#### KETOCONAZOLE- ketoconazole cream Rebel Distributors Corp

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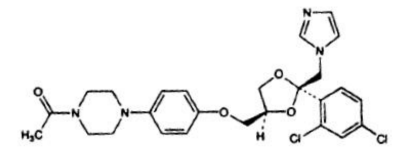
Ketoconazole Cream, 2%

Rx only

#### DESCRIPTION

Ketoconazole cream, 2% contains the broad-spectrum synthetic antifungal agent, ketoconazole 2%, formulated in an aqueous cream vehicle consisting of butylated hydroxyanisole (BHA), cetyl alcohol, isopropyl myristate, polysorbate 60, polysorbate 80, propylene glycol, purified water, sorbitan monostearate and stearyl alcohol.

Ketoconazole is *cis*-1-acetyl-4-[4-[[2-(2,4-dichlorophenyl)-2-(1*H*-imidazol-1-ylmethyl)-1,3-dioxolan-4-yl]methoxy]phenyl] piperazine and has the following structural formula:



Molecular Formula: C<sub>26</sub>H<sub>28</sub>Cl<sub>2</sub>N<sub>4</sub>O<sub>4</sub> Molecular Weight: 531.43

### **CLINICAL PHARMACOLOGY**

When ketoconazole cream, 2% was applied dermally to intact or abraded skin of beagle dogs for 28 consecutive days at a dose of 80 mg, there were no detectable plasma levels using an assay method having a lower detection limit of 2 ng/mL.

After a single topical application to the chest, back and arms of normal volunteers, systemic absorption of ketoconazole was not detected at the 5 ng/mL level in blood over a 72-hour period.

Two dermal irritancy studies, a human sensitization test, a phototoxicity study and a photoallergy study conducted in 38 male and 62 female volunteers showed no contact sensitization of the delayed hypersensitivity type, no irritation, no phototoxicity and no photoallergenic potential due to ketoconazole cream, 2%.

### Microbiology

Ketoconazole is a broad spectrum synthetic antifungal agent which inhibits the *in vitro* growth of the following common dermatophytes and yeasts by altering the permeability of the cell membrane: dermatophytes: *Trichophyton rubrum*, *T. mentagrophytes*, *T. tonsurans*, *Microsporum canis*, *M. audouini*, *M. gypseum* and *Epidermophyton floccosum*; *yeasts: Candida albicans*, *Malassezia ovale* (*Pityrosporum ovale*) and *C. tropicalis*; and the organism responsible for tinea versicolor, *Malassezia furfur* (*Pityrosporum orbiculare*). Only those organisms listed in the **INDICATIONS AND USAGE** section

have been proven to be clinically affected. Development of resistance to ketoconazole has not been reported.

# **Mode of Action**

*In vitro* studies suggest that ketoconazole impairs the synthesis of ergosterol, which is a vital component of fungal cell membranes. It is postulated that the therapeutic effect of ketoconazole in seborrheic dermatitis is due to the reduction of M. ovale, but this has not been proven.

# **INDICATIONS AND USAGE**

Ketoconazole cream, 2% is indicated for the topical treatment of tinea corporis, tinea cruris and tinea pedis caused by *Trichophyton rubrum*, *T. mentagrophytes* and *Epidermophyton floccosum*; in the treatment of tinea (pityriasis) versicolor caused by *Malassezia furfur (Pityrosporum orbiculare)*; in the treatment of cutaneous candidiasis caused by *Candida spp*. and in the treatment of seborrheic dermatitis.

### CONTRAINDICATIONS

Ketoconazole cream, 2% is contraindicated in persons who have shown hypersensitivity to the active or excipient ingredients of this formulation.

### WARNINGS

Ketoconazole cream, 2% is not for ophthalmic use.

# PRECAUTIONS

### General

If a reaction suggesting sensitivity or chemical irritation should occur, use of the medication should be discontinued. Hepatitis (1:10,000 reported incidence) and, at high doses, lowered testosterone and ACTH induced corticosteroid serum levels have been seen with orally administered ketoconazole; these effects have not been seen with topical ketoconazole.

### Carcinogenesis, Mutagenesis, Impairment of Fertility

A long-term feeding study in Swiss Albino mice and in Wistar rats showed no evidence of oncogenic activity. The dominant lethal mutation test in male and female mice revealed that single oral doses of ketoconazole as high as 80 mg/kg produced no mutation in any stage of germ cell development. The Ames' *salmonella* microsomal activator assay was also negative.

### Pregnancy

Teratogenic effects

### Pregnancy Category C

Ketoconazole has been shown to be teratogenic (syndactylia and oligodactylia) in the rat when given orally in the diet at 80 mg/kg/day, (10 times the maximum recommended human oral dose). However, these effects may be related to maternal toxicity, which was seen at this and higher dose levels.

There are no adequate and well-controlled studies in pregnant women. Ketoconazole should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

### **Nursing Mothers**

It is not known whether Ketoconazole cream, 2% administered topically could result in sufficient

systemic absorption to produce detectable quantities in breast milk. Nevertheless, a decision should be made whether to discontinue nursing or discontinue the drug, taking into account the importance of the drug to the mother.

# **Pediatric Use**

Safety and effectiveness in children have not been established.

# **ADVERSE REACTIONS**

During clinical trials 45 (5.0%) of 905 patients treated with ketoconazole cream, 2% and 5 (2.4%) of 208 patients treated with placebo reported side effects consisting mainly of severe irritation, pruritus and stinging. One of the patients treated with ketoconazole cream developed a painful allergic reaction.

In worldwide postmarketing experience, rare reports of contact dermatitis have been associated with ketoconazole cream or one of its excipients, namely propylene glycol.

# DOSAGE AND ADMINISTRATION

### Cutaneous candidiasis, tinea corporis, tinea cruris, tinea pedis, and tinea (pityriasis) versicolor

It is recommended that ketoconazole cream, 2% be applied once daily to cover the affected and immediate surrounding area. Clinical improvement may be seen fairly soon after treatment is begun; however, candidal infections and tinea cruris and corporis should be treated for two weeks in order to reduce the possibility of recurrence.

Patients with tinea versicolor usually require two weeks of treatment. Patients with tinea pedis require six weeks of treatment.

### Seborrheic dermatitis

Ketoconazole cream, 2% should be applied to the affected area twice daily for four weeks or until clinical clearing.

If a patient shows no clinical improvement after the treatment period, the diagnosis should be redetermined.

### HOW SUPPLIED

Ketoconazole cream, 2% is supplied in tubes of:

15gm NDC 21695-554-15,

30gm NDC 21695-555-30,

60gmNDC 21695-556-60

Store at 20°-25°C (68°-77°F) [see USP Controlled Room Temperature].

Manufactured by: Taro Pharmaceuticals Inc., Brampton, Ontario, Canada L6T 1C1

Manufactured for: **TIBER LABORATORIES**, Suwanee, GA 30024

Issued: April, 2010 REV. 04/10 054-10

PK-6519-0 97

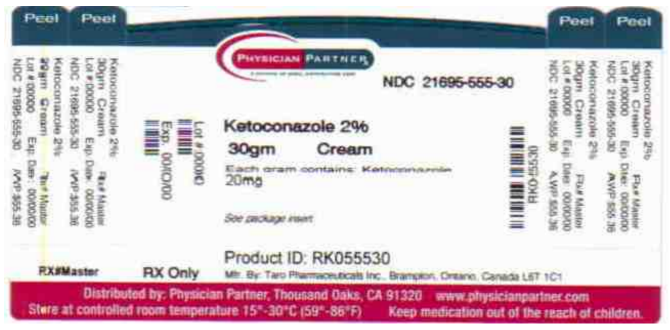
# Repackaged by: **REBEL DISTRIBUTORS CORP**

Thousand Oaks, CA 91320

# **Principal Display Panel**



# **Principal Display Panel**



**Principal Display Panel** 



KETOCONAZOLE					
ketoconazole cream					
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	G Item Code (Se	ource)	NDC:21695-554(N	NDC:51672-1298)
Route of Administration	TOPICAL				
Active Ingredient/Active Moi	etv				
0	redient Name		Bas	sis of Strength	Strength
Ketoconazole (UNII: R9400W927I) (Ko		I)		onazole	20 mg in 1 g
Inactive Ingredients					
	Ingredient Name				Strength
Butylated Hydroxyanisole (UNII: REK	4960K2U)				
cetyl alcohol (UNII: 936JST6JCN)					
isopropyl myristate (UNII: 0RE8K4LN	JS)				
polysorbate 60 (UNII: CAL22UVI4M)					
polysorbate 80 (UNII: 6OZP39ZG8H)					
propylene glycol (UNII: 6DC9Q167V3	)				
water (UNII: 059QF0KO0R)					
sorbitan monostearate (UNII: NVZ4I0	H58X)				
stearyl alcohol (UNII: 2KR8914H1Y)					
Product Characteristics					
Color	WHITE Sco	re			
Shape	Size				
Flavor		rint Code			

Contains								
Packaging								
# Item Code	Pacl	kage Description	Marketin	o Start I	Date	Marketin	ig End Date	
1 NDC:21695-554-15	1 in 1 CAF		What is the	ig oturt i	Juic	Whit is the	ig Liid Date	
1	15 g in 1 T							
-	8							
Marketing Info	rmation							
Marketing Category	Applicatio	on Number or Monog	raph Citation	Market	ting Star	t Date Mark	eting End Dat	
ANDA	ANDA075638				12/18/2002			
ketoconazole cream <b>Product Informati</b> Product Type Route of Administrat		HUMAN PRESCRIPTIO	N DRUG Item	Code (So	urce) N	IDC:21695-555(1	NDC:51672-1298	
Product Type	ion	TOPICAL	N DRUG Item	Code (So	urce) N	IDC:21695-555(I	NDC:51672-1298	
<b>Product Informati</b> Product Type Route of Administrat	ion 'Active Moi	TOPICAL	N DRUG	Code (So		IDC:21695-555(I of Strength		
<b>Product Informati</b> Product Type Route of Administrat	ion 'Active Moi Ing	TOPICAL e ty redient Name		Code (So		of Strength	NDC:51672-1298 Strength 20 mg in 1 g	
Product Informati Product Type Route of Administrat Active Ingredient/ Ketoconazole (UNII: RS	ion ( <b>Active Moi</b> Ing1 0400W927I) (Ka	TOPICAL e ty redient Name		Code (So	Basis	of Strength	Strength	
Product Informati Product Type Route of Administrat Active Ingredient/ Ketoconazole (UNII: RS	ion ( <b>Active Moi</b> Ing1 0400W927I) (Ka	TOPICAL e ty redient Name	0 W9 27 I)	Code (So	Basis	<b>of Strength</b> azole	Strength	
Product Informati Product Type Route of Administrat Active Ingredient/ Ketoconazole (UNII: R9 Inactive Ingredien	ion ' <b>Active Moi</b> Ing: 1400W927I) (Ko <b>Its</b>	TOPICAL ety redient Name etoconazole - UNII:R940 Ingredient Name	0 W9 27 I)	Code (So	Basis	<b>of Strength</b> azole	Strength 20 mg in 1 g	
Product Informati Product Type Route of Administrat Active Ingredient/ Ketoconazole (UNII: R9 Inactive Ingredien Butylated Hydroxyanis	ion (Active Moi Ing 1400 W9 27I) (Ka 145 145 101 Ka 101	TOPICAL ety redient Name etoconazole - UNII:R940 Ingredient Name	0 W9 27 I)	Code (So	Basis	<b>of Strength</b> azole	<b>Strength</b> 20 mg in 1 g	
Product Informati Product Type Route of Administrat Active Ingredient/ Ketoconazole (UNII: RS Inactive Ingredien Butylated Hydroxyanis cetyl alcohol (UNII: 936	ion (Active Moi Ing: 0400W927I) (Ko 0400W927I) (Ko 0400W927I) (Sole (UNII: REK 635T6JCN)	TOPICAL ety redient Name etoconazole - UNII:R940 Ingredient Name	0 W9 27 I)	Code (So	Basis	<b>of Strength</b> azole	<b>Strength</b> 20 mg in 1 g	
Product Informati Product Type Route of Administrat Active Ingredient/	ion (Active Moi Ing 400 W927I) (Ka 400 W927I) (Ka 501e (UNII: REK 501e (UNII: REK 515T6JCN) NII: 0 RE8K4LN	TOPICAL ety redient Name etoconazole - UNII:R940 Ingredient Name	0 W9 27 I)	Code (So	Basis	<b>of Strength</b> azole	<b>Strength</b> 20 mg in 1 g	
Product Informati Product Type Route of Administrat Active Ingredient/ Ketoconazole (UNII: R9 Inactive Ingredien Butylated Hydroxyanis cetyl alcohol (UNII: 936 isopropyl myristate (U	ion (Active Moi Ing: 0400 W927I) (Ko 0400 W927I) (Ko 1ts sole (UNII: REK 5JST6JCN) NII: 0 RE8K4LN CAL22UVI4M)	TOPICAL ety redient Name etoconazole - UNII:R940 Ingredient Name	0 W9 27 I)	Code (So	Basis	<b>of Strength</b> azole	Strength 20 mg in 1 g	

water (UNII: 059QF0KO0R) sorbitan monostearate (UNII: NVZ4I0H58X)

stearyl alcohol (UNII: 2KR8914H1Y)

Product Characteristics						
Color	WHITE	Score				
Shape		Size				
Flavor		Imprint Code				
Contains						

Packaging									
# Item Code	Pack	age Description	Μ	arketir	ig Start I	Date	Μ	arketin	g End Date
1 NDC:21695-555-30	1 in 1 CAF								
1	30 g in 17	TUBE							
Marketing Info	rmation								
Marketing Category	Applicatio	n Number or Monogra	aph Cit	ation	Marke	ting St	art Date	Mark	eting End Date
ANDA	ANDA075638				12/18/20	02			
KETOCONAZO	LE								
ketoconazole cream									
Product Information	on								
Product Type		HUMAN PRESCRIPTION	I DRUG	Ite m C	Code (So	urce)	NDC:2169	95-556(N	NDC:51672-1298
Route of Administration	on	TOPICAL							
Active Ingredient//		ety edient Name				Bas	is of Stro	ength	Strength
Ketoconazole (UNII: R94			) W9 27 I)				nazole	0	20 mg in 1 g
Inactive Ingredien	ts								
		Ingredient Name							Strength
Butylated Hydroxyaniso		4960K2U)							
cetyl alcohol (UNII: 936									
isopropyl myristate (UN		(S)							
polysorbate 60 (UNII: C.									
polysorbate 80 (UNII: 60									
propylene glycol (UNII: water (UNII: 059QF0KO)									
sorbitan monostearate		458 X )							
stearyl alcohol (UNII: 2)									
Product Character	istics								

Color		WHITE	Score		
Shape	2		Size		
Flavo	r		Imprint Code		
Conta	ins				
Pack	aging				
#	Item Code	Package Description	Marketing Start Date	Marketing End	d Date

1 NDC:21695-556-60	1 in 1 CARTON		
1	60 g in 1 TUBE		
Marketing Info	rmation		
Marketing Info Marketing Category	r <b>mation</b> Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
0		Marketing Start Date	Marketing End Date

Labeler - Rebel Distributors Corp (118802834)

Establishment							
Name	Address	ID/FEI	<b>Business Operations</b>				
Rebel Distributors Corp		118802834	RELABEL, REPACK				

Revised: 1/2011

Rebel Distributors Corp