PSORIASIS REAL RELIEF- sulphur, arsenicum album, alumen, magnesia sulphuricum, natrum sulphuricum tablet, chewable HOMEOLAB USA INC

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.

DRUG FACTS

ACTIVE INGREDIENTS HPUS

Sulphur (Sublimed sulfur) 8X Arsenicum album (Arsenous oxide) 8X Alumen (Potassium alum) 8X Magnesia sulphurica (Magnesium sulfate) 8X Natrum sulphuricum (Disodium sulfate) 8X

The letters 'HPUS' indicate that the components in this product are officially monographed in the Homeopathic Pharmacopoeia of the United States.

PURPOSE

burning itching peeling dry skin dry and cracked skin, reddish spots, burning itching (fish-scales), itchy pimples itching and redness stinging itching spots on skin itching pimples (moist and oozing)

USES

This homeopathic medicine helps relieve symptoms of psoriasis:

- dry peeling skin
- stinging
- burning itching
- redness
- itching spots

WARNINGS

Stop use and ask a doctor if symptoms worsen or persist for more than 7 days.

If pregnant or breast-feeding, ask a health professional before use.

Keep out of reach of children.

DIRECTIONS

Chew tablets and let dissolve in mouth. Do not use more than directed. Do not take with food.

Repeat 3 times daily and reduce with improvement or as directed by a health professional.

Age	Dose
Adults and children 12 years of age and older	2 tablets
Children 2 to 11 years of age	1 tablet
Children under 2 years of age	Ask a doctor

OTHER INFORMATION

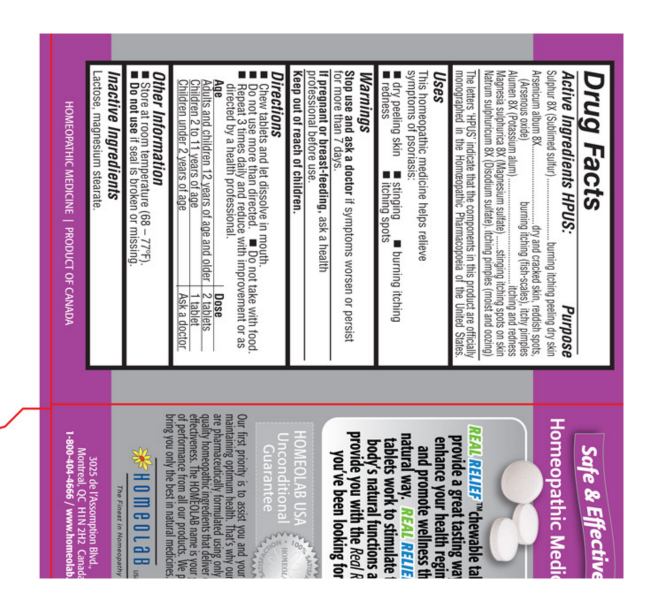
Store at room temperature (68 - 77F).

Do not use if seal is broken or missing.

INACTIVE INGREDIENTS

Lactose, magnesium stearate.

CARTON





PSORIASIS REAL RELIEF

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Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source) NDC:60512-6010		0 10	
Route of Administration	ORAL				
Active Ingredient/Active Moi	ety				
Ing	gredient Name		Basis of	fStrength	Strength
SULFUR (UNII: 70 FD1KFU70) (SULFUR - UNII:70 FD1KFU70)			SULFUR		8 [hp_X]
ARSENIC TRIOXIDE (UNII: S7V92P67HO) (ARSENIC CATION (3+) - UNII:C96613F5AV)			ARSENIC CA	ATION (3+)	8 [hp_X]
POTASSIUM ALUM (UNII: 1L24V9R23S) (ALUMINUM HYDROXIDE - UNII:5QB0T2IUN0)			ALUMINUM	HYDRO XIDE	8 [hp_X]
		MAGNESIUM SULFATE HEPTAHYDRATE (UNII: SK47B8698T) (MAGNESIUM CATION - UNII:T6 V3LHY838)			

UNII:36 KC S0 R750	DIUM SULFATE (UNII: 0 YPR65R21J) (SODIUM SULFATE ANHYDROUS - SODIUM SUL II:36 KCS0 R750) ANHYDROUS					8 [hp_X]	
Inactive Ingre	edients						
Ingredient Name					Sti	Strength	
LACTOSE (UNII:	J2B2A4N98C	i)					
MAGNESIUM STI	E ARATE (UN	II: 70097M6I30)					
Product Char	acteristics	5					
Color		white	Score no scor				
Shape		ROUND	Size 8mm				
Flavor			Imprint Code				
Contains							
Packaging							
	ode	Package Description	n Marketing	g Start Date	Marketing	End Date	
# Item Co		Package Descriptio	n Marketing	g Start Date	Marketing	End Date	
# Item Co		U	n Marketing	g Start Date	Marketing	End Date	
# Item Co		U	n Marketin _i	g Start Date	Marketing	End Date	
Packaging # Item Co 1 NDC:60512-601 Marketing	10-0 9	00 in 1 BOTTLE	n Marketin _i	g Start Date	Marketing	End Date	
# Item C 1 NDC:60512-601	10-0 9 Informa	00 in 1 BOTTLE		g Start Date Marketing Start D		End Date ing End Dat	

Labeler - HOMEOLAB USA INC (202032533)

Establishment					
Name	Address	ID/FEI	Business Operations		
HOMEOLAB USA INC		202032533	manufacture(60512-6010)		

Revised: 7/2012

HOMEOLAB USA INC