SODIUM CHLORIDE HYPERTONICITY- sodium chloride ointment CVS PHARMACY

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Drug Facts

Active ingredient

Sodium chloride, 50 mg (5%)

Purpose

Hypertonicity agent

Uses

temporary relief of corneal edema

Warnings

For external use only

Do not use

except under the advice and supervision of a doctor

When using this product

- it may cause temporary burning and irritation
- replace cap after use
- to avoid contamination do not touch tip of container to any surface

Stop use and ask a doctor if

- condition worsens or persists for more than 72 hours
- you experience eye pain, changes in vision, continued redness or irritation of the eye

Keep out of reach of children.

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

Directions

- pull down the lower lid of the affected eye
- apply a small amount (1/4 inch) of ointment to the inside of eyelid
- apply every 3 or 4 hours or as directed by a doctor

Other information

- store at 15°-25°C (59°-77°F)
- keep tightly closed
- DO NOT FREEZE
- see crimp of tube or carton for Lot Number and Expiration Date
- do not use if difficult to dispense or visible particles are seen in the product
- serious side effects associated with use of the product may be reported to the phone number below

Inactive ingredients

lanolin, mineral oil, purified water, white petrolatum

Questions

[phone icon] Call 1-866-767-9161

Package/Label Principal Display Panel - Carton

[heart icon] \mathbf{CVS} Health_{TM}

Compare to the active ingredient in Muro $128^{\$}*$

NDC 69842-285-35

Sodium Chloride Hypertonicity ophthalmic ointment, 5% Temporary relief of corneal edema

NET WT 0.125 OZ (3.5 g)

STERILE



SODIUM CHLORIDE HYPERTONICITY

LANOLIN (UNII: 7EV65EAW6H)

sodium chloride ointment						
Product Information						
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:6984	NDC:69842-285	
Route of Administration	OPHTHALMIC					
A -4' T 1'4/A -4' B.F'						
Active Ingredient/Active Moi	ету					
Ingredient Name			Basis of S	Strength	Strength	
SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37)			SODIUM CH	LORIDE	50 mg in 1 g	
Inactive Ingredients						
Ingredient Name				Strength		

MINERAL OIL (UNII: T5L8T28FGP)	
WATER (UNII: 059QF0KO0R)	
PETROLATUM (UNII: 4T6H12BN9U)	

Packaging				
3	# Item Code	Package Description	Marketing Start Date	Marketing End Date
:	NDC:69842-285-35	1 in 1 CARTON	05/01/2020	
:	L	3.5 g in 1 TUBE; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph final	part349	05/01/2020		

Labeler - CVS PHARMACY (062312574)

Establishment				
Name	Address	ID/FEI	Business Operations	
Bausch & Lomb Incorporated		079587625	MANUFACTURE(69842-285)	

Revised: 1/2021 CVS PHARMACY