4141 FIRST AID KIT- 4141 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.

4141 First Aid Kit (Eye Wash, Hand Sanitizer, FABC, Neomycin, alcohol wipe- FAKREFU-B)

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

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

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 Wipe 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

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

Alcohol Wipe

Inactive ingredient

water

Alcohol Wipe

Questions

1-800-430-5490

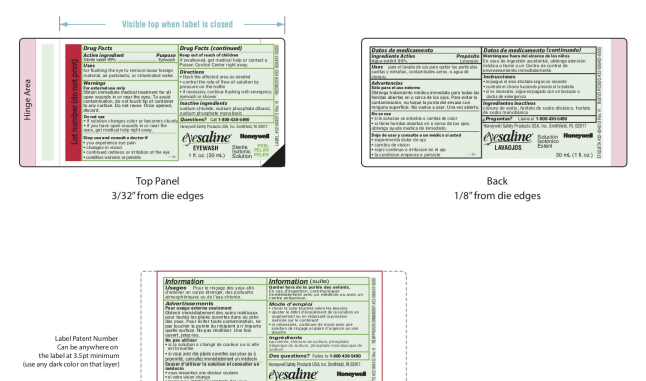
4141

FAKREFU-B Kit Contents 1 HAND SANITIZER 10/PER 1 NEOMYCIN OINT 0.9 GM, UNTZD 25/BX 1 EYE DRESS PKT W/4 ADH STRIPS 1 GAUZE BANDAGE, 4" X 6 YD **1 TOURNIQUET** 2 TRIANGULAR BDG, NON-STERILE **1 WIRE SPLINT 1 PER** 1 GAUZE PADS, 3" X 3", 4 PER 1 ADH TAPE, .5" X 2.5 YD, 2 PER 2 BANDAGE COMP 24" X 72", UNTZD 2/BX 1 FORCEPS & SCISSORS, 1 EA 1 GAUZE BANDAGE, 2" X 6 YD,2 PER 2 INSTANT COLD PACK 4" X 6" 1 1" X 3" PLAS STRIP BAN, UNITZD 50/BX **1 BURN CREAM POUCHES 25/EA** 2 BURN-STOP BURN DRESSING 4 X 4 1 ALCOHOL WIPES, UNITIZED 50/BX 2 NITRILE GLOVES 2PR BBP

1 FIRST AID GUIDE ASHI 1 CPR MICROSHIELD DOUBLE UNIT 1 4OZ BFS EYEWASH TRILINGUAL BOTTLE LBL STOCK 6-3/8"X4" 1 LBL STOCK 3"x1-7/8" 1 ZIP LOCK BAG 14 X 20 1.5 MIL

Eye Wash Package label

#32-004513 Rev. H



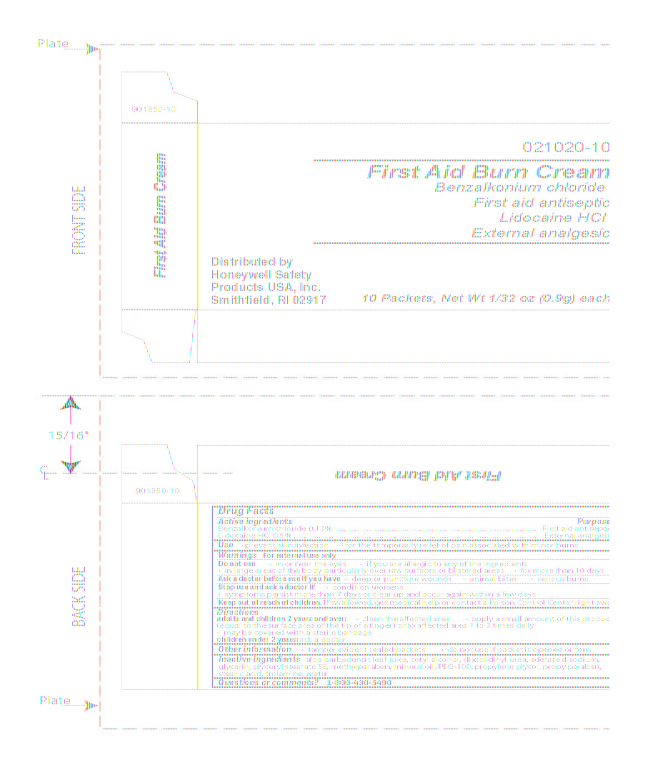
Base 3/32" from die edges

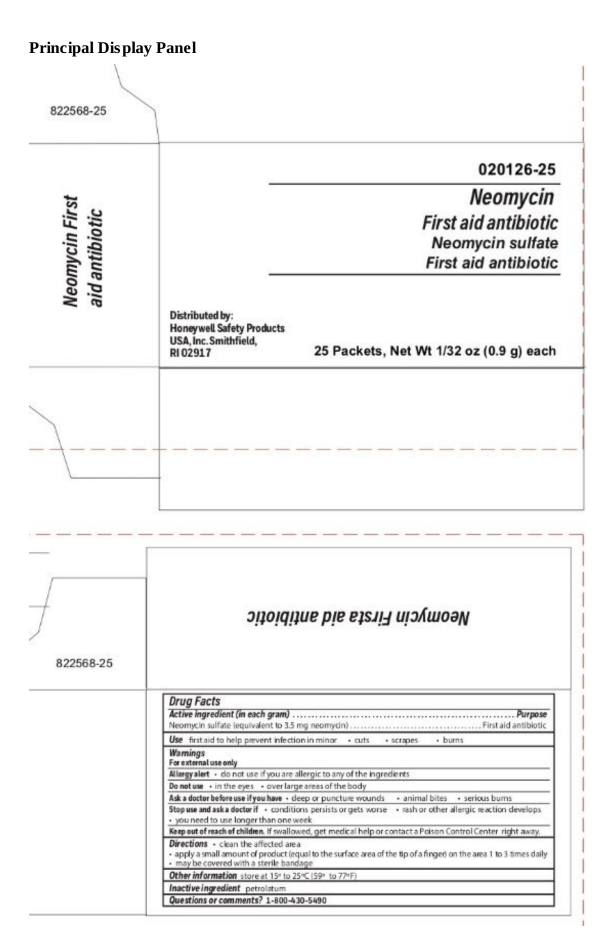
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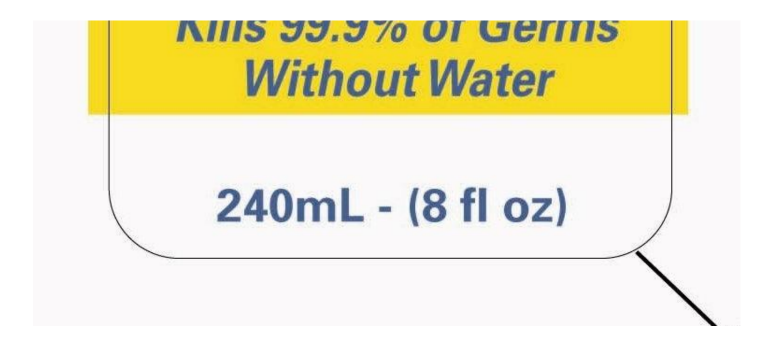
First Aid Burn Cream Principal Display Panel



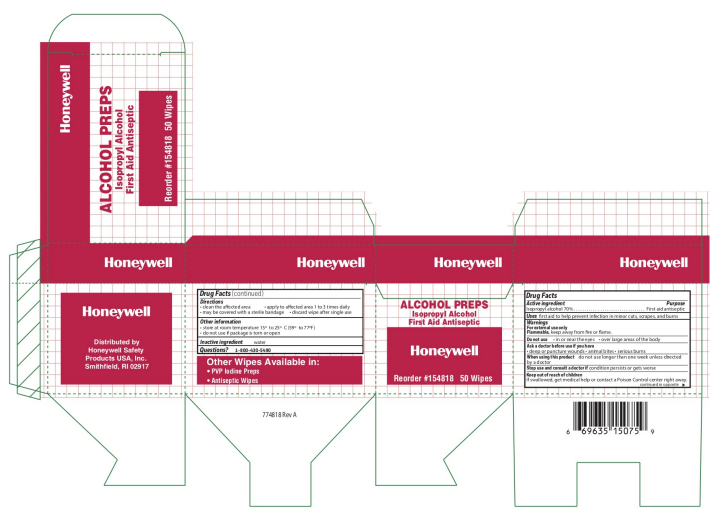


Hand Sanitizer Principal Display Panel

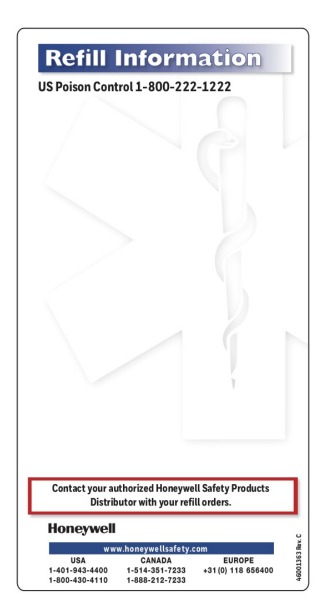




Alcohol Wipe Principal Display Panel



4141 Kit Label FAKREFU-B 46001363 Rev.C Prints 3 colors Black, Red (PMS 186) and Blue (PMS 072)



Product Informat	ion					
Product Type	HUMAN O	TC DRUG	Item Co	de (Source)	N	DC:0498-4141
Packaging						
# Item Code	Pa	ackage Descriptio	on	Marketing Start D	ate	Marketing End Date
1 NDC:0498-4141-01		0: Not a Combinatio		10/18/2018		
Quantity of Parts						
Part #	Dackage Oue	ntitu		Total Produc	+ O u av	
Part 1 1 BOTTLE	Package Qua	intity	118 mL	Total Produc	t Quai	litity
Part 2 50 POUCH			20 mL			
Part 3 25 PACKET			20 III. 22.5 g			
Part 4 25 PACKET			22.5 g			
Part 5 10 PACKET			9 mL			
EYESALINE E	EMERGEN	CY EYEWA	SH			
EYESALINE E purified water liquid Product Informat Item Code (Source)	ion	NDC:0498-0100	SH			
EYESALINE E purified water liquid Product Informat Item Code (Source)	ion		SH			
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat	ion tion	NDC:0498-0100 OPHTHALMIC	SH			
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat	ion tion /Active Moie Ingredie	NDC:0498-0100 OPHTHALMIC ety nt Name		Basis of Stren	gth	Strength
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat	ion tion /Active Moie Ingredie	NDC:0498-0100 OPHTHALMIC ety nt Name		Basis of Stren WATER	-	Strength 98.6 mL in 100 mL
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat Active Ingredient WATER (UNII: 059QF0	ion tion /Active Moie Ingredie KOOR) (WATER	NDC:0498-0100 OPHTHALMIC : ty nt Name - UNII:059QF0KO01	3)		-	98.6 mL in 100 mL
EYESALINE E ourified water liquid Product Informat Item Code (Source) Route of Administrat Active Ingredient WATER (UNII: 059QF0	ion tion /Active Moie Ingredie KOOR) (WATER nts	NDC:0498-0100 OPHTHALMIC ety nt Name - UNII:059QF0KO0I	3)		-	
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat Active Ingredient WATER (UNII: 059QF0 Inactive Ingredie	ion tion /Active Moie Ingredie KOOR) (WATER nts UNII: 451W47IQ8	NDC:0498-0100 OPHTHALMIC ety nt Name - UNII:059QF0KO01 Ingredien X)	3)		-	98.6 mL in 100 mL
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat Active Ingredient WATER (UNII: 059QF0 Inactive Ingredie SO DIUM CHLORIDE (SO DIUM CHLORIDE (ion tion /Active Moie Ingredie KOOR) (WATER nts UNII: 451W47IQ8 2, DIBASIC (UNI	NDC:0498-0100 OPHTHALMIC ety nt Name - UNII:059QF0KO0I Ingredien X) : GR686LBA74)	۲) ۲) ۲ Name	WATER	-	98.6 mL in 100 mL
EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat Active Ingredient WATER (UNII: 059QF0 Inactive Ingredie SO DIUM CHLORIDE (SO DIUM CHLORIDE (ion tion /Active Moie Ingredie KOOR) (WATER nts UNII: 451W47IQ8 2, DIBASIC (UNI	NDC:0498-0100 OPHTHALMIC ety nt Name - UNII:059QF0KO0I Ingredien X) : GR686LBA74)	۲) ۲) ۲ Name	WATER	-	98.6 mL in 100 mL
Part 1 of 5 EYESALINE E purified water liquid Product Informat Item Code (Source) Route of Administrat Active Ingredient WATER (UNII: 059QF0 Inactive Ingredie SODIUM CHLORIDE (SODIUM CHLORIDE (SODIUM PHO SPHATE SODIUM PHO SPHATE	ion tion /Active Moie Ingredie KOOR) (WATER nts UNII: 451W47IQ8 2, DIBASIC (UNI	NDC:0498-0100 OPHTHALMIC ety nt Name - UNII:059QF0KO0I Ingredien X) : GR686LBA74)	۲) ۲) ۲ Name	WATER	-	98.6 mL in 100 mL

Marketing Infor	mation					
Marketing Category	Applicatio	n Number or Monograph Citation	Marketing	Start Date	Marke	ting End Date
OTC monograph final	part349		12/18/2018			
Part 2 of 5						
ALCOHOL WIP)F					
isopropyl alcohol swal						
- I IJ	-					
Product Informatio	n					
Item Code (Source)		NDC:0498-0143				
Route of Administratio	n	TOPICAL				
Active Ingredient/A	Active Moie	tv				
		redient Name		Basis of Str	rength	Strength
ISOPROPYL ALCOHOL UNII:ND2M416302)	C	416302) (ISOPROPYL ALCOHOL -		ISOPROPYL ALCOHOL	0	0.7 mL in 1 mL
,						
Inactive Ingredients		agradiant Nama			Stron	ath
Inactive Ingredients WATER (UNII: 059QF0KC	Ir	ngredient Name			Stren	gth
	Ir	ngredient Name			Stren	gth
	Ir	ngredient Name			Stren	gth
WATER (UNII: 059QF0KC	Ir DOR)					
WATER (UNII: 059QF0KC Packaging # Item Code	Ir DOR)	Package Description	Marketing	Start Date		gth ting End Date
WATER (UNII: 059QF0KC Packaging # Item Code	Ir DOR)		Marketing	Start Date		
WATER (UNII: 059QF0KC Packaging # Item Code	Ir DOR)	Package Description	Marketing	Start Date		
WATER (UNII: 059QF0KC Packaging # Item Code 1 NDC:0498-0143-00 0.	Ir DOR) 4 mL in 1 POU	Package Description	Marketing	Start Date		
WATER (UNII: 059QF0KC Packaging # Item Code	In DOR) 4 mL in 1 POU *mation	Package Description		Start Date	Market	
WATER (UNII: 059QF0KC Packaging # Item Code 1 NDC:0498-0143-00 0.4 Marketing Infor	In DOR) 4 mL in 1 POU *mation Applicati	Package Description CH; Type 0: Not a Combination Product			Market	ting End Date
WATER (UNII: 059QF0KC Packaging # Item Code 1 NDC:0498-0143-00 0. Marketing Infor Marketing Category	In DOR) 4 mL in 1 POU *mation Applicati	Package Description CH; Type 0: Not a Combination Product	Marketing		Market	ting End Date
WATER (UNII: 059QF0KC Packaging # Item Code 1 NDC:0498-0143-00 0. Marketing Infor Marketing Category	In DOR) 4 mL in 1 POU *mation Applicati	Package Description CH; Type 0: Not a Combination Product	Marketing		Market	ting End Date
WATER (UNII: 059QF0KC Packaging # Item Code 1 NDC:0498-0143-00 0. Marketing Infor Marketing Category OTC monograph not final	Ir DOR) 4 mL in 1 POU *mation Applicati part333A	Package Description CH; Type 0: Not a Combination Product	Marketing		Market	ting End Date
WATER (UNII: 059QF0KC Packaging # Item Code 1 1 NDC:0498-0143-00 0. Marketing Infor Marketing Category OTC monograph not final Part 3 of 5 FIRST AID BUF	In COR) 4 mL in 1 POU mation Applicati part333A	Package Description CH; Type 0: Not a Combination Product Con Number or Monograph Citation	Marketing		Market	ting End Date
WATER (UNII: 059QF0KC Packaging # Item Code 1 NDC:0498-0143-00 0. Marketing Infor Marketing Category OTC monograph not final	In COR) 4 mL in 1 POU mation Applicati part333A	Package Description CH; Type 0: Not a Combination Product Con Number or Monograph Citation	Marketing		Market	ting End Date

Item Code (Sou	mation					
nem coue (oou	rce)	NDC:0498-0903				
	·					
Route of Admini	istration	TOPICAL				
Active Ingred	lient/Activ	ve Moiety				
		Ingredient Name		Basis of Stre	ength	Strength
BENZALKONIUM UNII:7N6JUD5X6Y		E (UNII: F5UM2KM3W7) (BENZALKONIUM -		BENZALKONIUM CHLORIDE	0	0.13 g in 100 g
L IDO CAINE HYD	ROCHLOR	IDE (UNII: V13007Z41A) (LIDOCAINE - UNII:981	91200987)	LIDOCAINE HYDROCHLORIDE		0.5 g in 100
Inactive Ingre	edients					
		Ingredient Name			5	Strength
PROPYLENE GL	YCOL (UNII	: 6DC9Q167V3)				
ALOE VERA LEA	F (UNII: ZY8	1Z8 3H0 X)				
WATER (UNII: 059	9QF0KO0R)					
STEARIC ACID (U	UNII: 4ELV7Z	265AP)				
METHYLPARABI	E N (UNII: A21	18 C 7 HI9 T)				
CETYL ALCOHO	L (UNII: 936	JST6JCN)				
GLYCERYL MON	NOSTEARAT	ГЕ (UNII: 230ОU9ХХЕ4)				
PEG-100 STEAR	ATE (UNII: Y	D0 1N19 9 9 R)				
LIGHT MINERAL	OIL (UNII:	N6K5787QVP)				
EDETATE DISOD	IUM (UNII: 7	'FLD91C86K)				
TROLAMINE (UN	NII: 903K93S	3TK)				
GLYCERIN (UNII:	PDC6A3C0	OX)				
PROPYLPARABE	E N (UNII: Z81	X2SC1OH)				
DIAZOLIDINYL						
		•				
Packaging						
		Package Description	Marketi	ing Start Date	Market	ing End Date
# Item Code	0.9 g in 1 PA	· ·	Market	ing Start Date	Market	ing End Date
# Item Code	0.9 g in 1 PA	Package Description CKET; Type 0: Not a Combination Product	Marketi	ing Start Date	Market	ing End Date
# Item Code 1		CKET; Type 0: Not a Combination Product	Market	ing Start Date	Market	ing End Date
# Item Code 1 Marketing	Informa	CKET; Type 0: Not a Combination Product		ng Start Date		
# Item Code 1	Informa	CKET; Type 0: Not a Combination Product		ing Start Date		ing End Date ting End Date
# Item Code 1 Marketing Marketing Cat	Informa tegory A	CKET; Type 0: Not a Combination Product		seting Start Date		
# Item Code 1 Marketing Cat	Informa tegory A	CKET; Type 0: Not a Combination Product Ition Application Number or Monograph Citation	n Marl	seting Start Date		
# Item Code 1 Marketing Cat	Informa tegory A	CKET; Type 0: Not a Combination Product Ition Application Number or Monograph Citation	n Marl	seting Start Date		
# Item Code 1 Marketing Marketing Cat OTC monograph n Part 4 of 5	Informa tegory A not final pa	CKET; Type 0: Not a Combination Product Ition Application Number or Monograph Citation	n Marl	seting Start Date		
Marketing	Informa tegory A not final pa	CKET; Type 0: Not a Combination Product Ition Application Number or Monograph Citation	n Marl	seting Start Date		
# Item Code 1 Marketing Marketing Cat OTC monograph n Part 4 of 5	Informa tegory A not final pa	CKET; Type 0: Not a Combination Product Ition Application Number or Monograph Citation	n Marl	seting Start Date		

Product Information			
Item Code (Source)	NDC:0498-0730		
Route of Administration	TOPICAL		
Active Ingredient/Active	•		
	Ingredient Name	Basis of Str	0
NEOMYCIN SULFATE (UNII: 057	Y626693) (NEOMYCIN - UNII:116QD7X297)	NEOMYCIN SUL	FATE 3.5 mg in 1 g
Inactive Ingredients			
0	Ingredient Name		Strength
PETROLATUM (UNII: 4T6H12BN	9 U)		
Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Da
	PACKET; Type 0: Not a Combination Product		
Marketing Information	D N Ication Number or Monograph Citation	Marketing Start Date	Marketing End Dat
OTC monograph final part333E	3	0 3/3 1/20 10	
Part 5 of 5			
INSTANT HAND SA	NITIZER		
Product Information			
Item Code (Source)	NDC:59898-420		
Route of Administration	TOPICAL		
	TOTIONE		
Active Ingredient/Active	•		
	gredient Name	Basis of Strength	
ALCOHOL (UNII: 3K9958V90M)	(ALCOHOL - UNII:3K9958V90M)	ALCOHOL	62 mL in 100 mL
Inactive Ingredients			
	Ingredient Name		Strength
ALOE VERA LEAF (UNII: ZY8 1Z8			
ALPHATOCOPHEROL ACETA			

TRIISOPROPANOLAMINE (UNII: W9EN9DLM98)							
CARBOMER COPOLYMER TYPE A (UNII: 71DD5V995L)							
WATER (UNII: 059QF0K00R)							
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)							
Packaging							
# Item Code	Package Description	Marketing Start Date	Marketing End Date				
1 NDC:59898-420-36	0.9 mL in 1 PACKET; Type 0: Not a Combination Product						
Marketing Info	rmation						
Marketing Category	y Application Number or Monograph Citation	Marketing Start Date	Marketing End Date				
OTC monograph not fina	al part333E	04/15/2011					
Marketing Information							
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date				
unapproved drug other		10/18/2018					

Labeler - Honeywell Safety Products USA, INC (079287321)

Establishment				
Name	Address	ID/FEI	Business Operations	
Honeywell Safety Products USA, INC		079287321	pack(0498-4141)	

Establishment

Name	Address	ID/FEI	Business Operations
Water-Jel Technologies		155522589	manufacture(0498-0903, 0498-0730, 59898-420)

Establishment

Name	Address	ID/FEI	Business Operations
Honeywell Safety Products USA, Inc		167518617	manufacture(0498-0100)

Establishment				
Name	Address	ID/FEI	Business Operations	
Changzhou Maokang Medical		421317073	manufacture(0498-0143)	

Revised: 11/2019

Honeywell Safety Products USA, INC