

END-ZIT- sulfur lotion, augmented
ABBE Laboratories, Inc.

END-ZIT® ACNE CONTROL DRYING LOTION

Intended for use on blemishes only. This advanced formulation is designed to dry and aid in elimination of pimples. It is make-up quality for excellent coverage of the blemish. It is unbelievably effective, clinically tested, dermatologist approved and recommended.

DIRECTIONS: Shake bottle well before each use. Apply directly to blemish only, using applicator. Wait a few moments until slightly dry. Pat with fingertip or clean cotton swab to blend color to skin. Make-up may be applied if desired. Should remain on skin all day. Re-apply at night to clean skin to remain on skin overnight.

NOTE: PERSONS WITH KNOWN SENSITIVITY TO SULFUR SHOULD AVOID USE OF THIS PRODUCT. KEEP FROM EYES.

DRUG FACTS

ACTIVE INGREDIENT SULFUR 5%

PURPOSE ACNE TREATMENT

WARNINGS:

- For external use only.
- Using other topical acne medications at the same time or immediately following use of this product may increase dryness or irritation of the skin. If this occurs, only one medication should be used unless directed by a doctor.
- Do not get into eyes. If excessive skin irritation develops or increases, discontinue use and consult a doctor.
- Keep out of reach of children.

OTHER INGREDIENTS: Isopropyl Alcohol, Water, Zinc Oxide, Propylene Glycol, Camphor, Talc, Sodium Laureth Sulfate, Titanium Dioxide, Diazolidinyl Urea, Methylparaben, Propylparaben. May contain Iron Oxides.

NDC #'s

Light/Medium 68605-2001-2 Medium/Dark 68605-2002-2 Acne Control Mask 68605-2010-2

——PACKAGE LABEL.PRINCIPAL DISPLAY PANEL——

DIRECTIONS: SHAKE BOTTLE WELL.

APPLY TWICE DAILY DIRECTLY TO ACNE BREAKOUT. ALLOW TO DRY FOR 10 SECONDS. PAT TO BLEND. APPLY MAKE-UP IF DESIRED.

ACTIVE INGREDIENT: SULFUR 5%.

WARNINGS:

AVOID IF ALLERGIC TO SULFUR

AVOID CONTACT WITH EYES
FOR EXTERNAL USE ONLY
KEEP FROM CHILDREN

Manufactured by ABBE Laboratories, Inc.
Farmingdale, NY 11735
Made in the U.S.A.

END-ZIT®

ACNE CONTROL
DRYING LOTION

ABBE

0.5 OZ. (14.78 g)



———PACKAGE LABEL.PRINCIPAL DISPLY PANEL———

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Farmingdale, NY 11735
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END-ZIT®

ACNE CONTROL
DRYING LOTION

ABBE

2.5 OZ. (70 g)

THE ZIT STOPS HERE!®

END-ZIT®

ACNE CONTROL MASK

**GREAT AS
A SPOT
TREATMENT!**



NET WT. 2.5 OZ. (70g)

DRUG FACTS

Active Ingredients Sulfur 10%
Purpose Acne-Treatment

USE: Controls Acne Breakouts

WARNINGS:
 -For External Use Only
 -Avoid Getting Into Eyes
 -Avoid if Allergic to Sulfur
 -Keep From Children

No Animal Testing

Stop use and consult physician if condition worsens or rash develops.

Directions: Apply End-Zit® Mask generously, spreading evenly over skin, avoiding eye area. Allow mask to remain on 7-10 minutes. Rinse thoroughly with warm water to remove all traces of mask. Pat dry. Use once or twice each week. May also be used as a spot treatment for acne pimples.

Other Ingredients: Water, Bentonite, Titanium Dioxide, Talc, Kaolin, Ahnfeltia Concinna Extract (Red Marine Algae), Butylene Glycol, Propylene Glycol, Diazolidinyl Urea, Methylparaben, Propylparaben.

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Manufactured by ABBE Laboratories, Inc.
 Farmingdale, NY 11735
Made in the USA

END-ZIT

sulfur lotion, augmented

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68605-2001
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SULFUR (UNII: 70FD1KFU70) (SULFUR - UNII:70FD1KFU70)	SULFUR	.05 g in 1 mL

Inactive Ingredients

Ingredient Name	Strength
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
WATER (UNII: 059QF0KO0R)	
ZINC OXIDE (UNII: SOI2LOH54Z)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)	
TALC (UNII: 7SEV7J4R1U)	
SODIUM LAURETH SULFATE (UNII: BPV390UAP0)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
DIAZOLIDINYL UREA (UNII: H5RIZ3MPW4)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	

Product Characteristics

Color	brown (Light/Medium)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:68605-2001-2	14.78 mL in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	04/20/2004	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	04/20/2004	

END-ZIT

sulfur lotion, augmented

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68605-2002
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SULFUR (UNII: 70FD1KFU70) (SULFUR - UNII:70FD1KFU70)	SULFUR	.05 g in 1 g

Inactive Ingredients

Ingredient Name	Strength
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
WATER (UNII: 059QF0KO0R)	
ZINC OXIDE (UNII: SOI2LOH54Z)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)	
TALC (UNII: 7SEV7J4R1U)	
SODIUM LAURETH SULFATE (UNII: BPV390UAP0)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
DIAZOLIDINYL UREA (UNII: H5RIZ3MPW4)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	

Product Characteristics

Color	brown (Medium/Dark)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:68605-2002-2	17.57 g in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	04/20/2004	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	04/20/2004	

END-ZIT

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Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68605-2010
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SULFUR (UNII: 70FD1KFU70) (SULFUR - UNII:70FD1KFU70)	SULFUR	.05 g in 1 g

Inactive Ingredients

Ingredient Name	Strength
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
WATER (UNII: 059QF0KO0R)	
ZINC OXIDE (UNII: SOI2LOH54Z)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET)	
TALC (UNII: 7SEV7J4R1U)	
SODIUM LAURETH SULFATE (UNII: BPV390UAP0)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
DIAZOLIDINYL UREA (UNII: H5RIZ3MPW4)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	

Product Characteristics

Color	white (Mask)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:68605-2010-2	70 g in 1 TUBE; Type 0: Not a Combination Product	04/20/2004	10/31/2023

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	04/20/2004	10/31/2023

Labeler - ABBE Laboratories, Inc. (781745286)

Establishment

Name	Address	ID/FEI	Business Operations
ABBE Laboratories, Inc.		781745286	manufacture(68605-2001, 68605-2002, 68605-2010)

Revised: 1/2026

ABBE Laboratories, Inc.