

MINERAL LP FOUNDATION SANDY BEIGE- mineral lp foundation cream
JENTRY KELLEY

JENTRY KELLEY MINERAL LIQUID POWDER FOUNDATION BROAD SPECTRUM
SPF 15 -Sandy Beige

Actives

Titanium Dioxide 6%

Zinc Oxide 3%

Purpose

Sunscreen

Use

Helps prevent sunburn. If used as directed with other sun protection measures (see directions), decreases the risk of skin cancer and early skin aging caused by the sun.

Warnings

For external use only. Do not use on damaged or broken skin. Stop use and ask doctor if rash occurs.

When using this product keep out of eyes. Rinse with water to remove.

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

Directions

Apply liberally 15 minutes before sun exposure. Use a water-resistant sunscreen if swimming or sweating. Reapply at least every 2 hours, Children under 6 months: ask a doctor.

Sun Protection Measures: Spending time in the sun increases risk of skin cancer and early aging. To decrease

this risk, regularly use a sunscreen with a Broad-Spectrum SPF value of 15 or higher and other sun protection measures including: limit time in the sun especially from 10:00 a.m. - 2:00 p.m. Wear long sleeved shirts, pants, hats, and sunglasses.

Inactive Ingredients

Cyclopentasiloxane, Aqua/Water/Eau, Dimethicone Crosspolymer, Octyldodecyl Neopentanoate, Butylene Glycol, Cetyl PEG/PPG-10/1 Dimethicone, Polyglyceryl-4

Isostearate, Aluminum Hydroxide, Stearic Acid, Dimethicone, Hexyl Laurate, Silica, Tocopherol, Tetrahexyldecyl Ascorbate, Sodium Hyaluronate, PEG/PPG-18/18 Dimethicone, Phytantriol, Nylon-12, Sodium Chloride, Nylon-12 Fluorescent Brightener 230 Salt, Polyvinylalcohol Crosspolymer, Octyldodecanol, Magnesium Chloride, Potassium Chloride, Zinc Chloride , Lysine, Methicone, Triethoxycaprylylsilane, Disodium EDTA, Hexylene Glycol, Caprylyl Glycol, Phenoxyethanol, Potassium Sorbate, Mica , Titanium Dioxide (CI 77891), Iron Oxides (CI 77492), Iron Oxides (CI 77491), Iron Oxides (CI 77499)

Other Information

Protect this product from excessive heat and direct sun

JENTRY KELLEY

Mineral Liquid
Powder Foundation

**BROAD SPECTRUM
SPF 15**

1 fl. oz. (30ml)

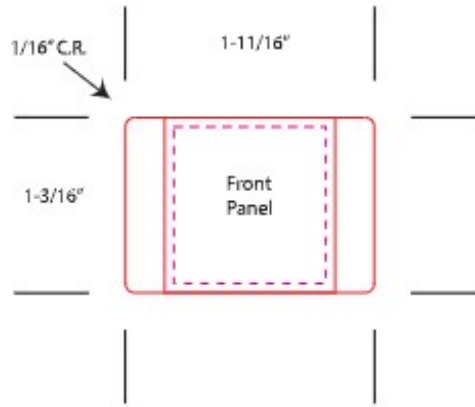


Kroger Packaging Inc.

7 Panel Fold over Label DieLine
 Date: 2/16/12
 Size: 1-11/16" x 1-3/16", 1/16" C.R. Fold over (7 panels)

- DieLine
- CLEAR/FACE AREA (no copy)

View After Folding



This Panel is behind the Front Panel
 Panels 4, 5, 6, 7 - Prints as 1 panel across the folds

Mineral Liquid Powder Foundation (and actual) see to

Drug Facts

Active Ingredients: Purpose: The active ingredients are listed on the label. The purpose of the product is to provide coverage and protection.

Warnings: Use only as directed. Do not use if you are allergic to any of the ingredients. Do not use if you are pregnant or breastfeeding. Do not use if you have a skin condition.

Directions: Apply to clean, dry skin. Use a brush or sponge to apply. Blend into skin. Reapply as needed.

Other Information: For more information, see the label. © 2012 Kroger Co.

GLUE

Drug Facts

Inside

Outside

DIE LINES/DECO AREA RULES/TUBE TEMPLATES NEVER PRINT, BUT SHOULD BE SHOWN ON PROOFS

Brand	Job #	Description	Ingredient List Formula	Component #
YOUR NAME	YN-LBL 416F	MINERAL LIQUID POWDER FOUNDATION SPF 15 DRUG FACTS PEEL OFF LABEL	MINERAL LIQUID POWDER FOUNDATION - 10/12/20	MAN-070-LPMF0000F
Version & Date Routed		Colors	UPC	Item / F6 #
R2 11/4/20		PANELS 1, 2, & 3 PRINT BLACK W/KNOCKOUT TO WHITE COPY/ART. PANELS 4 - 7 PRINTS BLACK	N/A	YNC-LPMFxxD
Date Released		Stock / Finish / Materials	Final Size	Production Method
12/1/20		WHITE LABEL STOCK		OUTSIDE SCREEN

F.P.O. MEANS "FOR POSITION ONLY", DO NOT PRINT ANY ITEM THAT HAS F.P.O. SUPERIMPOSED ON IT

MINERAL LP FOUNDATION SANDY BEIGE

mineral lp foundation cream

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:84179-210
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ZINC OXIDE (UNII: SOI2LOH54Z) (ZINC OXIDE - UNII:SOI2LOH54Z)	ZINC OXIDE	3 g in 30 mL
TITANIUM DIOXIDE (UNII: 15FIX9V2JP) (TITANIUM DIOXIDE - UNII:15FIX9V2JP)	TITANIUM DIOXIDE	6 g in 30 mL

Inactive Ingredients

Ingredient Name	Strength
FERRIC OXIDE YELLOW (UNII: EX438O2MRT)	
FERROSFERRIC OXIDE (UNII: XM0M87F357)	
BUTYLENE GLYCOL (UNII: 3XUS85K0RA)	
HYALURONATE SODIUM (UNII: YSE9PPT4TH)	
ALUMINUM HYDROXIDE (UNII: 5QB0T2IUN0)	
POTASSIUM CHLORIDE (UNII: 660YQ98I10)	
OCTYLDODECYL NEOPENTANOATE (UNII: X8725R883T)	
DIMETHICONE (UNII: 92RU3N3Y1O)	
DISODIUM 4,4'-BIS((4-ANILINO-6-((2-CARBAMOYLETHYL)(2-HYDROXYETHYL)AMINO)-S-TRIAZIN-2-YL)AMINO)-2,2'-STILBENEDISULFONATE (UNII: 30I2S866LK)	
MICA (UNII: V8A1AW0880)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)	
POLYVINYL ALCOHOL, UNSPECIFIED (UNII: 532B59J990)	
MAGNESIUM CHLORIDE (UNII: 02F3473H9O)	
ZINC CHLORIDE (UNII: 86Q357L16B)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
LYSINE (UNII: K3Z4F929H6)	
CYCLOMETHICONE 5 (UNII: 0THT5PCI0R)	
POLYGLYCERYL-4 ISOSTEARATE (UNII: 820DPX33S7)	
WATER (UNII: 059QF0KO0R)	
HEXYL LAURATE (UNII: 4CG9F9W01Q)	
DIMETHICONE CROSSPOLYMER (450000 MPA.S AT 12% IN CYCLOPENTASILOXANE) (UNII: UF7620L1W6)	
METHICONE (20 CST) (UNII: 6777U11MKT)	
TRIETHOXYCAPRYLYLSILANE (UNII: LDC331P08E)	
CAPRYLYL GLYCOL (UNII: 00YIU5438U)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TETRAHEXYLDECYL ASCORBATE (UNII: 9LBV3F07AZ)	
PEG/PPG-18/18 DIMETHICONE (UNII: 9H0AO7T794)	
NYLON-12 (UNII: 446U8J075B)	
CETYL PEG/PPG-10/1 DIMETHICONE (HLB 3) (UNII: G300307ZXP)	
TOCOPHEROL (UNII: R0ZB2556P8)	
PHYTANTRIOL (UNII: 8LVI07A72W)	
OCTYLDODECANOL (UNII: 461N1O614Y)	
EDETATE DISODIUM (UNII: 7FLD91C86K)	
HEXYLENE GLYCOL (UNII: KEH0A3F75J)	

Product Characteristics

Color	brown	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:84179-210-01	1 in 1 CARTON	08/01/2011	
1		30 mL 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 Drug	M020	08/01/2011	

Labeler - JENTRY KELLEY (040036679)

Registrant - MANA Products, Inc. (078870292)

Establishment

Name	Address	ID/FEI	Business Operations
MANA Products, Inc		078870292	manufacture(84179-210)

Establishment

Name	Address	ID/FEI	Business Operations
MANA Products, Inc		032870270	manufacture(84179-210)

Revised: 4/2024

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