

**ELTAMD UV CLEAR SPF46- zinc oxide and octinoxate sunscreen lotion**  
**CP Skin Health Group, Inc.**

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**EltAMD UV Clear SPF46**

**Warnings**

For external use only Do not use on damaged or broken skin When using the product keep out of eyes. Rinse with water to remove Stop use and ask a physician if rash occurs Keep out of reach of children. If product is swallowed, get medical help or contact a Poison Control Center right away.

**Active Ingredients**

Zinc Oxide 9.0% Sunscreen

Octinoxate 7.5% Sunscreen

**Uses**

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.

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**Directions**

apply liberally to face and neck 15 minutes before sun exposure use a water-resistant sunscreen if swimming or sweating reapply at least every 2 hours 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 of 15 or higher and other sun protection measures including: limit time in the sun, especially from 10 a.m.- 2 p.m. wear long-sleeve shirts, pants, hats and sunglasses children under 6 months: Ask a physician

**Inactive Ingredients**

Purified water, Cyclopentasiloxane, Niacinamide, Octyldodecyl Neopentanoate, Hydroxyethyl Acrylate/Sodium Acryloyldimethyl Taurate Copolymer, Polyisobutene, PEG-7 Trimethylolpropane Coconut Ether, Sodium Hyaluronate, Tocopheryl Acetate, Lactic Acid, Oleth-3 Phosphate, Phenoxyethanol, Butylene Glycol, Iodopropynyl Butylcarbamate, Triethoxycaprylylsilane

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## Labeling



# ELTAMD UV CLEAR SPF46

zinc oxide and octinoxate sunscreen lotion

## Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:72043-2500
<b>Route of Administration</b>	TOPICAL		

## Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
<b>ZINC OXIDE</b> (UNII: SOI2LOH54Z) (ZINC CATION - UNII:13S1S8SF37)	ZINC CATION	90 g in 1000 g
<b>OCTINOXATE</b> (UNII: 4Y5P7MUD51) (OCTINOXATE - UNII:4Y5P7MUD51)	OCTINOXATE	75 g in 1000 g

## Inactive Ingredients

Ingredient Name	Strength
<b>BUTYLENE GLYCOL</b> (UNII: 3XUS85K0RA)	
<b>IODOPROPYNYL BUTYLCARBAMATE</b> (UNII: 603P14DHEB)	
<b>PHENOXYETHANOL</b> (UNII: HIE492ZZ3T)	
<b>WATER</b> (UNII: 059QF0KO0R)	
<b>LACTIC ACID</b> (UNII: 33X04XA5AT)	
<b>HYALURONATE SODIUM</b> (UNII: YSE9PPT4TH)	
<b>NIACINAMIDE</b> (UNII: 25X51I8RD4)	
<b>CYCLOMETHICONE 5</b> (UNII: 0THT5PCI0R)	
<b>.ALPHA.-TOCOPHEROL ACETATE, DL-</b> (UNII: WR1WPI7EW8)	
<b>OCTYLDODECYL BENZOATE</b> (UNII: R04N7AS5EA)	
<b>OLETH-3 PHOSPHATE</b> (UNII: 8Q0Z18J1VL)	
<b>TRIETHOXYCAPRYLYLSILANE</b> (UNII: LDC331P08E)	
<b>HYDROXYETHYL ACRYLATE/SODIUM ACRYLOYLDIMETHYL TAURATE COPOLYMER (100000 MPA.S AT 1.5%)</b> (UNII: 86FQE96TZ4)	
<b>POLYISOBUTYLENE (1000 MW)</b> (UNII: 5XB3A63Y52)	

## Product Characteristics

<b>Color</b>	white	<b>Score</b>	
<b>Shape</b>		<b>Size</b>	
<b>Flavor</b>		<b>Imprint Code</b>	
<b>Contains</b>			

## Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72043-2500-1	48 g in 1 BOTTLE; Type 0: Not a Combination Product	01/10/2018	
2	NDC:72043-2500-5	14 g in 1 BOTTLE; Type 0: Not a Combination Product	12/17/2020	
3	NDC:72043-2500-2	2 g in 1 PACKET; Type 0: Not a Combination Product	07/06/2022	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M020	01/10/2018	

**Labeler** - CP Skin Health Group, Inc. (611921669)

**Registrant** - Swiss-American CDMO, LLC (080170933)

## Establishment

Name	Address	ID/FEI	Business Operations
Swiss-American CDMO, LLC		080170933	manufacture(72043-2500)

Revised: 11/2023

CP Skin Health Group, Inc.