

**NEUTROGENA SUN RESCUE AFTER SUN MEDICATED RELIEF- camphor
(synthetic) gel
Kenvue Brands LLC**

Neutrogena Sun Rescue After Sun Medicated Relief Gel

Drug Facts

Active ingredient

Camphor 0.45%

Purpose

External analgesic

Use

temporarily relieves pain and itching associated with sunburn

Warnings

For external use only.

Flammable: Keep away from fire or flame

When using this product avoid contact with eyes

Stop use and ask a doctor if

- condition worsens
- symptoms persist for more than 7 days or clear up and occur again within a few days

Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- adults and children 2 years of age and older: apply to affected area not more than 3 to 4 times daily
- children under 2 years of age: ask a doctor

Other information

- Store at 20°C to 25°C (68°F to 77°F)

Inactive ingredients

Water, Alcohol Denat., Triethanolamine, Carbomer, EDTA

Questions?

Call toll-free 800-299-4786 or 215-273-8755 (collect) or visit www.neutrogena.com

Distributed by:

Kenvue Brands LLC
Summit, NJ 07901

PRINCIPAL DISPLAY PANEL - 85 g Tube Label

NEW

Neutrogena®

DERMATOLOGIST RECOMMENDED BRAND

NEUTROGENA®

SUN

RESCUE

AFTER SUN

MEDICATED RELIEF GEL

CAMPBOR EXTERNAL ANALGESIC

pain relief for sunburn

cools, calms and soothes

skin on contact

NET WT 3.0 OZ (85 g)

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Kenvue Brands LLC
Summit, NJ 07901
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Active Made in India

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NEUTROGENA SUN RESCUE AFTER SUN MEDICATED RELIEF

camphor (synthetic) gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69968-0721
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET) (CAMPHOR (SYNTHETIC) - UNII:5TJD82A1ET)	CAMPHOR (SYNTHETIC)	4.5 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
CARBOXYPOLYMETHYLENE (UNII: 0A5MM307FC)	

EDETIC ACID (UNII: 9G34HU7RV0)	
TROLAMINE (UNII: 9O3K93S3TK)	
WATER (UNII: 059QF0KO0R)	
ALCOHOL (UNII: 3K9958V90M)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69968-0721-3	85 g in 1 TUBE; Type 0: Not a Combination Product	10/01/2021	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	10/01/2021	

Labeler - Kenvue Brands LLC (118772437)

Revised: 7/2025

Kenvue Brands LLC