

**LIP SCREEN BROAD SPECTRUM SPF 40- avobenzene, homosalate, octisalate, octocrylene liquid  
Supergoop, LLC**

*Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.*

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**Lip Screen Broad Spectrum SPF 40**

Avobenzene 3% Sunscreen

Homosalate 5% Sunscreen

Octisalate 5% Sunscreen

Octocrylene 10% 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.

**Keep out of reach of children.** If product is swallowed, get medical help or contact a poison Control Center right away.

**Stop use and ask a doctor if** rash occurs

**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

**Directions**

Apply generously and evenly 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 value of 15 or higher and other sun protection measures including: • limit your 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.

**Inactive Ingredients**

Hydrogenated Polyisobutene, Diisostearyl Malate, Caprylic/Capric Triglyceride, Butyloctyl Salicylate, C12-15 Alkyl Benzoate, Polyurethane-79, Polyester-7, Butyrospermum Parkii (Shea) Butter, Cocos Nucifera (Coconut) Oil, Isopropyl Palmitate, Sesamum Indicum (Sesame) Seed Oil, Neopentyl Glycol Diheptanoate, Diethylhexyl Syringlidenemalonate, Caprylyl Glycol, Glycine Soja (Soybean) Oil, Tocopherol, 2-Methylbutyric Acid, Barosma Betulina Leaf Oil, Benzaldehyde, Dimethylhydroxy Furanone, Trans-2-Hexenal, Helianthus

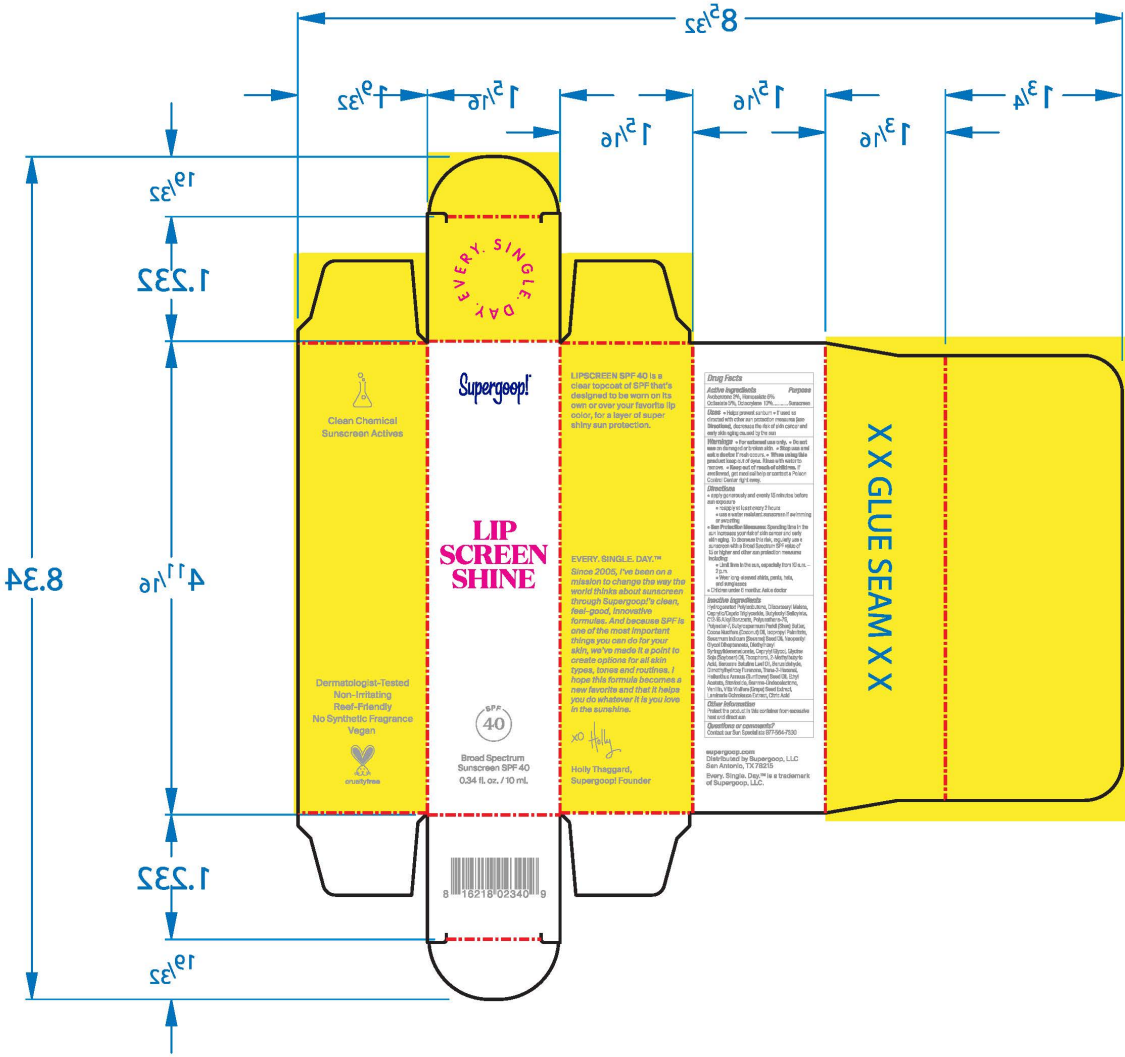
Annus (Sunflower) Seed Oil, Ethyl Acetate, Stevioside, Gamma-Undecalactone, Vanillin, Vitis Vinifera (Grape) Seed Extract, Laminaria Ochroleuca Extract, Citric Acid

Lip Screen

SPF 40

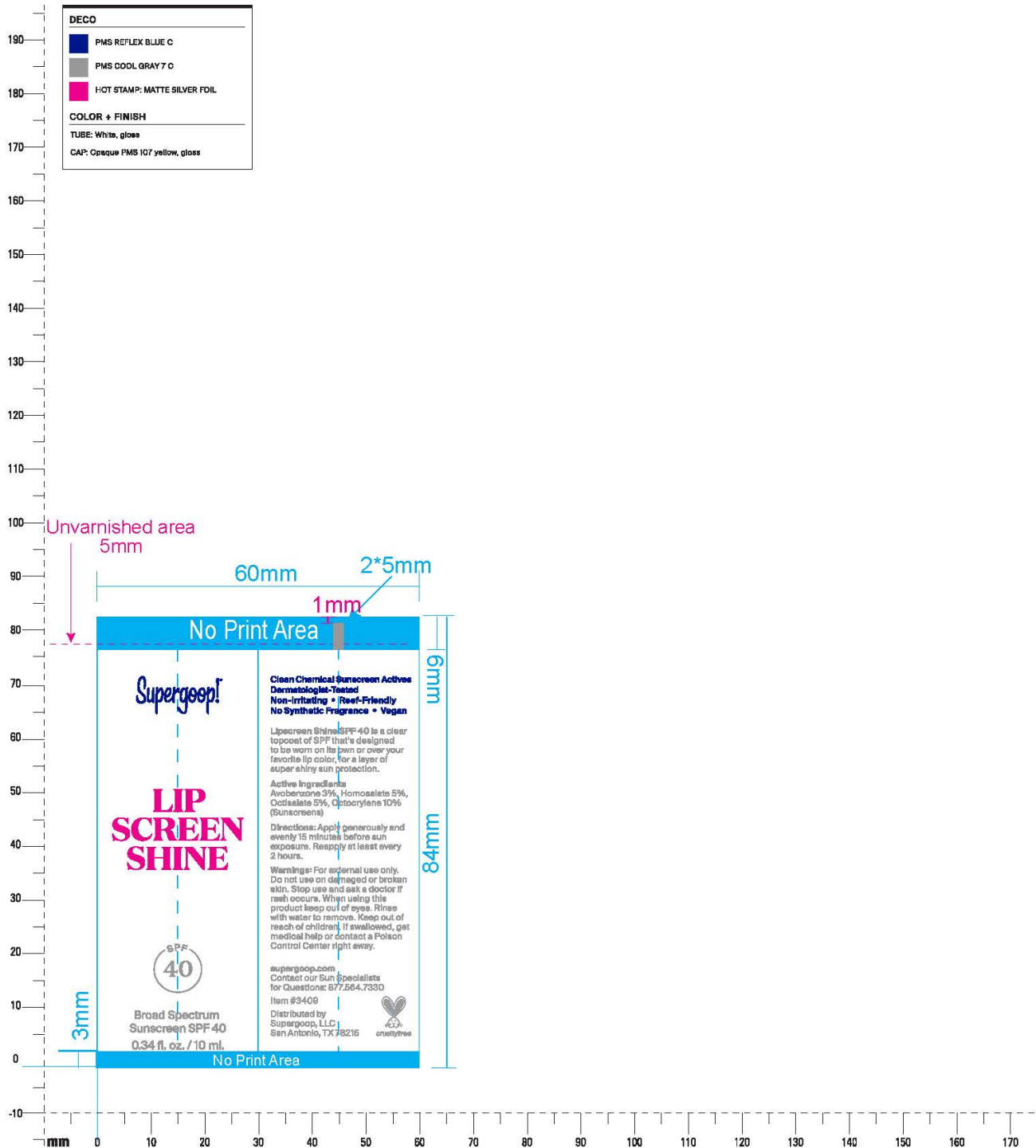
Broad Spectrum SPF 40

0.34 fl. oz. / 10 ml.



- Matte silver foil
- Reflex Blue C
- Cool Gray 7
- PMS 102 U

210  200	ARTWORK RELEASE STATUS
	<span style="background-color: green; display: inline-block; width: 15px; height: 10px; vertical-align: middle;"></span> PRODUCTION
	COMPONENT SPECS



	FILE INFORMATION		PROJECT INFORMATION		CLIENT APPROVAL CHECKLIST	
	FILE NAME		TUBE DIAMETER	∅ 19 mm	<b>Hot Stamping</b> <input type="checkbox"/> Silver <input type="checkbox"/> Gold <input type="checkbox"/> Other Metallic Color	
	CLIENT	B	TUBE LENGTH	84 mm	<input type="checkbox"/> Die Line <input type="checkbox"/> Copy <input type="checkbox"/> Colors <input type="checkbox"/> UPC <input type="checkbox"/> Photography <input type="checkbox"/> Illustration <input type="checkbox"/> Eye Mark	
	STARTED	CGP	PRINT INFORMATION		client signature _____ Date _____	
	MODIFIED	00/00/00 CGP	OFFSET	COLOR	Design: <i>Chel</i> Proof _____ UPC Test _____ Verify _____	
SOFTWARE	ai by Illustrator CS cdr by Coreldraw 9	SILK-SCREEN	COLOR	RELEASE DATE _____ <input checked="" type="checkbox"/> DIELINE DOES NOT PRINT		
PROOF NUMBER	PROOF SIZE	EXPORT	TUBE COLOR	IMPORTANT! This art has been checked and proofed for accuracy. It is the responsibility of the client to make all final approvals before the release of this art. PLEASE DOUBLE CHECK FOR ACCURACY.		
1	100%	PDF				

**LIP SCREEN BROAD SPECTRUM SPF 40**  
 avobenzone, homosalate, octisalate, octocrylene liquid

**Product Information**

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:75936-257
<b>Route of Administration</b>	TOPICAL		

**Active Ingredient/Active Moiety**

<b>Ingredient Name</b>	<b>Basis of Strength</b>	<b>Strength</b>
<b>AVOBENZONE</b> (UNII: G63QQF2NOX) (AVOBENZONE - UNII:G63QQF2NOX)	AVOBENZONE	3 g in 100 mL
<b>OCTOCRYLENE</b> (UNII: 5A68WGF6WM) (OCTOCRYLENE - UNII:5A68WGF6WM)	OCTOCRYLENE	10 g in 100 mL
<b>OCTISALATE</b> (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	5 g in 100 mL
<b>HOMOSALATE</b> (UNII: V06SV4M95S) (HOMOSALATE - UNII:V06SV4M95S)	HOMOSALATE	5 g in 100 mL

**Inactive Ingredients**

<b>Ingredient Name</b>	<b>Strength</b>
<b>ETHYL ACETATE</b> (UNII: 7684508NMZ)	
<b>CAPRYLYL GLYCOL</b> (UNII: 00YIU5438U)	
<b>CITRIC ACID MONOHYDRATE</b> (UNII: 2968PHW8QP)	
<b>POLYESTER-7</b> (UNII: 0841698D2F)	
<b>TOCOPHEROL</b> (UNII: R0ZB2556P8)	
<b>DIMETHYLHYDROXY FURANONE</b> (UNII: 20PI8YZP7A)	
<b>2-HEXENAL, (2E)-</b> (UNII: 69JX3AIR1I)	
<b>DIETHYLHEXYL SYRINGYLIDENEMALONATE</b> (UNII: 3V5U97P248)	
<b>LAMINARIA OCHROLEUCA</b> (UNII: 4R2124HE76)	
<b>AGATHOSMA BETULINA LEAF OIL</b> (UNII: KOS935A04V)	
<b>VANILLIN</b> (UNII: CHI530446X)	
<b>2-METHYLBUTYRIC ACID</b> (UNII: PX7ZNN5GXX)	
<b>SESAME OIL</b> (UNII: QX10HYY4QV)	
<b>.GAMMA.-UNDECALACTONE</b> (UNII: QB1T0AG2YL)	
<b>DIISOSTEARYL MALATE</b> (UNII: QBS8A3XZGQ)	
<b>ALKYL (C12-15) BENZOATE</b> (UNII: A9EJ3J61HQ)	
<b>MEDIUM-CHAIN TRIGLYCERIDES</b> (UNII: C9H2L21V7U)	
<b>BUTYLOCTYL SALICYLATE</b> (UNII: 2EH13UN8D3)	
<b>HYDROGENATED POLYBUTENE (1300 MW)</b> (UNII: 7D1YQ9Y5EZ)	
<b>STEVIOSIDE</b> (UNII: 0YON5MXJ9P)	
<b>VITIS VINIFERA SEED</b> (UNII: C34U15ICXA)	
<b>SHEA BUTTER</b> (UNII: K49155WL9Y)	
<b>COCONUT OIL</b> (UNII: Q9L0O73W7L)	
<b>ISOPROPYL PALMITATE</b> (UNII: 8CRQ2TH63M)	
<b>NEOPENTYL GLYCOL DIHEPTANOATE</b> (UNII: 5LKW3C543X)	
<b>SOYBEAN OIL</b> (UNII: 241ATL177A)	
<b>BENZALDEHYDE</b> (UNII: TA269SD04T)	

**Packaging**

<b>#</b>	<b>Item Code</b>	<b>Package Description</b>	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
1	NDC:75936-257-01	10 mL in 1 TUBE; Type 0: Not a Combination Product	04/22/2020	

## Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part352	04/22/2020	

**Labeler** - Supergoop, LLC (117061743)

Revised: 7/2023

Supergoop, LLC