### OXY SENSITIVE FACE WASH MAXIMUM SOOTHING- benzoyl peroxide lotion The Mentholatum Company

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.

-----

#### **Drug Facts**

#### **Active ingredient**

Benzoyl peroxide 5%

### **Purpose**

Acne treatment

#### Uses

treats and helps prevent 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 bleached by this product
- skin irritation may occur, characterized by redness, burning, itching, peeling, or possibly swelling. Irritation may be reduced by using this product less frequently or in a lower concentration.

### Stop use and ask a doctor if

irritation becomes severe

### If pregnant or breast-feeding

ask a health professional before use.

#### Keep Out of Reach of Children

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

#### **Directions**

- wet face
- apply to hands then work into a lather and massage gently onto face
- rinse thoroughly and pat dry
- because excessive drying of the skin may occur, start with 1 use daily, then gradually increase to 2 or 3 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.
- Sensitivity Test: Apply product sparingly to a small affected area for the first 3 days. If no discomfort occurs, follow directions above.

#### Other information

- THIS PRODUCT MAY BLEACH HAIR OR DYED FABRICS
- KEEP TIGHTLY CLOSED
- avoid storing at temperatures above 100°F (38°C)

### **Inactive ingredients**

water, cetostearyl alcohol, sodium C14-16 olefin sulfonate, disodium laureth sulfosuccinate, capryl/capramidopropyl betaine, xanthan gum, butylene glycol, citric acid, diazolidinyl urea, hydrolyzed soy protein, methylparaben, PEG-8 dimethicone, portulaca oleracea extract, propylparaben, sodium citrate, sodium hydroxide, sodium lauroyl sarcosinate

#### Questions?

**1-877-636-2677** MON-FRI 9 AM-5 PM (EST)

### **Principal Display Panel**



#### OXY SENSITIVE FACE WASH MAXIMUM SOOTHING

benzoyl peroxide lotion

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:10742-1313
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
BENZOYL PEROXIDE (UNII: W9WZ N9A0GM) (BENZ OYL PEROXIDE - UNII: W9WZ N9A0GM)	BENZOYL PEROXIDE	50 mg in 1 mL	

Inactive Ingredients	
Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)	
SODIUM C14-16 OLEFIN SULFONATE (UNII: O9W3D3YF5U)	
DISODIUM LAURETH SULFOSUCCINATE (UNII: D6DH1DTN7E)	
CAPRYL/CAPRAMIDOPROPYL BETAINE (UNII: 231H3ZT9NE)	
XANTHAN GUM (UNII: TTV12P4NEE)	
BUTYLENE GLYCOL (UNII: 3XUS85K0RA)	
ANHYDROUS CITRIC ACID (UNII: XF417D3PSL)	
DIAZOLIDINYL UREA (UNII: H5RIZ3MPW4)	
SOY PROTEIN (UNII: R44IWB3RN5)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
PEG-8 DIMETHICONE (UNII: GIA7T7640D)	
PURSLANE (UNII: M6S840WXG5)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	
SODIUM CITRATE, UNSPECIFIED FORM (UNII: 1Q73Q2JULR)	
SODIUM HYDROXIDE (UNII: 55X04QC32I)	
SODIUM LAUROYL SARCOSINATE (UNII: 632GS99618)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:10742- 1313-1	148 mL in 1 TUBE; Type 0: Not a Combination Product	08/01/2016	
2	NDC:10742- 1313-2	185 mL in 1 TUBE; Type 0: Not a Combination Product	08/01/2016	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph final	part333D	08/01/2016		

# Labeler - The Mentholatum Company (002105757)

## Registrant - The Mentholatum Company (002105757)

Establishment				
Name	Address	ID/FEI	<b>Business Operations</b>	
The Mentholatum Company		002105757	manufacture(10742-1313)	