

**ZILACTIN EARLY RELIEF COLD SORE - benzyl alcohol gel**  
**Applied Laboratories, Inc.**

*Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.*

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**Zilactin Red Applied**

**Drug Facts**

**Active ingredient**

Benzyl alcohol 10%

**Purpose**

Cold sore/fever blister treatment, Oral pain reliever

**Use**

temporarily relieves pain caused by:

- cold sores/fever blisters
- canker sores, mouth sores
- gum irritations

**Warnings**

**Flammable:**

- keep away from fire or flame

**Stop use and ask a physician if**

- sore mouth symptoms do not improve in 7 days
- condition worsens or symptoms clear up and occur again within a few days
- swelling, rash or fever develops
- irritation, pain or redness persists or worsens
- apply only to affected area
- do not exceed recommended dosage
- avoid contact with the eyes
- do not use for more than 7 days unless directed by a physician or dentist

**Keep out of reach of children.** If swallowed, get medical help or contact a poison control center immediately.

**Directions**

adults and children 2 years and older	dry affected area. Apply with cotton swab or clean finger up to 4 times daily. Allow to dry 30-60 seconds.
children under 12 years	adult supervision should be given in the use of this product
children under 2 years	do not use, consult a physician or dentist

### **Other information**

- do not peel off protective film. Attempting to peel off film may result in skin irritation or tenderness. To remove film, first apply another coat of Zilactin to film, and immediately wipe the area with a moist gauze pad or tissue.
- contains alcohol 68% by volume
- store at 15-30°C (59-86°F)

### **Inactive ingredients**

boric acid, hydroxypropylcellulose, propylene glycol, purified water, salicylic acid, SD alcohol 37, tannic acid

Manufactured for: Blairex Laboratories, Inc. P.O. Box 2127 Columbus, IN 47202 USA

**!** Questions or Comments?

Call TOLL FREE 1-800-252-4739

[www.zilactin.com](http://www.zilactin.com)

### **PROMOTES HEALING**

Cold Sore/Fever Blister/Oral Pain Treatment

Zilactin<sup>®</sup> Early Relief Cold Sore Gel

- Bioadhesive Promotes Healing
- Relieves Pain for up to 6 Hours

Net wt. 0.25 oz (7.1g)

PROMOTES HEALING

Cold Sore/  
Fever Blister/  
Oral Pain Treatment

# zilactin®

## Early Relief Cold Sore Gel

• **Bioadhesive  
Promotes Healing**

• **Relieves Pain  
for up to 6 Hours**

Net wt. 0.25 oz (7.1g)

**Dries Clear**

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# zilactin®

## Early Relief Cold Sore Gel



000-500027-03



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**Blair Laboratories, Inc.**  
P.O. Box 2127 Columbus, IN 47202 USA  
Questions or Comments?  
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[www.zilactin.com](http://www.zilactin.com)

**Tamper Evident Feature:** Do not use if glued carton has been opened or tampered with.

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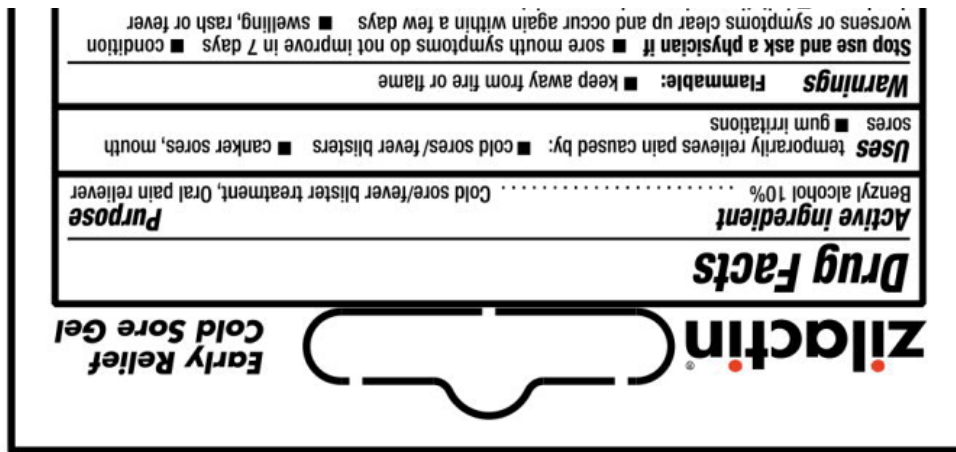
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**Keep out of reach of children.** If swallowed, get medical help or contact a poison control center immediately.

■ apply only to affected area ■ do not exceed recommended dosage ■ avoid contact with the eyes ■ do not use for more than 7 days unless directed by a physician or dentist

develops ■ irritation, pain or redness persists or worsens



## ZILACTIN EARLY RELIEF COLD SORE

benzyl alcohol gel

### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:72751-0301
<b>Route of Administration</b>	TOPICAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>BENZYL ALCOHOL</b> (UNII: LKG8494WBH) (BENZYL ALCOHOL - UNII:LKG8494WBH)	BENZYL ALCOHOL	0.1 g in 1 g

### Inactive Ingredients

Ingredient Name	Strength
<b>BORIC ACID</b> (UNII: R57ZHV85D4)	
<b>HYDROXYPROPYL CELLULOSE (1200000 WAMW)</b> (UNII: U3JF91U133)	
<b>PROPYLENE GLYCOL</b> (UNII: 6DC9Q167V3)	
<b>SALICYLIC ACID</b> (UNII: O414PZ4LPZ)	
<b>ALCOHOL</b> (UNII: 3K9958V90M)	
<b>EUCALYPTOL</b> (UNII: RV6J6604TK)	
<b>THYMOL</b> (UNII: 3J50XA376E)	
<b>MENTHOL, UNSPECIFIED FORM</b> (UNII: L7T10EIP3A)	
<b>TANNIC ACID</b> (UNII: 28F9E0DJY6)	
<b>WATER</b> (UNII: 059QF0KO0R)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72751-0301-1	1 in 1 CARTON	06/30/2005	
1		7.1 g in 1 TUBE; Type 0: Not a Combination Product		

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part356	06/30/2005	

**Labeler** - Applied Laboratories, Inc. (117337220)

Revised: 12/2021

Applied Laboratories, Inc.