4151 FIRST AID KIT- 4151 first aid kit 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.

0498-4151: First Aid Kit (Eye Wash, Hand Sanitizer, bagged components-SF00004420)

Active ingredient

Sterile Water 99%

Purpose

Eyewash

Uses

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

Warnings

For external use only- Obtain immediate medical treatment for all open wounds in or near the 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 ey
- 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
- 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

Inactive Ingredients

sodium chloride, sodium phosphate dibasic, sodium phosphate monobasic

Questions?

Call **1-800-430-5490**

Honeywell Safety Products USA, Inc. Smithfield, RI 02917

First Aid Burn Cream Active ingredient

Benzalkonium chloride o.13%

Lidocaine HCl 0.5%

First Aid Burn Cream *Purpose*

First aid antiseptic

External analgesic

First Aid Burn Cream *Uses*

- prevent skin infection
- for temporary relief of pain associated with minor burns

First Aid Burn Cream *Warnings*

For external use only

Do not use

- in or near the eyes
- if you are allergic to any of the ingredients
- in large areas of the body, particularly over raw surfaces or blistered areas
- for more than 10 days

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
- symptoms persist for more than 7 days or clear up and occurs again within a few days

First Aid Burn Cream Directions

- adults and children 2 years of age and older:
- clean the affected area
- apply a small amount of this product (equal to the surface area of the tip of a finger) onto affected area 1 to 3 times daily
- may be covered with a sterile bandage
- children under 2 years of age: consult a doctor

Other information

- tamper evident sealed packets
- do not use if packet is opened or torn

Inactive ingredients

aloe barbadensis juice, cetyl alcohol, diazolidinyl urea, edetate disodium, glycerin, glyceryl stearate SE, methylparaben, mineral oil, PEG-100, propylene glycol, propylparaben, stearic acid, trolamine, water

Questions

1-800-430-5490

BZK Antiseptic Wipe Active ingredient

Benzalkonium chloride 0.13%

BZK Purpose

First aid antiseptic

BZK Uses

Antiseptic cleansing of face, hands, and body without soap and water

BZK Warnings

For external use only

BZK

Do not use

- in the eyes or over large areas of the body
- on mucous membranes
- on irritated skin
- in case of deep puncture wounds, animal bites or serious burns, consult a doctor
- longer than 1 week unless directed by a doctor

Stop use and ask a doctor if

- if irritation, redness or other symptoms develop
- the condition persists or gets worse

Keep out of reach of children

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

BZK

Directions

• .tear open packet and use as a washcloth

BZK

Other information

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

BZK Inactive ingredients

water

BZK Questions

1-800-430-5490

Aypanal Active igredient

Acetaminophen 325 mg

Aypanal *Purpose*

Aypanal Uses

- temporarily relieves minor aches and pains due to the common cold and headache
- temporarily reduces fever

Ask a doctor before use if you have

liver disease

Aypanal *Warnings*

Liver warning: This product contains acetaminophen. Severe liver damage may occur if you take: more than 4,000 mg in 24 hours, which is the maximum daily amount - child takes

Allergy alert: Acetaminophen may cause severe skin reactions. Symptoms may include:

skin reddening blisters rash

Do not use

 with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen, ask a doctor or pharmacist.

Ask a doctor before use if you have

liver disease

Ask a doctor or pharmacist before use if

you are taking the blood thinning drug warfarin

if preganat or breast feeding

ask a health professional before use

Keep out of rech of children

Keep out of reach of children

Overdose Warning

Overdose warning: In case of accidental overdose, get medical help or contact a Poison Control Center right away. Prompt medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

Aypanal *Directions*

- do not take more than directed (see overdose warning) adults and children 12 years of age or older
- take two tablets every 4-6 hours while symptoms last
- do not take more than directed (see overdose warning)

adults and children 12 years of age or older

- take two tablets every 4-6 hours while symptoms last
- do not take more than 12 tablets in 24 hours
- children 6 to under 12 years of age
- take 1 tablet every 4-6 hours while symptoms last
- do not take more than 5 tablets in 24 hours
- children under 6 years
- consult a doctor

Aypanal Other information

store at room temperature 15 ° to 30 ° C (59 ° - 86 °F) TAMPER EVIDENT PACKETS- DO NOT USE IF OPEN OR TORN

Aypanal Inactive ingredients

corn starch, microcrystalline cellulose, povidone, sodium starch glycolate, stearic acid.

Aypanal Questions or Comments?

1-800-430-5490

Sting Relief Active ingredient

Ethyl alcohol 50.0%

Lidocaine HCl 2.0%

Sting Relief *Purpose*

Antiseptic Topical pain relief

Sting Relief

Uses

• prevent infection in minor scrapes, and temporary relief of itching of insect bites

Sting Relief *Warnings*

For external use only

Flammable, keep away from open fire or flame

Do not use

- over large areas of the body
- in eyes
- over raw or blistered areas

Stop use and ask a doctor

• if conditions worsen or persist for more than 7 days or clear up and occur again within a few days

Keep out of reach of children

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

Sting Relief Directions

- adults and children 2 years and older: Apply to cleaned affected area not more than 3 times daily.
- children under 2 years of age: consult a doctor.

Sting Relief Inactive ingredients

benzalkonium chloride, menthol, and purified water

Sting relief *Questions or Comments*

1-800-430-5490

Neomycin Antibiotic Ointment Active ingredient

Neomycin sulfate (5 mg equivalent to 3.5 mg Neomycin base)

Neomycin Antibiotic Ointment *Purpose*

First aid antibiotic

Neomycin Antibiotic Ointment Uses

first aid to help prevent infection in - minor cuts - scrapes - burns

Neomycin Antibiotic Ointment *Warnings*

For external use only

Do not use

- in the eyes
- over large areas of the body

Ask a doctor before use if you have

- deep or puncture wounds
- animal bites
- serious burns

Stop use and ask a doctor if

- the condition persists or gets worse
- a rash or other allergic reaction develops
- you need to use longer than 1 week

Keep out of reach of children

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

Neomycin Antibiotic Ointment Directions

- clean the affected area
- apply a small amount of the product (an amount equal to the surface area of the tip of a finger) on the area 1 to 3 times daily
- may be covered with a sterile bandage

Neomycin Antibiotic Ointment *Other information*

store at 15 ° to 25 °C (59 ° to 77 °F)

Neomycin Antibiotic Ointment Inactive ingredient

petrolatum

Neomycin Antibiotic Ointment *Questions*

1-800-430-5490

Hand Sanitizer Active ingredient

Ethyl alcohol 62%

Hand Sanitizer *Purpose*

Antiseptic handwash

Hand Sanitizer *Uses*

- for hand washing to decrease bacteria on skin
- recommended for repeated use

Hand Sanitizer *Warnings*

For external use only

Flammable, keep away from fire or flame

When using this product

- do not use in the eyes
- discontinue use if irritation and redness develops
- If condition persists for more than 72 hours consult a doctor.

Keep out of reach of children

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

Hand Sanitizer Directions

• wet hands thoroughly with product and allow to dry without wiping

Hand Santitizer Other information

- place a quarter size amount into one hand, spread over both hands to wrist and rub into skin until dry
- store at 15 ° to 25 ° C (59 ° to 77 ° F)

Hand Sanitizer Inactive ingredients

acrylates/C10-30 alkyl acrylate crosspolymer, aloe barbadensis leaf juice, dl-alpha tocopheryl acetate, fragrance, PEG-60 almond glycerides, propylene glycol, purified water .

Hand Sanitizer *Questions or Comments*

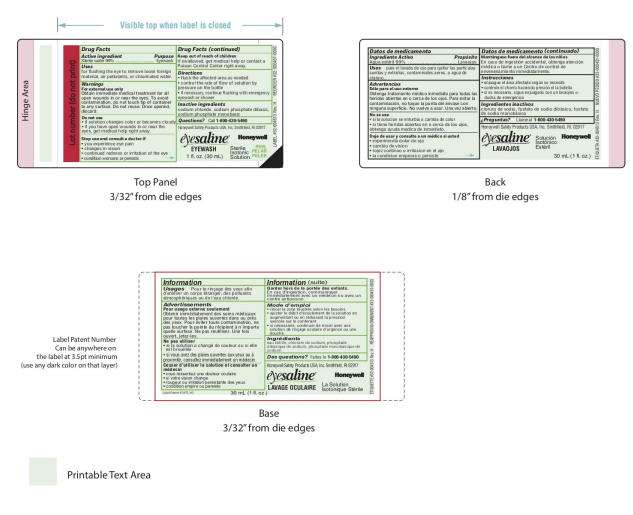
1-800-275-3433 info@waterjel.com

4151 SF00004420 Kit Contents

1 EYE DRESS PKT W/4 ADH STRIPS 1 TWEEZER PLASTICS 4" **1 FIRST AID GUIDE ASHI** 10 HAND SANITIZER 0.9G WJ BULK 2 GAUZE CLEAN-WRAP BDGE N/S 2" 1 GAUZE CLEAN-WRAP BDGE N/S 3" 2 ABD COMBINE PAD 5" X 9" 1 CPR FILTERSHIELD 77-100 **1 BAGGED COMP MISC** 11 OZ, BUFF EYEWASH 1 SCISSOR BDGE 4" RED PLS HDL LBL STOCK 6-3/8"X4" LBL STOCK 4"X2-7/8" 1 LBL STOCK 3"x1-7/8" 2 PR LRG NITRILE GLVES ZIP BAG 2 1" X 3" PLASTIC BANDS 16/BAG 2 TAPE ADHESIVE 1/2 X 2.5 125133 1 WATER-JEL BURN DRESSING 4 X 4 1 ADH BNDG PLASTIC EX-LG 4"X 2" 1 KIT, PP 16 UNIT FA 1 LBL CONTENTS ANSI 2015 CL A 1 TRI BNDG NON WOVEN 40"X40"X56" 1 COLD PACK UNIT 4"X6" BULK

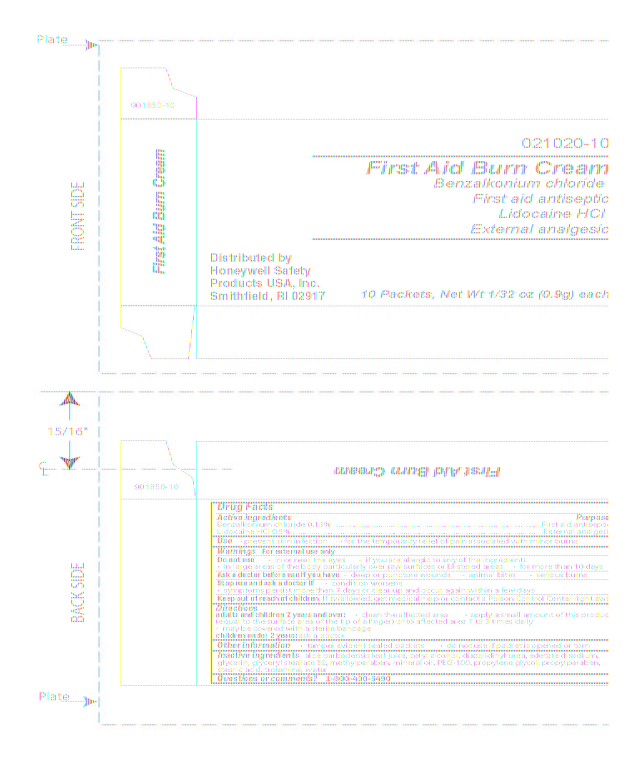
4 GAUZE PADS 3"X3" 12PLY 3 WOVEN FINGERTIP BANDAGE 2" 2 WOVEN KNUCKLE BANDAGE

Eye Wash Package label

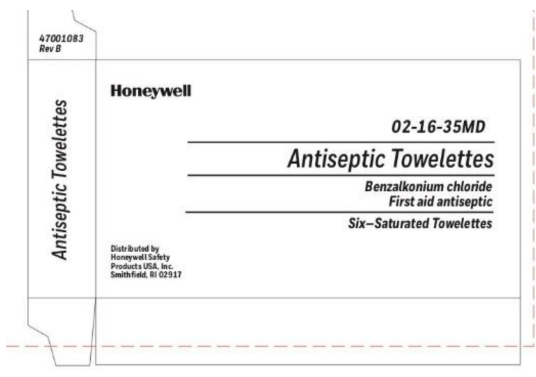


#32-004513 Rev. H

First Aid Burn Cream Principal Display Panel



Principal Display Panel

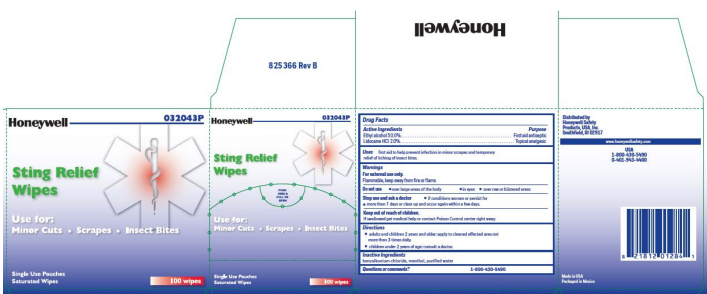


7001083 ev B	Antiseptic Towelettes
	Drug Facts
	Active Ingredient Purpose Benzalkonium chloride 0.133% w/v First ald antiseptic
	Uses • antiseptic cleaning of face, hands and body without soap and water. • air dries in seconds
	Warnings For external use only
	When using this product • do not use in the eyes or apply overlarge areas of the body
	Ask a doctor before use in case of deep or puncture wounds, animal bites, or serious burns
	Stop use and consult a doctor if irritation, mdness or other symptoms develop condition persists or gets worse
	Do not use I onger than 1 week unless directed by doctor
	Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.
	Directions • tear open packet, unfold and use as washcloth
	Other Information • store at room temperature 15° -30° C (59° -86° F) • do not reuse towe lotte
	•store at room temperature 15 - 50 C(58 - 66 F)

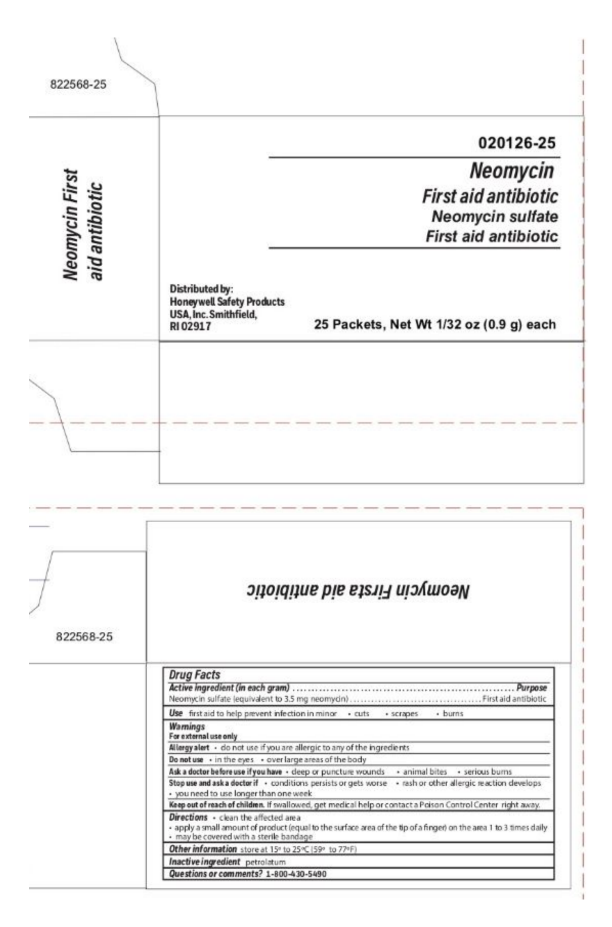
Aypanal Principal Display Panel



Sting Relief Principal Display Panel



Neomycin Antibiotic Ointment Principal Display Panel



Hand Sanitizer Principal Display Panel





4151 Kit Label SF00004420



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

4151 FIRST AID KIT

4151 first aid kit kit

Product Information

	uct Type	HUMAN	OTC DRUG	ltem Co	ode (Source)	NDC:0498-4151
Pack	aging					
# It	em Code	Рас	kage Description	ı	Marketing Start Date	Marketing End Date
1 NDC 01	2:0498-4151-	1 in 1 KIT; Ty Product	pe 0: Not a Combinat	ion	10/18/2018	
Quan	ntity of Pa	arts				
Part #	#	Package (Quantity		Total Product	Quantity
Part 1	1 BOTTLE			30 mL		
Part 2	3 PACKET			6		
Part 3	6 POUCH			2.4 mL		
Part 4	10 PACKET			9 g		
Part 5	10 PACKET			9 g		
Part 6	10 PACKET			14 mL		
Part 7	10 PACKET			13 mL		
Dart	t 1 of 7					
EYE	-	_	NCY EYEWA	SH		
EYE		_	NCY EYEWA	SH		
EYE S purifie	SALINE	uid	NCY EYEWA	SH		
EYE purifie Prod	SALINE ed water liqu	uid mation	NCY EYEWA	SH		
EYE: purifie Prod Item (SALINE ed water liqu	uid mation ce)		SH		
EYE purifie Prod Item (SALINE ed water liqu luct Inform Code (Sour	uid mation ce)	NDC:0498-0100	SH		
EYE purifie Prod Item (Route	SALINE ed water liqu luct Inform Code (Source e of Adminis	uid mation ce)	NDC:0498-0100 OPHTHALMIC	SH		
EYE purifie Prod Item (Route	SALINE ed water liqu luct Inform Code (Source e of Adminis	uid mation ce) stration	NDC:0498-0100 OPHTHALMIC Moiety	SH	Basis of Strength	Strength
EYE purifie Prod Item (Route Activ	SALINE ed water liqu luct Inform Code (Source e of Administry re Ingredie	uid mation ce) stration ent/Active Ingredien	NDC:0498-0100 OPHTHALMIC Moiety		Basis of Strength WATER	-
EYES purifie Prod Item (Route Activ	SALINE ed water liqu luct Inform Code (Source e of Administry re Ingredie	uid mation ce) stration ent/Active Ingredien	NDC:0498-0100 OPHTHALMIC Moiety t Name			-
EYE purifie Prod Item (Route Activ	SALINE ed water liqu luct Inform Code (Source e of Administry re Ingredie	uid mation ce) stration ent/Active Ingredien FOKOOR) (WATE	NDC:0498-0100 OPHTHALMIC Moiety t Name			Strength 98.6 mL in 100 mL

SODIUM CHLORIDE (UNII: 451W47IQ8X)

SODIUM PHOSPHATE, DIBASIC (UNII: GR686LBA74) SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE (UNII: 593YOG76RN)

Packaging

Item Code Package Description

1	NDC:0498-0100- 01	30 mL in 1 BOTTLE; Type 0: Not a Combinatio Product
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#			; Type 0: Not a (
	ltem Code	Pac	kage Descri	ption	Marketi Da	ng Start Ite		ting End ate
Pa	ackaging				N4	Chart	Marila	
	ontains							
	avor			Imprint Code			circle;U	
	nape	ROU	JND	Size			10mm	
	olor	whi		Score			2 pieces	
	roduct Charac							
-		,						
	DVIDONE (UNII: FZ 9		,U					
	ARCH, CORN (UNII			VII: AG9B05PV0B)				
	EARIC ACID (UNII: DDIUM STARCH GL							
C -		4511/22/25 42	Ingredier	nt Name			S	trength
In	active Ingred	ients						
_								
AC	CETAMINOPHEN (U	INII: 36209ITI	_9D) (ACETAMINO	PHEN - UNII:362C	9ITL9D)	ACETAMINOPH	IEN	325 mg
		-	edient Name			Basis of S		-
A	ctive Ingredie	nt/Active	Moiety					
Re	oute of Adminis	tration	ORAL					
lt	em Code (Sourc	e)	NDC:0498-2001	L				
Ρ	roduct Inform	ation						
	cetaminophen ta							
Δ	YPANAL NO		RIN					
Р	art 2 of 7							
σт	C Monograph Drug	M018	Citation		12/18/20		-	Juce
	Marketing Category	Applica	tion Number (Citatior	or Monograph		eting Start Date		eting End Date
	larketing Ir	nformat	ion					
N /								
•								

Market Catego		Applicat	tion Number or Monograph Citation	Marketing Start Date	Marketing End Date
inapproved d other	-			04/10/2012	
Part 3 c	of 7				
STING F	RELIEF	PAD			
ethyl alcoh	ol, lidoca	ine swab			
	_				
Product I	nforma	ition			
tem Code	(Source))	NDC:0498-0733		
Route of A	dministr	ation	TOPICAL		
Active Ing	redien	t/Active	Moiety		
	•	Ingredie	•	Basis of Streng	gth Strengt
JNII:98PI2009	IYDROCH	_	NII: V13007Z41A) (LIDOCAINE -	LIDOCAINE HYDROCHLOR ANHYDROUS	
ALCOHOL (U	NII: 3K995	8V90M) (ALC	COHOL - UNII:3K9958V90M)	ALCOHOL	0.5 mL in 1 mL
Inactive I	ngredie	ents			.
			Ingredient Name : F5UM2KM3W7)		Strength
MENTHOL (U					
NATER (UNII:					
Packagin	g				
# Item Code		Packa	age Description	Marketing Start Date	Marketing End Date
			; Type 0: Not a Combination		
1	Product	-			
1					
1					
	ng Inf	format	ion		
Marketi _{Market}	ing		ion tion Number or Monograph Citation	Marketing Start Date	Marketing End Date
Marketi	ing ory		tion Number or Monograph	-	Marketing End Date

Part 4 of 7

FIRST AID BURN

benzalkonium chloride, lidocaine hydrochloride cream

Pr	oduct In	formation					
lte	m Code (S	ource)	NDC:0498-0903				
Ro	ute of Adr	ninistration	TOPICAL				
٨	tive Ingr	edient/Active	Maiety				
AC	tive mgr		-		Decis of Ct		Cture in with
			dient Name		Basis of St	-	Strength
UN	I:7N6JUD5X6	Y)	: F5UM2KM3W7) (BENZALKONIUM	-	BENZ ALKONIUN CHLORIDE	4	0.13 g in 100 g
	OCAINE HY 1:98PI200987		NII: V13007Z41A) (LIDOCAINE -		LIDOCAINE HYDROCHLORIE	DE	0.5 g in 100 g
In	active Ing	gredients					
			Ingredient Name			S	trength
PR	OPYLENE GI	LYCOL (UNII: 6DC9	Q167V3)				
AL	DE VERA LE	AF (UNII: ZY81Z831	HOX)				
WA	TER (UNII: 0	59QF0KO0R)					
ST	EARIC ACID	(UNII: 4ELV7Z65AP					
ME	THYLPARAB	EN (UNII: A2I8C7HI	9Т)				
CE	TYL ALCOH	OL (UNII: 936JST6JC	N)				
GĽ	YCERYL MO	NOSTEARATE (UN	II: 2300U9XXE4)				
PE	G-100 STEA	RATE (UNII: YD01N	1999R)				
LIG	HT MINERA	LOIL (UNII: N6K578	37QVP)				
ED	ETATE DISO	DIUM (UNII: 7FLD9	1C86K)				
TR	DLAMINE (U	NII: 903K93S3TK)					
GĽ	YCERIN (UNI	I: PDC6A3C0OX)					
PR	OPYLPARAB	EN (UNII: Z8IX2SC	LOH)				
DI	ZOLIDINYL	UREA (UNII: H5RIZ	3MPW4)				
Pa	ckaging						
#	ltem Code	Packa	age Description		ting Start Date		eting End Date
1		0.9 g in 1 PACKET; Product	Type 0: Not a Combination				
Μ	arketin	g Informat	ion				
	Marketin Category	g Applica	tion Number or Monograpl Citation	n Mai	keting Start Date	Mar	keting End Date
	approved dru			12/20/			
oth	er			12/20/	2017		

Part 5 of 7						
NEOMYCIN						
antibiotic ointme	nt					
Product Inform	mation					
ltem Code (Sour	ce)	NDC:0498-0730				
Route of Adminis	stration	TOPICAL				
Active Ingredie	ent/Active	Moiety				
	Ingre	dient Name		Basis of Stre	ength	Strength
NEOMYCIN SULFAT	FE (UNII: 057Y6	526693) (NEOMYCIN - UNII:116QD7X	297)	NEOMYCIN SULFA	ATE	3.5 mg in 1 g
Inactive Ingree	dients					
		ngredient Name			Stre	ength
PETROLATUM (UNII	: 4T6H12BN9U)				
Packaging						
# Item Code	Pa	ckage Description	Mark	eting Start Date	Mar	keting End Date
	0.9 g in 1 PAC Product	KET; Type 0: Not a Combination				
Marketing I	nformat	ion				
Marketing Category	Applica	tion Number or Monograph Citation	Mai	rketing Start Date	Ma	rketing End Date
unapproved drug other			03/31/	2010		
Part 6 of 7						
	-	ETTE				
benzalkonium ch						
Product Inform	mation					
ltem Code (Sour	ce)	NDC:0498-0501				

Active Ingredien	t/Active	Moiety				
	Ingre	dient Name		Basis of Stre	ength	Strength
BENZALKONIUM CHL UNII: 7N6JUD5X6Y)	ORIDE (UNII	: F5UM2KM3W7) (BENZALKONIUM		BENZ ALKONIUM CHLORIDE		1.3 mg in 1 mL
Inactive Ingredi	ents					
j		redient Name			Streng	gth
WATER (UNII: 059QF0k	(00R)				-	
Packaging						
# Item Code	Packa	age Description		ing Start ate		eting End Date
1 1.4 mL Produc		F; Type 0: Not a Combination				
Marketing In	format	ion				
Marketing Category	Applicat	tion Number or Monograph Citation	Mark	eting Start Date	Mar	keting End Date
unapproved drug other			12/21/20	017		
Part 7 of 7						
INSTANT HAI		IITIZER				
alcohol liquid						
Product Informa	ation					
Item Code (Source)	NDC:59898-420				
Route of Administ	ration	TOPICAL				
Active Ingredien	t/Active	Moiety				
	Ingredie	nt Name	Basis	of Strength	9	Strength
ALCOHOL (UNII: 3K995	58V90M) (ALC	COHOL - UNII:3K9958V90M)	ALCOHO	L	62 m	L in 100 mL
Inactive Ingredie	ents					
		Ingredient Name				Strength
ALOE VERA LEAF (UNI	II: ZY81Z83H	10X)				

.ALPHATOCO				
TRIISOPROPA	NOLAMI	NE (UNII: W9EN9DLM98)		
CARBOMER C	OPOLYM	ER TYPE A (UNII: 71DD5V995L)		
WATER (UNII:	059QF0K0	DOR)		
PROPYLENE G	GLYCOL (UNII: 6DC9Q167V3)		
Packaging	I			
# Item Code		Package Description	Marketing Start Date	Marketing End Date
1	1.3 mL i Product	n 1 PACKET; Type 0: Not a Combination		
Marketir Marketin	•	ormation Application Number or Monograph	Marketing Start	Marketing End
Marketiı Categoı	ng ry		Marketing Start Date	Marketing End Date
Marketii	ng ry	Application Number or Monograph		-
Marketin Categor unapproved dr other	ng ry ^r ug	Application Number or Monograph Citation	Date	-
Marketin Categor unapproved dr other Marketin	ng ry ⁿ ug	Application Number or Monograph Citation	Date 04/15/2011	Date
Marketin Categor unapproved dr other Marketin Categor	ng ry ng Inf ng ry	Application Number or Monograph Citation	Date 04/15/2011	-
Marketin Categor unapproved dr other Marketin Marketin	ng ry ng Inf ng ry	Application Number or Monograph Citation Ormation Application Number or Monograph	Date 04/15/2011 Marketing Start	Date Marketing End

Labeler - Honeywell Safety Products USA, INC (118768815)

Revised: 1/2024

Honeywell Safety Products USA, INC