

LOTRIMIN ULTRA- butenafine hydrochloride cream
Bayer Healthcare LLC.

Lotrimin Ultra ®

Drug Facts

Active Ingredient

Butenafine hydrochloride 1%

Purpose

Antifungal

Uses

- cures most athlete's foot between the toes. Effectiveness on the bottom or sides of foot is unknown.
- relieves itching, burning, cracking, and scaling which accompany this condition

Warnings

For external use only

- on nails or scalp
- in or near the mouth or the eyes
- for vaginal yeast infections

When using this product do not get into the eyes. If eye contact occurs, rinse thoroughly with water.

Stop use and ask a doctor if too much irritation occurs or irritation gets worse

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

Directions

- **adults and children 12 years and older:**
 - wash the affected skin with soap and water and dry completely before applying
 - to open the applicator, twist the base to the ON position. Squeeze tube to dispense cream.
 - **for athlete's foot between the toes:** apply to affected skin between and around the toes twice a day for 1 week (morning and night), or once a day for 4 weeks, or as directed by a doctor. Wear well-fitting, ventilated shoes. Change shoes and socks at least once daily.
 - wipe applicator tip with a tissue and close the applicator by twisting the base to the OFF position

- wash hands after each use
- children under 12 years: ask a doctor

Other information

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

Inactive ingredients

benzyl alcohol, cetyl alcohol, glycerin, glyceryl monostearate SE, polyoxyethylene (23) cetyl ether, propylene glycol dicaprylate, purified water, sodium benzoate, stearic acid, trolamine, white petrolatum

Questions?

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

LOTRIMIN ULTRA ®

BUTENAFINE HYDROCHLORIDE CREAM 1% ANTIFUNGAL

NET WT 20g (0.70 OZ)



Actual Size 100%

FDA00439-3

PRODUCTS AND EQUIVALENTS		
CATEGORY	IBD	ACTUAL IN ILL
Drug Facts Table	at p.11	3 pt.
Drug Facts Text	at p.11	3 pt.
Directions	11.250000	3 pt.
Warnings	at p.11	3 pt.
Uses	at p.11	3 pt.
Side Effects	at p.11	3 pt.
Interactions	at p.11	3 pt.
Contraindications	at p.11	3 pt.
How and When to Use	at p.11	3 pt.
Other Information	at p.11	3 pt.
Questions?	at p.11	3 pt.

Approved by gebbl on 17-Dec-2020 18:31

LOTTRIMIN ULTRA butenafine hydrochloride cream

Product Information				
Product Type		HUMAN OTC DRUG	Item Code (Source)	NDC:11523-0082
Route of Administration		TOPICAL		
Active Ingredient/Active Moiety				
Ingredient Name			Basis of Strength	Strength
BUTENAFINE HYDROCHLORIDE (UNII: R8XA2029ZI) (BUTENAFINE - UNII:91Y494NLOX)			BUTENAFINE HYDROCHLORIDE	1 g in 100 g
Inactive Ingredients				
Ingredient Name				Strength
STEARIC ACID (UNII: 4ELV7Z65AP)				
TROLAMINE (UNII: 9O3K93S3TK)				
GLYCERIN (UNII: PDC6A3C0OX)				
WHITE PETROLATUM (UNII: B6E5W8RQJ4)				
BENZYL ALCOHOL (UNII: LKG8494WBH)				
CETYL ALCOHOL (UNII: 936JST6JCN)				
WATER (UNII: 059QF0KO0R)				
SODIUM BENZOATE (UNII: OJ245FE5EU)				
CETETH-23 (UNII: 495CTZ441V)				
GLYCERYL MONOSTEARATE (UNII: 230OU9XXE4)				
PROPYLENE GLYCOL DICAPRYLATE (UNII: 581437HWX2)				
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:11523-0082-1	1 in 1 CARTON	12/20/2021	
1		20 g in 1 TUBE, WTH APPLICATOR; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)		



Actual Size 100%

FDA00439-3

DRUG FACTS TABLE (continued)		
CATEGORY	REQUIRED	ACTUAL IN FILE
Drug Facts Table	all	all
Chemical Name	all	all
Indications	all	all
Contraindications	all	all
Warnings	all	all
Directions	all	all
How and how often	all	all
Other information	all	all
Questions	all	all

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Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA021307	12/20/2021	

Labeler - Bayer Healthcare LLC. (112117283)

Revised: 12/2024

Bayer Healthcare LLC.