

PENTOXIFYLLINE 0.5% / TRIAMCINOLONE ACETONIDE 0.1% - pentoxifylline 0.5% / triamcinolone acetonide 0.1% gel

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](#).

PENTOXIFYLLINE 0.5% / TRIAMCINOLONE ACETONIDE 0.1%

Directions for use



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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.

Office use only. Not for resale.



Sincerus Florida, LLC. Adverse reactions



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Sincerus Florida, LLC (800

3265 W McNab Rd, Pompano Beach

To report suspected adverse reactions

Sincerus Florida, LLC at (800) 604-503

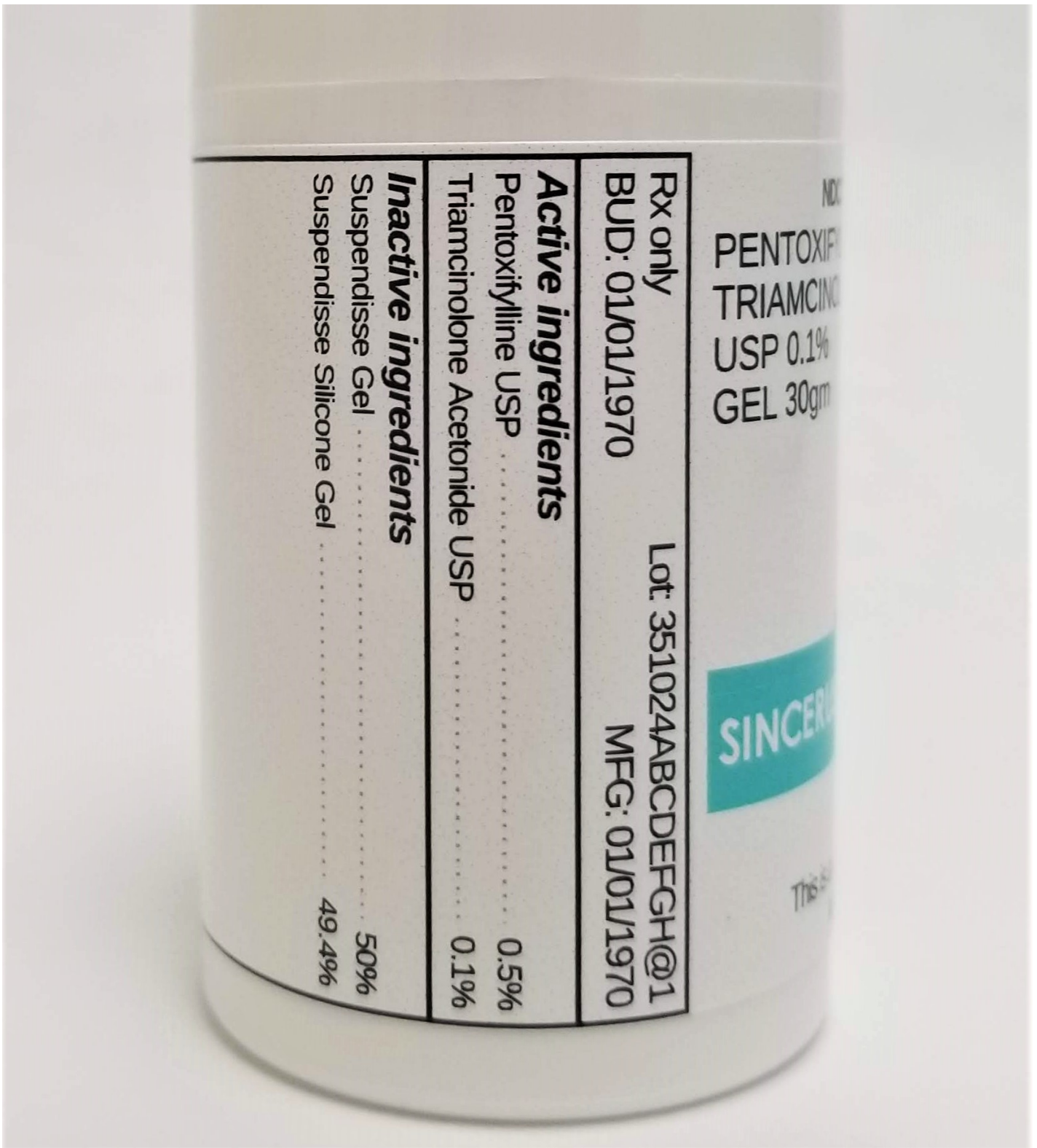
at www.FDA.gov/MedWatch or (800) F

Office use only. Not for resa



Active, inactive





NDC 72934- 1169-2 PENTOXIFYLLINE USP 0.5% / TRIAMCINOLONE ACETONIDE USP 0.1%. Gel 30gm.



NDC 72934-1169-2

PENTOXIFYLLINE USP 0.5%
TRIAMCINOLONE ACETONIDE
USP 0.1%
GEL 30gm



PENTOXIFYLLINE 0.5% / TRIAMCINOLONE ACETONIDE 0.1%

pentoxifylline 0.5% / triamcinolone acetonide 0.1% gel

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:72934-1169
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
PENTOXIFYLLINE (UNII: SD6QCT3TSU) (PENTOXIFYLLINE - UNII:SD6QCT3TSU)	PENTOXIFYLLINE	0.5 g in 100 g
TRIAMCINOLONE ACETONIDE (UNII: F446C597KA) (TRIAMCINOLONE ACETONIDE - UNII:F446C597KA)	TRIAMCINOLONE ACETONIDE	0.1 g in 100 g

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72934-1169-2	30 g in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	05/15/2019	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		05/15/2019	

Labeler - Sincerus Florida, LLC (080105003)

Establishment

Name	Address	ID/FEI	Business Operations
Sincerus Florida, LLC		080105003	manufacture(72934-1169)

Revised: 5/2019

Sincerus Florida, LLC