ULINE EYEWASH- eyewash solution Uline.

ULINE Eyewash

Active ingredient

Purified water 98.3%

Purpose

Eyewash

Warnings

For external use only

Do not use

- if you experience any open wounds in or near the eyes and obtain immediate medical treatment
- if solution changes color or becomes cloudy

When using this product

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

Use

For cleansing the eye to help relieve irritation or burning by removing loose foreign material

Stop use and ask a doctor if you have any of the following

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

Keep out of reach of children

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

Directions

- remove tamper evident seal and cap
- flush the affected eye as needed, controlling the rate of flow of solution by presssure

Other information

- lot number is printed on the bottle
- store at 20° to 25° C [68° to 77° F]
- for your protection, this bottle has an imprinted white seal with black printing "TAMPER EVIDENT SEAL"
- do not use if this seal is missing or broken
- use before expiration date marked on bottle

Inactive ingredients

boric acid, sodium borate, sodium chloride

Questions?

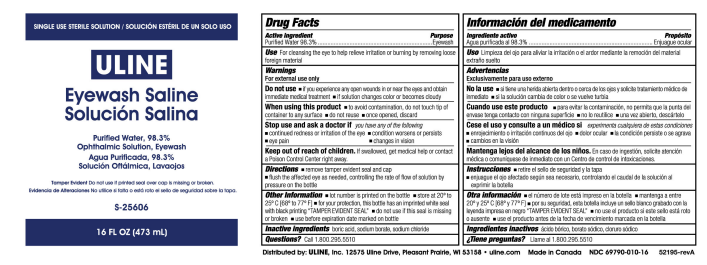
Tall 1.800.295.5510

Dosage & Administration

Remove Tamper evident seal and cap

flush the affected eye as needed, controlling the rate of flow of solution by pressure on the bottle

Label



ULINE EYEWASH eyewash solution						
Product Information	t Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69790-010			
Route of Administration	OPHTHALMIC					

Active Ingredient/Active Moiety										
		Ingredient Name	Basis of Stre	ngth	Strength					
W	ATER (UNII: 059QF0KO0R) (WATER - UNII:059QF0KO0R) WATER			465 mL in 473 mL						
In	active Ingr	edients								
			Strength							
	DRIC ACID (UNII									
		(UNII: 91MBZ8H3QO)								
sc	DIUM CHLORI	DE (UNII: 451W47IQ8X)		SODIUM CHLORIDE (UNII: 451W47IQ8X)						
Pa	ackaging									
	ackaging Item Code	Package Description	Marketin Dat	-	Marketing End Date					
#		Package Description 473 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a Combination Product		-	-					
# 1	Item Code NDC:69790-	473 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a	Dat	-	-					
# 1	Item Code NDC:69790- 010-16 NDC:69790-	473 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a Combination Product 946 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a	Dat 04/01/2024	-	-					
# 1 2	Item Code NDC:69790- 010-16 NDC:69790- 010-32	473 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a Combination Product 946 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a	Dat 04/01/2024	-	-					
# 1 2	Item Code NDC:69790- 010-16 NDC:69790- 010-32	473 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a Combination Product 946 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a Combination Product	04/01/2024 04/01/2024	e	-					

Labeler - Uline. (039612668)

Establishment

Name	Address	ID/FEI	Business Operations
Niagara Pharmaceuticals, Inc.		205477792	manufacture(69790-010)

Revised: 3/2024

Uline.