

ISLEAF TATTOO EYEBROW BROWN- methylparaben gel C3 Co., Ltd.

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Drug Facts

methylparaben

water, butylene glycol, glycerin, alcohol, xanthan gum, etc

for a tattoo effect

keep out of reach of the children

Shake the ball inside of the product before use, then press the tip of container and draw eyebrows. In about 5 hours, the eyebrows will look natural.

1. If the following symptoms occur after product use, stop using the product immediately and consult a dermatologist (continuous use can exacerbate the symptoms).

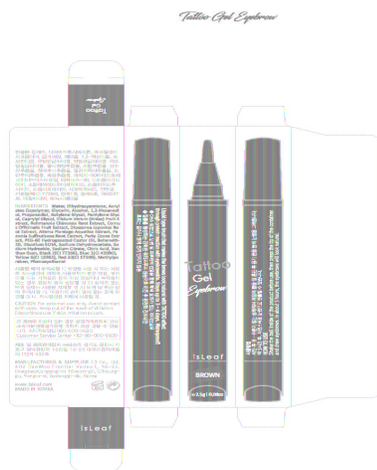
- 1) Occurrence of red spots, swelling, itchiness, and other skin irritation
 - 2) If the symptoms above occur after the application area is exposed to direct sunlight
2. Do not use on open wounds, eczema, and other skin irritations

3. Precaution for Storage and Handling

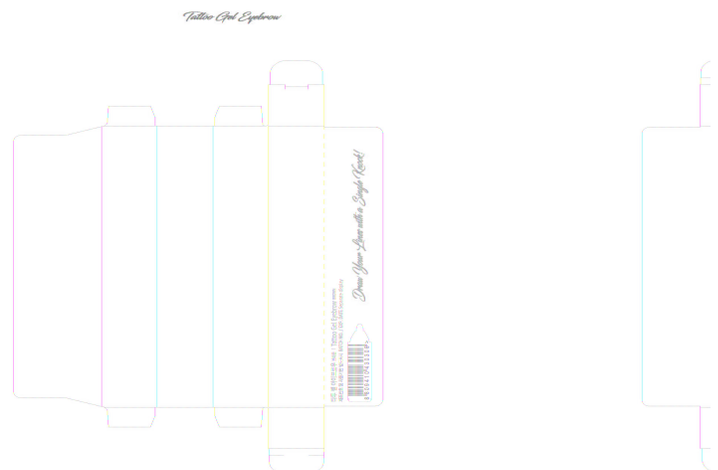
- 1) Close the lid after use
- 2) Keep out of reach of infants and children
- 3) Do not to store in a place with high/low temperature and exposed to direct sunlight

4. Use as avoiding eye areas.

for external use only



수량: 3,000개



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ISLEAF TATTOO EYEBROW BROWN

methylparaben gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70818-007	
Route of Administration	TOPICAL			
Active Ingredient/Active Moiety				
Ingredient Name		Basis of Strength	Strength	
METHYLPARABEN (UNII: A2I8C7HI9T) (METHYLPARABEN - UNII:A2I8C7HI9T)		METHYLPARABEN	0.3 g in 100 g	
Inactive Ingredients				
Ingredient Name			Strength	
WATER (UNII: 059QF0K00R)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:70818-007-01	2.5 g in 1 APPLICATOR; Type 0: Not a Combination Product	08/08/2017	
Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		08/08/2017		

Labeler - C3 Co., Ltd. (689846633)

Registrant - C3 Co., Ltd. (689846633)

Establishment

Name	Address	ID/FEI	Business Operations
C3 Co., Ltd.		689846633	label(70818-007) , pack(70818-007) , manufacture(70818-007)

Revised: 8/2017

C3 Co., Ltd.