerin USP, Dextran 40 USP, Methocel E4M,
Poloxamer 407 NF, Polysorbate 80 NF, Cremophor® EL,
Ethyl Alcohol in Balanced Salt Solution and
Sterile Water for injection.
Sodium Hydroxide may have been used to adjust pH.

Store at controlled room temperature 20-25°C (68 -77°F). This medicine was compounded for you at the direction of your prescriber. Protect from light. Rx Only - Not for resale



Lot#

Use By:

#### CYCLOSPORINE/CHONDROITIN SULFATE PF

cyclosporine/chondroitin sulfate pf emulsion

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Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:70261-514
Route of Administration	OPHTHALMIC		

#### **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
CYCLOSPORINE (UNII: 83HN0GTJ6D) (CYCLOSPORINE - UNII:83HN0GTJ6D)	CYCLOSPORINE	1 mg in 1 mL

# # Item Code Package Description Marketing Start Date 1 NDC:70261-514- 05 S.5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product 02/01/2018

Marketing Information								
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date					
unapproved drug other		02/01/2018						

### Labeler - Imprimis Rx NJ (931390178)

Revised: 2/2018 ImprimisRx NJ