

LOTRIMIN DAILY SWEAT AND ODOR CONTROL- topical starch powder
Bayer HealthCare LLC.

Lotrimin Daily Sweat and Odor Control Powder UI 1612345

Drug Facts

Active ingredient

Topical starch 83.7%

Purpose

Skin Protectant

Use

Use

temporarily protects and helps relieve minor skin irritation

Warnings

For external use only

Do not use on broken skin.

When using this product

- Do not get into eyes
- Keep away from face and mouth to avoid breathing it

Stop use and ask a doctor if

- Condition worsens
- Symptoms last more than 7 days or clear up and occur again within a few days

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

Directions

Directions

Apply as needed

Inactive ingredients

benzethonium chloride, camphor, eucalyptus globulus leaf oil, fragrance, kaolin, lemon oil, zinc oxide

Questions

Questions? 1-866-360-3266

Package display

LOTRIMIN

TOPICAL STARCH SKIN PROTECTANT

medicated foot powder

DAILY SWEAT

& ODOR CONTROL

6 odor-fighting

ingredients to control odor

- super sweat absorbent powder
- relieves irritation
- talc-free

NET WT 177 G (6.25 OZ)



Relieves irritation

Talc-free

C00011371-FL



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MEDICATED FOOT POWDER

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Questions? 1-866-360-3266 or visit us at www.lotrimin.com



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Dist. by:

Bayer HealthCare LLC
Whippany, NJ 07981



C00011371-BL

Bayer

LOTRIMIN DAILY SWEAT AND ODOR CONTROL

topical starch powder

Product Information

Product Type		HUMAN OTC DRUG	Item Code (Source)		NDC:11523-0012	
Route of Administration		TOPICAL				
Active Ingredient/Active Moiety						
Ingredient Name				Basis of Strength		Strength
STARCH, CORN (UNII: O8232NY3SJ) (STARCH, CORN - UNII:O8232NY3SJ)				STARCH, CORN		837 mg in 1 g
Inactive Ingredients						
Ingredient Name					Strength	
EUCALYPTUS OIL (UNII: 2R04ONI662)						
BENZETHONIUM CHLORIDE (UNII: PH41D05744)						
ZINC OXIDE (UNII: SOI2LOH54Z)						
LEMON OIL (UNII: I9GRO824LL)						
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)						
KAOLIN (UNII: 24H4NWX5CO)						
Product Characteristics						
Color		white	Score			
Shape			Size			
Flavor			Imprint Code			
Contains						
Packaging						
#	Item Code	Package Description		Marketing Start Date		Marketing End Date
1	NDC:11523-0012-1	177 g in 1 BOTTLE; Type 0: Not a Combination Product		02/17/2020		
Marketing Information						
Marketing Category		Application Number or Monograph Citation		Marketing Start Date		Marketing End Date
OTC Monograph Drug		M005		02/17/2020		

Labeler - Bayer HealthCare LLC. (112117283)