#### LOTRIMIN ULTRA- butenafine hydrochloride cream Bayer Healthcare LLC.

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#### 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)





CATEGORY	REQUIRED	ACTUAL IN FIL
Drug Facts Title	:8 # <sup>21</sup>	5 pt
Drug Facts Continued	8 pt.	
Headings?	8 pt-bold italic*	1 pt
Sabheedings	36 pt	š pt
Bady Text	36pt	li pt
Bulkts	Spt	5 pt
Leading	30.5 pt	15pt
Hairlines	0.5 pt nuile	1.5pt
Box and Barlines?	30.5 pt.mie	1 pt
Characters Per lach	No more than 39	<u>4</u> 33

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

## LOTRIMIN ULTRA

butenafine hydrochloride cream

Product In	ormation					
Product Typ	e	HUMAN OTC DRUG	Item Code (Source) NDC:11523-008		3-0082	
Route of Ad	ministration	TOPICAL				
Active Ingr	redient/Active	e Moiety				
	Ingr	edient Name		<b>Basis of Str</b>	ength	Strengt
BUTENAFINE   UNII:91Y494NLC		(UNII: R8XA2029ZI) (BUTEN		UTENAFINE YDROCHLORIDE		1 g in 100 g
Inactive In	aredients					
	9	Ingredient Name			St	rength
STEARIC ACID	(UNII: 4ELV7Z65AI	•				
TROLAMINE (L	JNII: 903K93S3TK)					
GLYCERIN (UN	II: PDC6A3C0OX)					
WHITE PETRO	LATUM (UNII: B6E	5W8RQJ4)				
BENZYL ALCO	HOL (UNII: LKG849	94WBH)				
CETYL ALCOH	<b>OL</b> (UNII: 936JST6J	CN)				
WATER (UNII: (	)59QF0KO0R)					
SODIUM BENZ	COATE (UNII: OJ245	FE5EU)				
CETETH-23 (U	NII: 495CTZ 441V)					
GLYCERYL MO	NOSTEARATE (UI	NII: 2300U9XXE4)				
PROPYLENE G	LYCOL DICAPRYL	ATE (UNII: 581437HWX2)				
	naracteristics					
Color	white (wh	nite to off white)	Sco	ore		
Shape			Siz	e		
Flavor			Imp	orint Code		
Contains						
Packaging						
# Item Code		Package Descriptio	'n	Marketir Start Da		arketing nd Date
<b>1</b> NDC:11523- 0082-1	1 in 1 CARTON			12/20/2021		
	20 g in 1 TURE W	ITH APPLICATOR; Type 2: Pr	ofilled Drug Delivery			



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

## **Marketing Information**

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

# Labeler - Bayer Healthcare LLC. (112117283)

Revised: 12/2024

Bayer Healthcare LLC.