

**TRIPLE ANTIBIOTIC PLUS PAIN RELIEF- bacitracin zinc, neomycin sulfate, polymyxin b sulfate, and pramoxine hydrochloride ointment
FRED'S, INC.**

**Triple Antibiotic
Plus Pain Relief**

Drug Facts

Active ingredients (in each gram)	Purpose
Bacitracin zinc USP 500 units	First aid antibiotic
Neomycin sulfate USP 3.5 mg	First aid antibiotic
Polymyxin B sulfate USP 10,000 units	First aid antibiotic
Pramoxine HCl USP 10 mg	External analgesic

Uses

first aid to help prevent infection and for the temporary relief of pain or discomfort 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 Center 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)
- see carton or tube crimp for lot number and expiration date

Inactive ingredient

white petrolatum

Questions?

Call **1-866-923-4914**

DISTRIBUTED BY: fred's, Inc.
4300 NEW GETWELL RD,
MEMPHIS, TN 38118

PRINCIPAL DISPLAY PANEL - 14.2 g Tube Carton

TRIPLE ANTIBIOTIC
OINTMENT
+ PAIN RELIEF

MAXIMUM STRENGTH

Bacitracin Zinc USP • Neomycin Sulfate USP • Polymyxin B
Sulfate USP • Pramoxine Hydrochloride USP
FIRST AID ANTIBIOTIC / PAIN RELIEVING OINTMENT

Helps Prevent Infection in Minor Cuts, Scrapes,
and Burns Plus Maximum Strength Pain Relief.

NET WT 0.5 oz (14.2 g)



NDC 55315-192-34



MAXIMUM STRENGTH

Compare to the Active Ingredients in Neosporin® + Pain Relief*

TRIPLE ANTIBIOTIC OINTMENT + PAIN RELIEF

Bacitracin Zinc USP • Neomycin Sulfate USP • Polymyxin B Sulfate USP • Pramoxine Hydrochloride USP

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FIRST AID ANTIBIOTIC / PAIN RELIEVING OINTMENT

Helps Prevent Infection in Minor Cuts, Scrapes, and Burns Plus Maximum Strength Pain Relief.

NET WT 0.5 oz (14.2 g)

100% satisfaction guaranteed

Questions or comments:
1-855-331-FRED (3733)



www.fredsinc.com
MEMPHIS, TN 38118
4300 NEW GETWELL RD,
DISTRIBUTED BY: fred's, Inc.
Made in Canada

*This product is not manufactured or distributed by Johnson & Johnson Corporation, owner of the registered trademark Neosporin® + Pain Relief.

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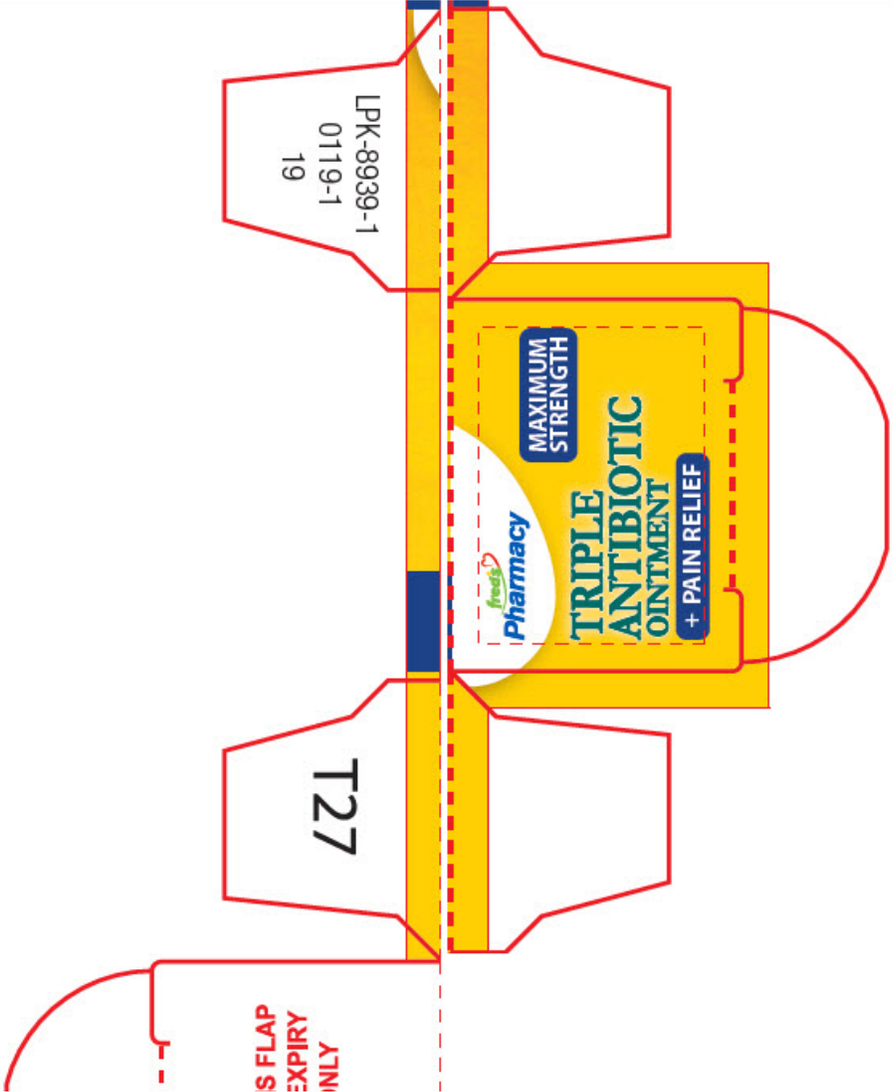
Inactive ingredient white petrolatum

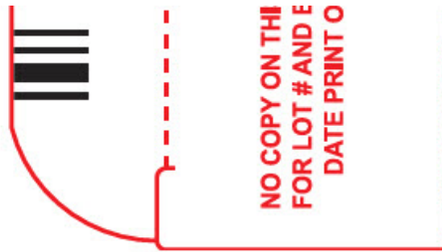
TRIPLE ANTIBIOTIC OINTMENT + PAIN RELIEF

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MAXIMUM STRENGTH

FIRST AID ANTIBIOTIC / PAIN RELIEF OINTMENT





TRIPLE ANTIBIOTIC PLUS PAIN RELIEF

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Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:55315-192
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BACITRACIN ZINC (UNII: 89Y4M234ES) (BACITRACIN - UNII:58H6RWO52I)	BACITRACIN	500 [USP'U] in 1 g
NEOMYCIN SULFATE (UNII: 057Y626693) (NEOMYCIN - UNII:I16QD7X297)	NEOMYCIN SULFATE	3.5 mg in 1 g
POLYMYXIN B SULFATE (UNII: 19371312D4) (POLYMYXIN B - UNII:J2VZ07J96K)	POLYMYXIN B	10000 [USP'U] in 1 g
PRAMOXINE HYDROCHLORIDE (UNII: 88AYB867L5) (PRAMOXINE - UNII:068X84E056)	PRAMOXINE HYDROCHLORIDE	10 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
PETROLATUM (UNII: 4T6H12BN9U)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:55315-192-34	1 in 1 CARTON	01/14/2019	
1		14.2 g in 1 TUBE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M004	03/31/2012	

Labeler - FRED'S, INC. (005866116)

Establishment

Name	Address	ID/FEI	Business Operations
Sun Pharma Canada Inc.		243339023	manufacture(55315-192)

Revised: 7/2025

FRED'S, INC.