4214 FIRST AID KIT- 4214 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.

0498-4214: First Aid Kit (Antiseptic Sp, EW, ASA, BZK wipe- 340420FP)

Eyewash Active ingredient

Sterile Water 99%

Eyewash *Purpose*

Eyewash

Eyewash

Uses

• for flushing the eye to remove loose foreign material, air pollutants or chlorinated water

Eyewash *Warnings*

For external use only Obtain immediate medical treatment for all open wounds in or near 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 eye
- condition worsens or persists

Keep out of reach of children

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

Eyewash 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

Eyewash Inactive ingredients

sodium chloride, sodium phosphate dibasic, sodium phosphate monobasic

Eyewash *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

Ammonia Active ingredient

Ammonia 15%

Ammonia *Purpose*

Respiratory stimulant

Ammonia

Uses

• to prevent or treat fainting

Ammonia Warnings

For external use only

Do not use

• if you have breathing problems such as asthma or emphysema

Stop use and ask a doctor if

• condition persists

Keep out of reach of children

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

Ammonia Directions

- hold inhalant away from face and crush ampoule between thumb and forefinger at position indicated on sleeve.
- hold near nostrils for inhalation of volatile vapor

Ammonia Other information

• store at room temperature away from light

Ammonia Inactive ingredient

alcohol USP, FD&C red #40, lavender oil, lemon oil fcc, nutmeg oil, purified water

Ammonia *Questions or Comments?*

1-800-430-5490

BZK Wipes Active ingredient

Benzalkonium chloride 0.13% w/v

BZK Wipe *Purpose*

First aid antiseptic

BZK Wipes *Uses*

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

BZK Wipes *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

Antiseptic Spray Active ingredient

Benzalkonium chloride 0.13%

Antiseptic Spray

Purpose

First aid antiseptic

Antiseptic Spray

Uses

• first aid to help prevent infection in minor cuts, scrapes and burns

Antiseptic Spray *Warnings*

For external use only

Do not use

- in or near the 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 one week unless directed by a doctor

Stop use and ask a doctor if:

• 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

Antiseptic Spray Directions

- clean the affected area
- spray a small amount of this product on the area 1 to 3 times daily
- may be covered with a sterile bandage
- if bandaged, let dry first

ntiseptic Spray Other information

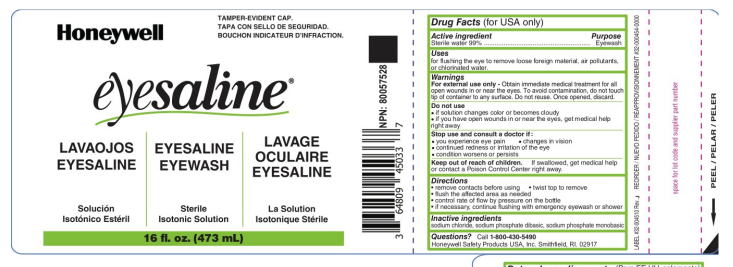
shake well

store at room temperature 15 0 -30 0 C (59 0 -86 0 F)

Antiseptic Spray Inactive ingredients diazolidinyl urea, edetate disodium, glycerin, hypromellose, methylparaben, octoxynol 9, propylene glycol, propylparaben, trolamine, water

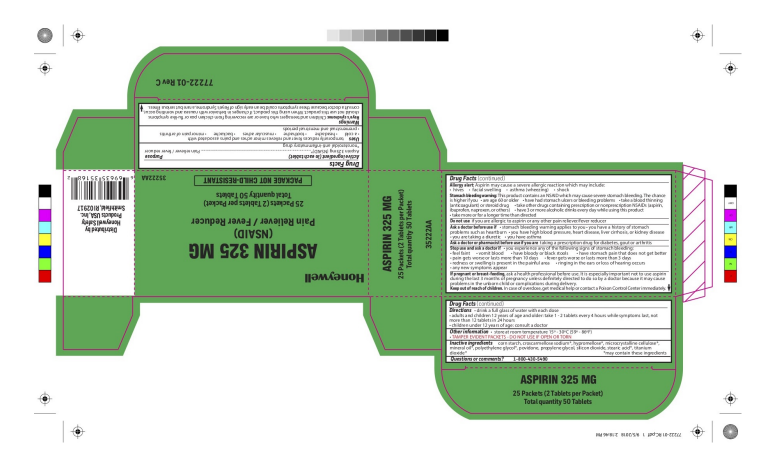
4214 340420FP Kit Contents

1 INSTANT COLD PACK 4" X 6" 1 ADHESIVE TAPE W/P 1/2"X 5 YD 1 ADH BDG, CLOTH, 1"X3", 16 PER 1 O/H PUMP ANTISEPTIC 2 OZ ID F **1 FIRST AID GUIDE ASHI** 1 GAUZE CLEAN-WRAP BDGE N/S 2" 1 40Z BFS EYEWASH TRILINGUAL BOTTLE 1 SCISSOR BDGE 4" RED PLS HDL 1 KIT TWEEZER 3 1/2" SLANTED 1 BANDAGE COMP 3" W/TELFA PAD 2 LBL STOCK 6-3/8"X4" LBL STOCK 4"X2-7/8" 1 LBL STOCK 3"x1-7/8" 4 BZK ANTISEPTIC WIPE, BULK **1 PR LRG NITRILE GLVES ZIP BAG** 1 KIT STL 16 UN (HORIZONTAL) 1 TRI BNDG NON WOVEN 40"X40"X56" 10 NON ADHERENT PAD 2" X 3" **1 WOVEN KNUCKLE 8'S** 1 FINGERTIP "T" 8/BX **3 ASPIRIN BULK 2/PK** 2 AMMONIA INHALANT, BULK

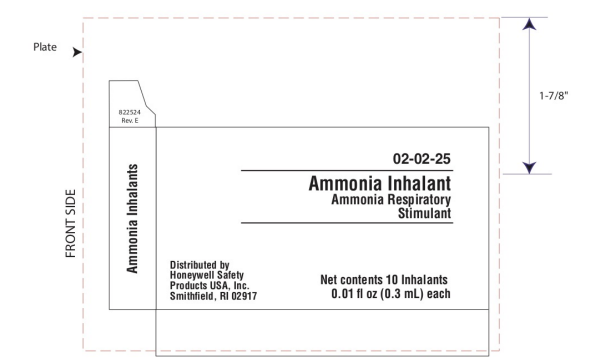


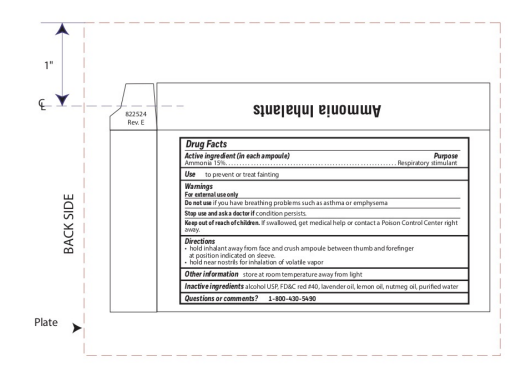
Ingrediente Activo Propósito
Agua estéril 99% Lavaojos
Usos para el lavado de ojo para quitar las particulas sueltas y extrañas, los contaminantes aeros, o agua de cloruro
Advertencias Para el uso externo sólo - Obtenga tratamiento médico inmediato para todas las heridas abiertas en o cerca de los ojos. Para evitar la contaminación, no toque la punta del envase con ninguna superficie. No vuelva a usar. Vez abierto, descarte.
No se use • si la solución se enturbia o cambia de color • si tiene heridas abiertas en o cerca del ojo, obtenga ayuda medica de inmediato
Deje de usar y consulte a un médico si: • experimenta dolor de ojo • cambio de visión • rojez continuo o irritación del ojo • la condición empeora o persiste
Manténgase fuera del alcance de los niños. En caso de ingestión accidental, obtenga atención médica o llame a un Centro de control de envenenamiento inmediatamente.
Instrucciones • quitese los lentes de contacto antes de usar la solución • tuerza la tapa para quitar • enjuague el área afectada según se necesite • controle el chorro haciendo presión el la botella • si es necesario, sigue enjuagado con un lavaojos o ducha de emergencia
Ingredientes inactivos cloruro de sodio, fosfato de sodio dibásico, fosfato de sodio monobásico
¿Preguntas? Llame al 1-800-430-5490 Honeywell Safety Products USA, Inc. Smithfield, RI. 02917
Information
Usages Pour le rinçage des yeux afin d'enlever un corps étranger,
des polluants atmospheriques oú de l'eau chlorée.
des polluarits atmospheriques où de l'eau chlorée . Advertissements Pour usage externe seulement - Obtenir immédiatement des soins médicaux pour toutes les plaies ouvertes dans ou près des yeux. Pour éviter toute contamination, ne pas toucher la pointe du récipient à n'importe quelle surface. Ne pas réutiliser. Une fois ouvert, jetz-les.
Advertissements Pour usage externe seulement - Obtenir immédiatement des soins médicaux pour toutes les plaies ouvertes dans ou près des yeux. Pour éviter toute contamination, ne pas toucher la pointe du récipient à n'importe quelle surface. Ne pas réutiliser. Une fois
Advertissements Pour usage externe seulement - Obtenir immédiatement des soins médicaux pour toutes les plaies ouvertes dans ou près des yeux. Pour éviter toute contamination, ne pas toucher la pointe du récipient à n'importe quelle surface. Ne pas réutiliser. Une tois ouvert, jetez-les. Ne pas utiliser • si la solution a changé de couleur ou si elle est brouillée • si vous avez des plaies ouvertes aux yeux ou à proximité, consultez immédiatement un médecin Cesser d'utiliser la solution et consulter un médecin • vous ressentez une douleur oculaire • si vous avez utrations change • vougeur ou intrâtion persistante des yeux
Advertissements Pour usage externe seulement - Obtenir immédiatement des sons médicaux pour toutes les plaies ouvertes dans ou près des précipient à n'imple outre contamination, ne pas toucher la pointe du ouvert, jetz-teis. Ne pas utiliser • si la solution a changé de couleur ou si elle est brouillée • si vous avez des plaies ouvertes aux yeux ou à proximité, consultez immédiatement un médecin • vous resert d'utiliser la solution et consulter un médecin • vous resertez une douleur coulaire
Advertissements Pour usage externe seulement - Obtenir immédiatement des sons médicaux pour toutes les plaies ouvertes dans ou près des yeux. Pour eviter toute contamination, ne pas toucher la pointe du touvert, jetez-teis. Ne pas utiliser i si la solution a changé de couleur ou si elle est brouillée • si vous avez des plaies ouvertes aux yeux ou à proximité, consultez immédiatement un médecin Cesser d'utiliser la solution et consulter un médecin • vous resentez une douleur coulaire • si vous coulaires • si vous douleur coulaire • si vous resentez une douleuro
Advertissements Pour usage externe seulement - Obtenir immédiatement des sons médicaux pour toutes les plates ouvertes dans ou près des yeux, Pour éviter toute contamination, ne pas toucher la pointe du récipient à rimporte quelle surface. Ne pas réutiliser. Une fois ouvert, jetez-les. Ne pas utiliser • si la solution a changé de couleur ou si elle est brouillée • si vous avez des plates ouvertes aux yeux ou à proximité, consultez immédiatement un médecin Cesser d'utiliser la solution et consulter un médecin • vous ressentez une douleur outaire • si vote vision change • condition empire ou persiste Garder hors de la portée des enfants. En cas d'ingestion, communiquer immédiatement avec un médecin ou avec un centre antipoion. Mode d'emploi • onicer les verres de contact avant l'utilisation • dévisser le besoins • ajuster le déti d'écoulement de la solution en augmentant ou en réduisant la pression exercée sur le contenant • si neisersie, continuer de incer avec unesolution de incage

Aspirin **Principal Display Panel**



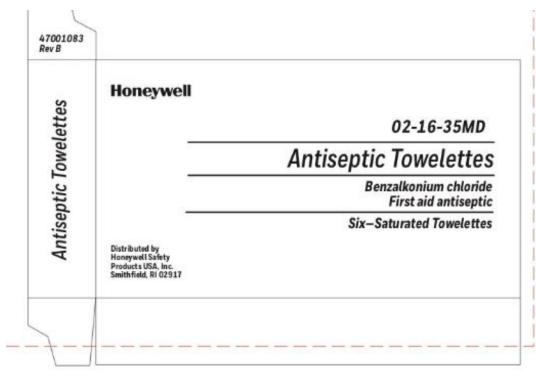






796006 Rev. E (page 3 of 3)

Principal Display Panel



7001083 ev B	səttələwoT citqəsitnA
	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
	Warnings For external use only
	When using this product • do not use in the eyes or apply overlarge 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 Ionger 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 towelotte
	•store acroom competature 15 - 50 C(58 - 50 P)

Antiseptic Spray Principal Display Panel



4214 Kit Label 340420FP

I.



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

AD14 EIDST AID VIT

Produ	ct Inform	ation					
Produc	t Type	HUMAN	OTC DRUG	ltem Co	de (Source)	NDC:0498-42	214
Packa	ging						
# Ite	m Code	Pac	kage Descriptio	n	Marketing Start Date	Marketii Dat	
1 NDC:0		1 in 1 KIT; Ty Product	pe 0: Not a Combina	tion	10/18/2018		
Quant	ity of Par	ts					
Part #	F	Package Q	Quantity		Total Product	Quantity	
Part 1	1 BOTTLE			118 mL			
Part 2	4 PACKET			5.6 mL			
	3 PACKET			6			
Part 4				0.6 mL			
Part 5	1 BOTTLE, SF	PRAY		59 mL			
Part	1 of 5		NCY FYFWA				
Part EYES	1 of 5	MERGE	NCY EYEWA				
Part EYES	1 of 5 ALINE E	MERGE	NCY EYEWA				
Part EYES purifiec	1 of 5 ALINE E	d	NCY EYEWA				
Part EYES purifiec Produ	1 of 5 ALINE E water liquid	MERGE	NCY EYEWA NDC:0498-0100				
Part EYES purifiec Produ	1 of 5 ALINE E water liquid	MERGE d ation					
Part EYES purified Produ Item Co Route	1 of 5 ALINE E water liquid ot Inform ode (Source of Administ	MERGE	NDC:0498-0100 OPHTHALMIC				
Part EYES purifiec Produ Item Co Route	1 of 5 ALINE E water liquid of Inform ode (Source of Administ	MERGE d ation ation a) tration	NDC:0498-0100 OPHTHALMIC Moiety		Pacia of Stronget	- Sture	
Part EYES purifiec Produ Item Co Route Active	1 of 5 ALINE E water liquid of Inform ode (Source of Administ	EMERGE d ation ation ation mt/Active Ingredient	NDC:0498-0100 OPHTHALMIC Moiety t Name	SH	Basis of Strengt		ngth
Part EYES purified Produ Item Co Route Active	1 of 5 ALINE E water liquid of Inform ode (Source of Administ	EMERGE d ation ation ation mt/Active Ingredient	NDC:0498-0100 OPHTHALMIC Moiety	SH	Basis of Strength WATER	1 Stre 98.6 mL in	-
Part EYES purifiec Produ Item Co Route Active	1 of 5 ALINE E water liquid of Inform ode (Source of Administ	MERGE d ation ation ation ration mt/Active Ingredient KOOR) (WATE	NDC:0498-0100 OPHTHALMIC Moiety t Name	SH	_		
Part EYES purifiec Produ Item Co Route Active WATER	1 of 5 ALINE E water liquid of Inform ode (Source of Administ Ingredier (UNII: 059QF01 ve Ingredi	EMERGE d ation ation a) ration mt/Active Ingredient KOOR) (WATE	NDC:0498-0100 OPHTHALMIC Moiety t Name R - UNII:059QF0K00P	SH	_	98.6 mL in	-
Part EYES purified Produ Item Ca Route Active WATER	1 of 5 ALINE E water liquid of Inform ode (Source of Administ (UNII: 059QF0 we Ingredi UCHLORIDE (EMERGE d ation ati	NDC:0498-0100 OPHTHALMIC Moiety t Name R - UNII:059QF0K00P	SH	_	98.6 mL in	100 mL

Packaging						
# Item Code	Pa	ckage Description	Marketiı Da		Mark	eting End Date
	18 mL in 1 BC roduct	OTTLE; Type 0: Not a Combination				
Marketing Ir	nformat	ion				
Marketing Category	Applica	tion Number or Monograph Citation		ing Start ate	Mar	keting End Date
OTC Monograph Drug	M018		12/18/2018	3		
Part 2 of 5						
ANTISEPTIC	TOWEL	ETTE				
benzalkonium chle	oride liquid					
Product Inform	ation					
Item Code (Source		NDC:0498-0501				
Route of Administ		TOPICAL				
Active Ingredie	nt/Active	Moiety				
		dient Name	Ba	asis of Str	ength	Strength
BENZALKONIUM CH UNII:7N6JUD5X6Y)	LORIDE (UNII	: F5UM2KM3W7) (BENZALKONIUM -		NZ ALKONIUM LORIDE		1.3 mg in 1 mL
Inactive Ingred	ients					
		redient Name			Streng	Jth
WATER (UNII: 059QF0	-				-	
Packaging						
# Item Code	Ра	ckage Description	Marketir Da		Mark	eting End Date
		CKET; Type 0: Not a Combination				
00	roduct					
Marketing Ir	nformat	ion				
Marketing Category	Applica	tion Number or Monograph Citation		ing Start ate	Mar	keting End Date

P	art 3 of 5						
	SPIRIN						
as	pirin tablet						
_							
P	roduct Infor	mation					
lte	em Code (Sour	ce)	NDC:0498-	0114			
Ro	oute of Admini	stration	ORAL				
A	tive Ingredi	ent/Active	e Moiety				
		Ingree	dient Name	e	Basis of	f Strength	Strength
AS	PIRIN (UNII: R16C	CO5Y76E) (AS	PIRIN - UNII:R1	L6CO5Y76E)	ASPIRIN		325 mg
In	active Ingre	dients					
			Ingred	lient Name			Strength
СЕ	LLULOSE, MICR	OCRYSTALL	-				<u>-</u>
	LYETHYLENE GI						
sт	EARIC ACID (UNI	I: 4ELV7Z65A	P)				
sт	ARCH, CORN (UN	NII: 08232NY3	BSJ)				
PO	VIDONE (UNII: FZ	Z989GH94E)					
SI		UNII: ETJ7Z6	XBU4)				
	OSCARMELLOSI						
	PROMELLOSE 2			1QE5P712K)			
	NERAL OIL (UNII:						
	FANIUM DIOXIDE		-				
PR	OPYLENE GLYC	DL (UNII: 6DC	9Q167V3)				
Pı	oduct Chara	cteristics	5				
Co	lor	W	hite	Score		2 piec	es
Sh	аре	R	DUND	Size		10mm	
Fla	avor			Imprint Code		FR21	
Co	ontains						
Pa	ackaging						
#	ltem Code	Pa	ackage De	scription	Marketing St Date	art Mai	keting End Date
1	NDC:0498-0114- 01		ET; Type 0: No	ot a Combination	Date		2400
÷.,		Product					

	Marketing	Applica	tion Number or Monograph	Marketing Start	Marketing End
	Category	Appred	Citation	Date	Date
ina oth	pproved drug er			09/18/2018	
Pā	art 4 of 5				
A	MMONIA I	NHALEN	т		
am	monia inhalen	it inhalant			
Pr	oduct Inform	mation			
lte	m Code (Sour	ce)	NDC:0498-3334		
Ro	ute of Adminis	stration	RESPIRATORY (INHALATION)		
Ac	tive Ingredie	ent/Active	Moiety		
		Ingredie	nt Name	Basis of Strength	Strength
٩M	MONIA (UNII: 513	38Q19F1X) (AM	MONIA - UNII:5138Q19F1X)	AMMONIA	0.045 g in 0.3 mL
	active Ingree	ulents			
ALC		Ing	gredient Name		Strength
AL(Ing	gredient Name		Strength
		Ing	gredient Name		Strength
Pa	C OHOL (UNII: 3KS	וחכ פאס איז	gredient Name ckage Description	Marketing Start Date	Strength Marketing End Date
Pa #	COHOL (UNII: 3K9 Ckaging Item Code NDC:0498-3334-	Ing 9958V90М) Ра			Marketing End
Pa #	COHOL (UNII: 3K9 Ckaging Item Code NDC:0498-3334-	Ing 9958V90М) Ра 0.3 mL in 1 АМ	ckage Description		Marketing End
Pa # 1	COHOL (UNII: 3K9 Ckaging Item Code NDC:0498-3334-	Ing 9958V90M) Pa 0.3 mL in 1 AM Product	ckage Description IPULE; Type 0: Not a Combination		Marketing End
Pa #	COHOL (UNII: 3K9 ckaging item Code NDC:0498-3334-	Ing 9958V90M) Pa 0.3 mL in 1 AM Product nformat	ckage Description IPULE; Type 0: Not a Combination		Marketing End Date
Pa # 1 ¦(COHOL (UNII: 3KG ockaging Item Code NDC:0498-3334- 00 arketing Category pproved drug	Ing 9958V90M) Pa 0.3 mL in 1 AM Product nformat	ckage Description IPULE; Type 0: Not a Combination ion tion Number or Monograph	Date Marketing Start	Marketing End Date Marketing End
Pa # 1	COHOL (UNII: 3KG ockaging Item Code NDC:0498-3334- 00 arketing Category pproved drug	Ing 9958V90M) Pa 0.3 mL in 1 AM Product nformat	ckage Description IPULE; Type 0: Not a Combination ion tion Number or Monograph	Date Marketing Start Date	Marketing End Date Marketing End Date
Pa # 1	COHOL (UNII: 3KG ockaging Item Code NDC:0498-3334- 00 arketing Category pproved drug	Ing 9958V90M) Pa 0.3 mL in 1 AM Product nformat	ckage Description IPULE; Type 0: Not a Combination ion tion Number or Monograph	Date Marketing Start Date	Marketing Enc Date Marketing Enc Date
Pa # 1	COHOL (UNII: 3K9 Ckaging Item Code NDC:0498-3334- Co Arketing Category Opproved drug er	Pa 0.3 mL in 1 AM Product nformat Applica	ckage Description IPULE; Type 0: Not a Combination ion tion Number or Monograph	Date Marketing Start Date	Marketing End Date Marketing End Date
Pa # 1 (M	COHOL (UNII: 3KG Ckaging Item Code NDC:0498-3334- 00 arketing Category Opproved drug er	Ing 2958V90M) Pa 0.3 mL in 1 AM Product nformat Applica	ckage Description PULE; Type 0: Not a Combination ion tion Number or Monograph Citation	Date Marketing Start Date	Marketing End Date Marketing End Date

	ormation					
ltem Code (So	urce)	NDC:0498-0402				
Route of Admi	nistration	TOPICAL				
Active Ingre	dient/Active	Moiety				
	Inare	dient Name		Basis of Stre	nath	Strength
BENZALKONIUM UNII:7N6JUD5X6Y)	CHLORIDE (UNII	: F5UM2KM3W7) (BENZALKONIUM	-	BENZ ALKONIUM CHLORIDE	5	0.13 g in 100 mL
Inactive Ingi	redients					
J		Ingredient Name				Strength
GLYCERIN (UNII:	PDC6A3C0OX)					
EDETATE DISOD		1С86К)				
TROLAMINE (UNI	I: 903K93S3TK)					
PROPYLPARABEI	N (UNII: Z8IX2SC1	.OH)				
WATER (UNII: 059	9QF0KO0R)					
DIAZOLIDINYL U	REA (UNII: H5RIZ:	3MPW4)				
DIPROPYLENE G	LYCOL (UNII: E10	71 85 (40)				
		7205040)				
METHYLPARABE						
METHYLPARABE HYPROMELLOSE	N (UNII: A2I8C7HI , UNSPECIFIED	9T) (UNII: 3NXW29V3WO)				
METHYLPARABE	N (UNII: A2I8C7HI , UNSPECIFIED	9T) (UNII: 3NXW29V3WO)				
METHYLPARABE HYPROMELLOSE	N (UNII: A2I8C7HI , UNSPECIFIED	9T) (UNII: 3NXW29V3WO)				
METHYLPARABE HYPROMELLOSE	N (UNII: A2I8C7HI , UNSPECIFIED	9T) (UNII: 3NXW29V3WO)				
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (N (UNII: A218C7HI , UNSPECIFIED UNII: 7JPC6Y25QS	9T) (UNII: 3NXW29V3WO)	Ma	rketing Start Date	Ma	rketing Enc Date
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa	9T) (UNII: 3NXW29V3WO)) nckage Description TLE, SPRAY; Type 0: Not a	Ma	-	Ma	-
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging # Item Code 1 NDC:0498-	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT	9T) (UNII: 3NXW29V3WO)) nckage Description TLE, SPRAY; Type 0: Not a	Ma	-	Ma	-
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging # Item Code 1 NDC:0498-	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT Combination Pre	9T) (UNII: 3NXW29V3WO)) ackage Description TLE, SPRAY; Type 0: Not a oduct	Ma	-	Ma	-
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1) Packaging # Item Code 1 NDC:0498- 0402-59 Marketing Category	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT Combination Pro- Informat	9T) (UNII: 3NXW29V3WO)) ackage Description TLE, SPRAY; Type 0: Not a oduct		-		-
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging # Item Code 1 NDC:0498- 0402-59 Marketing Marketing	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT Combination Pro- Informat	9T) (UNII: 3NXW29V3WO))		Date Date		Date Date
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging # Item Code 1 NDC:0498- 0402-59 Marketing Category unapproved drug	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT Combination Pro- Informat	9T) (UNII: 3NXW29V3WO))	Mai	Date Date		Date Date
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging # Item Code 1 NDC:0498- 0402-59 Marketing Category unapproved drug	N (UNII: A2I8C7HI UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT Combination Pro Informat Applica	9T) (UNII: 3NXW29V3WO)) ackage Description TLE, SPRAY; Type 0: Not a oduct ion tion Number or Monograph Citation	Mai	Date Date		Date Date
METHYLPARABE HYPROMELLOSE OCTOXYNOL 9 (1 Packaging # Item Code 1 NDC:0498- 0402-59 Marketing Category unapproved drug	N (UNII: A2I8C7HI , UNSPECIFIED UNII: 7JPC6Y25QS Pa 59 mL in 1 BOT Combination Pro Informat Applica	9T) (UNII: 3NXW29V3WO)) ackage Description TLE, SPRAY; Type 0: Not a oduct ion tion Number or Monograph Citation	Ma 09/18/	Date Date	Ma	Date Date

Labeler - Honeywell Safety Products USA, INC (118768815)

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