

**LOTRIMIN DAILY PREVENTION- tolnaftate powder**  
**Bayer HealthCare LLC.**

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**Lotrimin Daily Prevention Powder UI 1612510**

*Drug Facts*

**Active ingredient**

Tolnaftate 1%

**Purpose**

Antifungal

**Use**

**Use**

- clinically proven to prevent most athlete's foot with daily use

**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

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

**Directions**

**Directions**

- to prevent athlete's foot, wash the feet and dry thoroughly.
- apply a thin layer of the product on the feet once or twice daily (morning and/or night).
- supervise children in the use of this product.
- pay special attention to spaces between the toes; wear well-fitting ventilated shoes and socks at least once daily.

**Other information**

**Other information**

store between 20° to 25°C (68°- 77°F)

**Inactive ingredients**

benzethonium chloride;corn starch, kaolin;sodium bicarbonate

## **Questions**

**Questions?** 1-866-360-3266

## **Package Display 3 oz. label**

LOTRIMIN® AF

TOLNAFTATE ANTIFUNGAL

medicated foot powder

DAILY

PREVENTION

clinically proven to

prevent most

athlete's foot

- stops the growth of most athlete's foot fungus
- absorbs sweat & keeps feet dry
- destroys odor

NET WT 90g (3 OZ)

**LOTRIMIN<sup>®</sup> AF**  
TOLNAFTATE **ANTIFUNGAL**  
MEDICATED FOOT POWDER

**DAILY  
PREVENTION**

CLINICALLY PROVEN TO  
**PREVENT MOST  
ATHLETE'S  
FOOT**



**3 IN 1**

- stops the growth of most athlete's foot fungus
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C00011377



NET WT 90g (3 OZ)



## LOTRIMIN DAILY PREVENTION

tolnaftate powder

### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:11523-0011
<b>Route of Administration</b>	TOPICAL		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
TOLNAFTATE (UNII: 06KB629TKV) (TOLNAFTATE - UNII:06KB629TKV)	TOLNAFTATE	10 mg in 1 g

**Inactive Ingredients**

Ingredient Name	Strength
KAOLIN (UNII: 24H4NWX5CO)	
SODIUM BICARBONATE (UNII: 8MDF5V39QO)	
STARCH, CORN (UNII: O8232NY3SJ)	
BENZETHONIUM CHLORIDE (UNII: PH41D05744)	

**Product Characteristics**

Color	white	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:11523-0011-1	90 g in 1 BOTTLE; Type 0: Not a Combination Product	02/14/2020	
2	NDC:11523-0011-2	28 g in 1 BOTTLE; Type 0: Not a Combination Product	11/01/2022	

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M005	02/14/2020	

**Labeler** - Bayer HealthCare LLC. (112117283)

Revised: 1/2026

Bayer HealthCare LLC.