

**LANCOME PARIS TEINT IDOLE ULTRA WEAR BREATHABLE COVERAGE  
FOUNDATION RENO BROAD SPECTRUM SPF 25 SUNSCREEN- octinoxate and  
titanium dioxide liquid  
L'Oreal USA Products Inc**

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**Drug Facts**

**Active ingredients**

Octinoxate 6.7%

Titanium dioxide 11.5%

**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

For sunscreen use:

- shake well
- 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. - 2 p.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, dimethicone, isododecane, alcohol denat., trimethylsiloxy silicate, butylene glycol, PEG-10 dimethicone, perlite, synthetic fluorophlogopite, moringa oleifera seed extract, polymnia sonchifolia root juice, calcium aluminum borosilicate, glycerin, dipentaerythrityl tetrahydroxystearate/tetraisostearate, alpha-glucan oligosaccharide, sodium hyaluronate, silica, silica silylate, HDI/trimethylol hexyllactone crosspolymer, cellulose, aluminum hydroxide, magnesium sulfate, nylon-12, disodium phosphate, disodium stearyl glutamate, isopropyl lauroyl sarcosinate, hydrogen dimethicone, citric acid, diisopropyl sebacate, bis-PEG/PPG-14/14 dimethicone, lactobacillus, maltodextrin, disteardimonium hectorite, BHT, tocopherol, phenoxyethanol; may contain: titanium dioxide, iron oxides, chromium oxide greens

## Questions or comments?

**1-800-LANCOME**

**(1-800-526-2663)**

Monday - Friday

(9 a.m. - 5 p.m. EST)

LANCÔME

TEINT  
IDOLE  
ULTRA  
WEAR

SUNSCREEN  
BROAD SPECTRUM  
SPF 25

UP TO 24H WEAR FOUNDATION  
BREATHABLE COVERAGE

TRANSFER-RESISTANT - PARABEN-FREE  
NON-COMEDOGENIC - NON ACNEGENIC

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1-800-LANCOME (1-800-626-2663) Monday - Friday (9 a.m. - 5 p.m. EST)

6992804

TESTER, NOT FOR SALE.

V293548/1

LANCÔME, LUXURY PRODUCTS LLC, 10 HUDSON YARDS, NEW YORK, NY 10001

IMP. BY LANCÔME CANADA, MONTREAL, HAT 1K5

TS# 750000 93684 ST-QUIEN CEDEX FR

Lancome.com Made in France



100 ml - 3.38 FL. OZ.

63.0

126.0

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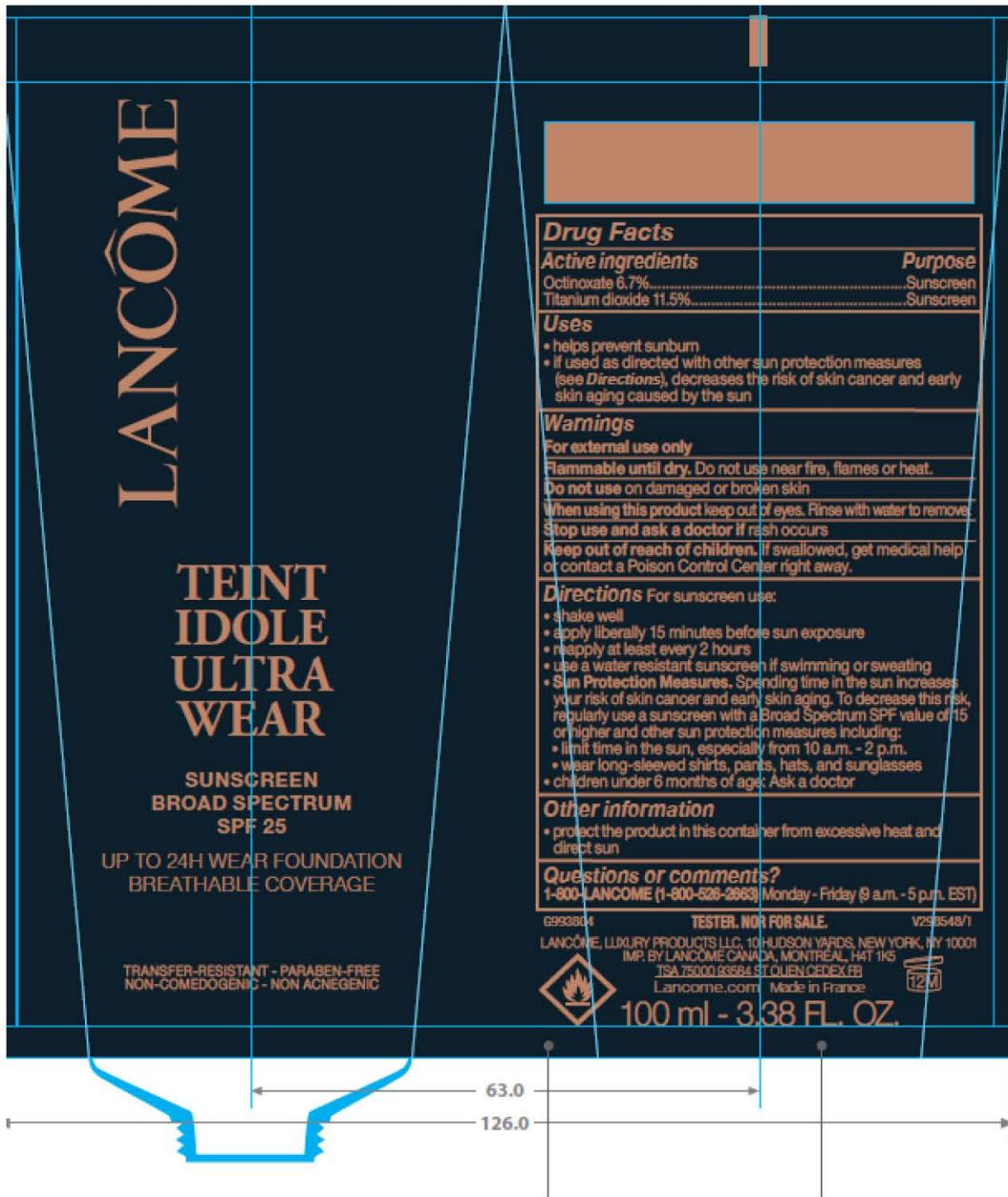
Lancome.com Made in France



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63.0

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C1 - Internal use

**LANCÔME PARIS TEINT IDOLE ULTRA WEAR BREATHABLE**

# COVERAGE FOUNDATION RENO BROAD SPECTRUM SPF 25 SUNSCREEN

octinoxate and titanium dioxide liquid

## Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:49967-303
<b>Route of Administration</b>	TOPICAL		

## Active Ingredient/Active Moiety

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
<b>OCTINOXATE</b> (UNII: 4Y5P7MUD51) (OCTINOXATE - UNII:4Y5P7MUD51)	OCTINOXATE	67 mg in 1 mL
<b>TITANIUM DIOXIDE</b> (UNII: 15FIX9V2JP) (TITANIUM DIOXIDE - UNII:15FIX9V2JP)	TITANIUM DIOXIDE	115 mg in 1 mL

## Inactive Ingredients

<b>Ingredient Name</b>	<b>Strength</b>
<b>WATER</b> (UNII: 059QF0KO0R)	
<b>DIMETHICONE</b> (UNII: 92RU3N3Y1O)	
<b>ISODODECANE</b> (UNII: A8289P68Y2)	
<b>ALCOHOL</b> (UNII: 3K9958V90M)	
<b>TRIMETHYLSILOXYSILICATE (M/Q 0.6-0.8)</b> (UNII: 5041RX63GN)	
<b>BUTYLENE GLYCOL</b> (UNII: 3XUS85K0RA)	
<b>PEG-10 DIMETHICONE (600 CST)</b> (UNII: 8PR7V1SVM0)	
<b>PERLITE</b> (UNII: 0SG101ZGK9)	
<b>MAGNESIUM POTASSIUM ALUMINOSILICATE FLUORIDE</b> (UNII: YK3DC63Y5M)	
<b>MORINGA OLEIFERA SEED</b> (UNII: TIX5482832)	
<b>SMALLANTHUS SONCHIFOLIUS ROOT JUICE</b> (UNII: M9S7HX36CT)	
<b>CALCIUM ALUMINUM BOROSILICATE</b> (UNII: 3JRB8A35M0)	
<b>GLYCERIN</b> (UNII: PDC6A3C0OX)	
<b>DIPENTAERYTHRITYL TETRAHYDROXYSTEARATE/TETRAISOSTEARATE</b> (UNII: 230K0823CE)	
<b>.ALPHA.-GLUCAN OLIGOSACCHARIDE</b> (UNII: S95658MI3W)	
<b>HYALURONATE SODIUM</b> (UNII: YSE9PPT4TH)	
<b>SILICON DIOXIDE</b> (UNII: ETJ7Z6XBU4)	
<b>HEXAMETHYLENE DIISOCYANATE/TRIMETHYLOL HEXYLLACTONE CROSSPOLYMER</b> (UNII: WB5K9Y35Y9)	
<b>POWDERED CELLULOSE</b> (UNII: SMD1X3XO9M)	
<b>ALUMINUM HYDROXIDE</b> (UNII: 5QB0T2IUN0)	
<b>MAGNESIUM SULFATE, UNSPECIFIED FORM</b> (UNII: DE08037SAB)	
<b>NYLON-12</b> (UNII: 446U8J075B)	
<b>SODIUM PHOSPHATE, DIBASIC, ANHYDROUS</b> (UNII: 22ADO53M6F)	
<b>DISODIUM STEAROYL GLUTAMATE</b> (UNII: 45ASM2L11M)	
<b>ISOPROPYL LAUROYL SARCOSINATE</b> (UNII: LYR06W430J)	
<b>HYDROGEN DIMETHICONE (13 CST)</b> (UNII: 4QGR4P2YOI)	
<b>CITRIC ACID MONOHYDRATE</b> (UNII: 2968PHW8QP)	
<b>DIISOPROPYL SEBACATE</b> (UNII: J8T3X564IH)	
<b>BIS-PEG/PPG-14/14 DIMETHICONE</b> (UNII: X2I70H0QJE)	
<b>EMMENTAL LACTOBACILLUS</b> (UNII: B253LSV04X)	

<b>MALTODEXTRIN</b> (UNII: 7CVR7L4A2D)	
<b>DISTEARDIMONIUM HECTORITE</b> (UNII: X687XDK09L)	
<b>BUTYLATED HYDROXYTOLUENE</b> (UNII: 1P9D0Z171K)	
<b>TOCOPHEROL</b> (UNII: R0ZB2556P8)	
<b>PHENOXYETHANOL</b> (UNII: HIE492ZZ3T)	
<b>FERRIC OXIDE RED</b> (UNII: 1K09F3G675)	

<b>Packaging</b>				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:49967-303-01	1 in 1 CARTON	12/12/2022	
1		30 mL in 1 BOTTLE; Type 0: Not a Combination Product		
2	NDC:49967-303-02	100 mL in 1 TUBE; Type 0: Not a Combination Product	12/12/2022	
3	NDC:49967-303-03	1 mL in 1 PACKET; Type 0: Not a Combination Product	12/12/2022	01/22/2027
4	NDC:49967-303-04	0.4 mL in 1 PACKAGE; Type 0: Not a Combination Product	12/12/2022	12/28/2026

<b>Marketing Information</b>			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M020	12/12/2022	

**Labeler** - L'Oreal USA Products Inc (002136794)

<b>Establishment</b>			
Name	Address	ID/FEI	Business Operations
SICOS ET CIE		276993581	manufacture(49967-303) , pack(49967-303)

<b>Establishment</b>			
Name	Address	ID/FEI	Business Operations
BPS60		272259304	pack(49967-303)

<b>Establishment</b>			
Name	Address	ID/FEI	Business Operations
Socoplan		276221405	pack(49967-303)