4386 FIRST AID KIT- 4386 first aid Honeywell Safety Products USA, Inc.

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

4386 First Aid Kit (PVP wipes, NaCl irr, EW, BZK wipe, alcohol wipe, PAWS- Z346100)

Eyewash Active ingredient

Sterile Water 99%

Eyewash

Purpose

Eyewash

Eyewash

Uses

• for flushing the eye to remove loose foreign material, air pollutants or chlorinated water

Eyewash *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 becomes cloudy
- if you have open wounds in or near the eyes, get medical help right away.

Stop use and ask a doctor if

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

Keep out of reach of children

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

Eyewash

Directions

- remove contacts before using
- twist top to remove
- flush the affected area as needed

- control rate of flow by pressure on the bottle
- if necessary, continue flushing with emergency eyewash or shower

Eyewash Inactive ingredients

sodium chloride, sodium phosphate dibasic, sodium phosphate monobasic

Eyewash

Questions

Call 1-800-430-5490 Honeywell Safety Products USA, Inc. Smithfield, RI 02917

Isotonic Solution for Irrigation.

For Irrigation Only.

Not for Injection.

Description NaCL Irrigation

Each 100 mL contains: Sodium Chloride USP 0.9 g; Water for Injection USP qs

pH adjusted with Hydrochloric Acid NF pH: 5.0 (4.5–7.0) Calculated Osmolarity: 310 mOsmol/liter

Concentration of Electrolytes (mEq/liter): Sodium 154; Chloride 154 0.9% Sodium Chloride Irrigation USP is sterile, nonpyrogenic, isotonic and contains no bacteriostatic or antimicrobial agents.

The formula of the active ingredient is: Ingredient Molecular Formula Molecular Weight Sodium Chloride USP NaCl 58.44

The plastic container is a copolymer of ethylene and propylene formulated and developed for parenteral drugs. The copolymer contains no plasticizers and exhibits virtually no leachability. The plastic container is also virtually impermeable to vapor transmission and, therefore, requires no overwrap to maintain the proper drug concentration. The safety of the plastic container has been confirmed by biological evaluation procedures. The material passes Class VI testing as specified in the U.S. Pharmacopeia for Biological Tests — Plastic Containers. These tests have shown that the container is nontoxic and biologically inert.

The PIC[™] Container is PVC-free and DEHP-free.

Ingredient Molecular Formula Molecular Weight Sodium Chloride USP NaCl 58.44

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Pharmacopeia for Biological Tests — Plastic Containers. These tests have shown that the container is nontoxic and biologically inert.

The PIC[™] Container is PVC-free and DEHP-free.

Clinical Pharmacology NaCL Irrigant

0.9% Sodium Chloride Irrigation USP is utilized for a variety of clinical indications such as sterile irrigation of body cavities, tissues or wounds, indwelling urethral catheters, surgical drainage tubes, and for washing, rinsing or soaking surgical dressings, instruments and laboratory specimens. It also serves as a diluent or vehicle for drugs used for irrigation or other pharmaceutical preparations.

0.9% Sodium Chloride Irrigation USP provides an isotonic saline irrigation identical in composition with 0.9% Sodium Chloride Injection USP (normal saline).

Physiological irrigation solutions are considered generally compatible with living tissues and organs.

Sodium, the major cation of the extracellular fluid, functions primarily in the control of water distribution, fluid balance, and osmotic pressure of body fluids. Sodium is also associated with chloride and bicarbonate in the regulation of the acid-base equilibrium of body fluid.

Chloride, the major extracellular anion, closely follows the metabolism of sodium, and changes in the acid-base balance of the body are reflected by changes in the chloride concentration.

Indication and Usage NaCl Irrigant

0.9% Sodium Chloride Irrigation USP is indicated for all general irrigation, washing, rinsing and dilution purposes which permit use of a sterile, nonpyrogenic electrolyte solution.

Contraindications NaCl Irrigant

0.9% Sodium Chloride Irrigation USP is not for injection by usual parenteral routes.

An electrolyte solution should not be used for irrigation during electrosurgical procedures.

Warnings NaCl Irrigant

FOR IRRIGATION ONLY. NOT FOR INJECTION.

Irrigating fluids have been demonstrated to enter the systemic circulation in relatively large volumes; thus, irrigation solutions must be regarded as systemic drugs. Absorption of large amounts can cause fluid and/or solute overload resulting in dilution of serum electrolyte concentrations, overhydration, congested states or pulmonary edema.

The risk of dilutional states is inversely proportional to the electrolyte concentrations of the administered parenteral solutions. The risk of solute overload causing congested states with peripheral and pulmonary edema is directly proportional to the electrolyte concentration.

Do not warm above 150°F (66°C).

After opening container, its contents should be used promptly to minimize the possibility of bacterial growth or pyrogen formation.

Discard unused portion of irrigating solution since it contains no preservatives.

Precautions NaCl Irrigant

General

Use aseptic technique when preparing and administering sterile irrigation solutions.

Use only if solution is clear and container and seal are intact.

Do not use for irrigation that may result in absorption of large amounts of fluid into the blood.

Caution should be observed when the solution is used for continuous irrigation or allowed to "dwell" inside body cavities because of possible absorption into the blood stream and the production of circulatory overload.

When used for irrigation via appropriate irrigation equipment, the administration set should be attached promptly. Unused portions should be discarded and a fresh container of appropriate size used for the start up of each cycle or repeat procedure. For repeated irrigations of urethral catheters, a separate container should be used for each patient.

Laboratory Tests

Clinical evaluation and periodic laboratory determinations are necessary to monitor changes in fluid balance, electrolyte concentrations, and acid-base balance after prolonged irrigation, when fluid absorption is suspected, or whenever the condition of the patient warrants such evaluation.

Drug Interactions

Some additives may be incompatible. Consult with pharmacist. When introducing additives, use aseptic technique. Mix thoroughly.

Do not store.

Carcinogenesis, Mutagenesis, Impairment of Fertility

Studies with 0.9% Sodium Chloride Irrigation USP have not been performed to evaluate carcinogenic potential, mutagenic potential, or effects on fertility.

Pregnancy

Teratogenic Effects

Animal reproduction studies have not been conducted with 0.9% Sodium Chloride Irrigation USP. It is also not known whether 0.9% Sodium Chloride Irrigation USP can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. 0.9% Sodium Chloride Irrigation USP should be given to a pregnant woman only if clearly needed.

Labor and Delivery

Safety and effectiveness of 0.9% Sodium Chloride Irrigation USP during labor and delivery have not been established. Caution should be exercised, and the fluid balance, glucose and electrolyte concentrations, and acid-base balance, of both mother and fetus should be evaluated periodically or whenever warranted by the condition of the patient or fetus.

Nursing Mothers

It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when 0.9% Sodium Chloride Irrigation USP is administered to a nursing woman.

Pediatric Use

The safety and effectiveness of 0.9% Sodium Chloride Irrigation USP in pediatric patients have not been established. Its limited use in pediatric patients has been inadequate to fully define proper dosage and limitations for use.

Geriatric Use

Clinical studies of 0.9% Sodium Chloride Irrigation USP did not include a sufficient number of patients age 65 years and over to determine whether they respond differently from younger subjects. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy.

This drug is known to be substantially excreted by the kidney, and the risk of toxic reactions to this drug may be greater in patients with impaired renal function. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selection, and it may be useful to monitor renal function. Frequent laboratory determinations and clinical evaluations are recommended to monitor changes in blood glucose, electrolyte concentrations, and renal function.

Adverse Reactions

Possible adverse effects arising from the irrigation of body cavities, tissues, or indwelling catheters and tubes can be minimized when proper procedures are followed. Displaced catheters or drainage tubes can lead to irrigation or infiltration of unintended structures or cavities. Excessive volume or pressure during irrigation of closed cavities may cause undue distension or disruption of tissues. Accidental contamination from careless technique may transmit infection.

If an adverse reaction does occur, discontinue administration of the irrigant, evaluate the patient, institute appropriate therapeutic countermeasures, and save the remainder of the fluid for examination if deemed necessary.

Overdosage

In the event of overhydration or solute overload, reevaluate the patient's condition, and institute appropriate corrective treatment. Intravasular volume overload may respond to hemodialysis. See WARNINGS, PRECAUTIONS, and ADVERSE REACTIONS.

Dosage and Administration

As required for irrigation.

When used as a diluent, or vehicle for other drugs, the drug manufacturer's recommendations should be followed.

Some additives may be incompatible. Consult with pharmacist. When introducing additives, use aseptic techniques. Mix thoroughly. Do not store.

Solutions should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permits.

How Supplied

0.9% Sodium Chloride Irrigation USP is supplied sterile and nonpyrogenic in PIC[™] (Plastic Irrigation Container). The 1000 mL and 500 mL containers are packaged 16 per case, the 2000 mL containers are packaged 8 per case, and the 4000 mL containers are packaged 4 per case.

0.9% Sodium Chloride Irrigation USP

NDC Cat. No. REF SIZE

0264-2201-00 R5200-01 1000 mL 0264-2201-10 R5201-01 500 mL 0264-2201-50 R5205-01 2000 mL 0264-2201-70 R5207 ,,,,4000 mL

Exposure of pharmaceutical products to heat should be minimized. Avoid excessive heat. Protect from freezing. It is recommended that the product be stored at room temperature (25°C); however, brief exposure up to 40°C does not adversely affect the product.

Do not warm above 150°F (66°C).

SPL Unclassified Section

Rx only Revised: March 2009 PIC is a trademark of B. Braun Medical Inc.

DIRECTIONS FOR USE OF PIC[™] (PLASTIC IRRIGATION CONTAINER)

Not for injection.

Aseptic technique is required.

Caution – Before use, perform the following checks:

(a) Read the label. Ensure solution is the one ordered and is within the expiration date.

(b) Invert container and inspect the solution in good light for cloudiness, haze, or particulate matter; check the

container for leakage or damage. Any container which is suspect should not be used.

Use only if solution is clear and container and seal are intact

Single unit container. Discard unused portion.

Outer Closure Removal – Grasp the container with one hand and turn the breakaway ring counterclockwise with the other hand until slight resistance is felt. Then, twisting the container in the opposite direction, turn the breakaway ring sharply until the entire outer cap is loose and can be lifted off.

Figure 1

Connect the administration set through the sterile set port according to set instructions or remove screw cap and pour.

Figure 2

Do not warm above 150°F (66°C) to assure minimal bottle distortion. Keep bottles upright.

SPL Unclassified Section

B. Braun Medical Inc.

Irvine, CA 92614-5895 USA Made in USA

Y36-002-699

PVP Wipe *Active ingredient*

Povidone-iodine 10% (equivalent to 1% titratable iodine)

PVP Wipe

Purpose

First aid antiseptic

PVP Wipe

Uses

• first aid antiseptic to help prevent infection in minor cuts, scrapes and burns

PVP Wipe

Warnings

For external use only.

Ask a doctor before use if you have

- deep or puncture wounds
- animal bites
- serious burns

Stop use and ask a doctor if

- condition worsens or persists for more than 72 hours
- irritation and redness develops

Keep out of reach of children.

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

PVP Wipe

Directions

- clean the affected area
- apply1 to 3 times daily
- may be covered with a sterile bandage
- if bandaged, let dry first
- discard wipe after single use

PVP Wipe

Other information

- do not use on individuals who are allergic or sensitive to iodine
- store at controlled temperature 59-86°F (15-30°C)
- do not use if pouch is open or torn

PVP Wipes Inactive ingredients nonoxynol 9, water

PVP Wipe *Questions*

1-800-430-5490

PAWS

Active ingredient

Ethyl alcohol 66.5%

PAWS Purpose

Antiseptic

PAWS

Uses

• for handwashing to decrease bacteria on skin whenever soap and water is not readily available

PAWS

Warnings

For external use only

Flammable: keep away from fire or flame

Do not use in the eyes.

If this happens, rinse thoroughly with water.

Stop use and ask a doctor if

• irritation or redness develop and 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

PAWS

Directions

- wet hands and wrists thoroughly for 15 seconds and allow to air dry
- always reseal after use
- children under 6 years of age should be supervised when using this product

PAWS

Inactive ingredients

aloe vera, fragrance, purified water, triethanolamine

PAWS Questions

Alcohol Wipe Active ingredient

Isopropyl alcohol 70%

Alcohol Wipe

Purpose First aid antiseptic

Alcohol Wipe

Uses

first aid to help prevent infection in minor cuts, scrapes, and burns

Alcohol Wipe *Warnings*

For external use only

Flammable, keep away from fire and flame

Do not use

- in or near eyes
- over large areas of the body

Ask a doctor before use if you have

- deep or puncture wounds
- animal bites
- serious burns

When using this product

• do not use longer than 1 week unless directed by a doctor

Stop use and consult a doctor if

• condition persists or gets worse

Keep out of the reach of children

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

Alcohol Wipe

Directions

- clean the affected area
- may be covered with a sterile bandage
- apply wipe to affeted are 1 to 3 times daily
- discard wipe after single use

Alcohol Wipe Directions

- clean the affected area
- may be covered with a sterile bandage
- apply wipe to affeted are 1 to 3 times daily
- discard wipe after single use

Alcohol Wipe

Other information

- store at room temperature 15 o to 25 o C (59 o to 77 oF)
- do not use if packet is torn or opened

Alcohol ipe Inactive ingredient

water

Alcohol Wipe *Questions*

1-800-430-5490

4386

Z346100 Kit Contents

1 ALCOHOL PREP PADS 10P

1 PVP IODINE WIPES 10 PER

1 NITRILE GLOVES 2PR BBP

1 O/H TAPE ADHESIVE TRI-CUT

1 BK GZ 4.5"X4.1YD6PLY RL ST MSO

1 FIRST AID GUIDE ASHI

1 EMERGENCY SURVIVAL BLANKET

2 GAUZE CLEAN-WRAP BDGE N/S 2"

1 BLOODSTOPPER

2 ABD COMBINE PAD 5" X 9"

2 ABD PADS 8"X10" STERILE

1 SOD. CHLORIDE 0.9% 500ML EA

1 4OZ BFS EYEWASH TRILINGUAL BOTTLE

1 EMPTY BAG RED 8X8X6

1 LBL STOCK 6-3/8"X4"

1 LBL STOCK 4"X2-7/8"

1 LBL STOCK 3"x1-7/8"

1 BANDAGE PACK FOR KIT

1 ZIP LOCK BAG FOR KIT #3

1 SELF-ADH WRAP 3 X 5 YDS NORTH REV E

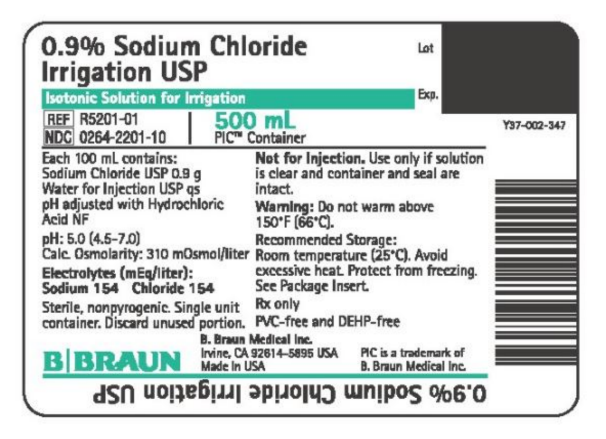
1 WATER-JEL BURN DRESSING 4 X 4 1 CORRUGATED 24PK 01-0810 RSC 1 TRI BNDG NON WOVEN 40"X40"X56" 1 COLD PACK UNIT 4"X6" BULK 1 CPR MSK,WPS,GLVS 1

Eyewash Principal Display Panel

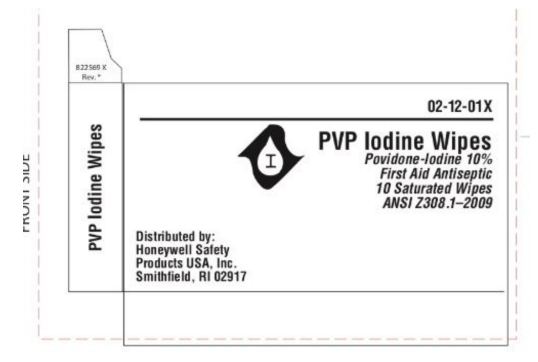


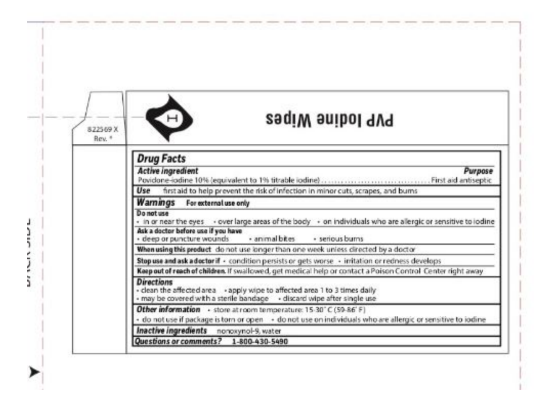


Principal Display Panel 500 ml Container



PVP Wipe Principal Display Panel





PAWS Principal Display Panel





Kills 99.99% of Germs!

- Contains 66.5% Ethyl Alcohol
 - Enriched with Aloe Vera

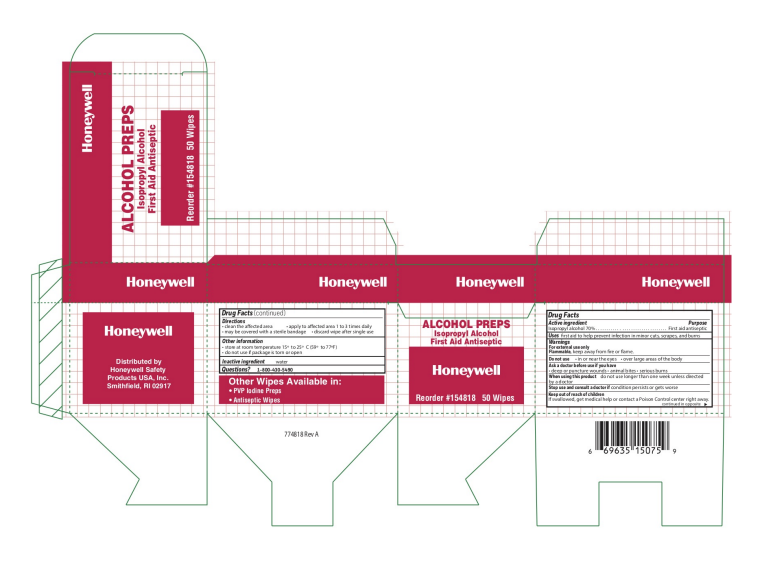
Fresh Scent

Drug Facts	
Active ingredients Ethyl Alcohol 66.5%	Purpose Antiseptic
Uses for handwashing to decrease bacteria on skin whenever soap and water is not re-	adily available
<i>Warnings</i> For external use only. Flammable, keep away from fire or flame	
Do not use in the eyes. If this happens, rinse thoroughly with water.	
Stop use and ask a doctor if irritation or redness develop and 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 wet hands and wrists thoroughly for 15 seconds and allow to air dry always reseal after use children under 6 years of age should be supervised when using this product 	
Inactive ingredients aloe vera, fragrance, purified water, triethanolamine	

SAFETEC OF AMERICA, Inc.

887 Kensington Ave. Buffalo, NY 14215 800-456-7077 www.safetec.com Reorder no. 34409 275 Gal. (1,041 L)

Alcohol Wipe Principal Display Panel



4386 Kit Label Z346100



Distributed by Honeywell Safety Products USA, Inc. Smithfield, RI 02917

438	6 FIRST AID	KIT		
4386	5 first aid kit			
Pro	duct Information	l		
Pro	duct T yp e	HUMAN OTC DRUG	Item Code (Source)	NDC:0498-4386
Pac	kaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NI	DC:0498-4386-01	1 in 1 KIT	09/13/2018	
Qua	ntity of Parts			

Part # Package Qu Part 1 2 PACKET	antity	Total Product Q	uantity	
FAILE Z FAGNET	0.0038 L	I otal I louact Q	auntity	
Part 2 10 POUCH	3 mL			
Part 3 1BOTTLE	118 mL			
Part 4 10 POUCH	4 mL			
Part 5 1 CONTAINER	500 mL			
Part 1 of 5				
PAWS				
ethyl alcohol liquid				
Product Information				
Item Code (Source)	NDC:0498-3111			
Route of Administration	TOPICAL			
Active Ingredient/Active Mo				
	edient Name	Basis of Stre		
ALCOHOL (UNII: 3K9958V90M) (AL	COHOL - UNII:3K9958 V90M)	ALCOHOL	665 mL in 1 L	
Inactive Ingredients				
	Ingredient Name		Strength	
WATER (UNII: 059QF0KO0R)				
TROLAMINE (UNII: 903K93S3TK)				
ALOE VERA LEAF (UNII: ZY81Z83H0	X)			
Packaging				
	ckage Description	Marketing Start Dat	e Marketing End Date	
1 0.0019 L in 1 PACKE	T; Type 0: Not a Combination Product			
Marketing Information				
Marketing Information				
	tion Number or Monograph Citation	on Marketing Start Date Marketing 12/21/2017		
		12/21/201/		
OTC monograph not final part333E				
OTC monograph not final part333E Part 2 of 5				
OTC monograph not final part333E Part 2 of 5 PVP IODINE WIPE				
OTC monograph not final part333E Part 2 of 5				

Product Informatio	11					
Item Code (Source)		NDC:0498-0121				
Route of Administration TOPICAL						
Active Ingredient/A						
	_	edient Name		Basis of Stre	ength	Strength
PO VIDO NE-IO DINE (UN	II: 85H0HZU95	9M) (IODINE - UNII:9679TC07X4)		IODINE		10 mg in 1 mL
Inactive Ingredient	S					
		Ingredient Name			5	Strength
NONOXYNOL-9 (UNII: 4						
WATER (UNII: 059QF0KC	D0R)					
Packaging						
# Item Code		Package Description	Marke	ting Start Dat	e Mar	keting End Da
1 NDC:0498-0121-00 0.3		CH; Type 0: Not a Combination Product		5		0
Marketing Category		n Number or Monograph Citation	Marke 09/18/20	e ting Start Date	Mar	keting End Dat
Marketing Category		n Number or Monograph Citation		-	Mar	keting End Dat
Marketing Infor Marketing Category unapproved drug other Part 3 of 5		n Number or Monograph Citation		-	Mar	keting End Dat
Marketing Category unapproved drug other	Applicatio			-	Mar	keting End Dat
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN	Applicatio			-	Mar	keting End Dat
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN	Applicatio /IERGEN			-	Mar	keting End Dat
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid	Applicatio /IERGEN			-	Mar	keting End Dat
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid Product Informatio	Applicatio /IERGEN	ICY EYEWASH		-	Mar	keting End Dat
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid Product Informatio Item Code (Source) Route of Administratio	Applicatio	NDC:0498-0100 OPHTHALMIC		-	Mar	keting End Dat
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid Product Informatio Item Code (Source) Route of Administratio	Applicatio /IERGEN	NDC:0498-0100 OPHTHALMIC	09/18/20	018	Mar	
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid Product Informatio Item Code (Source) Route of Administratio	Applicatio	ICY EYEWASH NDC:0498-0100 OPHTHALMIC ety nt Name	09/18/20	-		Strength
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid Product Informatio Item Code (Source) Route of Administratio	Applicatio	ICY EYEWASH NDC:0498-0100 OPHTHALMIC ety nt Name	09/18/20	018		
Marketing Category unapproved drug other Part 3 of 5 EYESALINE EN purified water liquid Product Informatio Item Code (Source)	Applicatio MERGEN In In Active Moie Ingredie DOR) (WATER	ICY EYEWASH NDC:0498-0100 OPHTHALMIC ety nt Name	09/18/20	018		Strength

	JNII: 451W47IQ	8 X)				
	лип. 4 010047 IQ					
Packaging						
# Item Code		Package Description	Marketing	g Start Date	Market	ting End Date
1 NDC:0498-0100-02	118 mL in 1 BO	TTLE; Type 0: Not a Combination Product	t			
Marketing Info	rmation					
Marketing Category	Applicatio	on Number or Monograph Citation	Marketing	Start Date	Market	ting End Date
OTC monograph final	part349		12/18/2018			
Part 4 of 5						
ALCOHOL WI	PE					
isopropyl alcohol sw	ab					
Product Informati	on					
Item Code (Source)		NDC:0498-0143				
Route of Administrat	ion	TOPICAL				
Active Ingredient/	Active Moi	ety				
		gredient Name		Basis of Str	rength	Strength
		gredient Name (416302) (ISOPROPYL ALCOHOL -		Basis of Str ISOPROPYL ALCOHOL	rength	Strength
ISOPROPYL ALCOHO				ISOPROPYL	rength	
ISOPROPYL ALCOHO UNII:ND2M416302)	L (UNII: ND2M			ISOPROPYL	rength	
ISOPROPYL ALCOHO UNII:ND2M416302)	L (UNII: ND2M			ISOPROPYL	rength Strenş	0.7 mL in 1 m
ISOPROPYL ALCOHO UNIEND2M416302) Inactive Ingredien	L (UNII: ND2M hts I	- 416302) (ISOPROPYL ALCOHOL -		ISOPROPYL		0.7 mL in 1 m
ISOPROPYL ALCOHO	L (UNII: ND2M hts I	- 416302) (ISOPROPYL ALCOHOL -		ISOPROPYL		0.7 mL in 1 m
ISOPROPYL ALCOHO UNIEND2M416302) Inactive Ingredien	L (UNII: ND2M hts I	- 416302) (ISOPROPYL ALCOHOL -		ISOPROPYL		0.7 mL in 1 m
ISOPROPYL ALCOHO UNII:ND2M416302) Inactive Ingredien WATER (UNII: 059QF0F Packaging	L (UNII: ND2M Its I KOOR)	- 416302) (ISOPROPYL ALCOHOL -		ISOPROPYL ALCOHOL	Strenş	0.7 mL in 1 m
ISOPROPYL ALCOHO UNII:ND2M416302) Inactive Ingredien WATER (UNII: 059QF0F Packaging # Item Code	L (UNII: ND2M hts I (OOR)	416302) (ISOPROPYL ALCOHOL - ngredient Name		ISOPROPYL ALCOHOL	Strenş	0.7 mL in 1 m
ISOPROPYL ALCOHO UNII:ND2M416302) Inactive Ingredien WATER (UNII: 059QF0F Packaging # Item Code	L (UNII: ND2M hts I (OOR)	416302) (ISOPROPYL ALCOHOL - ngredient Name Package Description		ISOPROPYL ALCOHOL	Strenş	0.7 mL in 1 m
ISOPROPYL ALCOHO UNII:ND2M416302) Inactive Ingredien WATER (UNII: 059QF0F Packaging # Item Code 1 NDC:0498-0143-04 (L (UNII: ND2M hts I (OOR)	416302) (ISOPROPYL ALCOHOL - ngredient Name Package Description		ISOPROPYL ALCOHOL	Strenş	0.7 mL in 1 m
ISOPROPYL ALCOHO UNII:ND2M416302) Inactive Ingredien WATER (UNII: 059QF0F Packaging # Item Code	L (UNII: ND2M hts I KOOR)).4 mL in 1 POU	416302) (ISOPROPYL ALCOHOL - ngredient Name Package Description	Marketing	ISOPROPYL ALCOHOL	Streng	0.7 mL in 1 m

Part 5 of 5

SODIUM CHLORIDE

sodium chloride irrigant

Product Informati	on				
Item Code (Source)	NDC:0264-2201				
oute of Administration IRRIGATION					
Active Ingredient/	Active Moiety				
	Ingredient Name			asis of crength	Strength
SODIUM CHLORIDE (U - UNII:Q32ZN48698)	DDIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37, CHLORIDE ION SODIUM				
Inactive Ingredien	its				
	Ingredient Name			Str	ength
WATER (UNII: 059QF0k	(OOR)				
HYDROCHLORIC ACIE) (UNII: QTT17582CB)				
Packaging					
# Item Code	Package Description	Marketing Start	Date	Marketi	ng End Date
1 500 mL	. in 1 CONTAINER; Type 0: Not a Combination Product				
Marketing Info	rmation				
Marketing Category	Application Number or Monograph Citation	Marketing Start	Date	Marketi	ng End Date
NDA	NDA016733	09/14/2009			-
Marketing Info	rmation				
Marketing Category	Application Number or Monograph Citation	Marketing Start	Date	Marketi	ng End Date
unapproved drug other		09/13/2018			

Labeler - Honeywell Safety Products USA, Inc. (079287321)

Registrant - Honeywell Safety Products USA, Inc. (079287321)

Establishment

Name	Addres	s	ID/	FEI		Business Operations
B. Braun Medical Inc.		0374	25308		label(0	264-2201)
Establishment						
Name		Addre	ess	ID/F	EI	Business Operations
Honeywell Safety Products USA, Inc				07928732	1	pack(0498-4386)
Establishment						
Name		Addre	ss	ID/FE	I	Business Operations
Honeywell Safety Products USA, Inc.				167518617		manufacture(0498-0100)
Establishment						
Name		Address		ID/FEI	Business Operations	
Changzhou Maokang Medical			4213	817073	manufacture(0498-0143)	
Establishment						
NT	Address		ID/F	EI	Business Operations	
Name	muncsa		10/1			-
Sion Medical Biotext	nuures	53277		r	nanufac	ture(0498-0121)
	nuics			I	nanufac	-
				I	nanufac	-
Sion Medical Biotext	Addre	53277	75194	r /FEI	nanufac	-

Revised: 10/2019

Honeywell Safety Products USA, Inc.