

LOTRIMIN AF JOCK ITCH- miconazole nitrate powder
Bayer HealthCare LLC.

Lotrimin ® AF Jock Itch Powder

Drug Facts

Active ingredient

Miconazole nitrate 2%

Purpose

Antifungal

Uses

- Cures most jock itch (tinea cruris)
- relieves Itching, burning, scaling, discomfort, and chafing associated with jock itch

Warnings

For external use only

Do not use on children under 2 years of age unless directed by a doctor.

When using this product

avoid contact with the eyes

Stop use and ask a doctor if

- irritation occurs
- if there is no improvement within 2 weeks

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- wash affected area and dry thoroughly
- sprinkle 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 condition persists longer, ask a doctor
- this product is not effective on the scalp or nails

Other information

store between 20° to 25°C (68° to 77°F)

Inactive ingredients

benzethonium chloride, corn starch, kaolin, sodium bicarbonate, zinc oxide

PDP**LOTRIMIN ® AF**

miconazole nitrate **ANTIFUNGAL**

MEDICATED POWDER**JOCK ITCH****CLINICALLY PROVEN**

to cure most

jock itch

RELIEVES

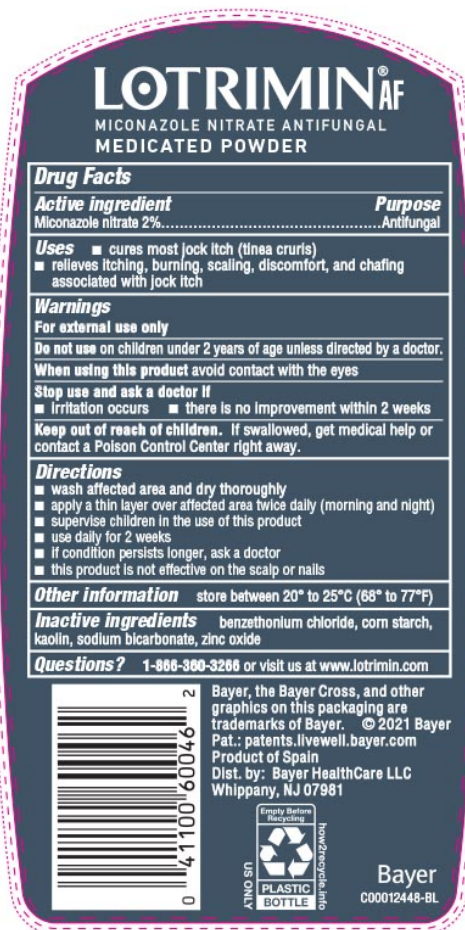
- Itching
- Burning
- Scaling
- Chafing

NET WT 177g (6.25 OZ)

FRONT



BACK

**LOTRIMIN® AF JOCK ITCH**

miconazole nitrate powder

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:11523-0150
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
MICONAZOLE NITRATE (UNII: VW4H1CYW1K) (MICONAZOLE - UNII:7NNO0D7S5M)	MICONAZOLE NITRATE	20 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength

BENZETHONIUM CHLORIDE (UNII: PH41D05744)

STARCH, CORN (UNII: 08232NY3SJ)

KAOLIN (UNII: 24H4NWX5CO)

SODIUM BICARBONATE (UNII: 8MDF5V39QO)

ZINC OXIDE (UNII: SOI2LOH54Z)

Product Characteristics

Color	white (White to off-white)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:11523-0150-1	177 g in 1 CAN; Type 0: Not a Combination Product	11/06/2023	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M005	11/06/2023	

Labeler - Bayer HealthCare LLC. (112117283)

Revised: 1/2026

Bayer HealthCare LLC.