CANDIDA- candida parapsilosis liquid USPharmaCo

Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.

CANDIDA DROPS 4X

Homeopathic Medicine

Indications

For temporary relief of external anal or vaginal itching and irritation

Dosage

5-10 drops, three times daily.

Active Ingredient

Candida parapsilosis 4X

Inactive Ingredients

Purified water, sodium chloride, potassium sorbate.

Warning

If symptoms persist more than a few days, contact a licensed practitioner. As with any drug, if you are pregnant or nursing, seek the advice of a health care professional before using this product.

Keep this and all medications out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.

Protect from light and heat.

Tamper Evident

Do not use product if the tamper evident strip is broken or removed from the base of the cap.

To report serious adverse events, call: 1-877-557-4276 or 1-623-582-3110

Manufactured for and distributed by:

USPharmaCo Distribution Ltd. 2205 W. Lone Cactus Drive #19, Phoenix, AZ 85027 www.uspharmaco.com info@uspharmaco.com

Made in Canada

PRINCIPAL DISPLAY PANEL - 10 mL Label CANDIDA DROPS 4X

Homeopathic Medicine 0.34 fl. oz. (10ml)

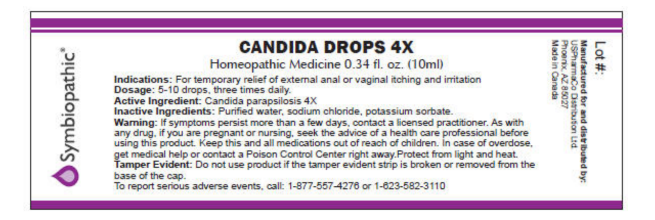
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CANDIDA

candida parapsilosis liquid

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:49693-1101
Route of Administration	ORAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
Candida parapsilosis (UNII: 0KZ676D44N) (Candida parapsilosis - UNII:0KZ676D44N)	Candida parapsilosis	4 [hp_X] in 10 mL	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:49693-1101-1	1 in 1 BOX		
1		10 mL in 1 BOTTLE		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED HOMEOPATHIC		12/15/2009	

Labeler - USPharmaCo (145322622)

Establishment				
Name	Address	ID/FEI	Business Operations	
DermaMed		256799461	MANUFACTURE	

Revised: 12/2009 USPharmaCo