SUNSCREEN SETTING SPRITZ, SPF 45- sunscreen setting spray liquid Cardinal Health, inc.

AVOBENZONE G63QQF2NOX 2.9% OCTISALATE 4X49Y0596W 4.9% OCTOCRYLENE 5A68WGF6WM 9.5%

WATER
BHT
ALCOHOL
PEG-400
PROPYLENE GLYCOL LAURATE

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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.lf swallowed, get medical help or contacta Poison Control Center right away

Flammable. Do not spray near heat sparks, sources of ignition, or flames

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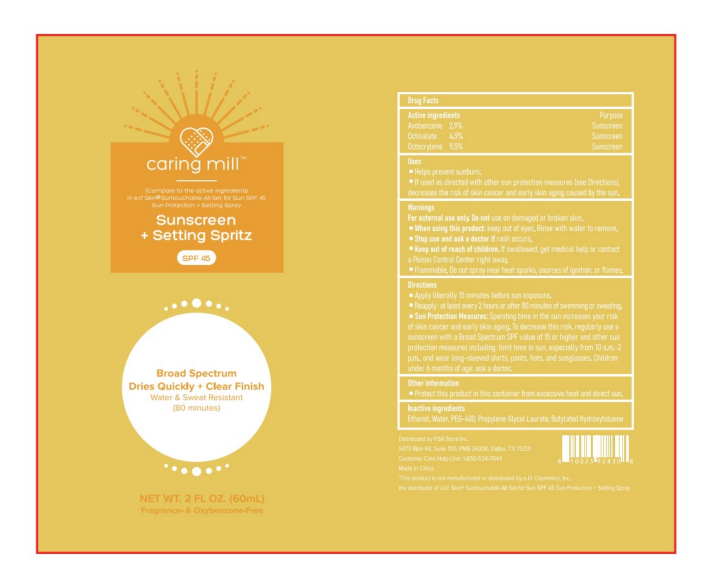
Directions

Apply liberally 15 minutes before sun exposure. Reaooly. at least every 2 hours or after 80 minutes ofswimming or sweating. Sun Protection Measures, Spending time in the sun increases your riskof skan cancer and early skin aging. lo decrease this risk, regularly use asunscreen with a Broad Spectrum SPF value of 15 or higher and other sunprotection measures including, limit time in sun, especially from 10 a.m.-2p.m, and wear long-sleeved shirts, pants, hats, and sunalasses. Chitdrenunder 6 months of age:

ask a doctor.

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SUNSCREEN SETTING SPRITZ, SPF 45 sunscreen setting spray liquid Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:70000-1543 Route of Administration TOPICAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
OCTISALATE (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	2.94 g in 60 mL		
OCTOCRYLENE (UNII: 5A68WGF6WM) (OCTOCRYLENE - UNII:5A68WGF6WM)	OCTOCRYLENE	5.7 g in 60 mL		
AVOBENZONE (UNII: G63QQF2NOX) (AVOBENZONE - UNII:G63QQF2NOX)	AVOBENZONE	1.74 g in 60 mL		

Inactive Ingredients			
Ingredient Name	Strength		
PROPYLENE GLYCOL LAURATE (UNII: 668Z5835Z3)			
ALCOHOL (UNII: 3K9958V90M)			
BHT (UNII: 1P9D0Z171K)			
PEG-400 (UNII: B697894SGQ)			
AQUA (UNII: 059QF0KO0R)			

Packaging				
# Item Code	Package Description	Marketing Start Date	Marketing End Date	
1 NDC:70000- 1543-1	60 mL in 1 BOTTLE; Type 0: Not a Combination Product	12/15/2024		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M020	12/15/2024		

Labeler - Cardinal Health, inc. (063997360)

Revised: 12/2024 Cardinal Health, inc.