TOLNAFTATE- tolnaftate jock itch powder spray talc free aerosol, spray Topco Associates LLC

TopcareTolnaftate Jock Itch Powder Spray

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

TopCare health

MEDICATED

Jock Itch

Powder Spray

TOLNAFTATE 1% ANTIFUNGAL

- Cures Most Jock Itch
- Relieves Itching, Chafing and Burning
- Talc- Free

NET WT 4.6 OZ (130 g)



TOLNAFTATE

tolnaftate jock itch powder spray talc free aerosol, spray

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:36800-877

Route of Administration TOPICAL

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
BUTYLATED HYDROXYTOLUENE (UNII: 1P9D0Z171K)	
ISOBUTANE (UNII: BXR49TP611)	
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
		NDC:36800-877- 46	130 g in 1 CAN; Type 0: Not a Combination Product	10/31/2017	

Marketing In	eting Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
OTC Monograph Drug	M005	10/31/2017				

Labeler - Topco Associates LLC (006935977)

Revised: 2/2024 Topco Associates LLC