

BUTENAFINE HYDROCHLORIDE 1%- butenafine hydrochloride cream
CVS Pharmacy

Butenafine Hydrochloride Cream 1%

Drug Facts

Active ingredient

Butenafine hydrochloride 1%

Purpose

Antifungal

Uses

- cures most jock itch
- relieves itching, burning, cracking, and scaling which accompany this condition

Warnings

For external use only

Do not use

- 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:
 - use the tip of the cap to break the seal and open the tube
 - wash the affected skin with soap and water and dry completely before applying
 - apply once a day to affected skin for 2 weeks or as directed by a doctor
 - wash hands after each use
- children under 12 years: ask a doctor

Other information

- **TAMPER EVIDENT:** DO NOT USE IF THE SEAL ON THE TUBE IS PUNCTURED OR

NOT VISIBLE

- 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?

Call **1-866-923-4914**

Distributed by: CVS Pharmacy, Inc.

One CVS Drive, Woonsocket, RI 02895

PRINCIPAL DISPLAY PANEL - 12 g Tube Carton

CVS
Health®

Compare to the active
ingredient in Lotrimin
Ultra® Jock Itch*

NDC 69842-989-08

Prescription Strength
Butenafine
Hydrochloride
Cream 1%
ANTIFUNGAL CREAM

Relieves itching, burning & chafing

Clinically Proven
to Cure Most
JOCK ITCH

Contains the Drug: Butenafine Hydrochloride

NET WT 12 g (0.42 OZ)

NO VARNISH
ON THIS FLAP



Compare to the active
ingredient in Lotrimin
Ultra® Jock Itch*

NDC 69842-989-08

Prescription Strength
**Butenafine
Hydrochloride
Cream 1%**

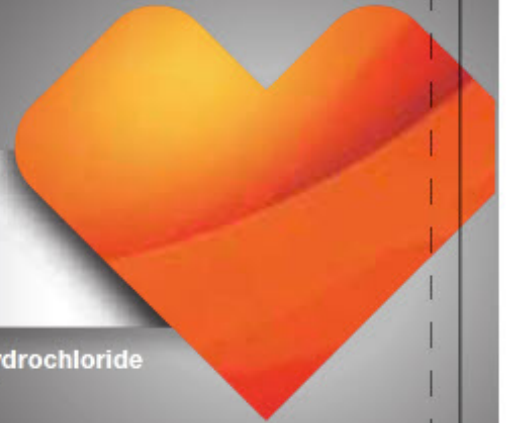
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guaranteed
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T302
B275.0
ENG 19.68



Prescription Strength
**Butenafine
Hydrochloride
Cream 1%**
ANTIFUNGAL CREAM



Compare to the

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Drug Facts (continued)

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Questions? Call
1-866-923-4914

*All trademarks are property of their respective owners. This product is not affiliated with the makers/owners of Lotrimin Ultra® Jock Itch.

CVS
Health.

active ingredient
in Lotrimin Ultra®
Jock Itch®

NDC 69842-989-08

Prescription Strength

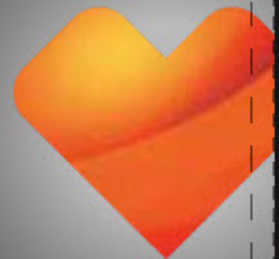
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ANTIFUNGAL CREAM

Relieves itching, burning & chafing

Contains the Drug:
Butenafine Hydrochloride

Clinically Proven to Cure Most
JOCK ITCH



NET WT
12 g (0.42 OZ)

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0323
16

NO VARNISH/NO AQ
NO COPY / NO COLOR
THIS FLAP FOR LOT #
AND EXP DATE PRINT

BUTENAFINE HYDROCHLORIDE 1%

butenafine hydrochloride cream

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69842-989
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
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Butenafine Hydrochloride (UNII: R8XA2029ZI) (Butenafine - UNII:91Y494NLOX)	Butenafine Hydrochloride	10 mg in 1 g
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Inactive Ingredients	
Ingredient Name	Strength
benzyl alcohol (UNII: LKG8494WBH)	
cetyl alcohol (UNII: 936JST6JCN)	
glycerin (UNII: PDC6A3C0OX)	
Glyceryl Stearate Se (UNII: FCZ5MH785I)	
ceteth-23 (UNII: 495CTZ441V)	
propylene glycol dicaprylate (UNII: 581437HWX2)	
water (UNII: 059QF0KO0R)	
sodium benzoate (UNII: OJ245FE5EU)	
stearic acid (UNII: 4ELV7Z65AP)	
trolamine (UNII: 9O3K93S3TK)	
petrolatum (UNII: 4T6H12BN9U)	

Product Characteristics			
Color	WHITE	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69842-989-08	1 in 1 CARTON	10/18/2018	
1		12 g in 1 TUBE; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA205181	10/18/2018	

Labeler - CVS Pharmacy (062312574)