ACNE CLEANSER- benzoyl peroxide cream Meijer Distribution, 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.

Daily Acne Control Cleanser 264.005/264AI rev 1-264AJ

Active ingredient

Benzoyl peroxide 10%

Purpose

Acne medication

use

for the treatment of acne

Warnings

For external use only

Do not use if you

- have very sensitive skin
- 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 the eyes, lips and mouth
- avoid contact with hair and dyed fabrics, which may be bleaced by this product
- skin irritation may occur, characterized by redness, burning, itching, peeling or possible 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 a Poison Control Center right away. Avoid contact with eyes. If contact occurs, flush thoroughly with water.

Directions

- wet face. Gently massage all over face for 20-30 seconds avoiding the eyes. Rinse thoroughly and pat dry.
- 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 daily 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
- Sensitivity Test for a New User. Apply product sparingly to one or two small affected areas, during the first 3 days. If no discomfort occurs, follow the directions stated above.

Other information

- keep tightly closed
- store at room temperature (59°-77°)

Inactive ingredients

water, cetyl alcohol, petrolatum, acrylates/C10-30 alkyl acrylate crosspolymer, zinc lactate, steareth-2, glycerin, potassium cetyl phosphate, xanthan gum, benzyl alcohol, fragrance, disodium EDTA, laureth-4, BHT, sodium hydroxide, lactic acid, menthol

*This product is not manufactured or distributed by Johnson & Johnson Consumer Products Company, distributor ofClean & Clear Continuous Control Acne Cleanser

Questions 1-888-593-0593

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GRAND RAPIDS, MI 49544

www.meijer.com

OUR QUALITY GUARANTEE

Principal display panel

Compare to CLEAN & CLEAR Continuous Control Acne Cleanser*

meijer

Acne

Cleanser

10% Benzoyl Peroside

Acne Medication Helps Fight Blemishes DAILY CONTROL Keeps fighting Acne after you wash Daily treatment for daily cleaning Paraben free NET WT 5 OZ (141 g)



ACNE CLEANSER benzoyl peroxide cream				
Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:41250-264	
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 g

Inactive Ingredie	ents			
Ingredient Name			Strength	
water (UNII: 059QF0K0	D0R)			
CETYL ALCOHOL (UN	CETYL ALCOHOL (UNII: 936JST6JCN)			
PETROLATUM (UNII: 4	T6H12BN9U)			
CARBOMER INTERPO	LYMER TYPE A (ALLYL SUCROSE CROSSLI	NKED) (UNII: 59TL3WG5C	O)	
ZINC LACTATE (UNII:	2GXR25858Y)			
STEARETH-2 (UNII: V5	6DFE46J5)			
GLYCERIN (UNII: PDC6	GLYCERIN (UNII: PDC6A3C0OX)			
POTASSIUM CETYL P	HOSPHATE (UNII: 03KCY6P7UT)			
XANTHAN GUM (UNII: TTV12P4NEE)				
BENZYL ALCOHOL (U	NII: LKG8494WBH)			
EDETATE DISODIUM	ANHYDROUS (UNII: 8NLQ36F6MM)			
LAURETH-4 (UNII: 6HC	Q855798J)			
BUTYLATED HYDROX	YTOLUENE (UNII: 1P9D0Z171K)			
SODIUM HYDROXIDE	(UNII: 55X04QC32I)			
LACTIC ACID (UNII: 33	3X04XA5AT)			
menthol (UNII: L7T10E	EIP3A)			
Packaging				
# Item Code	Package Description	Marketing Start Date	Marketing End Date	

 I
 NDC:41250-264-56
 141 g in 1 TUBE; Type 0: Not a Combination Product
 02/18/2014
 Date
 Date

 Marketing Category
 Application Number or Monograph Citation
 Marketing Start Date
 Marketing End Date

02/18/2014

Labeler - Meijer Distribution, INC (006959555)

Registrant - Vi-Jon, LLC (790752542)

OTC monograph final part333D

Establishment			
Name	Address	ID/FEI	Business Operations
Vi-Jon, LLC		790752542	manufacture(41250-264)
Establishment			

Name	Address	ID/FEI	Business Operations
Vi-Jon, LLC		088520668	manufacture(41250-264)

Revised: 5/2023

Meijer Distribution, INC