DR. B ANTI POLLUTION BUBBLE MASK- glycerin liquid Ddoruroo Co., Ltd.

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.

Drug Facts

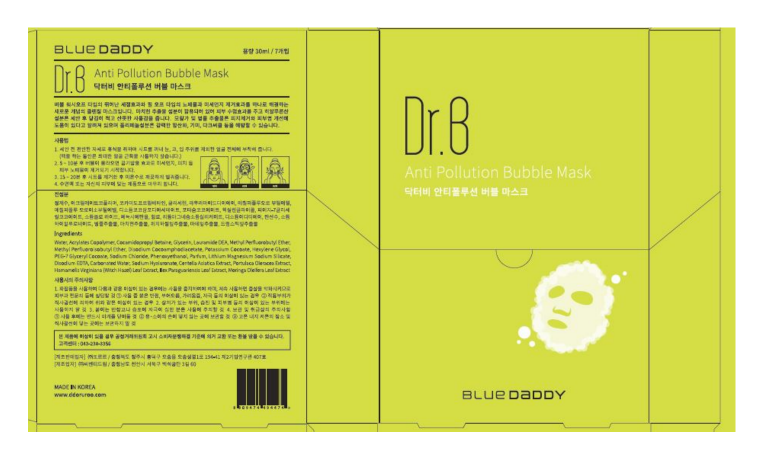
glycerin

water, sodium chloride, carbonated water, etc

SKIN PROTECTANT

KEEP OUT OF REACH OF THE CHILDREN

- ☐ Put the sheet on the face in a comfortable posture without cleansing
- $\ \square$ After 5 to 10 minutes, bubbles begin to remove skin wastes, sebum and fine dust
- Unit 5 to 10 more minutes, remove the sheet then wash it with lukewarm water or take a shower
- ☐ Finish with appropriate toner
- 1. Do not use in the following cases(Eczema and scalp wounds)
- 2.Side Effects
- 1)Due to the use of this druf if rash, irritation, itching and symptopms of hypersnesitivity occur dicontinue use and consult your phamacisr or doctor
- 3.General Precautions
- 1)If in contact with the eyes, wash out thoroughty with water If the symptoms are servere, seek medical advice immediately
- 2)This product is for exeternal use only. Do not use for internal use
- 4. Storage and handling precautions
- 1)If possible, avoid direct sunlight and store in cool and area of low humidity
- 2)In order to maintain the quality of the product and avoid misuse
- 3)Avoid placing the product near fire and store out in reach of children for external use only



DR. B ANTI POLLUTION BUBBLE MASK

glycerin liquid

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:71484-0006

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name

Basis of Strength

GLYCERIN (UNII: PDC6A3C0OX) (GLYCERIN - UNII:PDC6A3C0OX)

GLYCERIN (4 g in 100 mL

Inactive Ingredients

Ingredient Name Strength

WATER (UNII: 059QF0KO0R)

Packaging

	l	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
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1 NDC:71484-0006-1 210 mL in 1 PACKAGE; Type 0: Not a Combination Product 12/19/2017

Marketing Information

Marketing Category Application Number or Monograph Citation Marketing Start Date Marketing End Date

OTC monograph not final	part347	12/19/2017	

Labeler - Ddoruroo Co., Ltd. (694209410)

Registrant - Ddoruroo Co., Ltd. (694209410)

Establishment						
Name	Address	ID/FEI	Business Operations			
Ddoruroo Co., Ltd.		694209410	manufacture(71484-0006), pack(71484-0006), label(71484-0006)			

Revised: 12/2017 Ddoruroo Co., Ltd.