

**ZEASORB ATHLETES FOOT- miconazole nitrate powder**  
**Crown Laboratories**

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**Zeasorb Athletes Foot**

***Active ingredient***

Miconazole nitrate 2%

***Purpose***

Antifungal

***Use***

- Proven clinically effective in the treatment of most athlete's foot

***Warnings***

**For external use only.**

**Avoid contact with eyes.**

**Do not use**

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

If irritation occurs or if there is no improvement within 4 weeks, discontinue use and consult a doctor.

**Keep out of reach of children.**

If swallowed, get medical help or contact a poison control center right away.

***Directions***

Clean the affected area and dry thoroughly. Apply a thin layer of the product over affected area twice daily (morning and night) or as directed by a doctor. Supervise children in the use of this product. Pay special attention to spaces between the toes; wear well fitting, ventilated shoes, and change shoes and socks at least once daily. Use daily for 4 weeks. If condition persists longer, consult a doctor. This product is not effective on the scalp or nails.

***Other information***

- store at 20 ° - 25 °C (68 ° - 77 °F) [see USP Controlled Room Temperature]. Product settles during shipment. Package contains full net weight.

### ***Inactive ingredients***

1,2-Hexanediol, Aloe Barbadensis Leaf Juice Powder, Beta-Glucan, Caprylyl Glycol, Fragrance, Glycerin, Lactobacillus Ferment, Potassium Sorbate, Sodium Benzoate, Tapioca Starch, Tricalcium Phosphate, Water, Zea Mays (Corn) Starch

### ***Questions?***

call **1-833-279-6522**

### **Principal Display**

Zeasorb® AF

Antifungal Powder

with 2% Miconazole Nitrate

Cures Most Athlete's Foot

Relieves Itching, Burning, and Scaling

Patented Odor Control Technology

Attacks and Absorbs Moisture

Talc-Free, Paraben Free, Aluminum Free

Dermatologist Recommended

Net wt. 2.5 oz (71g)

P12320.01



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## **ZEASORB ATHLETES FOOT**

miconazole nitrate powder

### **Product Information**

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:0316-8000
<b>Route of Administration</b>	TOPICAL		

### **Active Ingredient/Active Moiety**

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

### **Inactive Ingredients**

<b>Ingredient Name</b>	<b>Strength</b>
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<b>TRICALCIUM PHOSPHATE</b> (UNII: K4C08XP666)	
<b>STARCH, TAPIOCA</b> (UNII: 24SC3U704I)	
<b>WATER</b> (UNII: 059QF0KO0R)	
<b>STARCH, CORN</b> (UNII: O8232NY3SJ)	
<b>CAPRYLYL GLYCOL</b> (UNII: 00YIU5438U)	
<b>1,2-HEXANEDIOL</b> (UNII: TR046Y3K1G)	
<b>ALOE VERA LEAF</b> (UNII: ZY81Z83H0X)	
<b>YEAST .BETA.-D-GLUCAN</b> (UNII: 44FQ49X6UN)	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>POTASSIUM SORBATE</b> (UNII: 1VPU26JZZ4)	
<b>SODIUM BENZOATE</b> (UNII: OJ245FE5EU)	
<b>LACTOBACILLUS ACIDOPHILUS</b> (UNII: 1PRR1V42V5)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0316-8000-01	71 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	10/19/2023	

### Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M005	10/19/2023	

**Labeler** - Crown Laboratories (119508400)

Revised: 1/2026

Crown Laboratories