PARASOL DAILY MINERAL SUNSCREEN- zinc oxide cream Bodi & Wax, LLC



WHOLE PERSON SKINCARE

Parasol Daily Mineral Sunscreen



NET WT. 1.75 OZ / 50 G

DRUG FACTS

ACTIVE INGREDIENTS:

PURPOSE:

Zinc Oxide 16%

Sunscreen

USES:

Helps prevent sunburn. If used as directed with other sun protection measures (see Directions), decrease 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 a 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 product 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 your risk of skin cancer and early skin 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 10am-2pm
- · Wear long-sleeved shirts, pants, hats, and sunglasses.

INACTIVE INGREDIENTS:

Capric/Caprylic Triglyceride, Ceramide 3, Cyclohexasiloxane,
Cyclopentasiloxane, Dimethicone, Dimethicone Crosspolymer, Dimethicone/
Vinyl Dimethicone Crosspolymer, Dimethicone, Iron Oxide, PEG-10 Dimethicone, Polyhydroxystearic Acid, Tetrahexyldecyl
Ascorbate, Tocopheryl Acetate, Vinyl Dimethicone/Hydrogen Dimethicone
Silsesquioxane Crosspolymer

OTHER INFORMATION:

- Protect this product from excessive heat and direct sun.
- · May stain some fabrics.



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MANUFACTURED FOR MEG BODI SKINCARE, AUSTIN TEXAS 78751

PARASOL DAILY MINERAL SUNSCREEN

zinc oxide cream

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Prod	uct	Inform	nation

Product Type HUMAN OTC DRUG Item Code (Source) NDC:85373-208

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name		Basis of Strength	Strength
	ZINC OXIDE (UNII: SOI2LOH54Z) (ZINC OXIDE - UNII:SOI2LOH54Z)	ZINC OXIDE	160 mg in 1 g

Inactive	

Ingradiant Nama	Strongth
Ingredient Name	Strength

VINYL DIMETHICONE/METHICONE SILSESQUIOXANE CROSSPOLYMER (UNII: 9NH1UDD2RR) ALPHA-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0) **CERAMIDE 3** (UNII: 4370DF050B) **DIMETHICONE CROSSPOLYMER** (UNII: UF7620L1W6) **CI 77492** (UNII: EX438O2MRT) PEG-10 DIMETHICONE (600 CST) (UNII: 8PR7V1SVM0) **DIMETHICONE** (UNII: 92RU3N3Y10) **DIMETHICONE/VINYL DIMETHICONE CROSSPOLYMER (SOFT PARTICLE)** (UNII: 9E4CO0W6C5) CI 77499 (UNII: XM0M87F357) POLYHYDROXYSTEARIC ACID (2300 MW) (UNII: YXH47AOU0F) TETRAHEXYLDECYL ASCORBATE (UNII: 9LBV3F07AZ) CAPRYLIC/CAPRIC TRIGLYCERIDE (UNII: C9H2L21V7U) CYCLOHEXASILOXANE (UNII: XHK3U310BA) HYDROGEN DIMETHICONE (20 CST) (UNII: 12Z59IF64N) **DIMETHICONOL (2000 CST)** (UNII: T74O12AN6Y) CYCLOPENTASILOXANE (UNII: 0THT5PCI0R) CI 77491 (UNII: 1K09F3G675)

Packaging				
# Item Code	Package Description	Marketing Start Date	Marketing End Date	
1 NDC:85373-208-50	50 g in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	05/05/2025		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M020	05/05/2025	

Labeler - Bodi & Wax, LLC (013179703)

Establishment			
Name	Address	ID/FEI	Business Operations
Fragrance Manufacturing INC		793406000	manufacture(85373-208)

Establishment				
Name	Address	ID/FEI	Business Operations	
Custom Analytics LLC		144949372	analysis (85373-208)	

Revised: 5/2025 Bodi & Wax, LLC