DULCAMARA COMP.- dulcamara comp. liquid Uriel Pharmacy 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.

Dulcamara comp.

Directions: FOR ORAL USE ONLY.

Take 3-4 times daily. Ages 12 and older: 10 drops. Ages 2-11: 5 drops. Under age 2: Consult a doctor.

Active Ingredients: Glandulae suprarenales (Bovine adrenal glands) 5X, Arsenicum album (White arsenic) 6X, Dulcamara (Bittersweet) 6X, Lobelia 6X

Inactive Ingredients: Distilled water, Organic cane alcohol

Use: Temporary relief of skin rash.

KEEP OUT OF REACH OF CHILDREN.

Warnings: Claims based on traditional homeopathic practice, not accepted medical evidence. Not FDA evaluated. Do not use if allergic to any ingredient. Contains traces of lactose. Consult a doctor before use for serious conditions or if conditions worsen or persist. If pregnant or nursing, consult a doctor before use. Do not use if safety seal is broken or missing.

Questions? Call 866.642.2858 Made with care by Uriel, East Troy, WI 53120 www.urielpharmacy.com



DULCAMARA COMP. dulcamara comp. liquid					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Sour	ce)	NDC:48951-4010	
Route of Administration	ORAL				
Active Ingredient/Active Moi	ety				
Ing	Basis of Strength		Strength		
BOS TAURUS ADRENAL GLAND (UNII: M2776SWB29) (BOS TAURUS ADRENAL GLAND - UNII:M2776SWB29)			BOS TAURUS ADRENAL GLAND		5 [hp_X] in 1 mL
					6 [bp V]

ARSENIC TRIOXIDE (UNII: S7V92P67HO) (ARSENIC CATION (3+) - UNII:C96613F5AV) ARSENIC TR					OXIDE	ο μιρ_λι in 1 mL	
SOLANUM DULCAMARA TOP (UNII: KPS1B1162N) (SOLANUM DULCAMARA TOP - UNII: KPS1B1162N)SOLANUM D TOP					JLCAMARA	6 [hp_X] in 1 mL	
LOBELIA SPICATA LEAF (UNII: 1G4GK01F67) (LOBELIA SPICATA LEAF - UNII: 1G4GK01F67)				LOBELIA SPI	CATA LEAF	6 [hp_X] in 1 mL	
I	nactive Ingred	ients					
			Ingredient Name			Stre	ngth
W	WATER (UNII: 059QF0KO0R)						
S	O DIUM CHLO RID	E (UNI	: 451W47IQ8X)				
Packaging							
#	Item Code		Package Description	Package Description Marketing Sta Date		t Marketing End Date	
1	NDC:48951-4010- 3	60 mI Produ	in 1 BOTTLE, DROPPER; Type 0: Not a Combination ct	09/0	01/2009		
Marketing Information							
	Marketing Categ	ory	Application Number or Monograph Citation	Marke	ting Start Date	e Marketi	ng End Date
uı	happroved homeopa	athic		09/01/20	09		

Labeler - Uriel Pharmacy Inc. (043471163)

Establishment						
Name	Address	ID/FEI	Business Operations			
Uriel Pharmacy Inc.		043471163	manufacture(48951-4010)			

Revised: 4/2018

Uriel Pharmacy Inc.