TOLNAFATE- tolnaftate jock itch powder spray - talc free aerosol, spray Chain Drug Consortium, LLC

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Premier Value Tolnaftate Jock Itch Powder Spray - Talc Free

Active ingredient

Tolnaftate 1%

Purpose

Antifungal

Uses

- cures most jock itch (tinea cruis)
- relieves itching, burning and crafing associated with jock itch

For external use only.

Flammable:

Do not use while smoking or near heat or flame. Do not puncture or incinerate. Contents under pressure. Do not store at temperature above 120°F

When using this product

- do not get into eyes or mouth, if products get into eyes, rinse eyes thoroughly with water
- use only as directed

Intentional misuse by deliberately concentrating and inhaling contents cans be harmful or fatal.

Stop use and ask a doctor if

- irritation occurs
- no improvement within 2 weeks

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away. Do not use on children under 2 years of age unless directed by a doctor.

Directions

• wash affected area and dry thoroughly

- shake can well and spray a thin layer over affected area twice daily (morning and night)
- supervise children in the use of this product
- use daily for 2 weeks; if conditions persist, consult a doctor
- if nozzle clogs, clean with a pin

Other information

store between 20° and 30°C (68°F and 86°F)

Inactive ingredients

BHT, Isobutane, Kaolin, PPG-12-Buteth-16, SD Alcohol 40-B, Zea Mays (Corn) Starch

Questions?

call 1-866-964-0939

Principal Display Panel
Premier Value
Antifungal
Jock Itch Powder Spray

Tolnaftate 1%

Cures most jock itch

Relieves itching, chafing and burning



Talc- Free

NET WT 4.6 OZ (130 g)

BUTYLATED HYDROXYTOLUENE (UNII: 1P9D0Z171K)

ISOBUTANE (UNII: BXR49TP611)

TOLNAFATE tolnaftate jock itch powder spray - talc free aerosol, spray **Product Information Product Type** HUMAN OTC DRUG **Item Code (Source)** NDC:68016-653 **TOPICAL Route of Administration Active Ingredient/Active Moiety Ingredient Name Basis of Strength** Strength TOLNAFTATE (UNII: 06KB629TKV) (TOLNAFTATE - UNII:06KB629TKV) **TOLNAFTATE** 1.3 g in 130 g **Inactive Ingredients Ingredient Name** Strength

KAOLIN (UNII: 24H4NWX5CO)	
PPG-12-BUTETH-16 (UNII: 58CG7042J1)	
ALCOHOL (UNII: 3K9958V90M)	
ZEA MAYS SUBSP. MAYS WHOLE (UNII: 1G5HNE09V8)	

l	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:68016-653- 46	130 g in 1 CAN; Type 0: Not a Combination Product	10/31/2017	

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph final	part333C	10/31/2017			

Labeler - Chain Drug Consortium, LLC (101668460)

Revised: 1/2023 Chain Drug Consortium, LLC