SPERIAN EYESALINE EMERGENCY EYEWASH- purified water liquid Sperian Eye & Face Protection Inc

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.

Sperian Eyesaline Emergency Eyewash

Active ingredient

Purified Water 98.6%

Purpose

Emergency eyewash

Uses

• For flushing or irrigating the eye to reduce chances of severe injury caused by acid, alkali, or particulate contamination

Warnings

For external use only- Obtain immediate medical treatment for all open wounds in or near eyes. To avoid contamination, do not touch tip of container to any surface. Do not reuse. Once opened, discard.

Do not use

- if solution changes color or gets cloudy
- with contact lenses
- if seal is broken or tampered with
- if the eye is lacerated or object is embedded

Stop use and ask a doctor if you have

- changes in vision
- eye pain
- continued redness or irritation of the eye or if the condition worsens or persists

Keep out of reach of children

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

Directions

- remove contacts before using
- flush the affected area as needed
- if necessary, continue flushing with emergency eyewash or shower

Other information

- store at room temperature, 59 ° to 86 °F (15 ° to 30 °C)
- do not freeze

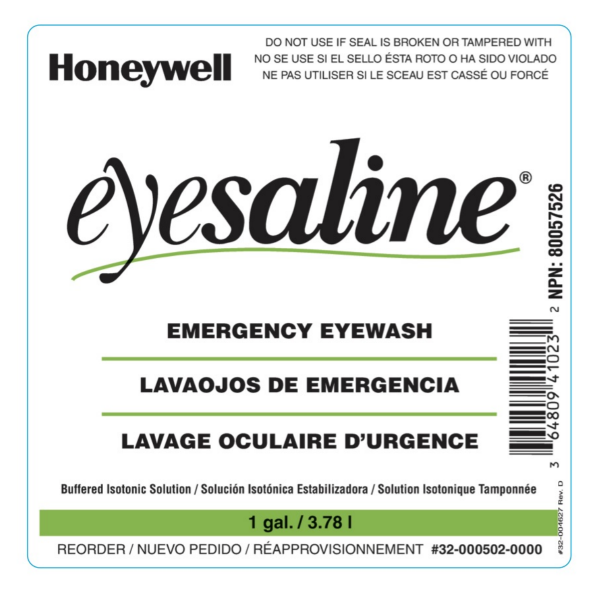
Inactive Ingredients

benzalkonium chloride, edatate disodium, sodium chloride, sodium phosphate dibasic, sodium phosphate monobasic

Questions?

Call **1-800-430-5490** Sperian Eye & Face Protection, Inc. (a Honeywell Company) 825 East Highway 151 Platteville, WI 53818 USA

Package label



Product T ype		HUMAN OTC DRUG	Item Co	n Code (Source) ND		NDC:6	DC:64809-100	
Route of Administ	ration	OPHTHALMIC						
Active Ingredie	nt/Active Moi	ety						
Ingredient Name				Basis of Strength			Strength	
WATER (UNII: 059QF0KO0R) (WATER - UNII:059QF0KO0R)				WATER		98.6 n	98.6 mL in 100 mL	
Inactive Ingred	ients							
Ingredient Name						Strength		
SODIUM CHLORIDE (UNII: 451W47IQ8X)								
SO DIUM PHO SPHATE, MO NO BASIC, MO NO HYDRATE (UNII: 593YO G76 RN)								
SO DIUM PHO SPHA	TE, MONOBASIC	C, MONOHYDRATE (UNII: 593)	YOG76RN)				
			YOG76 RN)				
SO DIUM PHO SPHA EDETATE DISO DIU	TE, DIBASIC (UN M (UNII: 7FLD91C	II: GR686LBA74) 86K)	YOG76RN)				
SO DIUM PHO SPHA EDETATE DISO DIU	TE, DIBASIC (UN M (UNII: 7FLD91C	II: GR686LBA74) 86K)	YOG76RN)				
SO DIUM PHO SPHA SO DIUM PHO SPHA EDETATE DISO DIU BENZALKO NIUM C	TE, DIBASIC (UN M (UNII: 7FLD91C	II: GR686LBA74) 86K)	YO G76 RN)				
SODIUM PHOSPHA EDETATE DISODIU BENZALKONIUM C	TE, DIBASIC (UN M (UNII: 7FLD91C	II: GR686LBA74) 86K)	YO G76 RN)				
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SODIUM PHOSPHA EDETATE DISODIU BENZALKONIUM C Packaging # Item Code	TE, DIBASIC (UN M (UNII: 7FLD91C HLORIDE (UNII: 1	II: GR686LBA74) 86K) F5UM2KM3W7)			-	rt N	-	
SODIUM PHO SPHA EDETATE DISODIU BENZALKONIUM C Packaging # Item Code 1 NDC:64809-100- 10	TE, DIBASIC (UN M (UNII: 7FLD91C HLORIDE (UNII: 1 14384 mL in 1 BA	II: GR686LBA74) 86K) F5UM2KM3W7) Package Description	Pro duct	0	Date	rt N	-	
SODIUM PHO SPHA EDETATE DISODIU BENZALKONIUM C Packaging I tem Code NDC:64809-100- 10 NDC:64809-100-	TE, DIBASIC (UN M (UNII: 7FLD91C HLORIDE (UNII: 1 14384 mL in 1 BA 3785 mL in 1 BO	II: GR686LBA74) 86K) F5UM2KM3W7) Package Description AG; Type 0: Not a Combination F	Pro duct	0	Date	rt N	-	
SODIUM PHO SPHA EDETATE DISODIU BENZALKONIUM C I Item Code NDC:64809-100- 10 NDC:64809-100- 11	TE, DIBASIC (UN M (UNII: 7FLD91C HLORIDE (UNII: 1 14384 mL in 1 BA 3785 mL in 1 BO Product	II: GR686LBA74) 86K) F5UM2KM3W7) Package Description AG; Type 0: Not a Combination F	Pro duct	0	Date	rt N	-	
SO DIUM PHO SPHA EDETATE DISODIU BENZALKO NIUM C Packaging I tem Code NDC:64809-100- 10 NDC:64809-100-	TE, DIBASIC (UN M (UNII: 7FLD91C HLORIDE (UNII: 1 14384 mL in 1 BA 3785 mL in 1 BO Product	II: GR686LBA74) 86K) F5UM2KM3W7) Package Description AG; Type 0: Not a Combination F	Product Combinatio	on 0	Date		-	

Labeler - Sperian Eye & Face Protection Inc (013435034)

Establishment								
Name	Address	ID/FEI	Business Operations					
Sperian Eye & Face Protection Inc		013435034	manufacture(64809-100)					

Revised: 3/2018

Sperian Eye & Face Protection Inc