ACNE MED 10%- benzoyl peroxide gel Face Reality, LLC

Acne Med 10%

Drug Facts

Active ingredient

Benzoyl Peroxide 10%

Purpose

Acne Treatment

Use

for the treatment of acne

Warnings

For external use only

Do not use

- if you have very sensitive skin
- if you are sensitive to benzoyl peroxide

When using this product

- skin irritation and dryness is 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.
- avoid unnecessary sun exposure and use a sunscreen
- avoid contact with eyes, lips and mouth
- avoid contact with hair and dyed fabrics, which may be bleached by this product
- skin irritation may occur, characterized by redness, burning, itching, peeling, or possibly swelling. Irritation may be reduced by using the product less frequently or in a lower concentration.

Stop use and ask a doctor if

irritation becomes severe

Keep out of reach of children.

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

Directions

- 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
- if going outside, apply sunscreen after using this product. If irritation or sensitivity develops, stop use of both products and ask a doctor.

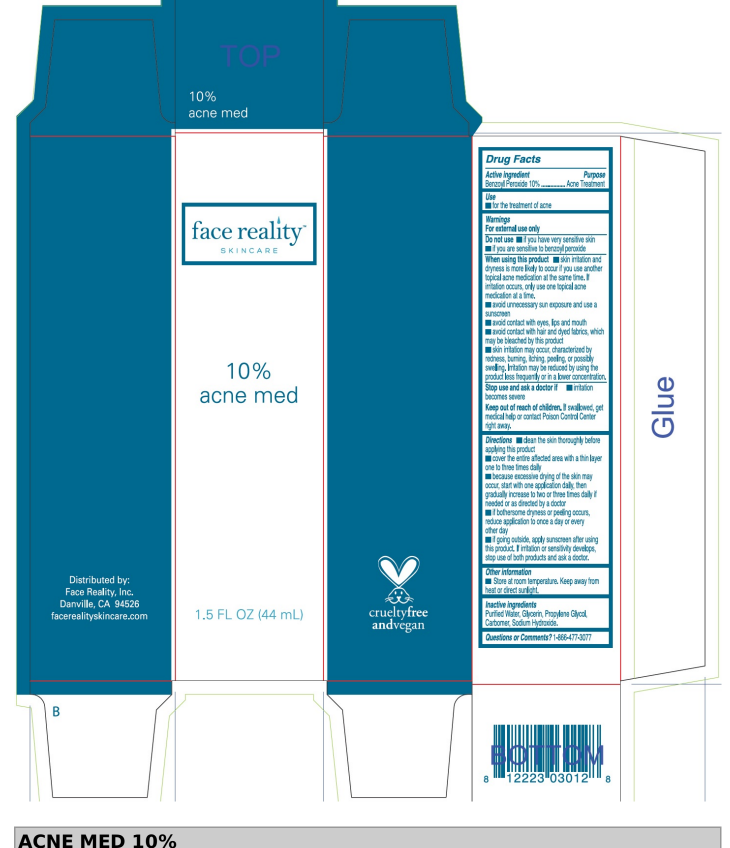
Other information

• Store at room temperature. Keep away from heat or direct sunlight.

Inactive ingredients

Purified Water, Glycerin, Propylene Glycol, Carbomer, Sodium Hydroxide.

Package Labeling:



ACIVE MED 10%

benzoyl peroxide gel

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:70707-212

Route of Administration TOPICAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
BENZOYL PEROXIDE (UNII: W9WZN9A0GM) (BENZOYL PEROXIDE - UNII: W9WZN9A0GM)	BENZOYL PEROXIDE	100 mg in 1 mL		

Inactive Ingredients			
Ingredient Name	Strength		
WATER (UNII: 059QF0KO0R)			
GLYCERIN (UNII: PDC6A3C0OX)			
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)			
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)			
SODIUM HYDROXIDE (UNII: 55X04QC32I)			

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:70707- 212-15	1 in 1 CARTON	01/01/2023			
1		44 mL in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product				

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
OTC Monograph Drug	M006	01/01/2023				

Labeler - Face Reality, LLC (602958071)

Revised: 12/2023 Face Reality, LLC