ALOE VERA GEL- lidocaine hcl, menthol gel Retail Business Services, 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.

After Sun Gel with Aloe 747.001/747AB

Active ingredient

Lidocaine HCI 0.7%

Menthol 0.2%

Purpose

Topical analgesic

Uses

for the temporary relief of pain and itching associated with

- minor burns
- sunburn
- minor cuts
- scrapes
- insect bites
- minor skin irritations

Warnings

For external use only

Do not use

in large quantities, particularly over raw surfaces or blistered areas

When using this product

avoid contact with the eyes

Stop use and ask a doctor if

condition worsens, or if 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

Inactive ingredients

water, alcohol denat., polysorbate 20, glycerin, Aloe barbadensis leaf juice, carbomer, benzophenone-4, triethanolamine, benzyl alcohol, phenoxyethanol, blue 1

DISTRIBUTED BY:

AUDSA DISTRIBUTION, LLC

SALISBURY, NC 28147

For product questions or concerns, contact us at 1-833-992-3872

Quality guaranteed or your money back.

DSP-TN-21091

DSP-MO-20087

principal display panel

CAREONE

AFTER SUN

Gel with Aloe

Pain Relieving Gel

- with Lidocaine and Menthol
- Soothing

NET 16 FL OZ (473 mL)



idocaine hcl, menthol gel					
Product Information					
Product Type	HUMAN OTC DRUG	Item C	Code (Source)	NDC:72	2476-747
Route of Administration	TOPICAL				
Active Ingredient/Activ	e Moiety				
Ingredient Name			Basis of Stren	ngth	Strength
LIDOCAINE HYDROCHLORIDE (UNII: V13007Z41A) (LIDOCAINE - UNII:98PI200987)			LIDOCAINE HYDROCHLORIDE ANHYDROUS		7 mg in 1 mL
0111.301 (200307)			ANHYDROUS		
MENTHOL (UNII: L7T10EIP3A) (M	ENTHOL - UNII:L7T10EIP3A)		MENTHOL		2 mg in 1 mL
MENTHOL (UNII: L7T10EIP3A) (M	ENTHOL - UNII:L7T10EIP3A)				
	ENTHOL - UNII:L7T10EIP3A)				
MENTHOL (UNII: L7T10EIP3A) (M	ENTHOL - UNII:L7T10EIP3A) Ingredient Name			S	
MENTHOL (UNII: L7T10EIP3A) (M				S	in 1 mL
MENTHOL (UNII: L7T10EIP3A) (M				S	in 1 mL
MENTHOL (UNII: L7T10EIP3A) (M Inactive Ingredients water (UNII: 059QF0K00R)	Ingredient Name			S	in 1 mL
MENTHOL (UNII: L7T10EIP3A) (M Inactive Ingredients water (UNII: 059QF0K00R) ALCOHOL (UNII: 3K9958V90M) POLYSORBATE 20 (UNII: 7T1F3	Ingredient Name			S	in 1 mL
MENTHOL (UNII: L7T10EIP3A) (M Inactive Ingredients water (UNII: 059QF0K00R) ALCOHOL (UNII: 3K9958V90M) POLYSORBATE 20 (UNII: 7T1F3 glycerin (UNII: PDC6A3C00X)	Ingredient Name 0V5YH)			S	in 1 mL
MENTHOL (UNII: L7T10EIP3A) (M Inactive Ingredients water (UNII: 059QF0K00R) ALCOHOL (UNII: 3K9958V90M)	Ingredient Name OV5YH) 33H0X)			S	in 1 mL

BE	NZYL ALCOHO	L (UNII: LKG8494WBH)		
	enoxyethanol			
•		L (UNII: H3R47K3TBD)		
Pa	ackaging			
#	ltem Code	Package Description	Marketing Start Date	Marketing End Date
1		473 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	11/24/2021	
Marketing Information				
	Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OT fin	C monograph no al	part348	11/24/2021	

Labeler - Retail Business Services, LLC (967989935)

Registrant - Vi-Jon, LLC (790752542)

Establishment				
Name	Address	ID/FEI	Business Operations	
Vi-Jon, LLC		790752542	manufacture(72476-747)	

Establishment

Name	Address	ID/FEI	Business Operations
Vi-Jon, LLC		088520668	manufacture(72476-747)

Revised: 1/2023

Retail Business Services, LLC