REDNESS RELIEVER AND LUBRICANT- dextran 70, polyethylene glycol 400, povidone, tetrahydrozoline hcl liquid Promex, 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.

Colirio Manzanill A+

Active Ingredient

Dextran 70 0.1%

Polyethylene Glycol 400 1%

Povidone 1%

Tetrahydrozoline HCI 0.5%

Purpose

Lubricant

Lubricant

Lubricant

Redness reliever

Use

- for the relief of redness of the eye due to minor eye irritations
- for use as a protectant against further irritation or to relieve dryness of the eye

Warnings

Ask a doctor before use if you have narrow angle glaucoma

When using this product

- pupils may become enlarged temporarily
- overuse may cause more eye redness
- remove contact lenses before using
- do not use if this solution changes color or become cloudy
- do not touch tip of container to any surface to avoid contamination
- replace cap after each use

Stop use and ask a doctor if

- you feel eye pain
- changes in vision occur

- redness or irritation of the eye lasts
- condition worsens or lasts more than 72 hours

If pregnant or breast-feeding,

ask a health professional before use.

Keep out of reach of children.

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

Directions

- Put 1 to 2 drops in the affected eye(s) up to 4 times daily
- children under 6 years of age: ask a doctor

Other information

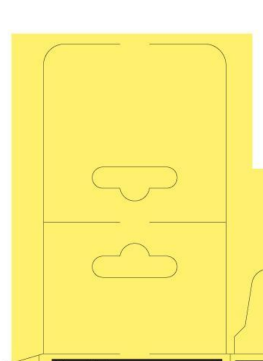
• store at 15-30 °C (59-86 °F)

Inactive ingredients

benzalkonium chloride, boric acid, edetate disodium, purified water, sodium borate, and sodium chloride

package label

Redness Reliever and Lubricant



- · Tamper Evident. Do not use this product if neckband is missing or broken.
- Prueba de Manipulación . No utilice este producto si la banda para el cuello falta o está roto.
- RETAIN THIS CARTON FOR FUTURE REFERENCE
- GUARDE ESTE ESTUCHE PARA FUTURAS REFERENCIAS

Drug Facts

Active ingredients Dextran 70 0.1%..... Purpose Lubricant Polyethylene Glycol 400 1%....Lubricant Povidone 1% ..Lubricant Tetrahydrozoline HCl 0.5%...Redness reliever

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Información del Medicamento

Ingredientes activos Dextrano 70 0 1% Lubricante Polietilenglical 400 1% Lubricante Povidona1% Lubricante Tetrahidrozolina HCI 0,5% .Alivia Enrojecimiento

Usos alivia el enrojecimiento de los ojos debido a irritaciones oculares menores e protector contra irrita ción adicionales o para aliviar la seguedad en los ojo Precauciones Consulte con un médico antes de usarlo si usted tiene glaucoma de ángulo estrecho Al utilizar este producto

pupilas pueden agrandarse temporalmente

Drug Facts(continued)

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Colirio





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Información del Medicamento (continuado)

- uso excesivo puede causar más enrojecimiento de los ojos quitar lentes de contacto antes
- No usar si la solución cambia. de color o se enturbie
- No toque la punta del envase con ninguna superficie para evitar la contaminación
- tapar después de cada uso

Pare el uso y pregúntele al doctor si misiente dolor en los ojos

- cambios en la visión ocurrer enrojecimiento o irritación de
- los ojos dura a condición empeora o dura más de 72 horas Si está embarazada o en periodo de lactancia, consulte a un profesional de la salud antes de usar.

Mantener fuera del alcance de los niños. En caso de ingestión, obtenga avuda médica o contactar con un centro de información toxicológica.

Indicaciones ■ Coloque 1 a 2 ootas en el oio afectado hasta 4 veces al día Menores de 6 años de edad: consulte un médico Otra información guarde entre

15-30°C (59-86 °F) Ingredientes inactivos cloruro benzalconio, ácido bórico, edetat disódico, agua, borato sodico, cloruro sodico

REDNESS RELIEVER AND LUBRICANT

dextran 70, polyethylene glycol 400, povidone, tetrahydrozoline hcl liquid

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:58988-0018
Route of Administration	OPHTHALMIC		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
TETRAHYDROZOLINE HYDROCHLORIDE (UNII: 0YZT43HS7D) (TETRAHYDROZOLINE - UNII:S9U025Y077)	TETRAHYDROZOLINE HYDROCHLORIDE	0.5 mg in 1 mL	
DEXTRAN 70 (UNII: 7SA290YK68) (DEXTRAN 70 - UNII:7SA290YK68)	DEXTRAN 70	1 mg in 1 mL	
POLYETHYLENE GLYCOL 400 (UNII: B697894SGQ) (POLYETHYLENE GLYCOL 400 - UNII: B697894SGQ)	POLYETHYLENE GLYCOL 400	10 mg in 1 mL	
POVIDONE (UNII: FZ989GH94E) (POVIDONE - UNII:FZ989GH94E)	POVIDONE	10 mg in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
BORIC ACID (UNII: R57ZHV85D4)			
SODIUM BORATE (UNII: 91MBZ8H3QO)			
EDETATE DISODIUM (UNII: 7FLD91C86K)			
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)			
SODIUM CHLORIDE (UNII: 451W47IQ8X)			
WATER (UNII: 059QF0KO0R)			

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:58988- 0018-1	1 in 1 BOX	12/06/2013		
1		15 mL in 1 BOTTLE; 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	12/06/2013		

Labeler - Promex, LLC (789974388)

Revised: 7/2022 Promex, LLC