4299 FIRST AID KIT- 4299 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.

0498-4299: First Aid Kit (Triple, Burn Jel, BZK wipe, sting relief- 011551-4340)

Burn Jel Active ingredient

Lidocaine HCI 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

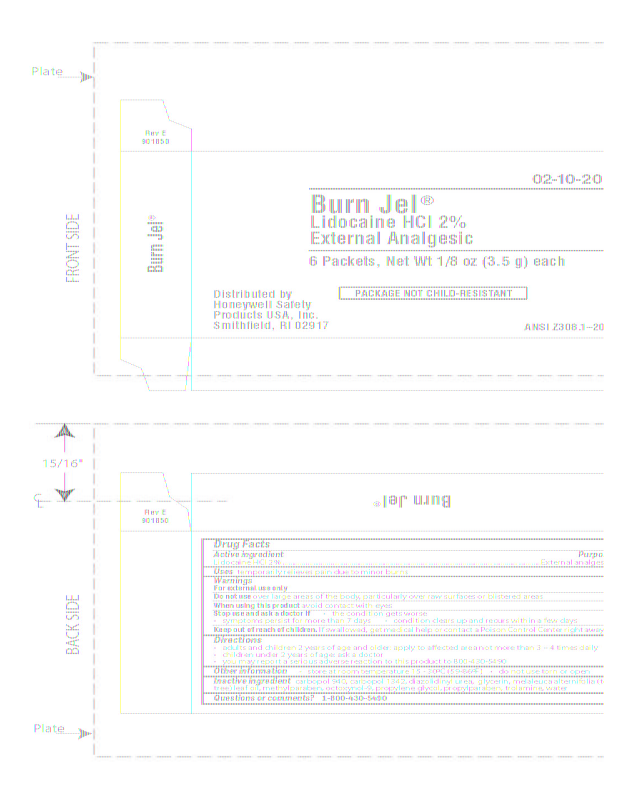
4299 011551-4340 kit contents

- 1 TRIPLE ANTIBIOTIC 10 PER
- 1 TRIANGULAR BDG, NON-STERILE
- 1 GAUZE PADS, 3" X 3", 4 PER
- 1 ADH TAPE, .5" X 2.5 YD, 2 PER
- 2 ADHESIVE BDG,PLSTIC,1"X3"16PER
- 1 FINGERTIP BANDAGE, 10 PER
- 1 BURN JEL 1/8 OZ, 6 PER
- 1 IVYX CLEANSER TOWEL 5 PER
- 1 NITRILE GLOVES 2PR BBP
- 1 ANTIMCRBL ANTSPTC TWLETTS
- 1 COMPRESS, 8" X 10", 1 PER
- 1 CPR MICROSHIELD DOUBLE UNIT

LBL STOCK 6-3/8"X4"

- 1 LBL STOCK 3"x1-7/8"
- 1 KIT STL 16 UN (VERTICAL)
- 1 STING Relief WIPES 10

Burn Jel *Principal Display Panel*



Principal Display Panel

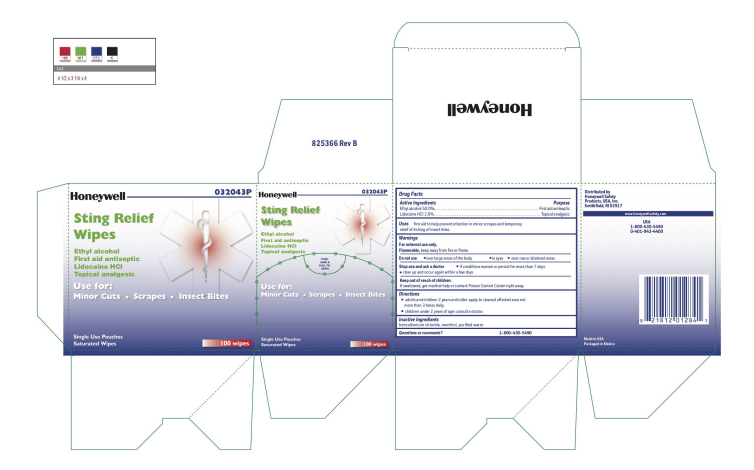


BZK Wipe Principal Display Panel

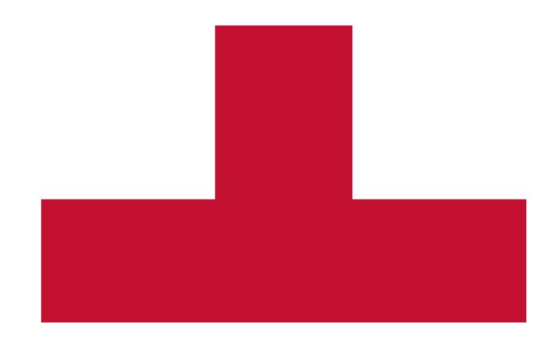
S	Honeywell	
lette		02-16-35MD
оме	-	Antiseptic Towelettes
Antiseptic Towelettes	A.	Benzalkonium chloride First aid antiseptic
tise		Six-Saturated Towelettes
An	Distributed by Honeywell Safety Products USA, Inc. Smithfield, RI 02917	

7001083 ev B	Antiseptic Towelettes
	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. • 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 ● 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
	Questions or comments 1-800-430-5490

Sting Relief Principal Display Panel



4299 Kit Label 011551-4340



FIRST AID KIT

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

4299 FIRST AID KIT

4299 first aid kit

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:0498-4299

Packaging

I	# Item Code	Package Description	Marketing Start Date	Marketing End Date
I	1 NDC:0498-4299-01	1 in 1 KIT	09/13/2018	

Quantity of Parts

4 44111	ny or runto	
Part #	Package Quantity	Total Product Quantity
Part 1	6 PACKET	21 g
Part 2	10 PACKET	9 g
Part 3	1 PACKET	1.4 mL

Part 1 of 4

BURN JEL

gel for burns gel

Product Information

Item Code (Source) NDC:0498-0203

Route of Administration TOPICAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
LIDOCAINE HYDROCHLORIDE (UNII: V13007Z41A) (LIDOCAINE - UNII:98PI200987)	LIDOCAINE HYDROCHLORIDE ANHYDROUS	2 g in 100 g		

Inactive Ingredients		
Ingredient Name	Strength	
TEA TREE OIL (UNII: VIF565UC2G)		
DIAZOLIDINYL UREA (UNII: H5RIZ3MPW4)		
METHYLPARABEN (UNII: A218C7HI9T)		
EDETATE DISODIUM (UNII: 7FLD91C86K)		
GLYCERIN (UNII: PDC6A3C0OX)		
TROLAMINE (UNII: 903K93S3TK)		
CARBOMER HOMOPOLYMER TYPE C (ALLYL PENTAERYTHRITOL CROSSLINKED) (UNII: 4Q93RCW27E)		
CARBOMER HOMOPOLYMER TYPE B (ALLYL PENTAERYTHRITOL CROSSLINKED) (UNII: HHT01Z NK31)		
PROPYLPARABEN (UNII: Z8IX2SC1OH)		
OCTOXYNOL-9 (UNII: 7JPC6Y25QS)		
DIPROPYLENE GLYCOL (UNII: E107L85C40)		

l	Packaging				
	# Item Code Package Description		Marketing Start Date	Marketing End Date	
	1	NDC:0498-0203- 00	3.5 g in 1 PACKET; Type 0: Not a Combination Product		

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
unapproved drug						

Part 2 of 4

TRIPLE ANTIBIOTIC

bacitracin zinc, polymyxin b sulfate, neomycin sulfate ointment

Product Information

Item Code (Source) NDC:0498-0750

Route of Administration TOPICAL

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
POLYMYXIN B SULFATE (UNII: 19371312D4) (POLYMYXIN B - UNII: J2VZ 07J96K)	POLYMYXIN B	5000 [iU] in 1 g			
BACITRACIN ZINC (UNII: 89Y4M234ES) (BACITRACIN - UNII:58H6RW052I)	BACITRACIN	400 [iU] in 1 g			
NEOMYCIN SULFATE (UNII: 057Y626693) (NEOMYCIN - UNII:116QD7X297)	NEOMYCIN	3.5 mg in 1 g			

Inactive Ingredients				
Ingredient Name	Strength			
PETROLATUM (UNII: 4T6H12BN9U)				

Product Characteristics			
Color white Score		Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

ı	Packaging					
	# Item Code Package Description		Package Description	Marketing Start Date	Marketing End Date	
	1	NDC:0498-0750- 35	0.9 g in 1 PACKET; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		09/19/2018		

Part 3 of 4

ANTISEPTIC TOWELETTE

benzalkonium chloride liquid

Product Information

Item Code (Source) NDC:0498-0501

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient NameBasis of StrengthStrengthDE (UNII: F5UM2KM3W7) (BENZALKONIUM -BENZALKONIUM1.3 mg

in 1 mL

BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)

BENZALKONIUM - BENZALKONIUM - CHLORIDE

Inactive Ingredients

Ingredient Name Strength

WATER (UNII: 059QF0KO0R)

Packaging

#	tem Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0498-0501- 00	1.4 mL in 1 PACKET; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug		12/22/2017		

Part 4 of 4

STING RELIEF PAD

ethyl alcohol, lidocaine swab

Product	Inform	ation
FIUUUCL		ativii

Item Code (Source) NDC:0498-0733

Route of Administration TOPICAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
LIDOCAINE HYDROCHLORIDE (UNII: V13007Z41A) (LIDOCAINE - UNII:98PI200987)	LIDOCAINE HYDROCHLORIDE ANHYDROUS	20 mg in 1 mL		
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	0.5 mL in 1 mL		

Inactive Ingredients			
Ingredient Name	Strength		
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)			
MENTHOL (UNII: L7T10EIP3A)			
WATER (UNII: 059QF0KO0R)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0498-0733- 00	0.4 mL in 1 POUCH; Type 0: Not a Combination Product		

Marketing Information				
Marketi Catego		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved dr other	ug		12/23/2017	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		09/13/2018		

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

Revised: 1/2024 Honeywell Safety Products USA, Inc.