

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

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

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

Active Ingredient/Active Moiety

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

Inactive Ingredients

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

Product Characteristics

Color	white	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

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:72043-2500-3	106 g in 1 TUBE; Type 0: Not a Combination Product	06/04/2025	
3	NDC:72043-2500-5	14 g in 1 BOTTLE; Type 0: Not a Combination Product	12/17/2020	

4	NDC:72043-2500-2	2 g in 1 PACKET; Type 0: Not a Combination Product	07/06/2022	
5	NDC:72043-2500-1	48 g in 1 BOTTLE; Type 0: Not a Combination Product	05/01/2025	

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: 6/2025

CP Skin Health Group, Inc.