4250 FIRST AID KIT- 4250 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.

0498-4250: First Aid Kit (BZK wipes, FABC, triple,alcohol wipe, ASA-68P10SCHW)

First Aid Burn Cream Active ingredient

Benzalkonium chloride o.13%

Lidocaine HCI 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

First Aid Burn Cream *Other information*

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

First Aid Burn Cream 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

First Aid Burn Cream *Questions*

1-800-430-5490

Triple Active ingredient (each gram contains)

Bacitracin zinc 400 units - Neomycin sulfate (5 mg equivalent to 3.5 mg Neomycin base) Polymyxin B sulfate 5000 units

Triple *Purpose*

First aid antibiotic First aid antibiotic First aid antibiotic

Triple *Uses*

first aid to help prevent infection in

- minor cuts
- scrapes
- burns

Triple *Warnings*

For external use only

Allergy alert do not use if you are allergic to any of the ingredients

Do not use

- in the eyes
- over large areas of the body

Ask a doctor before use if you have

- a 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 the reach of children

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

Triple 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

Triple Other information

store at 15 0 to 25 0 C (59 0 to 77 0 F) tamper evident sealed packets - do not use if packet is torn or opened

Triple Inactive ingredient

petrolatum

Triple *Questions*

1-800-430-5490

Aspirin Active ingredient (in each tablet)

Aspirin 325 mg (NSAID)* *nonsteroidal anti-inflammatory drug

Aspirin *Purpo*se

Pain reliever/fever reducer

Aspirin *Uses*

temporarily reduces fever and relieves minor aches and pains associated with:

- a cold
- headache
- toothache
- muscular aches
- backache
- minor pain of arthritis
- premenstrual and menstrual periods

Aspirin *Warnings*

Reye's syndrome: Children and teenagers who have or are recovering from chicken pox or flu-like symptoms should not use this product. When using this product, if changes in behavior with nausea and vomiting occur, consult a doctor because these symptoms could be an early sign of Reye's syndrome, a rare but serious illness.

Allergy alert: Aspirin may cause a severe allergic reaction which may include:

- hives
- facial swelling
- asthma (wheezing)
- shock

Stomach bleeding warning: This product contains an NSAID, which may cause severe stomach bleeding. The chance is higher if you:are:

- age 60 or older
- have had stomach ulcers or bleeding problems
- take a blood thinning (anticoagulant) or steroid drug
- take other drugs containing prescription or nonprescription NSAIDs (aspirin,

ibuprofen, naproxen, or others)

- have 3 or more alcoholic drinks every day while using this product
- take more or for a longer time than directed

Do not use

• if you are allergic to aspirin or any other pain reliever/fever reducer

Ask a doctor before use if

- stomach bleeding warning applies to you
- you have a history of stomach problems such as heartburn
- you have high blood pressure, heart disease, liver cirrhosis or kidney disease
- you are taking a diuretic
- you have asthma

Ask a doctor or pharmacist before use if you are

• taking a prescription drug for diabetes, gout or arthritis

Stop use and ask a doctor if

- you experience any of the following signs of stomach bleeding:
- feel faint
- vomit blood
- have bloody or black stools
- have stomach pain that does not get better
- pain gets worse or lasts more than 10 days
- fever gets worse or lasts more than 3 days
- redness or swelling is present in the painful area
- ringing in the ears or loss of hearing occurs
- any new symptoms appear

If pregnant or breast-feeding,

If pregnant or breat-feeding, ask a health professional before use. It is especially important not to use aspirin during the last three months of pregnancy unless definitely directed to do so by a doctor because it may cause problems in the unborn child or complications during delivery.

Keep out of reach of children.

• In case of overdose, get medical help or contact Poison Control Center right away.

Aspirin

Directions

- drink a full glass of water with each dose
- adults and children 12 years of age and older: take 1 or 2 tablets every 4 hours while symptoms last, not more than 12 tablets in 24 hours
- children under 12 years of age: consult a doctor

Aspirin Other information

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

Aspirin Inactive ingredients

corn starch, croscarmellose sodium*, hypromellose*, microcrystalline cellulose*, mineral oil*, polyethylene glycol*, povidone, propylene glycol, silicon dioxide, stearic acid*, titanium dioxide*

*may contain these ingredients

Aspirin *Questions or Comments*

1-800-430-5490

BZK Active ingredient

Benzalkonium chloride 0.13% w/v

BZK Purpose

First aid antiseptic

BZK Uses

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

BZK Warnings

For external use only

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

Keep out of reach of children

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

Stop use and ask a doctor if

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

BZK

Directions

• tear open packet and use as a washcloth

BZK

Other information

- store at room temperature 15 0 to 30 0 C (59 0 86 0 F)
- do not reuse towelette

BZK Inactive ingredients

water

BZK Questions

1-800-430-5490

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 Wipes *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 Other information

- store at room temperature 15 0 to 25 0 C (59 0 to 77 0 F)
- do not use if packet is torn or opened

Alcohol Wipe Inactive ingredient

water

Alcohol Wipe *Questions*

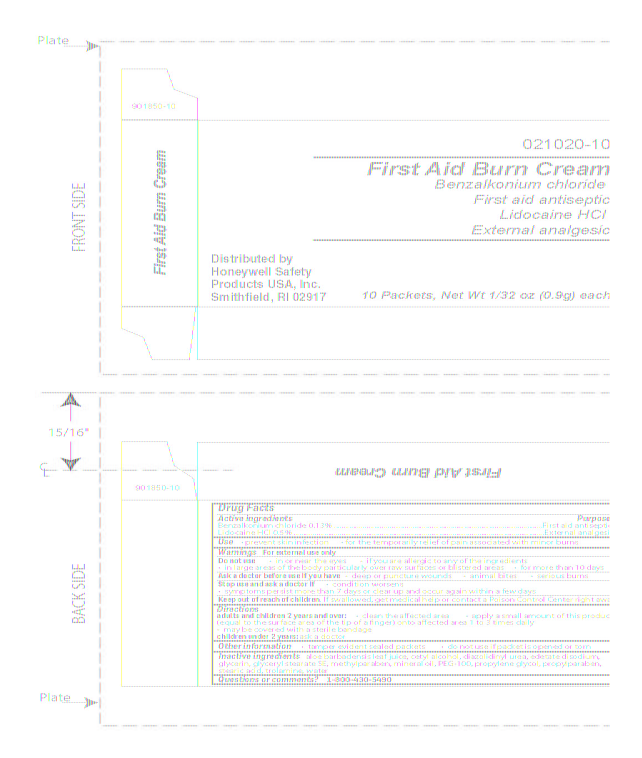
1-800-430-5490

4250 68P10SCHW Kit Contents

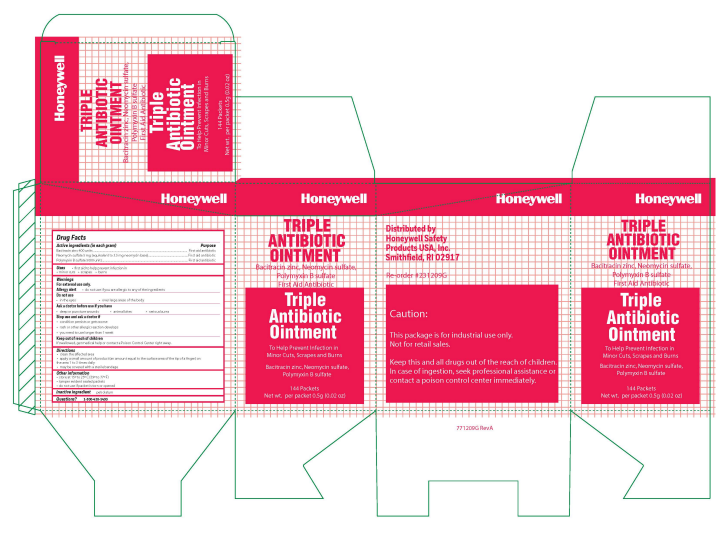
1 INSTANT COLD PACK 4" X 6"

1 ADH BDG, PLSTIC, 3/4"X3", 16 PER 1 TWEEZER PLASTICS 4" **1 FIRST AID GUIDE ASHI** 1 TAPE ADHESIVE 1"X 5 YD PLSTC 1 GAUZE CLEAN-WRAP BDGE N/S 2" 1 GAUZE CLEAN-WRAP BDGE N/S 3" 1 ABD COMBINE PAD 5" X 9" 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" 6 BZK ANTISEPTIC WIPE, BULK 2 PR LRG NITRILE GLVES ZIP BAG 6 FIRST AID CREAM 1.0GR PKT EACH **3 TRIPLE BIOTIC FOIL PACK EACH** 6 WIPE ALCOHOL PREP IPA 70% (DUKAL) 1 KIT, PP 10 UNIT FA 1 TRI BNDG NON WOVEN 40"X40"X56" 4 GAUZE PADS 2"X2" 12PLY **1 EYE PADS STD OVAL STERILE** 4 GAUZE PADS 3"X3" 12PLY 2 GAUZE PADS 4"X4" 12PLY **1 WOVEN FINGERTIP BANDAGE 2"** 1 HEAVY FLEX LARGE PATCH 2" X 3" 3 ASPIRIN BULK 2/PK

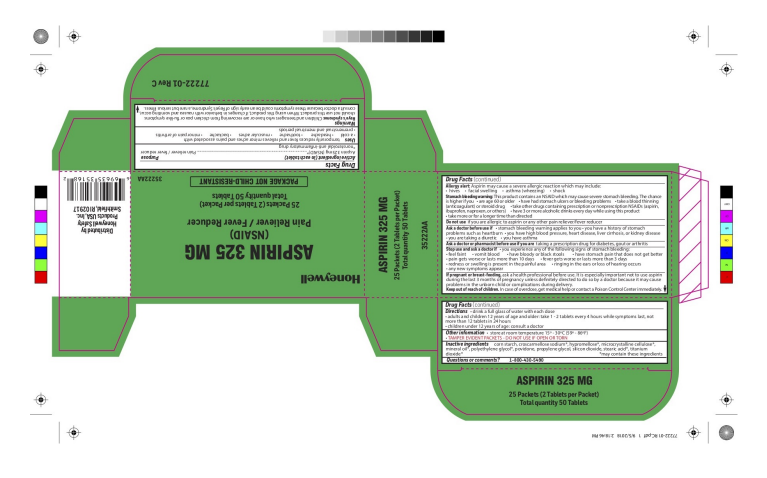
First Aid Burn Cream Principal Display Panel



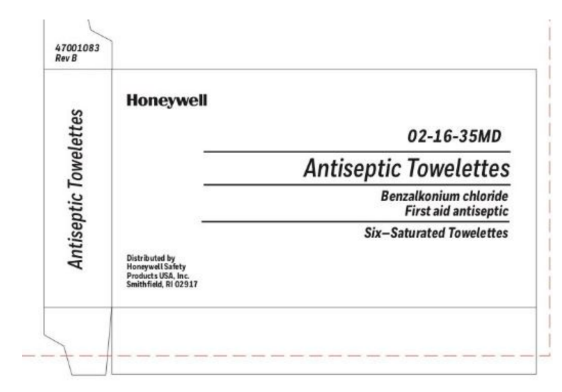
Principal Display Panel



Aspirin Principal Display Panel

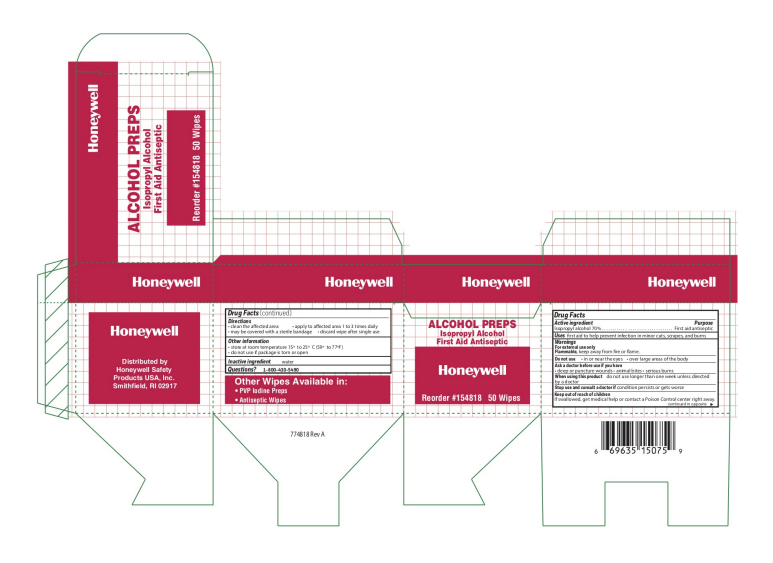






7001083 Rev B	səttələwoT oitqəsitnA
	Drug Facts
	Active Ingredient Purpose Benzalkonium chloride 0.133% w/v First ald antiseptic
	Uses antiseptic cleaning of face, hands and body without soap and water. air dries in seconds
	Wamings For external use only
	When using this product • do not use in the eyes or apply over large 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, redness or other symptoms develop condition persists or gets worse
	Do not use I longer 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 towelette
	Inactive ingredient water

Alcohol Wipe Principal Display Panel



4250 Kit Label 68FP10SCHW



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

Produ					
iiout	ict Inforr	nation			
Product Type HUMAN OTC DRUG			ltem Co	ode (Source)	NDC:0498-4250
Packa	ging				
# Ite	m Code	Package Des	scription	Marketing Start Date	Marketing End Date
1 NDC:	NDC:0498-4250- 011 in 1 KIT; Type 0: Not a Combinatio Product			10/18/2018	
Quant	tity of Pa	rts			
Part #		Package Quantity		Total Product	Quantity
Part 1	6 PACKET		5.4 g		
	6 PACKET		8.4 mL		
Part 2			6		
Part 2 Part 3	3 PACKET		2.7 g		
Part 2 Part 3	3 PACKET 3 PACKET 6 POUCH		2.7 g		

FIRST AID BURN

benzalkonium chloride, lidocaine hydrochloride cream

Dr/	duct In	formation				
Itei	n Code (S	ource)	NDC:0498-0903			
Rοι	ite of Adr	ninistration	TOPICAL			
۵ct	ive Inar	edient/Active	Moiety			
	ive mgi		dient Name	Basis of Str	rength Stren	nath
	ZALKONIU 7N6JUD5X6	M CHLORIDE (UNI	I: F5UM2KM3W7) (BENZALKONIUM ·		-	-
	DCAINE HY 98PI200987		NII: V13007Z41A) (LIDOCAINE -	LIDOCAINE HYDROCHLORID	0.5 g E in 100	g
	ative las					
ina		gredients				
TRA			Ingredient Name		Strengt	n
	•	NII: 903K93S3TK)				
	•	I: PDC6A3C0OX)	104)			
		EN (UNII: Z8IX2SC				
		UREA (UNII: H5RIZ				
		DIUM (UNII: 7FLD9				
		LYCOL (UNII: 6DC9				
		AF (UNII: ZY81Z83				
		59QF0KO0R)	N N			
		OL (UNII: 936JST6JC				
		NOSTEARATE (UN				
		RATE (UNII: YD01N				
LIGF	11 MINEKA	L OIL (UNII: N6K57	67QVP)			
Pa	ckaging					
#	ltem Code		age Description	Marketing Start Date	Marketing E Date	Ind
1		0.9 g in 1 PACKET; Product	Type 0: Not a Combination			
			_			
Ma	arketin	g Informat	ion			
	Marketin		tion Number or Monograph Citation	Marketing Start Date	Marketing Date	End
	Categor	-	Citation			
unap	oproved dru	-	Citation	12/20/2017		

Part 2 of 5						
ANTISEPTIC	TOWEL	ETTE				
benzalkonium chl	-					
Product Inform	nation					
ltem Code (Sourc	e)	NDC:0498-0501				
Route of Adminis	tration	TOPICAL				
Active Ingredie	nt/Active	Moietv				
<u> </u>		dient Name		Basis of Str	ength	Strength
	-	: F5UM2KM3W7) (BENZALKONIUM -		BENZALKONIUM	-	1.3 mg
UNII:7N6JUD5X6Y)				CHLORIDE		in 1 mL
Inactive Ingred	lients					
	-	redient Name			Streng	gth
WATER (UNII: 059QF	0KO0R)					
Packaging						
Packaging	Da	ckage Description	Mark	eting Start	Mark	ceting End
# Item Code		ckage Description	Mark	eting Start Date	Mark	ceting End Date
 # Item Code 1 NDC:0498-0501- 1 		ckage Description CKET; Type 0: Not a Combination	Mark		Mark	
 # Item Code 1 NDC:0498-0501- 1 	4 mL in 1 PA		Mark		Mark	
# Item Code 1 NDC:0498-0501- 00 1 F	4 mL in 1 PA Product	CKET; Type 0: Not a Combination	Mark		Mark	
# Item Code 1 NDC:0498-0501- 1 00 00 1 Marketing In 1	4 mL in 1 PA Product	CKET; Type 0: Not a Combination		Date		Date
# Item Code 1 NDC:0498-0501- 00 1 F	4 mL in 1 PA Product	CKET; Type 0: Not a Combination				
 # Item Code 1 NDC:0498-0501- 00 1 NDC:0498-0501- 00 1 Marketing Category Unapproved drug 	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph		Date keting Start Date		Date keting End
 # Item Code 1 NDC:0498-0501- 00 1 Marketing Category 	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
 # Item Code 1 NDC:0498-0501- 00 1 NDC:0498-0501- 00 1 Marketing Category Unapproved drug 	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
 # Item Code 1 NDC:0498-0501- 00 1 NDC:0498-0501- 00 1 Marketing Category Unapproved drug 	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 00 0 Item Code 1 Marketing Category Item Code Item Code Item Code Unapproved drug other Item Code Item Code Item Code Part 3 of 5 Item Code Item Code Item Code Item Code	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 00 0 1 Marketing Category 1 unapproved drug other 1	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 00 0 1 Marketing Category 1 unapproved drug other 1 Part 3 of 5 ASPIRIN	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 00 0 1 Marketing Category 1 unapproved drug other 1 Part 3 of 5 ASPIRIN	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 00 0 1 Marketing Category 1 unapproved drug other 1 Part 3 of 5 ASPIRIN	4 mL in 1 PA Product	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 10 00 1 Marketing Category 1 Unapproved drug other 1 Part 3 of 5 ASPIRIN as pirin tablet	nformat Applica	CKET; Type 0: Not a Combination ion tion Number or Monograph	Marl	Date keting Start Date		Date keting End
# Item Code 1 NDC:0498-0501- 1 00 Marketing 1 Marketing Category 1 Unapproved drug 00 1 Part 3 of 5 ASPIRIN 1 aspirin tablet 1 1	nformat Applica hation	CKET; Type 0: Not a Combination ion tion Number or Monograph Citation	Marl	Date keting Start Date		Date keting End

Active Ingredie		-				
		redient Nam		Basis of Stre	ngth	Strength
ASPIRIN (UNII: R16C	O5Y76E) (ASPIRIN - UNII:R	16CO5Y76E)	ASPIRIN		325 mg
Inactive Ingre	dients					
		Ingree	dient Name			Strength
CELLULOSE, MICR	OCRYSTA	LLINE (UNII: OP	1R32D61U)			
POLYETHYLENE GI	YCOL, U	NSPECIFIED (U	NII: 3WJQ0SDW1A)			
STEARIC ACID (UNI						
STARCH, CORN (UN						
POVIDONE (UNII: FZ						
SILICON DIOXIDE (
		-				
HYPROMELLOSE 2	-		DIQEDP/12K)			
MINERAL OIL (UNII:						
TITANIUM DIOXIDE		-				
PROPILENE GLICO		DC9Q107V3)				
Product Chara	cterist	ics				
Color		white	Score		2 pieces	
Shape		ROUND	Size		10mm	
Flavor			Imprint Code		FR21	
Contains						
Packaging						
				Marketing Start	Mar	keting End
# Item Code		Package De	escription	Date	Mai	Date
1 NDC:0498-0114-	2 in 1 PA	CKET; Type 0: N	ot a Combination			
01	Product					
Markating	nform	otion				
Marketing I				Maykating Ctart	N4-	ukating End
Marketing Category	Арр		ber or Monograph ation	Marketing Start Date	ма	rketing End Date
unapproved drug other				09/18/2018		
Part 4 of 5						
TRIPLE ANT	IBIOT	IC				

Product Information Item Code (Source) Route of Administration Active Ingredient/Active Ingre	NDC:0498-0750 TOPICAL					
Route of Administration Active Ingredient/Active Ingre	TOPICAL Moiety					
Ingre	-					
Ingre	-					
-	edient Name					
BACITRACIN ZINC (UNII: 89Y4M2				Basis o Strengt		Strength
	34ES) (BACITRACIN - U	UNII:58H6RWO52	21)	BACITRACIN		400 [iU] in 1 g
POLYMYXIN B SULFATE (UNII: 19	9371312D4) (POLYMY)	XIN B -		POLYMYXIN B		5000 [iU] in 1
UNII:J2VZ07J96K) NEOMYCIN SULFATE (UNII: 057Y	(626693) (NEOMYCIN -	- UNII:I16QD7X2	97)	NEOMYCIN		3.5 mg in 1 g
Inactive Ingredients						
I PETROLATUM (UNII: 4T6H12BN9U	ngredient Name				St	rength
Product Characteristics						
Color	white	Score				
Shape		Size				
Flavor Contains		Imprint Cod	le			
Packaging						
	ackage Description	on	Mark	eting Start	Ма	rketing End
NDC:0498-0750- 0.9 g in 1 PAC	• •			Date		Date
35 Product						
	.					
Marketing Informat Marketing Applica	LION ation Number or N	Ionograph	Mar	keting Start	Ma	rketing End
Category	Citation	ionograph	Fier	Date		Date
unapproved drug other			09/19/	2018		
Part 5 of 5						
ALCOHOL WIPE						
isopropyl alcohol swab						
Product Information						

Item Code (Sou	rce)	NDC:0498-0143			
Route of Admini	stration	TOPICAL			
Active Ingredi	ent/Active	Moiety			
	Ingr	edient Name		Basis Streng	Strendth
ISOPROPYL ALCO UNII:ND2M416302)	HOL (UNII: ND2)	M416302) (ISOPROPYL ALCOHOL	-	ISOPROPYL ALCOHOL	0.7 mL in 1 mL
Inactive Ingre	dients				
9		redient Name			Strength
WATER (UNII: 059Q	-				
Packaging					
	_		Market	ing Start	Marketing End
# Item Code	Ра	ckage Description		ate	Date
1 NDC:0498-0143- 04	0.4 mL in 1 PC Product	UCH; Type 0: Not a Combination			
Marketing	Informat	ion			
Marketing Category	Applica	tion Number or Monograp Citation		ting Start Date	Marketing End Date
unapproved drug other			09/18/203	18	
Marketing	Informat	ion			
Marketing Category	Applica	tion Number or Monograpl Citation		ting Start Date	Marketing End Date
unapproved drug			10/18/20	18	
other					

Labeler - Honeywell Safety Products USA, INC (118768815)

Revised: 1/2024

Honeywell Safety Products USA, INC