# **LEADER LUBRICANT EYE DROPS- carboxymethylcellulose sodium solution/** drops

**Cardinal Health** 

-----

## Leader Lubricant Eye Drops 70ct and 30ct (PLD)

## **Active ingredients**

Carboxymethylcellulose sodium 0.5%

## **Purpose**

Lubricant

#### Uses

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

## Warnings

For external use only

## Do not use this product if

solution changes color or becomes cloudy

## When using this product

- do not reuse
- once opened, discard
- to avoid contamination, do not touch tip of container to any surface
- 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 the reach of children.

If accidentally swallowed, get medical help or contact a Poison Control Center (1800-222-1222) immediately.

#### **Directions**

• to open, twist and pull tab to remove

- instill 1 or 2 drops in the affected eye(s) as needed and discard container
- if used for post-operative (e.g. LASIK) dryness and discomfort, follow your eye doctor's instructions

#### Other information

- store at 15°-25°C (59°-77°F)
- use only if single-use container is intact
- use before expiration date marked on container
- RETAIN THIS CARTON FOR FUTURE REFERENCE

## **Inactive ingredients**

calcium chloride, magnesium chloride, potassium chloride, purified water, sodium chloride and sodium lactate. May contain sodium hydroxide and/or hydrochloric acid to adjust pH.

#### Questions or comments?

Call 1-888-527-4276

Leader Lubricant Eye Drops Preservative-Free 70 ct



#### **LEADER LUBRICANT EYE DROPS**

carboxymethylcellulose sodium solution/ drops

carboxymethylcellulose sodium solution/ drops					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (S	Source)	NDC:70000	-0012
Route of Administration	OPHTHALMIC				
Active Ingredient/Active Moiety					
Ingredient Name			Basis of St	rength	Strength
CARBOXYMETHYLCELLULOSE SODIUM (UNII: K6790BS311) (CARBOXYMETHYLCELLULOSE - UNII:05 Z17B19X)  CARBOXYMETHYLCELLULOSE - UNII:05 Z17B19X)			CELLULOSE	0.5 g in 100 mL	

Inactive Ingredients				
Ingredient Name	Strength			
CALCIUM CHLORIDE (UNII: M4I0D6VV5M)				
WATER (UNII: 059QF0KO0R)				
SODIUM CHLORIDE (UNII: 451W47IQ8X)				
SODIUM LACTATE (UNII: TU7HW0W0QT)				
SODIUM HYDROXIDE (UNII: 55X04QC32I)				
HYDROCHLORIC ACID (UNII: QTT17582CB)				
MAGNESIUM CHLORIDE (UNII: 02F3473H9O)				
POTASSIUM CHLORIDE (UNII: 660YQ98I10)				

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:70000- 0012-1	70 in 1 BOX	07/13/2019	
1		0.4 mL in 1 VIAL, SINGLE-USE; Type 0: Not a Combination Product		
2	NDC:70000- 0012-2	30 in 1 BOX	07/13/2019	
2		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 Drug	M018	07/13/2019		

## Labeler - Cardinal Health (063997360)

# Registrant - KC Pharmaceuticals, Inc. (174450460)

Establishment				
Name	Address	ID/FEI	Business Operations	
KC Pharmaceuticals, Inc.		174450460	pack(70000-0012) , label(70000-0012)	

Establishment			
Name	Address	ID/FEI	Business Operations
Unimed		689852052	manufacture(70000-0012) , pack(70000-0012) , label(70000-0012)

Revised: 12/2023 Cardinal Health