

UPANDUP CLOTRIMAZOLE- clotrimazole cream

Target Corporation

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Up&Up

Clotrimazole 1%

Drug Facts

Active ingredient

Clotrimazole 1%

Purpose

Antifungal

Uses

- cures most athlete's foot, jock itch and ringworm
- relieves itching, burning, cracking, scaling and discomfort which accompany these conditions

Warnings

For external use only

Do not use on children under 2 years of age unless directed by a doctor.

When using this product avoid contact with eyes.

Stop use and ask a doctor if

- irritation occurs
- there is no improvement within 4 weeks (for athlete's foot or ringworm) or 2 weeks (for jock itch).

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

Directions

- wash affected area and dry thoroughly
- apply a thin layer over affected area twice daily (morning and night)
- supervise children in the use of this product
- for athlete's foot, pay special attention to spaces between the toes, wear well-fitting, ventilated shoes and change socks at least once daily
- for athlete's foot and ringworm, use daily for 4 weeks; for jock itch, use daily for 2 weeks
- if condition persists longer, ask a doctor
- this product is not effective on the scalp or nails

Other information

- To open: unscrew cap, use the pointed end of cap to puncture seal.
- store between 20° - 25°C (68° - 77°F)
- see carton or tube crimp for lot number and expiration date

Inactive ingredients

benzyl alcohol (1%), cetostearyl alcohol, cetyl esters wax, 2-octyldodecanol, polysorbate 60, purified water, sorbitan monostearate

Questions?

Call 1-800-910-6874

Dist. by Target Corp., Mpls., MN 55403

PRINCIPAL DISPLAY PANEL - 30 g Tube Carton

up&up

athlete's foot cream

clotrimazole cream USP, 1%

antifungal

helps cure most athlete's foot

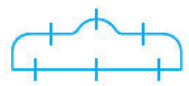
helps relieve burning and itching

NET WT 1 OZ (30 g)



NDC 11673-910-12

Compare to active ingredient in Lotrimin® AF
athlete's foot cream
clotrimazole cream USP, 1%
antifungal



LPK-3766-10
0515-10
M43



athlete's foot cream
clotrimazole cream USP, 1%
antifungal
helps cure most athlete's foot
helps relieve burning and itching

NET WT 1 OZ (30 g)



T30



Dist. by Target Corp., Minneapolis, MN 55403
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245060157 R00
IDC-000021-01-108

Drug Facts (continued)
Inactive ingredients Benzyl alcohol (1%), cetostearyl alcohol, cetyl esters wax, 2-octyldodecanol, polyorbate O, purified water, sorbitan monostearate
Questions? Call 1-800-910-6874

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

This product is not
manufactured or
distributed by MSP
Consumer Care, Inc.
owner of the registered
trademark Lotrimin® AF.

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UPANDUP CLOTRIMAZOLE

clotrimazole cream

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:11673-910
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
Clotrimazole (UNII: G07GZ97H65) (Clotrimazole - UNII:G07GZ97H65)	Clotrimazole	10 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
benzyl alcohol (UNII: LKG8494WBH)	
cetostearyl alcohol (UNII: 2DMT128M1S)	
cetyl esters wax (UNII: D072FFP9GU)	
octyldodecanol (UNII: 461N1O614Y)	
polysorbate 60 (UNII: CAL22UVI4M)	
water (UNII: 059QF0K00R)	
sorbitan monostearate (UNII: NVZ4I0H58X)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:11673-910-12	1 in 1 CARTON		
1		30 g in 1 TUBE; Type 0: Not a Combination Product		
2	NDC:11673-910-16	1 in 1 CARTON		
2		40 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
OTC MONOGRAPH FINAL	part333C	08/31/1993	

Labeler - Target Corporation (006961700)

Revised: 6/2015

Target Corporation