CANTHARIDIN 0.7% - cantharidin 0.7% liquid Sincerus Florida, LLC

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

CANTHARIDIN 0.7%

Directions for use

Active ingredients Cantharidin Inactive ingredients Acetone NF Hexible Collodion USP Hydroxypropyl Cellulose NF 0.7% 0.7%

Directions for use

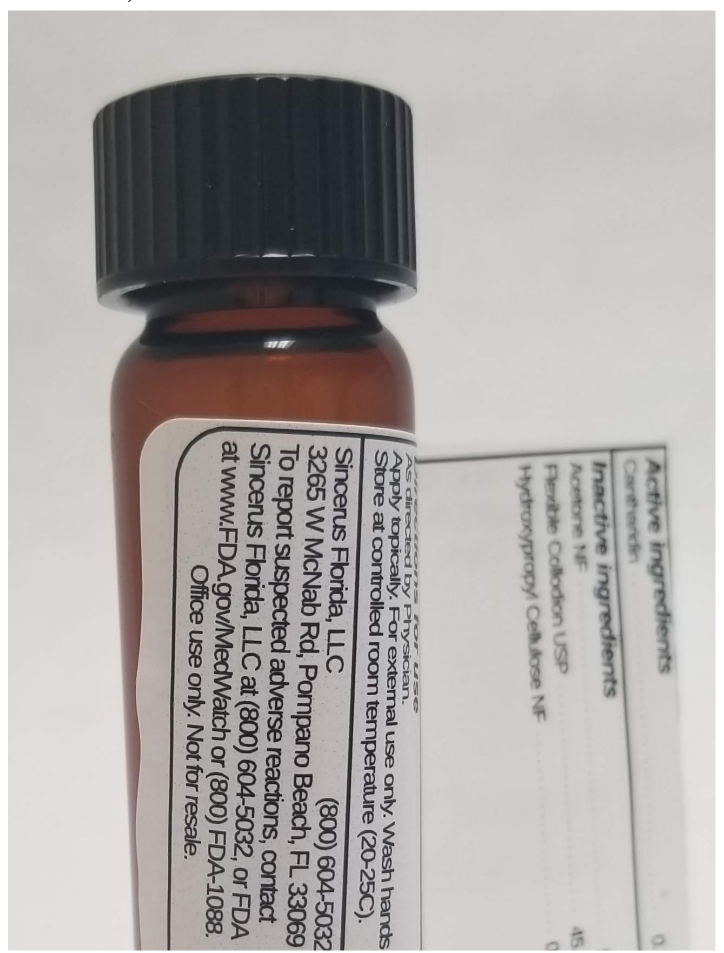
As directed by Physician.

Apply topically. For external use only. Wash hands after use. Store at controlled room temperature (20-25C).

Sincerus Florida, LLC (800) 604-5032
3265 W McNab Rd, Pompano Beach, FL 33069
To report suspected adverse reactions, contact Sincerus Florida, LLC at (800) 604-5032, or FDA at www.FDA.gov/Medwatch or (800) FDA-1088.



Sincerus Florida, LLC. Adverse reactions





Active, inactive

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CANTHARIDIN 0.7%

cantharidin 0.7% liquid

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:72934-9035
Route of Administration	TOPICAL		

ı	Active Ingredient/Active Moiety			
l	Ingredient Name	Basis of Strength	Strength	
ı	CANTHARIDIN (UNII: IGL471WQ8P) (CANTHARIDIN - UNII:IGL471WQ8P)	CANTHARIDIN	0.7 g in 100 g	

Product Characteristics			
Color	white (clear liquid)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging				
# Item Code Package Description		Marketing Start Date	Marketing End Date	
ı	1 NDC:72934-9035-9	15 g in 1 VIAL; Type 0: Not a Combination Product	05/17/2019	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		05/17/2019		

Labeler - Sincerus Florida, LLC (080105003)

Establishment					
Name	Address	ID/FEI	Business Operations		
Sincerus Florida, LLC		080105003	manufacture(72934-9035)		

Revised: 5/2019 Sincerus Florida, LLC