

SKINSUIT LIP SPF 30- zinc oxide gel
Skin Authority LLC

Skin Authority - SKINSUIT LIP SPF-30 (46007-205)

ACTIVE INGREDIENT

ZINC OXIDE 10%

PURPOSE

SUNSCREEN

USES

- HELPS PREVENT SUNBURN
- IF USED WITH OTHER SUN PROTECTION MEASURES (SEE DIRECTIONS), DECREASES THE RISK OF SKIN CANCER AND EARLY SKIN AGING CAUSED BY SUN.

WARNINGS

FOR EXTERNAL USE ONLY.

DO NOT USE ON

- BROKEN SKIN
- SERIOUS BURNS

WHEN USING THIS PRODUCT

- KEEP OUT OF EYES. RINSE WITH WATER TO REMOVE.

STOP USE AND ASK A DOCTOR IF

- RASH OR IRRITATION DEVELOPS AND LASTS

KEEP OUT OF REACH OF CHILDREN.

- IF SWALLOWED, GET MEDICAL HELP OR CONTACT A POISON CONTROL CENTER RIGHT AWAY.

SUN PROTECTION MEASURES:

SPENDING TIME IN THE SUN INCREASES YOUR RISK OF SKIN CANCER AND EARLY SKIN AGING. TO DECREASE RISK, REGULARLY USE A SUNSCREEN WITH BROAD SPECTRUM SPF VALUE OF 15+ AND OTHER SUN PROTECTION MEASURES

- LIMIT TIME IN THE SUN FROM 10AM - 2PM
- WEAR LONG SLEEVED SHIRT, PANTS, HATS, AND SUNGLASSES

DIRECTIONS

APPLY EVENLY TO UPPER AND LOWER LIP 15 MINUTES BEFORE EXPOSURE. NATURAL TINT WILL ABSORB AND BECOME SHEER.

- REAPPLY EVERY 2 HOURS
- REAPPLY AS NEEDED AFTER SWIMMING OR PERSPIRING
- CHILDREN UNDER 6 MONTHS OF AGE: ASK A DOCTOR

HYDROGENATED POLY(C6-14 OLEFIN), SQUALANE, BUTYLOCTYL SALICYLATE, CETEARYL DIMETHICONE, ISODODECANE, ETHYLENE/PROPYLENE/STYRENE COPOLYMER, CAPRYLYL METHICONE, SYNTHETIC FLUORPHLOGOPITE, IRON OXIDE, LECITHIN, BUTYLENE/ETHYLENE/STYRENE COPOLYMER, CAPRYLYL GLYCOL, DIMETHICONE, DIMETHICONOL/PROPYLSILSESQUIOXANE/SILICATE CROSSPOLYMER, PENTAERYTHRITYL TETRA-DI-T-BUTYL HYDROXYHYDROCINNAMATE, PHENYLPROPANOL, PROPANEDIOL, TIN OXIDE, TITANIUM DIOXIDE (CI 77891), TOCOPHEROL.

OTHER INFORMATION

PROTECT THIS PRODUCT FROM EXCESSIVE HEAT AND DIRECT SUN. STORE AT 40° - 95°F (5° - 35°C).

QUESTIONS?

1-866-325-7546

MONDAY - FRIDAY (9 A.M. TO 5 P.M. PST)



SkinSuit™ Labio

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zinc oxide gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:46007-205
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ZINC OXIDE (UNII: SOI2LOH54Z) (ZINC CATION - UNII:13S1S8SF37)	ZINC CATION	10 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
HYDROGENATED POLY(C6-14 OLEFIN; 2 CST) (UNII: POTX083987)	
SQUALANE (UNII: GW89575KF9)	
BUTYLOCTYL SALICYLATE (UNII: 2EH13UN8D3)	
CETEARYL METHICONE (15000 MW) (UNII: VY9RTR7MSY)	
ISODODECANE (UNII: A8289P68Y2)	
CAPRYLYL TRISILOXANE (UNII: Q95M2P1KJL)	
SYNTHETIC WAX (1200 MW) (UNII: Q3Z4BCH099)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
CAPRYLYL GLYCOL (UNII: 00YIU5438U)	
DIMETHICONE (UNII: 92RU3N3Y1O)	
DIMETHICONOL/PROPYLSILSESQUIOXANE/SILICATE CROSSPOLYMER (450000000 MW) (UNII: 9KB5R958PB)	
PENTAERYTHRITOL TETRAKIS(3-(3,5-DI-TERT-BUTYL-4-HYDROXYPHENYL)PROPIONATE) (UNII: 255PIF62MS)	
PHENYLPROPANOL (UNII: 0F897O3O4M)	
PROPANEDIOL (UNII: 5965N8W85T)	
STANNIC OXIDE (UNII: KM7N50LOS6)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
TOCOPHEROL (UNII: ROZB2556P8)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:46007-205-11	3.2 g in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product	01/02/2020	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M020	01/02/2020	

