

CONTROLLING CREAM- benzoyl peroxide cream CA-BOTANA INTERNATIONAL

Controlling Cream

Warnings:

For external use only.

Do not use on wounds or damaged skin

When using this product: use only as directed. Avoid contact with eyes. Do not bandage tightly

Stop use and ask a doctor if: redness is present. Irritation develops. Condition worsens or symptoms persist more than 7 days. Symptoms clear up and occur again within a few days.

Store at room temperature. Lot number and expiration date see crimp or see box.

cleanse the skin thoroughly with soft foam deep cleanse before applying. cover the entire affected area with a thin layer 1 to 2 times daily. because too much drying of the skin may occur, start with 1 application daily, then gradually increase to 2 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. Allow benzoyl peroxide to dry the follow directions on sunscreen label.

Alpha Oligo Peptide-6 TM
Calendula officinalis (Marigold) Extract
Cetearyl Alcohol
Cetearyl Isononanoate
Cynamomum Zeylanicum
Ethylhexylglycerin
Glycerine
Glycine
Hydrogen Peroxide
Mineral oil
Oenothera biennis (Evening Primrose) extract
Panicum Miliaceum (Millet) extract
Petrolatum
Phenoxyethanol
Phosphatidylcholine
Polysorbate 60
Potassium Sorbate
Rosa canina (Rosehip) extract
Rosmarinus officinalis (Rosemary) extract
Sorbitan Stearate
Tetra Sodium
Water (Aqua)

Keep out of reach of children. If swallowed get medical help or contact a Poison Control

Center immediately.

when using this product apply to affected areas only. avoid unnecessary sun exposure and use a sunscreen. do not use in or near eyes. using other topical acne drugs at the same time or right

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DOCTOR D.
SCHWAB®

CONTROLLING CREAM

CRÈME RÉGULATRICE
CREMA CONTROLANTE



**ACNE +
OILY SKIN**



1 FL OZ / 30 ML



Drug Facts

Active ingredient	Purpose
Benzoyl Peroxide 2.7%	Acne Treatment

Uses •cleas acne pimples •helps prevent new acne pimples •penetrates pores to eliminate most acne blemishes, blackheads, and whiteheads

Warnings For external use only

Do not use •on broken skin •on large areas of the body

When using this product •apply to affected areas only •avoid unnecessary sun exposure and use a sunscreen •do not use in or near eyes •using other topical acne drugs at the same time or right after use of this product may increase dryness or irritation of the skin. Only one drug should be used unless

NDC # 36192-017-05

DOCTOR D. SCHWAB®
www.DoctorSchwabCA.com
A Division of CA BOTANICA
San Diego, CA 92121, USA
Made in USA

REF #860

Peel Here

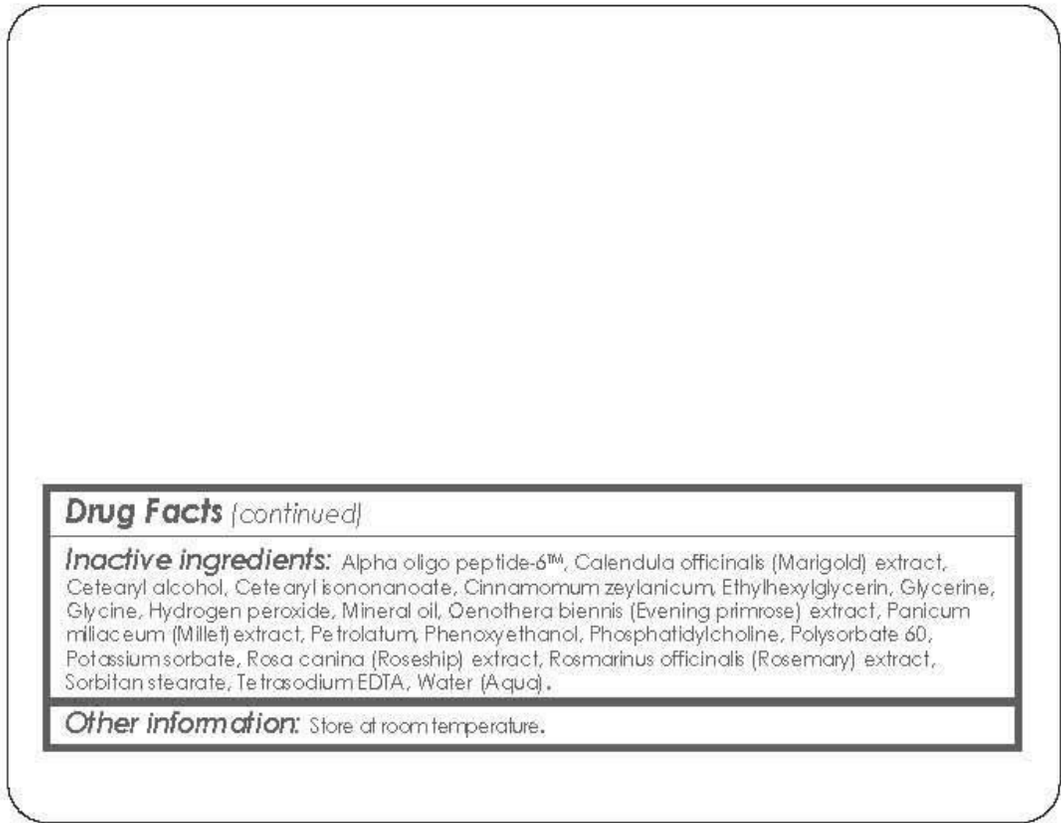
Drug Facts (continued)

directed by a doctor.

Stop use and ask a doctor if too much skin irritation or sensitivity develops or increases

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions •cleanse the skin thoroughly with Soft Foam Deep Cleanse before applying •cover the entire affected area with a thin layer 1 to 2 times daily •because too much drying of the skin may occur, start with 1 application daily, then gradually increase to 2 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, use a sunscreen. Allow benzoyl peroxide to dry, the follow directions on sunscreen label.



Package label display panel

CONTROLLING CREAM			
benzoyl peroxide cream			
Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:35192-017
Route of Administration	TOPICAL		
Active Ingredient/Active Moiety			
Ingredient Name		Basis of Strength	Strength
BENZOYL PEROXIDE (UNII: W9WZ N9A0GM) (BENZOYL PEROXIDE - UNII:W9WZ N9A0GM)		BENZOYL PEROXIDE	0.8 mg in 29.6 mg

Inactive Ingredients

Ingredient Name	Strength
CINNAMON (UNII: 5S29HWU6QB)	
CINNAMON LEAF OIL (UNII: S92U8SQ71V)	
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)	
GLYCERIN (UNII: PDC6A3C0OX)	
GLYCINE (UNII: TE7660XO1C)	
HYDROGEN PEROXIDE (UNII: BBX060AN9V)	
LIGHT MINERAL OIL (UNII: N6K5787QVP)	
OENOTHERA BIENNIS (UNII: 76UI55V071)	
MILLET (UNII: TJR6B3R47P)	
PETROLATUM (UNII: 4T6H12BN9U)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
HYDROGENATED SOYBEAN LECITHIN (UNII: H1109Z9J4N)	
POLYSORBATE 60 (UNII: CAL22UVI4M)	
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)	
ROSA CANINA LEAF (UNII: J3N2Z889QP)	
ROSEMARY (UNII: IJ67X351P9)	
SORBITAN MONOSTEARATE (UNII: NVZ4I0H58X)	
EDETATE SODIUM (UNII: MP1J8420LU)	
WATER (UNII: 059QF0KO0R)	
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)	
CETEARYL ISONONANOATE (UNII: P5O01U99NI)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:35192-017-05	29.6 mg in 1 TUBE; Type 0: Not a Combination Product	11/02/2017	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	11/09/2014	

Labeler - CA-BOTANA INTERNATIONAL (106276728)

Registrant - RODOLFO UGELSTAD (106276728)

Establishment

Name	Address	ID/FEI	Business Operations
CA-BOTANA INTERNATIONAL		106276728	manufacture(35192-017)