

KIEHLS DERMATOLOGIST SOLUTIONS BETTER SCREEN UV SERUM BROAD SPECTRUM SPF 50 PLUS SUNSCREEN- avobenzone, homosalate, octisalate and octocrylene lotion
L'Oreal USA Products Inc

Drug Facts

Active ingredients

Avobenzone 3%

Homosalate 7%

Octisalate 5%

Octocrylene 7%

Purpose

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

Warnings

For external use only

Flammable until dry. Do not use near fire, flame or heat.

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 swallowed, get medical help or contact a Poison Control Center right away.

Directions

- shake well before use
- apply liberally 15 minutes before sun exposure
- reapply at least every 2 hours
- use a water resistant sunscreen if swimming or sweating
- **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 10 a.m. - 2p.m.
 - wear long-sleeved shirts, pants, hats and sunglasses
 - children under 6 months of age: Ask a doctor

Other information

- protect the product in this container from excessive heat and direct sun

Inactive ingredients

water, glycerin, c15-19 alkane, propanediol, c12-22 alkyl acrylate/hydroxyethylacrylate copolymer, tocopherol, sodium stearyl glutamate, butylene glycol, cetearyl alcohol, sclerotium gum, hydroxyacetophenone, caprylyl glycol, sodium starch octenylsuccinate, glyceryl stearate, jojoba esters, helianthus annuus (sunflower) seed wax, hydroxypropyl starch phosphate, trisodium ethylenediamine disuccinate, carbomer, sodium lactate, sodium hydroxide, polysorbate 20, polyglycerin-3, palmitoyl tripeptide-1, palmitoyl tetrapeptide-7

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Directions

For sunscreen use:

- apply generously and evenly 15 minutes before sun exposure
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Questions or comments?

Call toll free 1-800-946-4453

Fmla 2068500
FJ.L. Code V70029421/1

KIEHL'S SINCE 1851 LLC,
NEW YORK, NY 10014

Ingredients sourced
worldwide, made in U.S.A.

Dist. Kiehl's Canada, Montreal H4T 1K5
14 rue Royale 75008 Paris • LONDON W6 8AZ
TSA 75000 93584 STOUEN CEDEX FR
www.kiehls.com • 3757918



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SINCE 1851

KIEHL'S
DERMATOLOGIST
SOLUTIONS™

Better Screen™⁺
UV Serum

SUNSCREEN

Broad Spectrum SPF 50+

with Collagen Peptide

*Helps Visibly Correct Fine
Lines, Uneven Skin Tone
and Dull Skin

1.7 fl. oz. - 50 ml

Kiehl's Dermatologist Solutions are highly advanced targeted treatments developed by Kiehl's Since 1851. Relying on our extensive skincare expertise and botanical knowledge, Kiehl's chemists partner with an international team of leading dermatologists to deliver powerful, yet safe skincare solutions.

Better Screen* UV Serum goes beyond our traditional sunscreens, helping to correct visible signs of aging as well as helping to protect against UVA and UVB rays. Formulated with Collagen Peptide, our formula visibly diminishes fine lines, uneven skin tone and dullness that can be caused by UV damage. With an invisible finish, this weightless serum is absorbed easily into skin to deliver protection and correction across all skin tones and types, including sensitive and acne prone.

*Compared to other Kiehl's sunscreen formulas.



UPC
80%



Learn More

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
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
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Ingredients sourced worldwide, made in U.S.A.
Dist. Kiehl's Canada, Montreal H4T 1K5
14 rue Royale 75008 Paris • LONDON W6 8AZ
TSA 75000 93584 STOUEN CEDEX FR
www.kiehls.com • 3757918



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UV Serum

SUNSCREEN
Broad Spectrum SPF 50+

with Collagen Peptide


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
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Learn More

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avobenzone, homosalate, octisalate and octocrylene lotion

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:49967-053
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
AVOBENZONE (UNII: G63QQF2NOX) (AVOBENZONE - UNII:G63QQF2NOX)	AVOBENZONE	30 mg in 1 mL
HOMOSALATE (UNII: V06SV4M95S) (HOMOSALATE - UNII:V06SV4M95S)	HOMOSALATE	70 mg in 1 mL
OCTISALATE (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	50 mg in 1 mL
OCTOCRYLENE (UNII: 5A68WGF6WM) (OCTOCRYLENE - UNII:5A68WGF6WM)	OCTOCRYLENE	70 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6A3C0OX)	

C15-19 ALKANE (UNII: C187N1IM01)	
PROPANEDIOL (UNII: 5965N8W85T)	
TOCOPHEROL (UNII: R0ZB2556P8)	
SODIUM STEAROYL GLUTAMATE (UNII: 65A9F4P024)	
BUTYLENE GLYCOL (UNII: 3XUS85K0RA)	
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)	
BETASIZOFIRAN (UNII: 2X51AD1X3T)	
HYDROXYACETOPHENONE (UNII: G1L3HT4CMH)	
CAPRYLYL GLYCOL (UNII: 00YIU5438U)	
GLYCERYL MONOSTEARATE (UNII: 230OU9XXE4)	
HYDROGENATED JOJOBA OIL/JOJOBA OIL, RANDOMIZED (IODINE VALUE 64-70) (UNII: 96YYQ5TK1K)	
HELIANTHUS ANNUUS SEED WAX (UNII: 42DG15CHXV)	
TRISODIUM ETHYLENEDIAMINE DISUCCINATE (UNII: YA22H34H9Q)	
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)	
SODIUM LACTATE (UNII: TU7HW0W0QT)	
SODIUM HYDROXIDE (UNII: 55X04QC32I)	
POLYSORBATE 20 (UNII: 7T1F30V5YH)	
POLYGLYCERIN-3 (UNII: 4A0NCJ6RD6)	
PALMITOYL TRIPEPTIDE-1 (UNII: RV743D216M)	
PALMITOYL TETRAPEPTIDE-7 (UNII: Q41S464P1R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:49967-053-01	1 in 1 CARTON	07/01/2023	
1		50 mL in 1 BOTTLE; Type 0: Not a Combination Product		
2	NDC:49967-053-02	1 in 1 CARTON	07/01/2023	
2		15 mL in 1 TUBE; Type 0: Not a Combination Product		
3	NDC:49967-053-03	2 mL in 1 PACKET; Type 0: Not a Combination Product	07/01/2023	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M020	07/01/2023	

Labeler - L'Oreal USA Products Inc (002136794)

Establishment

Name	Address	ID/FEI	Business Operations
L'Oreal USA, Inc.		185931458	manufacture(49967-053) , pack(49967-053)

