MICONAZOLE NITRATE- anti-fungal powder miconazole nitrate talc free powder

LEADER/ Cardinal Health 110, Inc.

Leader Miconazorb Antifungal Powder - Talc Free

Active ingredient

Miconazole nitrate 2%

Purpose

Antifungal

Uses

for the cure of most athlete's foot, jock itch and ringworm

Warnings

For external use only.

Do not use

on children 2 years of age unless directed by a doctor.

When using this product

avoid contact with the eyes

Stop and ask a doctor if

irritation occurs or there is no improvement within 4 weeks for athlete's foot and ringworm, or 2 weeks for jock itch.

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

- clean the affected area and dry thoroughly
- apply a thin layer of product over affected area twice daily (morning and night) or as directed by a doctor
- supervise children in the use of this product
- for athlete's foot: pay special attention to spaces between the toes; wear well-fitting, ventilated shoes, and change shoes and socks at least once daily
- for athlete's foot and ringworm, use daily for 4 weeks; for jock itch, use daily for 2

weeks

- if conditions persist longer, consult a doctor
- this product is not effective on the scalp or nails

Other information

- store between 59º 86ºF
- lightly shake bottle to loosen settled powder

Inactive ingredients

allantoin, chloroxylenol, fragrance, imidazolidinyl urea, microcrystalline cellulose, tricalicum phosphate, zea mays (corn) starch

Questions?

call 1-866-964-0939

Principal Display Panel

LEADER

Moisture Absorbing

Antifungal Powder

Miconzole Nitrate 2% | Antfungal

Cures Most Athlete's Foot, Jock Itch, and Ringworm

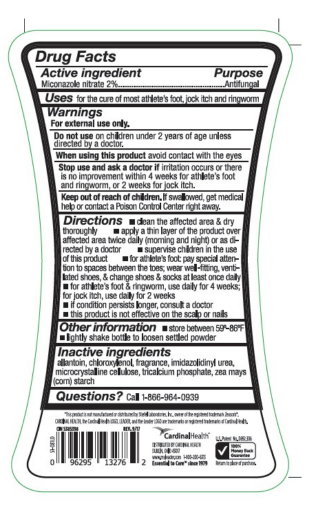
Relieves Chafing, Itching, Burning, and Scaling

Absorbs Moisture

Talc - Free

2.5 OZ (71 g)





MICONAZOLE NITRATE

anti-fungal powder miconazole nitrate talc free powder

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:70000-0323

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength			
MICONAZOLE NITRATE (UNII: VW4H1CYW1K) (MICONAZOLE - UNII:7NNO0D7S5M)	MICONAZOLE NITRATE	1.42 g in 71 g			

Inactive Ingredients			
Ingredient Name	Strength		
ALLANTOIN (UNII: 344S277G0Z)			
CHLOROXYLENOL (UNII: 0F32U78V2Q)			
IMIDUREA (UNII: M629807ATL)			
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)			
TRICALCIUM PHOSPHATE (UNII: K4C08XP666)			

ZEA MAYS SUBSP. MAYS WHOLE (UNII: 1G5HNE09V8)

l	P	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
	1	NDC:70000- 0323-1	71 g in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product	10/20/2017		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC Monograph Drug	M005	10/20/2017			

Labeler - LEADER/ Cardinal Health 110, Inc. (063997360)

Revised: 2/2024 LEADER/ Cardinal Health 110, Inc.