

REFRESH RELIEVA PF XTRA- carboxymethylcellulose sodium and glycerin solution/ drops
Allergan, Inc.

REFRESH RELIEVA® PF XTRA
Drug Facts

Active ingredients

Carboxymethylcellulose sodium 0.5%
Glycerin 0.9%

Purpose

Eye lubricant
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.**
- **To avoid contamination, do not touch tip of container to any surface. Replace cap after using.**
- **If solution changes color or becomes cloudy, do not use.**

Stop use and ask a doctor if

you experience eye pain, changes in vision, continued redness or irritation of the eye, or if the condition 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.

Directions

- Prior to first use, please read the "Instructions For Use" inside this carton.
- Instill 1 or 2 drops in the affected eye(s) as needed.

Other information

- Use only if tape seals on top and bottom flaps are intact.
- Use before expiration date marked on container.
- Discard 90 days after opening.

- Store at 59°-77°F (15°-25°C).
- RETAIN THIS CARTON AND THE INSERT FOR FUTURE REFERENCE.

Inactive ingredients

Boric acid; calcium chloride dihydrate; erythritol; levocarnitine; magnesium chloride hexahydrate; potassium chloride; purified water; sodium borate decahydrate; sodium citrate dihydrate; sodium hyaluronate, and trehalose. May contain hydrochloric acid and/or sodium hydroxide (to adjust pH).

Questions or comments?

1.800.678.1605

refresheyedrops.com

v2.0DFL3782

PRINCIPAL DISPLAY PANEL

NDC 0023-3782-10

PRESERVATIVE-FREE

Refresh[®]

RELIEVA[®] ***PF***

Xtra

Lubricant Eye Drops

With

HydroCell[®]

Fast-Acting Relief

Hydrates & Protects

Dry, Sensitive Eyes

0.33 fl oz (10 mL) Sterile

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SILHOUETTE OF BOTTLE IS ACTUAL SIZE

Drug Facts (continued)

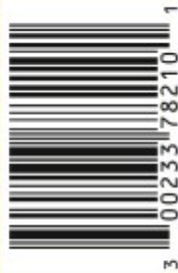
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Questions or comments?
 ☎ 1.800.678.1605
 refresheyedrops.com

PRESERVATIVE-FREE MULTIDOSE BOTTLE

BUILT-IN DOUBLE LOCKOUT SYSTEM KEEPS DROPS STERILE.




Distributed by:
 AbbVie, Inc.
 North Chicago, IL
 60064
 Product of US

PRESERVATIVE-FREE

Refresh RELIEVA PF Xtra

Lubricant Eye Drops

with **HydroCell®**



Fast-Acting Relief
Hydrates & Protects
Dry, Sensitive Eyes

0.33 fl oz (10 mL) Sterile

Refresh RELIEVA PF Xtra

Innovative preservative-free formula in a soft squeeze multidose bottle acts fast to relieve dry, burning, irritated eyes while providing a layer of protection that locks in hydration and prevents against further irritation.

REFRESH® RELIEVA® PF Xtra is the only lubricant eye drop enhanced with Trehalose, Sodium Hyaluronate, and HydroCell®—a proprietary NaCl-free, Glycerin-based technology.

TRUST YOUR EYES TO REFRESH®

The #1 Doctor-Recommended Preservative-Free Brand.*

abbvie

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REFRESH RELIEVA PF XTRA

carboxymethylcellulose sodium and glycerin solution/ drops

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:0023-3782
Route of Administration	OPHTHALMIC		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CARBOXYMETHYLCELLULOSE SODIUM (UNII: K679OBS311) (CARBOXYMETHYLCELLULOSE - UNII:05JZ17B19X)	CARBOXYMETHYLCELLULOSE SODIUM	5 mg in 1 mL
GLYCERIN (UNII: PDC6A3C0OX) (GLYCERIN - UNII:PDC6A3C0OX)	GLYCERIN	9 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
BORIC ACID (UNII: R57ZHV85D4)	
CALCIUM CHLORIDE (UNII: M410D6VV5M)	
ERYTHRITOL (UNII: RA96B954X6)	
LEVOCARNITINE (UNII: 0G389FZZ9M)	
MAGNESIUM CHLORIDE (UNII: 02F3473H9O)	
POTASSIUM CHLORIDE (UNII: 660YQ98I10)	

WATER (UNII: 059QF0KO0R)	
SODIUM BORATE (UNII: 91MBZ8H3QO)	
TRISODIUM CITRATE DIHYDRATE (UNII: B22547B95K)	
HYALURONATE SODIUM (UNII: YSE9PPT4TH)	
TREHALOSE (UNII: B8WCK70T7I)	
HYDROCHLORIC ACID (UNII: QTT17582CB)	
SODIUM HYDROXIDE (UNII: 55X04QC32I)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0023-3782-10	1 in 1 CARTON	03/01/2024	
1		10 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product		
2	NDC:0023-3782-35	1 in 1 CARTON	03/01/2024	
2		10 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product		
3	NDC:0023-3782-50	1 in 1 CARTON	03/01/2024	
3		3.5 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product		
4	NDC:0023-3782-20	2 in 1 CARTON	10/15/2025	
4		10 mL in 1 BOTTLE, DROPPER; 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	03/01/2024	

Labeler - Allergan, Inc. (144796497)

Revised: 9/2025

Allergan, Inc.