4150 FIRST AID KIT- 4150 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.

0498-4150: First Aid Kit (BZK wipes, FABC, triple, alcohol wipe- SF00003260)

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

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

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*

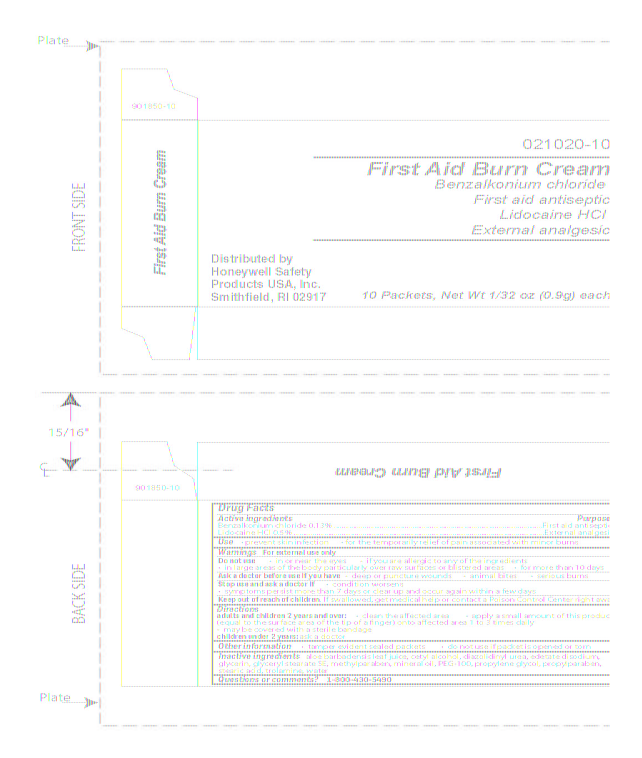
1-800-430-5490

4150 SF00003260 Kit Contents

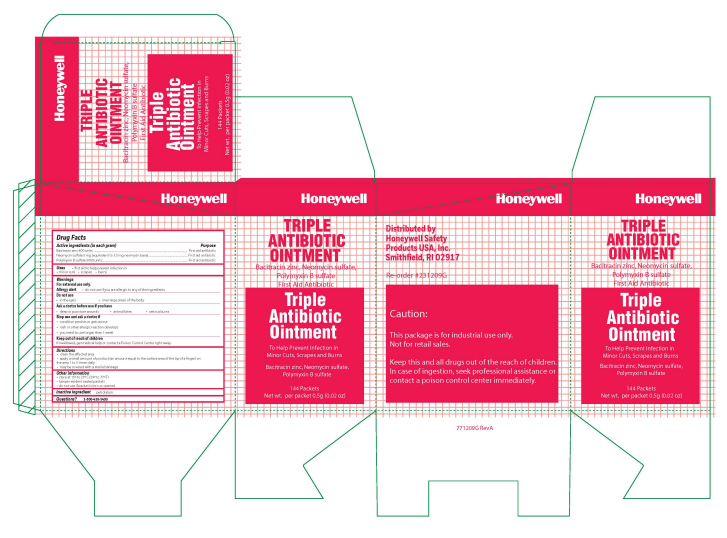
TRIPLE ANTIBIOTIC 10 PER
 FIRST AID BURN CREAM 6 PER
 TRIANGULAR BDG, NON-STERILE
 GAUZE PADS, 3" X 3", 4 PER
 ADH TAPE, .5" X 2.5 YD, 2 PER
 GAUZE COMP, 1 SQ YARD, 1 PER
 ADHESIVE BDG,PLSTIC,1"X3"16PER
 NITRILE GLOVES 2PR BBP
 ANTIMCRBL ANTSPTC TWLETTS
 FIRST AID GUIDE ASHI

1 ELASTIC BANDAGE 3" X 4.5YD 1 MICROSHIELD W/VNL GLV/ALCL 1 NORTH RESPONSE REFILL/KIT 1 SCISSOR BDGE 4" RED PLS HDL 1 KIT TWEEZER 3 1/2" SLANTED LBL STOCK 6-3/8"X4" LBL STOCK 4"X2-7/8" 1 LBL STOCK 3"x1-7/8" 1 KIT STL 24 UN WHITE 01 1 COLD PACK UNIT 4"X6" BULK

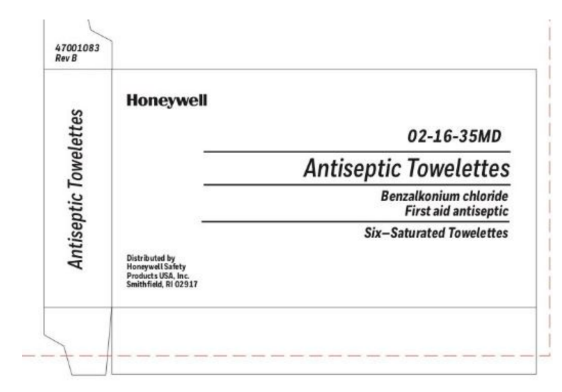
First Aid Burn Cream Principal Display Panel



Principal Display Panel

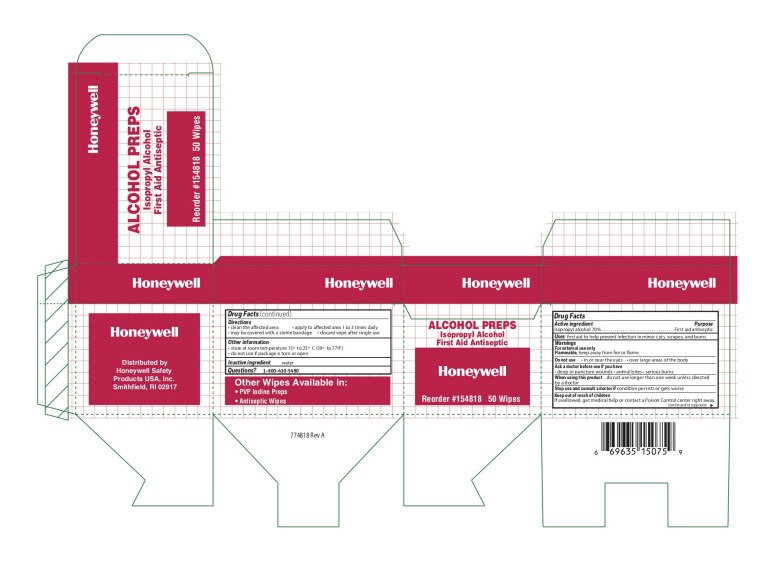


BZK Principal Display Panel



7001083 Rev B	səttələwoT oitqəsitnA
	Drug Facts
	Active Ingredient Purpose Benzelkonium 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 over large areas of the body
	Ask a doctor before use in case of deep or puncture wounds, animal bites, orserious 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 I 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



4150 Kit Label SF00003260

RADNOR

FIRST AID KIT



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

4150 FIRST AID	р кіт						
4150 first aid kit kit	4150 first aid kit kit						
Product Information							
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:0498-4150				

Pack	aging						
# Ite	em Code	Package Description		Marketing Start Date	Marketing End Date		
1 NDC 01	:0498-4150-	1 in 1 KIT; Type 0: Not a Combination Product		10/18/2018			
Quan	Quantity of Parts						
Part #	ŧ	Package Quantity		Total Product (Quantity		
Part 1	6 PACKET		5.4 g				
Part 2	1 PACKET		1.4 mL				
Part 3	10 PACKET		9 g				
Part 4	4 POUCH		1.6 mL				

Part 1 of 4

FIRST AID BURN

benzalkonium chloride, lidocaine hydrochloride cream

Product Information

ltem Code (Source)	NDC:0498-0903
Route of Administration	TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
/	BENZ ALKONIUM CHLORIDE	0.13 g in 100 g
,	LIDOCAINE HYDROCHLORIDE	0.5 g in 100 g

Inactive Ingredients	
Ingredient Name	Strength
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
WATER (UNII: 059QF0KO0R)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
CETYL ALCOHOL (UNII: 936JST6JCN)	
GLYCERYL MONOSTEARATE (UNII: 2300U9XXE4)	
PEG-100 STEARATE (UNII: YD01N1999R)	
LIGHT MINERAL OIL (UNII: N6K5787QVP)	
EDETATE DISODIUM (UNII: 7FLD91C86K)	
TROLAMINE (UNII: 903K93S3TK)	
GLYCERIN (UNII: PDC6A3C0OX)	

Code Jate Date 0.9 g in 1 PACKET; Type 0: Not a Combination Product Date Date Marketing Information Marketing Application Number or Monograph Citation Marketing Start Date Marketing End Date imapproved drug other 12/20/2017 Marketing Start Date Marketing End Date Part 2 of 4 ANTISEPTIC TOWELETTE beenzalkonium chloride liquid NDC:0498-0501 Route of Administration V Product Information Item Code (Source) NDC:0498-0501 Route of Administration Strength Strength Ingredient Name Basis of Strength Strength in 1 mL Active Ingredient/Active Moiety Ingredient Name Basis of Strength in 1 mL 1.3 mg in 1 mL Inactive Ingredients Ingredient Name Strength Package Description Marketing Start Date Marketing End Date	PROPYLPARABE	N (UNII: Z8IX2SCI	10H)		
Item Code Package Description Marketing Start Date Marketing End Date 0.9 g in 1 PACKET; Type 0: Not a Combination Product 0.9 g in 1 PACKET; Type 0: Not a Combination Marketing Start Date Marketing End Date Marketing Information Marketing Category Category anaproved drug other Application Number or Monograph Citation Marketing Start Date Marketing End Date Marketing Category Category Application Number or Monograph Category Numproved drug Marketing Start Date Marketing End Date Part 2 of 4 ANTISEPTIC TOWELETTE Desnzalkonium chloride liquid Variation Variation Product Information Item Code (Source) NDC:0498-0501 Route of Administration NDC:0498-0501 TOPICAL Basis of Strength Strength SenzakKonium Chloride liquid/ Date Ingredient Name Basis of Strength Strength SenzakKonium Chloride (UNII: F5UM2KM3W7) (BENZALKONIUM - BENZAKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - BENZAKONIUM 1: 3 mg (CHLORIDE 1.3 mg (CHLORIDE 1.3 mg Marketing Start Date Ingredient Name Strength Packaging Marketing Start Date Marketing Start Date Marketing End Date	DIAZOLIDINYL U	IREA (UNII: H5RIZ	3MPW4)		
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NDC:0498-0501- 1.4 mL in 1 PACKET; Type 0: Not a Combination		e Pa	ckage Description	-	
			CKET; Type 0: Not a Combination		Batt
		r roudet			

Marketing	Applicati	ion Number or M	lonograph	Mar	keting Start	Ma	rketing End
Category		Citation			Date		Date
unapproved drug other				09/18/	2018		
Part 3 of 4							
TRIPLE ANTI	BIOTIC						
bacitracin zinc, pol	ymyxin b su	ılfate, neomycin	sulfate oint	ment			
Product Inform	ation						
Item Code (Source	•)	NDC:0498-0750					
Route of Administ		TOPICAL					
Active Ingredier	nt/Active M	loiety					
	Ingred	lient Name			Basis of Strength		Strength
BACITRACIN ZINC (UNII: 89Y4M234ES) (BACITRACIN - UNII:58H6RW052I)					BACITRACIN		400 [iU] in 1 g
POLYMYXIN B SULFATE (UNII: 19371312D4) (POLYMYXIN B - UNII:J2VZ07J96K)				POLYMYXIN B		5000 [iU] in 1	
NEOMYCIN SULFATE	(UNII: 057Y62	26693) (NEOMYCIN -	UNII:I16QD7X	297)	NEOMYCIN		3.5 mg in 1 g
Inactive Ingredi	ents						
	Ing	gredient Name				Sti	rength
PETROLATUM (UNII: 4	T6H12BN9U)						
Product Charact	teristics						
Color		white	Score				
Shape			Size	_			
Flavor			Imprint Co	de			
Contains							
Packaging							
# Item Code	Pac	kage Descriptio	on	Mark	eting Start Date	Ма	rketing End Date
1 NDC:0498-0750- 35 Pr	9 g in 1 PACKI oduct	ET; Type 0: Not a C	ombination				

Marketing Category	Applica	tion Number or Monograph Citation		eting Start Date	Mar	keting End Date
unapproved drug other			09/19/20	18		
Part 4 of 4						
ALCOHOL W isopropyl alcohol						
Product Inform	nation					
ltem Code (Sourc	e)	NDC:0498-0143				
Route of Adminis	tration	TOPICAL				
• • • • • • • • • • •						
Active Ingredie		-		Basis	of	Charles and L
	-	edient Name		Streng		Strength
JNII:ND2M416302)	JL (UNII: ND2	M416302) (ISOPROPYL ALCOHOL -		ISOPROPYL ALCOHOL		0.7 mL in 1 mL
Inactive Ingred	lients					
	-	redient Name			Streng	Jth
WATER (UNII: 059QF	OKOOR)					
Packaging						
# Item Code	Pa	ckage Description		ing Start ate	Mark	eting End Date
).4 mL in 1 PC Product	OUCH; Type 0: Not a Combination				
Marketing I	nformat	ion				
Marketing Category	Applica	tion Number or Monograph Citation		eting Start Date	Mar	keting End Date
unapproved drug other			09/18/20	18		
Marketing I	nformat	ion				
Marketing II Marketing Category		ion tion Number or Monograph Citation		ting Start Date	Mar	keting End Date
Marketing		tion Number or Monograph		Date	Mar	_

Labeler - Honeywell Safety Products USA, INC (118768815)

Revised: 1/2024

Honeywell Safety Products USA, INC