JOHNSON AND JOHNSON ALL PURPOSE FIRST AID KIT- acetaminophen, diphenhydramine hydrochloride, zinc acetate, neomycin sulfate, polymyxin b sulfate, and pramoxine hydrochloride Kenvue Brands LLC

Johnson and Johnson All Purpose First Aid Kit

TYLENOL[®] Extra Strength Caplets

Drug Facts

Active ingredient (in each caplet)

Acetaminophen 500 mg

Purpose

Pain reliever/fever reducer

Uses

- temporarily relieves minor aches and pains due to:
 - the common cold
 - headache
 - backache
 - minor pain of arthritis
 - toothache
 - muscular aches
 - premenstrual and menstrual cramps
- temporarily reduces fever

Warnings

Liver warning

This product contains acetaminophen. Severe liver damage may occur if you take

- more than 4,000 mg of acetaminophen in 24 hours
- with other drugs containing acetaminophen
- 3 or more alcoholic drinks every day while using this product

Allergy alert: acetaminophen may cause severe skin reactions. Symptoms may include:

- skin reddening
- blisters
- rash

If a skin reaction occurs, stop use and seek medical help right away.

Do not use

- with any other drug containing acetaminophen (prescription or nonprescription). If you are not sure whether a drug contains acetaminophen, ask a doctor or pharmacist.
- if you are allergic to acetaminophen or any of the inactive ingredients in this product

Ask a doctor before use if you have liver disease

Ask a doctor or pharmacist before use if you are taking the blood thinning drug warfarin

Stop use and ask a doctor if

- pain gets worse or lasts more than 10 days
- fever gets worse or lasts more than 3 days
- new symptoms occur
- redness or swelling is present

These could be signs of a serious condition.

If pregnant or breast-feeding, ask a health professional before use.

Keep out of reach of children.

Overdose warning

In case of overdose, get medical help or contact a Poison Control Center right away. (1-800-222-1222) Quick medical attention is critical for adults as well as for children even if you do not notice any signs or symptoms.

Directions

do not take more than directed (see overdose warning)

adults and children 12 years and over	 take 2 caplets every 6 hours while symptoms last do not take more than 6 caplets in 24 hours, unless directed by a doctor do not use for more than 10 days unless directed by a doctor
children under 12 years	ask a doctor

Other information

- store between 20-25°C (68-77°F)
- do not use if pouch is torn or damaged

Inactive ingredients

carnauba wax ¹, corn starch ¹, FD&C red no. 40 aluminum lake, hypromellose, magnesium stearate, modified starch ¹, polyethylene glycol ¹, powdered cellulose, pregelatinized starch, propylene glycol, shellac, sodium starch glycolate, titanium dioxide

1 contains one or more of these ingredients

Questions or comments?

call 1-877-895-3665 (toll-free) or 215-273-8755 (collect)

BENADRYL[®] Extra Strength Itch Stopping Cream

Drug Facts

Active ingredients	Purpose	
Diphenhydramine hydrochloride 2%	Topical analgesic	
Zinc acetate 0.1%	Skin protectant	

Uses

- temporarily relieves pain and itching associated with:
 - insect bites
 - minor burns
 - sunburn
 - minor skin irritations
 - minor cuts
 - scrapes
 - rashes due to poison ivy, poison oak, and poison sumac
- dries the oozing and weeping of poison ivy, poison oak, and poison sumac

Warnings

For external use only

Do not use

- on large areas of the body
- with any other product containing diphenhydramine, even one taken by mouth

Ask a doctor before use

- on chicken pox
- on measles

When using this product avoid contact with eyes

Stop use and ask a doctor if

- condition worsens or does not improve within 7 days
- symptoms 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.

Directions

- do not use more than directed
- 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

Other information

protect from excessive heat (40°C/104°F)

Inactive ingredients

cetyl alcohol, diazolidinyl urea, methylparaben, polyethylene glycol monostearate 1000, propylene glycol, propylparaben, purified water

Questions?

call toll-free 800-524-2624 (English/Spanish) or 215-273-8755 (collect)

NEOSPORIN[®] + PAIN RELIEF First Aid Antibiotic/Pain Relieving Cream

Drug Facts

Active ingredients (in each gram)	Purpose
Neomycin Sulfate (3.5 mg)	First aid antibiotic
Polymyxin B Sulfate (10,000 units)	First aid antibiotic
Pramoxine HCI (10 mg)	External analgesic

Uses

first aid to help prevent infection and for the temporary relief of pain in minor:

- cuts
- scrapes
- burns

Warnings

For external use only.

Do not use

- if you are allergic to any of the ingredients
- 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

- you need to use longer than 1 week
- condition persists or gets worse
- symptoms persist for more than 1 week, or clear up and occur again within a few days
- rash or other allergic reaction develops

Keep out of reach of children.

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

Directions

- adults and children 2 years of age and older:
 - clean the affected area
 - apply a small amount of this 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
- children under 2 years of age: ask a doctor

Other information

store at 20° to 25°C (68° to 77°F)

Inactive ingredients

Water, Emulsifying Wax, Mineral Oil, Petrolatum, Propylene Glycol, Methylparaben, Sulfuric acid, Sodium Hydroxide

Questions?

call 800-223-0182 or 215-273-8755 (collect)

Distributed by: JOHNSON & JOHNSON CONSUMER INC. Skillman, NJ 08558

PRINCIPAL DISPLAY PANEL - Kit Package Label

Johnson & Johnson ®

ALL-PURPOSE

FIRST AID KIT

OUTDOORS ON-THE-GO AT HOME

INCLUDES \$20 VALUE 5 FULL-SIZE ITEMS

CUTS & SCRAPES MINOR BURNS

ITCH RELIEF PAIN RELIEF

SKIN RASHES INSECT BITES

140 ITEMS

SEE BACK PANEL



JOHNSON AND JOHNSON ALL PURPOSE FIRST AID KIT

acetaminophen, diphenhydramine hydrochloride, zinc acetate, neomycin sulfate, polymyxin b sulfate, and pramoxine hydrochloride kit

Produ	ct Informa	tion					
Produc	t Type	HUMAN O	TC DRUG	Item Code (Source)	NDC:69968-0)711
_							
Packa	ging						
	em Code	-	e Description	Marketing Start	Date	Marketing I	Ind Date
1 NDC:6	59968-0711-9	1 in 1 PACE	KAGE	08/16/2021			
Quant	ity of Parts	:					
Part #	-	, ackage Q	uantity	Tota	al Produ	ct Quantity	
	2 POUCH			4			
Part 2	1 TUBE			28.3 g			
Part 3	1 TUBE			14.2 g			
Part 4	6 PACKET			6			
Part	1 of 4						
TYLE	NOL EXTI	RA STR	ENGTH				
acetam	ninophen tabl	et, film co	ated				
Produ	ct Informa	tion					
ltem Co	ode (Source)		NDC:50580-449				
Route	of Administra	ation	ORAL				
Active	Ingredient	Active	Moiety				
		Ingre	dient Name		Basis	of Strength	Strength
ACETAM	IINOPHEN (UNII	: 36209ITL9	D) (ACETAMINOPHE	N - UNII:36209ITL9D)	ACETAN	MINOPHEN	500 mg
Inactiv	ve Ingredie	nts					
			Ingredient N	lame		S	Strength
CARNAU	BA WAX (UNII:	R12CBM0EI	Z)				
STARCH	, CORN (UNII: C)8232NY3SJ))				
	ED NO. 40 (UNI						
	UM OXIDE (UNI						
			UNII: 3NXW29V3WO)				
	SIUM STEARAT						
POLYET	HYLENE GLYC	DL, UNSPE	CIFIED (UNII: 3WJQ)	USDWIA)			

POWDERED CELLULO	DSE (UNII: SI	MD1X3XO9M)				
PROPYLENE GLYCOL	(UNII: 6DC90	Q167V3)				
SHELLAC (UNII: 46N10)7B71O)					
SODIUM STARCH GL	COLATE T	(PE A POTATO (UNII: 5856J3G2A	2)			
TITANIUM DIOXIDE (JNII: 15FIX9V	2JP)				
Product Charac	teristics					
Color	white	Score		no score		
Shape	OVAL	Size		19mm		
Flavor		Imprint Code		TYLENOL;	500	
Contains						
Packaging						
			Market	ing Start	Market	ing End
# Item Code	Pac	kage Description		ate		ate
	in 1 POUCH; roduct	Type 0: Not a Combination				
1	louuer					
Marketing In	format	ion				
Marketing		tion Number or Monograph	Marl	ceting Start	Marko	ting End
Category	Applica	Citation	i Mari	Date	Marketing End Date	
OTC Monograph Drug	M013		08/19/1	984		
Part 2 of 4						
	ΥΤΡΛ ς	TRENGTH ITCH ST		G		
		ide and zinc acetate cream		U		
upnennyurannine i	ryurucmur	ide and zinc acetate cream				
Product Inform	ation					
		NDC:69968-0223				
Item Code (Source		NDC.03300-0225				
Item Code (Source Route of Administ		TOPICAL				
Route of Administ	ration	TOPICAL				
Route of Administ	ration nt/Active	TOPICAL		Basis of St	trenath	Strengt
Route of Administ Active Ingredier	ration ht/Active Ingree	TOPICAL Moiety dient Name DRIDE (UNII: TC2D6JAD40)		Basis of St	-	Strengt 20 mg
Route of Administ Active Ingredier	ration ht/Active Ingree	TOPICAL Moiety dient Name DRIDE (UNII: TC2D6JAD40)			NE	20 mg in 1 g
Route of Administ Active Ingredier DIPHENHYDRAMINE I (DIPHENHYDRAMINE - U	ration ht/Active Ingree HYDROCHLO	TOPICAL Moiety dient Name DRIDE (UNII: TC2D6JAD40)	ł	DIPHENHYDRAMI	NE	20 mg in 1 g 1 mg
Route of Administ Active Ingredier DIPHENHYDRAMINE I (DIPHENHYDRAMINE - U	ration ht/Active Ingree HYDROCHLO	TOPICAL Moiety dient Name DRIDE (UNII: TC2D6JAD40) 83M)	ł	Diphenhydrami Hydrochloridi	NE	20 mg in 1 g
Route of Administ Active Ingredier DIPHENHYDRAMINE I (DIPHENHYDRAMINE - U	ration ht/Active Ingree HYDROCHLO	TOPICAL Moiety dient Name DRIDE (UNII: TC2D6JAD40) 83M)	ł	Diphenhydrami Hydrochloridi	NE	20 mg in 1 g 1 mg
Route of Administ Active Ingredier DIPHENHYDRAMINE I (DIPHENHYDRAMINE - U	ration ht/Active Ingree HYDROCHLC NII:8GTS82S FM5526K07A	TOPICAL Moiety dient Name DRIDE (UNII: TC2D6JAD40) 83M)	ł	Diphenhydrami Hydrochloridi	NE	in 1 g 1 mg

CETYL ALCOHOL DIAZOLIDINYL UR METHYLPARABEN		N)		
PROPYLENE GLYC				
PROPYLPARABEN	(UNII: Z8IX2SC1	.OH)		
NATER (UNII: 0590	QF0KO0R)			
Packaging				
# Item Code	Pa	ckage Description	Marketing Start Date	Marketing End Date
NDC:69968- 0223-1	1 in 1 CARTON			2 4 1 2
1	28.3 g in 1 TUBE; Type 0: Not a Combination			
	Product			
Marketing	Informat	ion		
Marketing Category	Applica	tion Number or Monograph Citation	Marketing Start Date	Marketing End Date
DTC Monograph Dr	ug M017		11/01/2009	
		AIN DELICE EIDET A		
RELIEVING	İ	AIN RELIEF FIRST A		/PAIN
RELIEVING	İ	AIN RELIEF FIRST A		/PAIN
RELIEVING neomycin sulfat	te, polymyxin			/PAIN
RELIEVING neomycin sulfat Product Infor	te, polymyxin rmation	b sulfate, and pramoxine hyd		/PAIN
RELIEVING neomycin sulfat Product Infor Item Code (Sou	te, polymyxin rmation urce)	b sulfate, and pramoxine hyd NDC:69968-0055		P AIN
RELIEVING neomycin sulfat Product Infor Item Code (Sou	te, polymyxin rmation urce)	b sulfate, and pramoxine hyd		P/PAIN
RELIEVING neomycin sulfat Product Infor Item Code (Sou	te, polymyxin rmation urce)	b sulfate, and pramoxine hyd NDC:69968-0055		P AIN
RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin	te, polymyxin rmation urce) histration	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL		P/PAIN
RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin	te, polymyxin rmation (rce) (istration lient/Active Ingredi	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL Moiety ent Name		
RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin Active Ingred	te, polymyxin rmation (rce) (istration lient/Active Ingredi	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL Moiety	rochloride cream	
RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin Active Ingred NEOMYCIN SULFA JNII:116QD7X297) POLYMYXIN B SU	te, polymyxin rmation (rce) (istration lient/Active Ingredi ATE (UNII: 057Ye	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL Moiety ent Name	rochloride cream Basis of Streng	th Strength
RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin Active Ingred NEOMYCIN SULFA UNII: I16QD7X297) POLYMYXIN B SU UNII: J2VZ 07J96K)	te, polymyxin rmation (rce) (istration lient/Active Ingredi ATE (UNII: 057Y6 (UNII: 19	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL Moiety ent Name 526693) (NEOMYCIN -	rochloride cream Basis of Streng NEOMYCIN	th Strength 3.5 mg in 1 g 10000 [USP'U]
RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin Active Ingred NEOMYCIN SULF/ UNII:116QD7X297) POLYMYXIN B SU UNII:12VZ 07J96K) PRAMOXINE HYDI UNII:068X84E056)	te, polymyxin rmation (rce) (istration lient/Active Ingredi ATE (UNII: 057YC LFATE (UNII: 19 ROCHLORIDE (U	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL Moiety ent Name 526693) (NEOMYCIN - 371312D4) (POLYMYXIN B -	rochloride cream Basis of Streng NEOMYCIN POLYMYXIN B PRAMOXINE	th Strength 3.5 mg in 1 g 10000 [USP'U] in 1 g 10000 [USP'U]
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RELIEVING neomycin sulfat Product Infor Item Code (Sou Route of Admin Active Ingred NEOMYCIN SULF/ JNII:116QD7X297) POLYMYXIN B SU JNII:J2VZ07J96K) PRAMOXINE HYDI JNII:068X84E056)	te, polymyxin te, polymyxin rmation irce) iistration lient/Active Ingredi ATE (UNII: 057Y6 LFATE (UNII: 19 ROCHLORIDE (U	b sulfate, and pramoxine hyd NDC:69968-0055 TOPICAL Moiety ent Name 526693) (NEOMYCIN - 371312D4) (POLYMYXIN B -	rochloride cream Basis of Streng NEOMYCIN POLYMYXIN B PRAMOXINE	th Strength 3.5 mg in 1 g 10000 [USP'U] in 1 g 10000 [USP'U]

MINERAL OIL (UNII:							
PETROLATUM (UNI							
	5L (UNII: 6DC9Q167V3)						
METHYLPARABEN							
SULFURIC ACID (UI							
	DE (UNII: 55X04QC32I)						
Packaging							
# Item Code	Package Description	Marketing Start Date	Marketing End Date				
1 NDC:69968-	1 in 1 CARTON	Dute					
0055-2	14.2 g in 1 TUBE; Type 0: Not a Combination	on					
1	Product						
Marketing I	nformation						
Marketing Category	Application Number or Monogr Citation	aph Marketing Start Date	Marketing End Date				
OTC Monograph Dru		12/01/2009	Date				
ore nonograph bra	g 11017	12,01,2000					
-	ND JOHNSON HAND CLEA y products] cloth	ANSING WIPES					
Product Infor	mation						
NOULE OF ADMINI	I UPICAL	Route of Administration TOPICAL					
Other Ingredie	ents						
		ient Name	Ouantity				
Other Ingredie Ingredient K		ient Name	Quantity				
Ingredient K	ind Ingred		Quantity				
Ingredient K	(ind Ingred WATER (UNII: 059QF0K00R)	2M416302)	Quantity				
Ingredient K INGR INGR	(ind Ingred WATER (UNII: 059QF0K00R) ISOPROPYL ALCOHOL (UNII: ND)	2M416302) III: F5UM2KM3W7)	Quantity				
Ingredient K INGR INGR INGR	Kind Ingred WATER (UNII: 059QF0K00R) ISOPROPYL ALCOHOL (UNII: ND: BENZALKONIUM CHLORIDE (UNI)	2M416302) III: F5UM2KM3W7)	Quantity				
Ingredient K INGR INGR INGR INGR	Kind Ingred WATER (UNII: 059QF0K00R) ISOPROPYL ALCOHOL (UNII: ND: BENZALKONIUM CHLORIDE (UNI)	2M416302) III: F5UM2KM3W7)	Quantity				
Ingredient K INGR INGR INGR INGR	Kind Ingred WATER (UNII: 059QF0K00R) ISOPROPYL ALCOHOL (UNII: ND: BENZALKONIUM CHLORIDE (UNI)	2M416302) III: F5UM2KM3W7) MDF5V39QO)					
Ingredient K INGR INGR INGR INGR	Kind Ingred WATER (UNII: 059QF0K00R) ISOPROPYL ALCOHOL (UNII: ND: BENZALKONIUM CHLORIDE (UNI)	2M416302) III: F5UM2KM3W7)	Quantity Quantity Marketing End Date				
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INGR INGR INGR INGR INGR INGR INGR INGR	Lingred WATER (UNII: 059QF0K00R) ISOPROPYL ALCOHOL (UNII: ND) BENZALKONIUM CHLORIDE (UNI) SODIUM BICARBONATE (UNII: 8	2M416302) III: F5UM2KM3W7) MDF5V39QO) Marketing Start	Marketing End				

Marketing In	formation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
Cosmetic		05/26/2021	
Marketing In	formation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	08/16/2021	

Labeler - Kenvue Brands LLC (118772437)

Revised: 9/2024

Kenvue Brands LLC