HEB BROAD SPECTRUM SPF 30 POMEGRANATE SCENTED SUNSCREEN LIP BALM- avobenzone, homosalate, octisalate, octocrylene stick H.E.B

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.

HEB BROAD SPECTRUM SPF 30 POMEGRANATE SCENTED SUNSCREEN LIP BALM

Active ingredients

Avobenzone 3.0%, Homosalate 10.0%, Octisalate 5.0%, Octocrylene 5.0%

Purpose

Sunscreen

□ Uses

helps prevent sunburn

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 I

rash occurs.

□Keep out of reach of children□.

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

□ Directions

Apply liberally 15 minutes before sun exposure

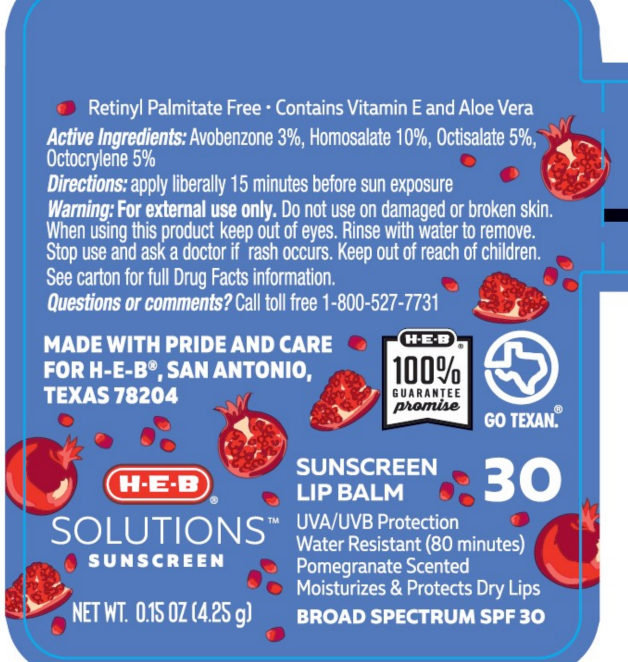
- Reapply:
- after 80 minutes of swimming or sweating
- immediately after towel drying
- at least every 2 hours
- Children under 6 months of age: Ask a doctor
- **Sun Protection Measures**. Spending time in the sun increases your risk of skin cancer and early skin aging. To decrease this risk, regularly use a sunscreen with a Broad Spectrum SPF value of 15 or higher and other sun protection measures including:
- limit time in the sun, especially from 10 a.m.-2 p.m.
- wear long-sleeved shirts, pants, hats, and sunglasses

Other information

- protect the product in this container from excessive heat and direct sun
- may stain or damage some fabrics, materials or surfaces

Inactive ingredients

C12-15 alkyl benzoate, paraffin, butyloctyl salicylate, synthetic wax, ozokerite, euphorbia cerifera (candelilla) wax, dimethicone, aloe barbadensis leaf extract, mineral oil, ascorbyl palmitate, tocopheryl acetate, dimethicone crosspolymer-3, cetyl dimethicone, cetyl alcohol, caprylyl glycol, flavor.







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Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:37808-337
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
OCTISALATE (UNII: 4X49Y0596W) (OCTISALATE - UNII:4X49Y0596W)	OCTISALATE	50 mg in 1 g	
OCTOCRYLENE (UNII: 5A68WGF6WM) (OCTOCRYLENE - UNII:5A68WGF6WM)	OCTOCRYLENE	50 mg in 1 g	
AVOBENZONE (UNII: G63QQF2NOX) (AVOBENZONE - UNII:G63QQF2NOX)	AVOBENZ ONE	30 mg in 1 g	
HOMOSALATE (UNII: V06SV4M95S) (HOMOSALATE - UNII:V06SV4M95S)	HOMOSALATE	100 mg in 1 g	

Inactive Ingredients	
Ingredient Name	Strength
CANDELILLA WAX (UNII: WL0328HX19)	
ALKYL (C12-15) BENZOATE (UNII: A9EJ3J61HQ)	
PARAFFIN (UNII: 1900E3H2ZE)	
BUTYLOCTYL SALICYLATE (UNII: 2EH13UN8D3)	
CERESIN (UNII: Q1LS2UJO3A)	
CETYL DIMETHICONE 25 (UNII: U4AS1BW4ZB)	

CAPRYLYL GLYCOL (UNII: 00YIU5438U)	
CETYL ALCOHOL (UNII: 936JST6JCN)	
MINERAL OIL (UNII: T5L8T28FGP)	
ASCORBYL PALMITATE (UNII: QN83US2B0N)	
.ALPHATOCOPHEROL ACETATE (UNII: 9E8X80D2L0)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
DIMETHICONE (UNII: 92RU3N3Y10)	

P	Packaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:37808- 337-89	2 in 1 CARTON	01/10/2020	
1	NDC:37808- 337-58	0.15~g in 1 CONTAINER; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part352	01/10/2020	

Labeler - H.E.B (007924756)

Revised: 8/2021 H.E.B