

**ULTA BEAUTY TINTED MOISTURIZER CREAM SPF 24 TAN- avobenzone, homosalate, octisalate, octocrylene cream
Cosmax Usa, Corporation**

Ulta Beauty Tinted Moisturizer Cream Spf 24 Tan

Drug Facts

Active ingredients

Avobenzone 3%

Homosalate 9.5%

Octisalate 4.5%

Octocrylene 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.

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 15 minutes before sun exposure
- reapply:
- after 80 minutes of swimming or sweating

- immediately after towel drying
- 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 doctor

Inactive ingredients

water, glycerin, butylene glycol, triisostearyl citrate, titanium dioxide, pentylene glycol, polyglycerin-4, cetearyl olivate, candelilla/jojoba/rice bran polyglyceryl-3 esters, glyceryl stearate, phenoxyethanol, sorbitan olivate, potassium cetyl phosphate, cetearyl alcohol, sodium stearoyl latylate, oleyl alcohol, carbomer, olea europaea (olive) fruit oil, tromethamine, iron oxides, acacia senegal gum, ethylhexylglycerin, galactoarabinan, xanthan gum, betaine, citric acid, disodium edta, potassium sorbate, tremella fuciformis sporocarp extract

Other information

- protect this product from excessive heat and direct sun

Questions?

1-866-983-8582

Package Labeling:

TT - top of top ply

TAN • HAVANE



Drug Facts

Active ingredients

Avobenzone 3%,
Homosalate 9.5%,
Octisalate 4.5%,
Octocrylene 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

Drug Facts (continued)

Warnings

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.

Drug Facts (continued)

Directions • apply 15 minutes before sun exposure • reapply: • after 80 minutes of swimming or sweating • immediately after towel drying • 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

BT - back of top ply

3.4

Drug Facts (continued)

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 doctor

Drug Facts (continued)

Inactive ingredients water, glycerin, butylene glycol, triisostearyl citrate, titanium dioxide, pentylene glycol, polyglycerin-4, cetearyl olivate, candelilla/jojoba/rice bran polyglyceryl-3 esters, glyceryl stearate, phenoxyethanol, sorbitan olivate, potassium cetyl phosphate, cetearyl

Drug Facts (continued)

alcohol, sodium stearyl lactylate, oleyl alcohol, carbomer, olea europaea (olive) fruit oil, tromethamine, iron oxides, acacia senegal gum, ethylhexylglycerin, galactoarabinan, xanthan gum, betaine, citric acid, disodium edta, potassium sorbate, tremella fuciformis sporocarp extract

Drug Facts (continued)

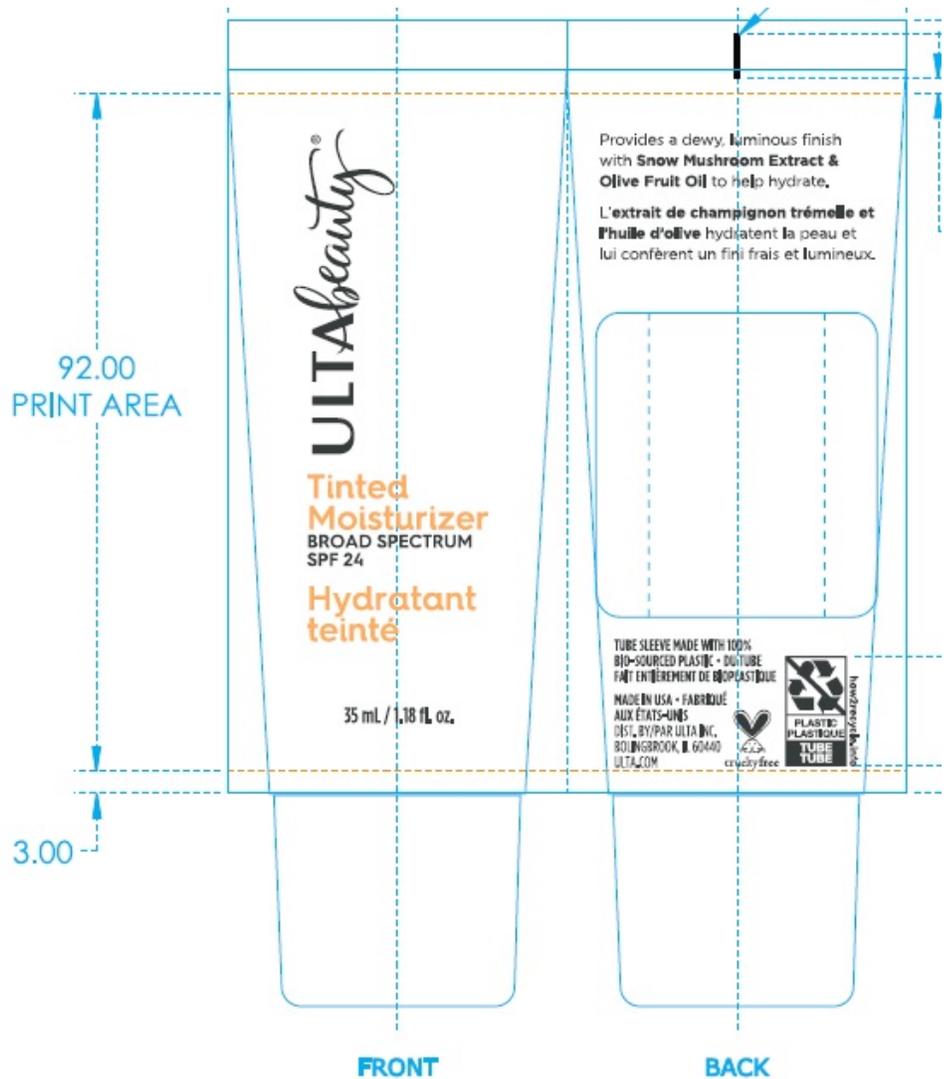
Other information • protect this product from excessive heat and direct sun

Questions?

1-866-983-8582



RENDERING DO NOT PRINT



ULTA BEAUTY TINTED MOISTURIZER CREAM SPF 24 TAN

avobenzone, homosalate, octisalate, octocrylene cream

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:68577-035
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	95 mg in 1 mL
OCTISALATE (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	45 mg in 1 mL
OCTOCRYLENE (UNII: 5A68WGF6WM) (OCTOCRYLENE - UNII:5A68WGF6WM)	OCTOCRYLENE	50 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
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GLYCERIN (UNII: PDC6A3C0OX)
BUTYLENE GLYCOL (UNII: 3XUS85KORA)
TRISOSTEARYL CITRATE (UNII: 50XT3250OH)
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)
PENTYLENE GLYCOL (UNII: 50C1307PZG)
POLYGLYCERIN-4 (UNII: YX76MGM96B)
CETEARYL OLIVATE (UNII: 58B69Q84JO)
GLYCERYL MONOSTEARATE (UNII: 230OU9XXE4)
PHENOXYETHANOL (UNII: HIE492ZZ3T)
SORBITAN OLIVATE (UNII: MDL271E3GR)
POTASSIUM CETYL PHOSPHATE (UNII: 03KCY6P7UT)
CETOSTEARYL ALCOHOL (UNII: 2DMT128M1S)
SODIUM STEAROYL LACTYLATE (UNII: IN99IT31LN)
OLEYL ALCOHOL (UNII: 172F2VN8DV)
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)
OLIVE OIL (UNII: 6UYK2W1W1E)
TROMETHAMINE (UNII: 023C2WHX2V)
FERRIC OXIDE RED (UNII: 1K09F3G675)
ACACIA (UNII: 5C5403N26O)
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)
GALACTOARABINAN (UNII: SL4SX1O487)
XANTHAN GUM (UNII: TTV12P4NEE)
BETAINE (UNII: 3SCV180C9W)
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)
EDETATE DISODIUM ANHYDROUS (UNII: 8NLQ36F6MM)
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)
TREMELLA FUCIFORMIS FRUITING BODY (UNII: GG8N28393G)
WATER (UNII: 059QF0KO0R)

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:68577-035-01	35 mL in 1 TUBE; Type 0: Not a Combination Product	06/01/2021	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M020	06/01/2021	

Labeler - Cosmax Usa, Corporation (010990210)

Establishment

Name	Address	ID/FEI	Business Operations
Cosmax Usa, Corporation		010990210	manufacture(68577-035)

