4152 FIRST AID KIT- 4152 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.

0498-4152: First Aid Kit (Triple, Burn Jel, BZK wipe, antiseptic hand gel, sting relief- SF00001774)

Burn Jel Active ingredient

Lidocaine HCl 2.0%

Burn Jel *Purpose*

External analgesic

Burn Jel

Uses

• temporarily relieves pain due to minor burns

Burn Jel *Warnings*

For external use only

Do not use

• on large areas of the body, particularly over raw surfaces or blistered areas

When using this product

• avoid contact with eyes

Stop use and ask a doctor if

- the condition gets worse
- symptoms persist for more than 7 days
- condition clears up and recurs within a few days

Keep out of reach of children

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

Burn JEI Directions

• adults and children 2 years of age and older; apply to affected area not more than 3

to 4 times daily

- children under 2 years of age: ask a doctor
- you may report a serious reaction to this product to 800-430-5490

Burn Jel Other information

• store at room temperature - do not use if opened or torn

Burn Jel Inactive ingredients

carbopol 940, carbopol 1342, diazolidinyl urea, glycerin, melaleuca alternifolia (tea tree) leaf oil, methylparaben, octoxynol-9, propylene glycol, propylparaben, trolamine, water

Burn Jel *Questions*

1-800-430-5490

Triple Active ingredient

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

BZK Wipe Active ingredient

Benzalkonium chloride 0.13% w/v

BZK Wipe Purpose

First aid antiseptic

BzK Wipe *Uses*

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

BZK Wipe *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

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 Wipe Directions

tear open packet and use as a washcloth

BZK Wipe Other information

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

BZK Wipe Inactive ingredient

water

BZK Wipe Questions

1-800-430-5490

Sting Relief Active ingredient (in each wipe)

Ethyl alcohol 50.0%

Lidocaine HCl 2.0%

Sting Relief *Purposse*

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

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 Sanitizer Other information

• store at 15 0 to 25 0 C (59 0 to 77 0 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, triisopropanolamine

Hand Sanitizer *Questions or Comments?*

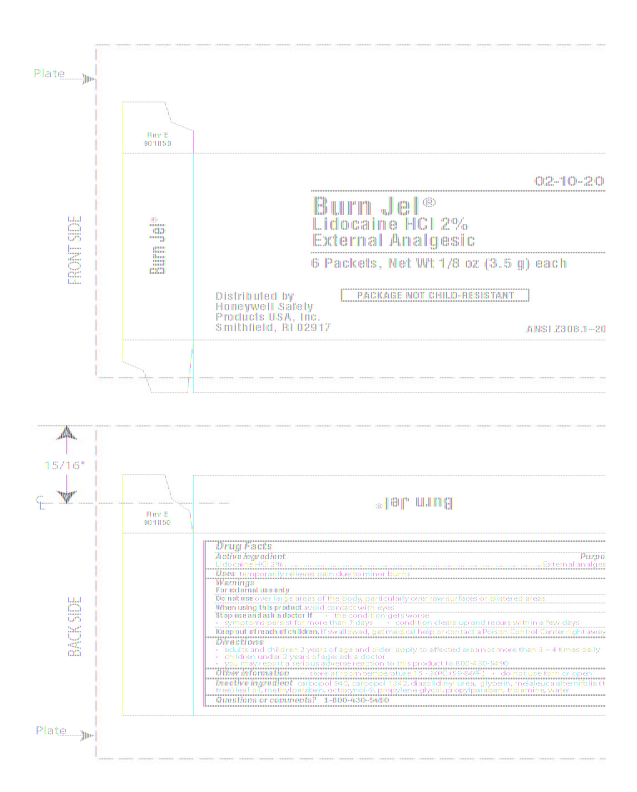
1-800-275-3433 info@waterjel.com www.waterjel.com

4152 SF00001774 Kit Contents

TRIPLE ANTIBIOTIC 10 PER
 GAUZE BANDAGE, 4" X 6 YD
 TRIANGULAR BDG, NON-STERILE
 GAUZE PADS, 3" X 3", 4 PER
 ADH TAPE, .5" X 2.5 YD, 2 PER
 FORCEPS & SCISSORS, 1 EA
 RESCUE BLANKET 1 PER
 BURN JEL 1/8 OZ, 6 PER
 ANTIMCRBL ANTSPTC TWLETTS
 ADH BDG, CLOTH, 1"X3", 16 PER
 FIRST AID GUIDE ASHI
 HAND SANITIZER 0.9G WJ 25/BX
 BANDAGE COMP 4" W/TELFA PAD 1
 LBL STOCK 6-3/8"X4"
 LBL STOCK 6-3/8"X4"

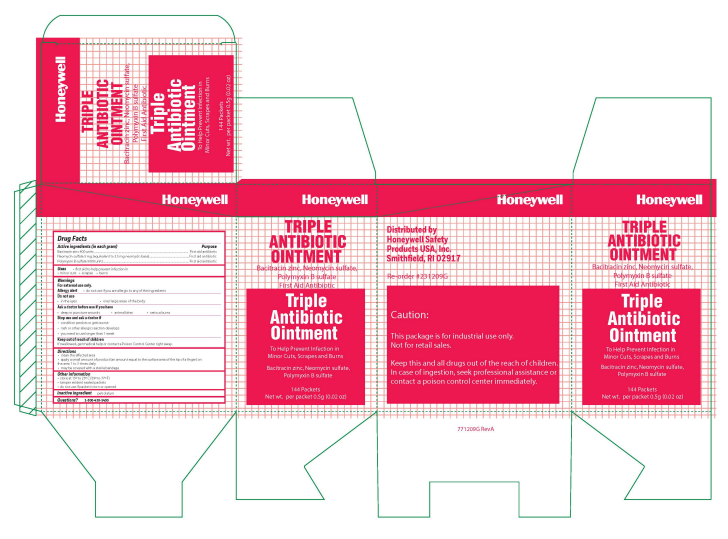
LBL STOCK 4"X2-7/8" 1 LBL STOCK 3"x1-7/8" 2 PR LRG NITRILE GLVES ZIP BAG 1 KIT STL 16 UN (VERTICAL) 1 COLD PACK UNIT 4"X6" BULK 2 ZIP LOCK BAG 5 X 8" 2 MIL 1 STING Relief WIPES 10

Burn Jel Principal Display Panel

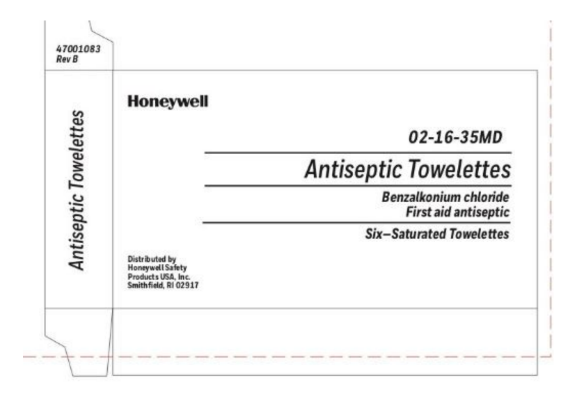


796353 Rev. E Unit Carton Printing Plate for "B" size cartor

Principal Display Panel

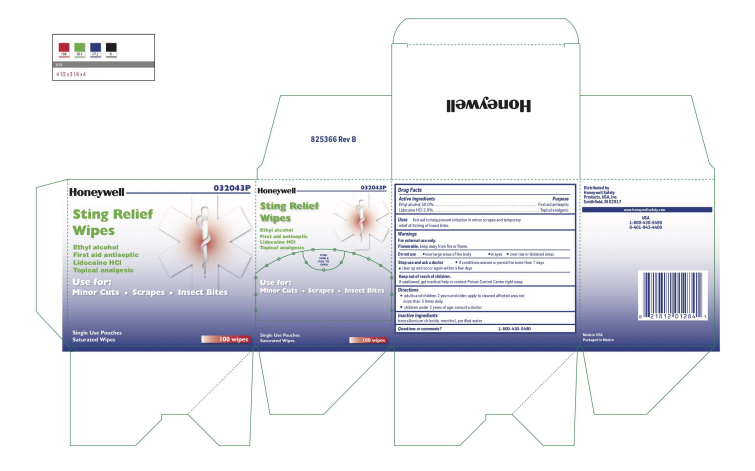


BZK Wipe Principal Display Panel



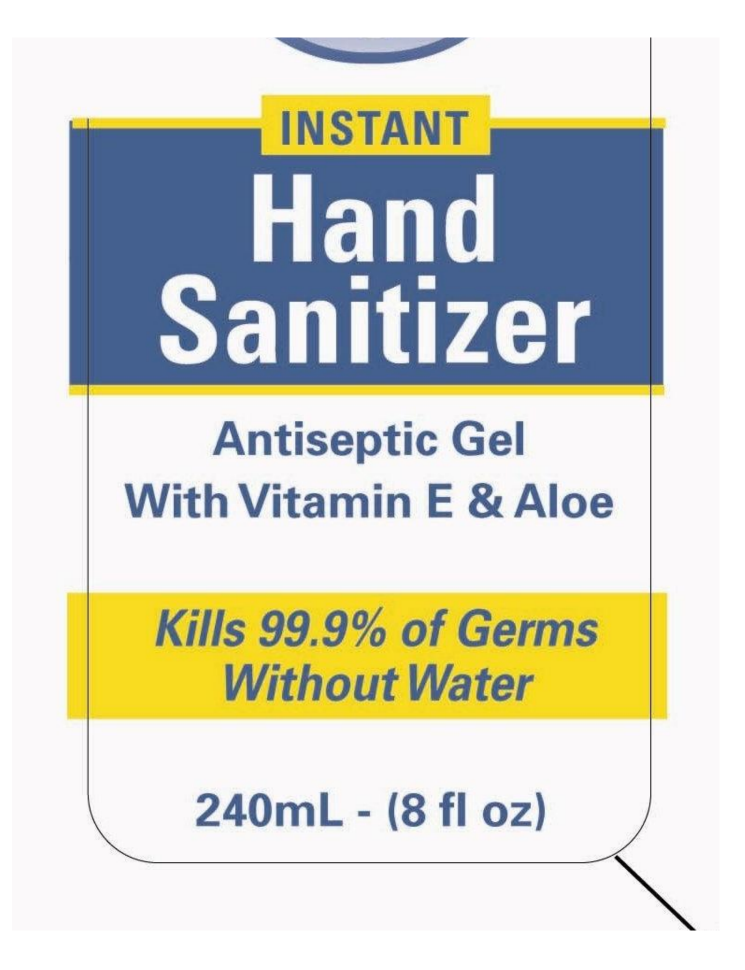
7001083 ev B	səttələwoT oitqəsitnA
	Drug Facts
	Active Ingredient Purpose Benzalkonium chloride 0.133% w/v First aid antiseptic
	Uses antiseptic cleaning of face, hands and body without soap and water. e air dries in seconds
	Warnings For external use only
	When using this product • do not use in the 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 towelatte
	•store at room temperature 15 - 50 ' G(35 - 86 ' F) • 40 interesting

Sting Relief Principal Display Panel



Hand Sanitizer Principal Display Panel





4152 Kit Label SF00001774



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

	FIRST AI	U KII		
4152 fir	st aid kit			
Produ	ct Informat	lion		
Produc	t Type	HUMAN OTC DRUG	ltem Code (Source)	NDC:0498-4152
Packa	ging			
# It	em Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:0	0498-4152-01	1 in 1 KIT	09/13/2018	
Quant	ity of Parts			
Part #	Pa	ckage Quantity	Total Produc	ct Quantity
Part 1	6 PACKET		21 g	
Part 2	10 PACKET		9 g	
i aic E	1 PACKET		1.4 mL	
Part 3	10 POUCH		4 mL	

Part 1 of 5					
BURN JEL					
gel for burns gel					
genor buills gel					
Product Inform	nation				
Item Code (Sourc	e)	NDC:0498-0203			
Route of Adminis	tration	TOPICAL			
Active Ingredie	ent/Active	Moiety			
J	Ingredie	•	Basis of Stre	nath	Strength
	-	III: V13007Z41A) (LIDOCAINE -	LIDOCAINE HYDROCHLO	•	2 g
UNII:98PI200987)			ANHYDROUS		in 100 g
Inactive Ingred	lients				
		Ingredient Name			Strength
TEA TREE OIL (UNII:	VIF565UC2G)				
DIAZOLIDINYL UREA					
METHYLPARABEN (U					
EDETATE DISODIUN		LC86K)			
GLYCERIN (UNII: PDC					
CARROMER HOMOS		E C (ALLYL PENTAERYTHRITOL			
4Q93RCW27E)		E B (ALLYL PENTAERYTHRITOL			
HHT01ZNK31)		(·····		
PROPYLPARABEN (L	JNII: Z8IX2SC1	OH)			
OCTOXYNOL-9 (UNI					
DIPROPYLENE GLYC	COL (UNII: E10	7L85C40)			
Packaging					
# Item Code	Pac	kage Description	Marketing Start Date	Marketi Da	-
	3.5 g in 1 PACH Product	ET; Type 0: Not a Combination			
Marketing I	nformat	ion			
Marketing Category		ion Number or Monograph Citation	Marketing Start Date		ing End ite
unapproved drug			09/19/2018		
other					

Part 2 of 5

TRIPLE ANTIBIOTIC

bacitracin zinc, polymyxin b sulfate, neomycin sulfate ointment

Product Inform	mation					
ltem Code (Sour	ce)	NDC:0498-0750				
Route of Adminis	stration	TOPICAL				
Active Ingredie	ent/Active I	Moiety				
	Ingred	lient Name			asis of trength	Strength
POLYMYXIN B SULI JNII:J2VZ 07J96K)	FATE (UNII: 193	71312D4) (POLYMY)	XIN B -	POLYMY	-	5000 [iU] in 1
BACITRACIN ZINC	(UNII: 89Y4M234	IES) (BACITRACIN - U	JNII:58H6RW052	2I) BACITRA	ACIN	400 [iU] in 1 g
NEOMYCIN SULFAT	FE (UNII: 057Y62	26693) (NEOMYCIN -	UNII:I16QD7X2	97) NEOMY	CIN	3.5 mg in 1 g
Inactive Ingre	dients					
		gredient Name			St	rength
PETROLATUM (UNII		-				-
Product Chara	cteristics					
Color		white	Score			
Shape			Size			
Flavor			Imprint Cod	е		
Contains						
Packaging						
# Item Code	Рас	kage Descriptio	on	Marketing S Date	itart Ma	rketing End Date
		ET; Type 0: Not a C	ombination	2410		2410
- 35	Product					
Marketing I	nformati	on				
Marketing Category	Applicat	ion Number or M Citation	lonograph	Marketing Date	Start Ma	arketing End Date
unapproved drug other				09/19/2018		
Part 3 of 5						

ANTISEPTIC						
benzalkonium ch	noriae liquia					
Product Infori	mation					
Item Code (Sour		NDC:0498-0501				
Route of Adminis		TOPICAL				
Active Ingredi	ent/Active	Moiety				
	Ingre	dient Name		Basis of Str	ength	Strength
BENZALKONIUM C UNII:7N6JUD5X6Y)	HLORIDE (UNI	I: F5UM2KM3W7) (BENZALKONIUM -		BENZ ALKONIUM CHLORIDE	I	1.3 mg in 1 mL
Inactivo Ingro	dianta					
Inactive Ingre		radiant Nama			Strong	•+h
WATER (UNII: 059QI	-	redient Name			Streng	yın
Packaging						
# Item Code	Ра	ckage Description	Mark	eting Start Date	Mark	ceting End Date
1 NDC:0498-0501- 00	1.4 mL in 1 PA Product	CKET; Type 0: Not a Combination				
Marketing I	nformat	ion				
Marketing Category	Applica	tion Number or Monograph Citation	Mar	keting Start Date	Mar	keting End Date
unapproved drug other			12/22/2	2017		
Part 4 of 5						
STING RELI	EF PAD					
ethyl alcohol, lido	ocaine swab					
Product Infor	mation					
ltem Code (Sour	ce)	NDC:0498-0733				
Route of Admini	stration	TOPICAL				
Active Ingradi	ont/Active	Mojoty				
Active Ingredie	ent/Active	molety				

IDOCAINE HYDRO		nt Name	Basis of Streng	th Strengt
UNII:98PI200987)	CHLORIDE (UI	NII: V13007Z41A) (LIDOCAINE -	LIDOCAINE HYDROCHLORIE ANHYDROUS	DE 20 mg in 1 mL
ALCOHOL (UNII: 3K9	958V90M) (ALC	COHOL - UNII:3K9958V90M)	ALCOHOL	0.5 mL in 1 mL
Inactive Ingred	dients			
		Ingredient Name		Strength
BENZALKONIUM CH		: F5UM2KM3W7)		
MENTHOL (UNII: L7T	-			
WATER (UNII: 059QF	UKUUK)			
Packaging				
# Item Code	Ра	ckage Description	Marketing Start Date	Marketing End Date
	0.4 mL in 1 PO Product	UCH; Type 0: Not a Combination		
Marketing I	nformat	ion		
Marketing Category	Applicat	tion Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other			12/23/2017	
Part 5 of 5				
INSTANT H	AND SAN	ITIZER		
-	AND SAN	ITIZER		
-	AND SAN	ITTIZER		
alcohol liquid		ITTIZER		
alcohol liquid Product Inform	nation			
alcohol liquid Product Inform Item Code (Sourc	mation ce)	NDC:59898-420		
INSTANT HA alcohol liquid Product Inform Item Code (Source Route of Adminis	mation ce)			
alcohol liquid Product Inform Item Code (Source Route of Adminis	mation ce) stration	NDC:59898-420 TOPICAL		
alcohol liquid Product Inform Item Code (Source Route of Adminis	mation ce) stration ent/Active	NDC:59898-420 TOPICAL Moiety	Basis of Strength	Strength
alcohol liquid Product Inform Item Code (Source Route of Adminis Active Ingredie	mation ce) stration ent/Active Ingredie	NDC:59898-420 TOPICAL	Basis of Strength ALCOHOL	Strength 62 mL in 100 mL
alcohol liquid Product Inform Item Code (Source Route of Adminis Active Ingredie	mation ce) stration ent/Active Ingredie	NDC:59898-420 TOPICAL Moiety nt Name		_
alcohol liquid Product Inform Item Code (Source Route of Adminis Active Ingredie	mation ce) stration ent/Active Ingredie 9958V90M) (ALC	NDC:59898-420 TOPICAL Moiety nt Name COHOL - UNII:3K9958V90M)		62 mL in 100 mL
alcohol liquid Product Inform Item Code (Source Route of Adminis Active Ingredie ALCOHOL (UNII: 3K9	mation ce) stration ent/Active Ingredie 9958V90M) (ALC dients	NDC:59898-420 TOPICAL Moiety nt Name COHOL - UNII:3K9958V90M)		-

FR	IISOPROPANOL	AMINE (UNII: W9EN9DLM98)		
		DLYMER TYPE A (UNII: 71DD5V995L)		
	ATER (UNII: 0590			
		· /		
Pa	ackaging			
#	ltem Code	Package Description	Marketing Start Date	Marketing Enc Date
	NDC:59898- 420-36	0.9 mL in 1 PACKAGE; Type 0: Not a Combination Product		
Μ	arketing	Information		
Μ	arketing Marketing Category	Information Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
	Marketing Category	Application Number or Monograph		Marketing End Date
una	Marketing Category	Application Number or Monograph	Date	
una oth	Marketing Category approved drug ler	Application Number or Monograph Citation	Date	
una oth	Marketing Category approved drug ler	Application Number or Monograph	Date	
una oth	Marketing Category approved drug ler	Application Number or Monograph Citation	Date	

Labeler - Honeywell Safety Products USA, Inc. (118768815)

Revised: 1/2024

Honeywell Safety Products USA, Inc.