# SUNMARK LUBRICATING PLUS- carboxymethylcellulose sodium solution/ drops

# Strategic Sourcing Services LLC

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.

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# McKesson Lubricating Plus Eye Drops Drug Facts

# Active ingredient (in each single-use container)

Carboxymethylcellulose sodium 0.5%

# Purpose

Eye lubricant

### Uses

- for the temporary relief of burning, irritation, and discomfort due to dryness of the eye or exposure to wind or sun
- may be used as a protectant against further irritation

# Warnings

#### For external use only

#### Do not use

if solution changes color or becomes cloudy

# When using this product

to avoid contamination

- do not touch tip of container to any surface
- do not reuse
- once opened, discard
- do not touch unit-dose tip to eye

# Stop use and ask a doctor if

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

#### Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away (1-800-222-1222).

#### Directions

- to open, twist and pull tab to remove
- instill 1 or 2 drops in the affected eye(s) as needed and discard container

### Other information

- store at 20-25°C (68-77°F)
- RETAIN THIS CARTON FOR FUTURE REFERENCE.

### Inactive ingredients

calcium chloride dihydrate, magnesium chloride hexahydrate, potassium chloride, sodium chloride, sodium lactate solution, water for injection. May also contain hydrochloric acid and/or sodium hydroxide to adjust pH.

### **Questions or comments?**

1-800-719-9260

# Package/Label Principal Display Panel

sunmark<sup>®</sup> COMPARE TO REFRESH PLUS<sup>®</sup> ACTIVE INGREDIENT lubricating plus Carboxymethylcellulose Sodium 0.5% Lubricant Eye Drops Moisturizing Relief Sterile PRESERVATIVE FREE Actual vial size 30 Sterile Single-Use Containers 0.01 FL OZ (0.4 mL) EACH



SUNMARK LUBRICA carboxymethylcellulose sodiu					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:49348-329	
Route of Administration	OPHTHALMIC				
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Active Ingredient/Active	Molety				
Ingredient Name			<b>Basis of Strength</b>		Strength
<b>CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM</b> (UNII: K6790BS311) (CARBOXYMETHYLCELLULOSE - UNII:05JZ17B19X)			CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM		0.5 g in 100 mL
Inactive Ingredients					
Ingredient Name			Strength		
CALCIUM CHLORIDE (UNII: M410D	6VV5M)				

# Item Code	Package Description	Marketing Start Date	Marketing End Date
Packaging			
HYDROCHLORIC ACI	<b>D</b> (UNII: QTT17582CB)		
SODIUM HYDROXIDE	(UNII: 55X04QC32I)		
WATER (UNII: 059QF0	KO0R)		
SODIUM LACTATE (U	NII: TU7HW0W0QT)		
	UNII: 451W47IQ8X)		
POTASSIUM CHLORI	DE (UNII: 660YQ98I10)		
MAGNESIUM CHLOR	<b>DE</b> (UNII: 02F3473H9O)		

1	5 in 1 POUCH					
	0.4 mL in 1 VIAL, SINGLE-USE; Type 0: Not a Combination Product					
Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
OTC monograph fir	al part349	06/06/2013				

6 in 1 CARTON

Revised: 11/2022

**1** NDC:49348-329-44

Strategic Sourcing Services LLC

06/06/2013