EXCHANGE SELECT MAXIMUM REDNESS RELIEF EYE DROPS- glycerin, naphazoline hcl solution/ drops Army and Air Force Exchange Service

Exchange Select Maximum Redness Relief Eye Drops 15mL (PLD)

Active ingredients

Glycerin 0.5% Naphazoline HCI 0.03%

Purposes

Lubricant

Redness reliever

Uses

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

Warnings

For external use only

Do not use

if solution changes color or becomes cloudy

Ask a doctor before use if you have

narrow angle glaucoma

When using this product

- to avoid contamination, do not touch tip of container to any surface
- replace cap after using
- overuse may produce increased redness of the eye
- pupils may become enlarged temporarily

Stop use and ask a doctor if

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

Keep out of reach of children.

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

Directions

instill 1 to 2 drops in the affected eye(s) up to 4 times daily.

Other information

- store at room temperature
- remove contact lenses before using

Inactive ingredients

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

Questions or comments?

Call 1-888-527-4276

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EXCHANGE SELECT MAXIMUM REDNESS RELIEF EYE DROPS glycerin, naphazoline hcl solution/ drops							
Product Information							
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:5	IDC:55301-998		
Route of Administration	OPHTHALMIC						
Active Ingredient/Active Moiety							
Ingredient Name			Basis of Stre	ngth	Strength		
GLYCERIN (UNII: PDC6A3C0OX) (GLYCERIN - UNII:PDC6A3C0OX)			GLYCERIN		0.5 g in 100 mL		
NAPHAZOLINE HYDROCHLORIDE (UNII: MZ1131787D) (NAPHAZOLINE -			NAPHAZ OLINE		0.03 g		

Inactive Ingredients				
Ingredient Name	Strength			
BORIC ACID (UNII: R57ZHV85D4)				
EDETATE DISODIUM (UNII: 7FLD91C86K)				
WATER (UNII: 059QF0K00R)				
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)				
SODIUM BORATE (UNII: 91MBZ8H3QO)				

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:55301- 998-01	1 in 1 BOX	01/01/2024				
1		15 mL in 1 BOTTLE, DROPPER; Type 0: Not a Combination Product					
Marketing Information							
Marketing Application Number or Monog Category Citation		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
01	ΓC Monograph Dru	ug M018	01/01/2024				

Labeler - Army and Air Force Exchange Service (001695568)

Registrant - KC Pharmaceuticals, Inc. (174450460)

Establishment

Name	Address	ID/FEI	Business Operations
KC Pharmaceuticals, Inc.		174450460	manufacture(55301-998) , pack(55301-998) , label(55301-998)

Revised: 1/2024

Army and Air Force Exchange Service