## CITY SUNSCREEN SERUM BROAD SPECTRUM SPF 30 SUPERGOOPhomosalate, octisalate, avobenzene, octocrylene cream Baxter Laboratories Pty. Ltd.

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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## City Sunscreen Serum Broad Spectrum SPF 30

Active ingredients: Purpose

Homosalate 8%, Octisalate 4%, Avobenzene 3%, Octocrylene 2% Sunscreen

Purpose

Uses Simultaneously protects skin, combats premature aging and moisturizes on contact

helps prevent sunburn

If used as directed with other sun protection measures (see directions), decreases the risk of skin cancer and early skin aging skin aging caused by the sun

Keep out of reach of children

Warnings

For external use only.

Do not use on damaged or broken skin

When using this product, keep out of of eyes.

Rinse with water to remove.

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

Directions: Apply liberally 15 minutes before sun exposure

Use a water resistant sunscreen if swimming or sweating

Reapply at least every 2 hours

children under 6 months: ask a doctor

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 sunscreenwith 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

Inactive Ingredients Purified water (Aqua), Cyclometicone, Isostearyl Neopentanoate, Glycerin, Ceteareth-20, Polypropylene, Cetearyl Alcohol, Xanthan Gum, d-Panthenol, Octanohydroxamic acid, Caprylyl Glycol, Silica, Triacontanyl PVP, Cetyl Dimethicone, Ammonium Acryloyldimethyltaurate/VP Copolymer, PEG-40 Stearate, Tocopheryl, Disodium EDTA, Pentylene Glycol, Pantheyl Triacetate, Sodium Lactate, Lactic Acid,

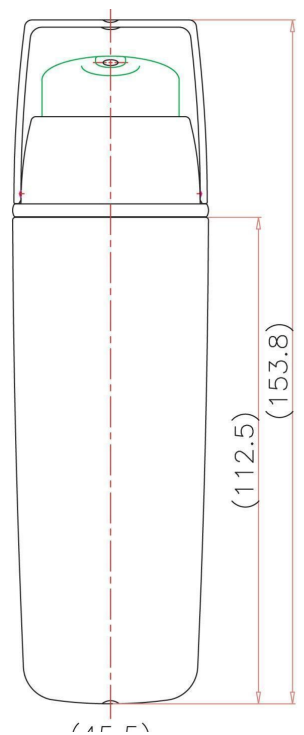
Serine, Urea, Sorbitol, Sodium Chloride, Allantoin, Oleyl Alcohol, Ethyl Linoleate.

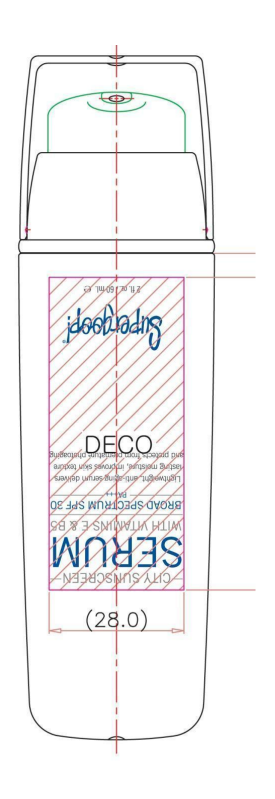
City Sunscreen

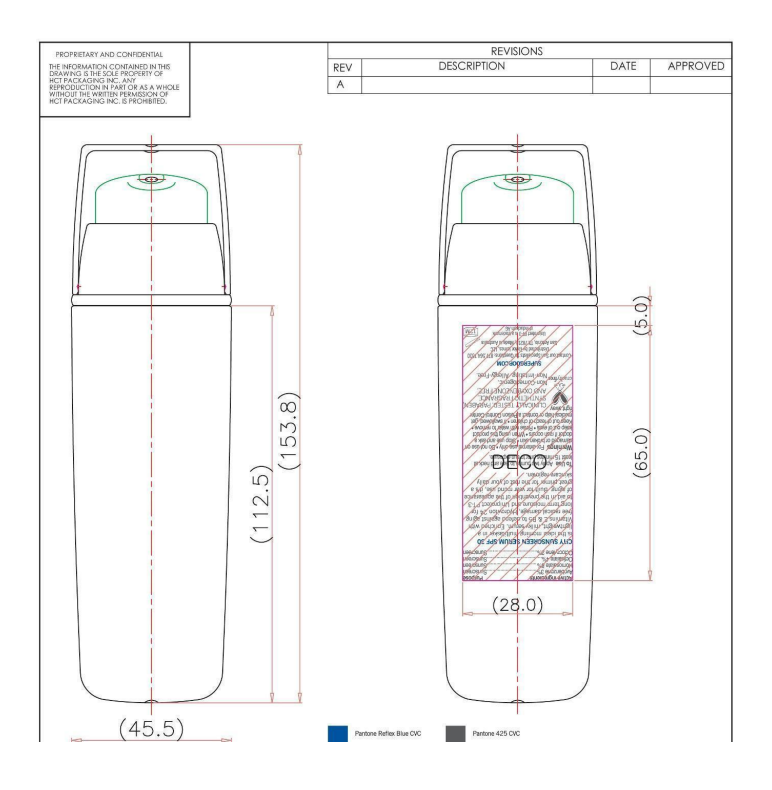
SERUM

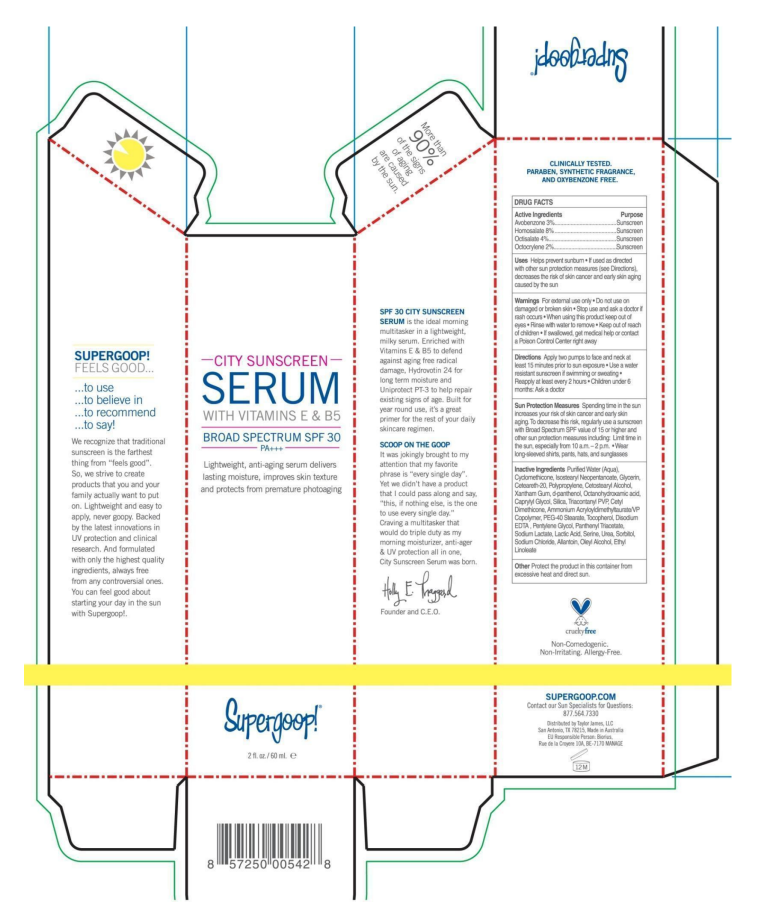
With Vitamins E & B5

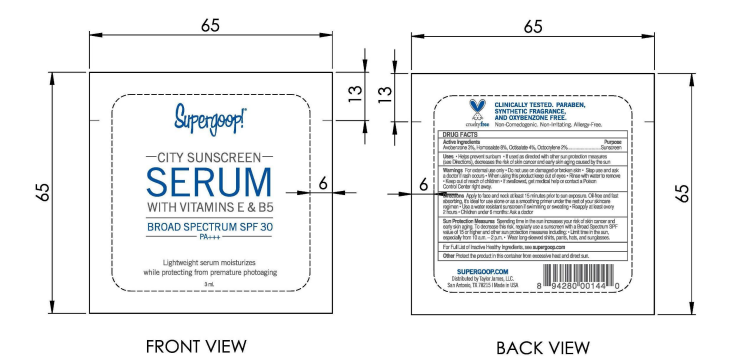
Supergoop!











## CITY SUNSCREEN SERUM BROAD SPECTRUM SPF 30 SUPERGOOP

homosalate, octisalate, avobenzene, octocrylene cream

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70157-004	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety			
Ingredient Name	<b>Basis of Strength</b>	Strength	
HOMOSALATE (UNII: V06SV4M95S) (HOMOSALATE - UNII:V06SV4M95S)	HOMOSALATE	8 g in 100 kg	
OCTISALATE (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	4 g in 100 kg	
AVOBENZONE (UNII: G63QQF2NOX) (AVOBENZONE - UNII:G63QQF2NOX)	AVOBENZONE	3 g in 100 kg	
OCTOCRYLENE (UNII: 5A68WGF6WM) (OCTOCRYLENE - UNII:5A68WGF6WM)	OCTOCRYLENE	2 g in 100 kg	

Inactive Ingredients			
Ingredient Name	Strength		
CYCLOMETHICONE (UNII: NMQ347994Z)			
GLYCERIN (UNII: PDC6A3C0OX)			
XANTHAN GUM (UNII: TTV12P4NEE)			
PANTHENOL (UNII: W/9CM0067Z)			
CAPRYLYL GLYCOL (UNII: 00YIU5438U)			
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)			
POLYOXYL 40 STEARATE (UNII: 13A4J4NH9I)			
EDETATE DISODIUM (UNII: 7FLD91C86K)			
PENTYLENE GLYCOL (UNII: 50C1307PZG)			
SODIUM LACTATE (UNII: TU7HW0W0QT)			
LACTIC ACID (UNII: 33X04XA5AT)			
SERINE (UNII: 452VLY9402)			
UREA (UNII: 8W8T17847W)			
SORBITOL (UNII: 506T60A25R)			
SODIUM CHLORIDE (UNII: 451W47IQ8X)			
ALLANTOIN (UNII: 344S277G0Z)			

OLEYL ALCOHOL (UNII: 172F2WN8DV)	
ETHYL LINOLEATE (UNII: MJ2YTT4 8M)	

l	P	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
	1	NDC:70157-004- 11	480 kg in 1 DRUM; Type 0: Not a Combination Product	12/07/2010		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph final	M020	12/07/2010		

**Labeler -** Baxter Laboratories Pty. Ltd. (740537709)

**Registrant -** Baxter Laboratories Pty. Ltd. (740537709)

Revised: 11/2022 Baxter Laboratories Pty. Ltd.