

SULFUR SPOT TREATMENT- sulfur acne treatment cream
Banuskín, Inc.

Sulfur Spot Treatment

Active Ingredients

Sulfur 10%

Purpose

Acne treatment

Uses

For the treatment of acne

Warnings

For external use only

When using this product

When using this product

- skin irritation and dryness more likely to occur if you use another topical acne medication at the same time. If irritation occurs, only use one topical acne medication at a time.

Do not use on

Do not use on

- broken skin
- large areas of the skin

When using this product

When using this product

- apply only to areas with acne

Keep out of reach of children.

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

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- Clean the skin thoroughly before applying this product.
- Cover the entire affected area with a thin layer one to three times daily.
- Because excessive drying of the skin may occur, start with one application daily, then gradually increase to two or three times daily if needed or as directed by a doctor.
- If bothersome dryness or peeling occurs, reduce application to once a day or every other day.

Inactive Ingredients

Aqua/Water/Eau, Glycerin, Kaolin, Bentonite, Propylene Glycol, Niacinamide, Sorbitol, Titanium Dioxide (CI 77891), Phenoxyethanol, Hamamelis Virginiana (Witch Hazel) Water, Acacia Senegal Gum, Xanthan Gum, Alpha-Glucan Oligosaccharide, Carbomer, Polyglyceryl-4 Caprate, Polyglyceryl-6 Caprylate, Chromium Oxide Green (CI 77288), Butylene Glycol, Potassium Sorbate, Sodium Benzoate, Glycyrrhiza Glabra (Licorice) Root Extract, Salix Nigra (Willow) Bark Extract, Camellia Sinensis (Green Tea) Leaf Extract, Benzoic Acid.

Questions or comments?

www.banuskin.com

855-215-2268

Principal Display Panel

banu

SULFUR SPOT TREATMENT

10% Sulfur Acne Treatment

SAFE FOR ACNE-PRONE SKIN

15 mL/0.5 fl.oz.

Topical Cream

Label

15 mL Tube Carton



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Questions or comments?

www.banuskincare.com
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Made in the U.S.A. with globally sourced ingredients. Please recycle this box.

Dist. By: BANUSKIN, INC.
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New York, NY 10004
855-215-BANU
www.banuskincare.com

SULFUR SPOT TREATMENT

10% Sulfur
Acne Treatment

SAFE FOR
ACNE-PRONE
SKIN

banu

15 mL/0.5 fl. oz.

Formulated with 10% sulfur, kaolin and bentonite clays, this spot treatment visibly reduces pimples and redness—without over-drying. The creamy formula forms a breathable barrier over breakouts to target problem areas, while skin-soothing ingredients protect the skin around the pimple to minimize redness, irritation, and flakiness caused by dryness.

Use overnight or as a 20-minute mask on problem areas. For best results, apply directly to clean, dry skin.

CLINICALLY TESTED ON
ACNE-PRONE SKIN

NON-COMEDOGENIC
& NON-ACNEGENIC

DERMATOLOGIST TESTED

NON-IRRITATING
& NON-DRYING

SAFE FOR
ALL SKIN TONES

VEGAN & CRUELTY-FREE

FRAGRANCE-FREE



15 mL Tube



SULFUR SPOT TREATMENT

sulfur acne treatment cream						
Product Information						
Product Type		HUMAN OTC DRUG	Item Code (Source)	NDC:85439-0102		
Route of Administration		TOPICAL				
Active Ingredient/Active Moiety						
Ingredient Name			Basis of Strength	Strength		
SULFUR (UNII: 70FD1KFU70) (SULFUR - UNII:70FD1KFU70)			SULFUR	10 g in 100 mL		
Inactive Ingredients						
Ingredient Name				Strength		
POLYGLYCERYL-6 CAPRYLATE (UNII: DGV8R54VG7)						
BUTYLENE GLYCOL (UNII: 3XUS85K0RA)						
SODIUM BENZOATE (UNII: OJ245FE5EU)						
BENZOIC ACID (UNII: 8SKN0B0MIM)						
CARBOMER (UNII: 0A5MM307FC)						
WATER (UNII: 059QF0KO0R)						
KAOLIN (UNII: 24H4NWX5CO)						
BENTONITE (UNII: A3N5ZCN45C)						
HAMAMELIS VIRGINIANA (WITCH HAZEL) LEAF WATER (UNII: 8FP93ED6H2)						
XANTHAN GUM (UNII: TTV12P4NEE)						
CHROMIUM OXIDE GREENS (UNII: X5Z09SU859)						
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)						
PHENOXYETHANOL (UNII: HIE492ZZ3T)						
WILLOW BARK (UNII: S883J9JDYX)						
CAMELLIA SINENSIS LEAF (UNII: W2ZU1RY8B0)						
ACACIA SENEGAL GUM (UNII: 5C5403N26O)						
ALPHA-GLUCAN OLIGOSACCHARIDE (UNII: S95658MI3W)						
GLYCERIN (UNII: PDC6A3C0OX)						
SORBITOL (UNII: 506T60A25R)						
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)						
NIACINAMIDE (UNII: 25X51I8RD4)						
POLYGLYCERYL-4 CAPRATE (UNII: 3N873UN885)						
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)						
GLYCYRRHIZA GLABRA (LICORICE) ROOT POWDER (UNII: 2788Z9758H)						
Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:85439-0102-1	15 mL in 1 TUBE; Type 0: Not a Combination Product	04/15/2025			

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	04/15/2025	

Labeler - Banuskin, Inc. (119398280)

Registrant - Banuskin, Inc. (119398280)

Revised: 4/2025

Banuskin, Inc.