

**FULL SPECTRUM MATTE AMBITION SKIN PRIMER SPF 20- ensulizole liquid  
Noxell Corporation**

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**Covergirl Full Spectrum Matte Ambition Primer SPF 20**

**Active ingredient**

Ensulizole 3%

**Purpose**

Sunscreen

**Uses**

helps prevent sunburn

**Warnings**

Skin Cancer/Skin Aging Alert: Spending time in the sun increases your risk of skin cancer and early skin aging. This product has been shown only to help prevent sunburn, not skin cancer or early skin aging.

For external use only.

Do not use on damaged or broken skin

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

Stop use and ask a doctor if rash occurs

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

**Directions**

- apply liberally 15 minutes before sun exposure
- reapply at least every 2 hours
- use water resistant sunscreen if swimming or sweating
- children under 6 months: ask a doctor

**Other information**

Protect this product from excessive heat and direct sun

**Inactive ingredients**

Aqua/Water/Eau, Cyclopentasiloxane, Talc, Propylene Glycol, Dimethicone, Aluminum

Starch Octenylsuccinate, PEG/PPG-18/18 Dimethicone, Sodium Chloride, PVP, Phenoxyethanol, Sodium Hydroxide, Acrylonitrile/Methyl Methacrylate/Vinylidene Chloride Copolymer, Synthetic Beeswax, Trihydroxystearin, Methicone, 1,2-Hexanediol, Caprylyl Glycol, Silica, Sodium Benzoate, Synthetic Wax, Polyglyceryl-4 Isostearate, Cetyl PEG/PPG-10/1 Dimethicone, Hexyl Laurate, Isopropyl Titanium Triisostearate, Ethylene Brassylate, Polyethylene.

## Questions?

1-800-426-8374

full spectrum

COVERGIRL

MATTE AMBITION

MATTIFYING

SKIN PRIMER +

ENSULIZOLE

SUNSCREEN

SPF 20

30 mL (1.0 FL OZ)



# FULL SPECTRUM MATTE AMBITION SKIN PRIMER SPF 20

ensulizole liquid

## Product Information

|                                |                |                           |               |
|--------------------------------|----------------|---------------------------|---------------|
| <b>Product Type</b>            | HUMAN OTC DRUG | <b>Item Code (Source)</b> | NDC:22700-201 |
| <b>Route of Administration</b> | TOPICAL        |                           |               |

## Active Ingredient/Active Moiety

| <b>Ingredient Name</b>                                       | <b>Basis of Strength</b> | <b>Strength</b> |
|--|--------------------------|-----------------|
| ENSULIZOLE (UNII: 9YQ9DI1W42) (ENSULIZOLE - UNII:9YQ9DI1W42) | ENSULIZOLE               | 3 g in 100 mL   |

## Inactive Ingredients

| <b>Ingredient Name</b>                                    | <b>Strength</b> |
|---|-----------------|
| ALUMINUM STARCH OCTENYLSUCCINATE (UNII: I9PJ0O6294)       |                 |
| PEG/PPG-18/18 DIMETHICONE (UNII: 9H0AO7T794)              |                 |
| PHENOXYETHANOL (UNII: HIE492ZZ3T)                         |                 |
| SODIUM HYDROXIDE (UNII: 55X04QC32I)                       |                 |
| HEXYL LAURATE (UNII: 4CG9F9W01Q)                          |                 |
| ETHYLENE BRASSYLATE (UNII: 9A87HC7ROD)                    |                 |
| DIMETHICONE (UNII: 92RU3N3Y1O)                            |                 |
| SODIUM CHLORIDE (UNII: 451W47IQ8X)                        |                 |
| POLYGLYCERYL-4 ISOSTEARATE (UNII: 820DPX33S7)             |                 |
| TALC (UNII: 7SEV7J4R1U)                                   |                 |
| WATER (UNII: 059QF0KO0R)                                  |                 |
| CETYL PEG/PPG-10/1 DIMETHICONE (HLB 2) (UNII: V2W71V8T0X) |                 |
| ISOPROPYL TITANIUM TRIISOSTEARATE (UNII: 949E3KB1I)       |                 |

## Packaging

| # | Item Code        | Package Description                                | Marketing Start Date | Marketing End Date |
|---|------------------|--|----------------------|--------------------|
| 1 | NDC:22700-201-30 | 30 mL in 1 TUBE; Type 0: Not a Combination Product | 02/28/2019           |                    |

## Marketing Information

| <b>Marketing Category</b> | <b>Application Number or Monograph Citation</b> | <b>Marketing Start Date</b> | <b>Marketing End Date</b> |
|---------------------------|---|-----------------------------|---------------------------|
| OTC Monograph Drug        | M020  | 02/28/2019                  |                           |

**Labeler** - Noxell Corporation (003082997)

**Establishment**

| <b>Name</b>        | <b>Address</b> | <b>ID/FEI</b> | <b>Business Operations</b> |
|--------------------|----------------|---------------|----------------------------|
| Noxell Corporation |                | 003082997     | manufacture(22700-201)     |

Revised: 11/2024

Noxell Corporation