## ANY MI NATURAL MAGIC BB NO.5 - allantoin cream K.N.Life 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.

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#### **Drug Facts**

allantoin

water, cyclomethicone, titanium dioxide, zinc oxide, ethylhexyl methoxycinnamate, panax ginseng root ext, peg-7 dimethicone, propylene glycol, arbutin, bentonite, cetyl peg/ppg-10/1 dimethicone, hexyl laurate, betaine, polymethyl methacrylate, sodium chloride, etc

whitening anti-wrinkle sun block

keep out of reach of the children

after foundation, apply small amounts to whole face by tapping until it is absorbed to skin completely

- 1. if you have any abnormal symptoms as followings, you should discontinue to use this cream. In case of using continuously, you have to consult to dermatologist because it may make symptoms worse.
- a. red macule, swelling, urtication, and irritation during use
- b. symptoms like above by direct ray
- 2. do not use if you have a wound, eczema, and dermatitis to the area where you apply this cream
- 3. caution during storing and handling
- a. close the cap after use
- b. keep it to the area where babies and infants cannot reachto
- c. keep it to the area where has not high or low temperature and has no direct ray

for external use only

#### BB 5호 케이스



#### **ANY MI NATURAL MAGIC BB NO.5**

allantoin cream

#### **Product Information**

Product Type HUMAN OTC DRUG Item Code (Source) NDC:62695-1001

Route of Administration TOPICAL

#### **Active Ingredient/Active Moiety**

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Ingredient Name	Basis of Strength	Strength
<b>ALLANTO IN</b> (UNII: 344S277G0Z) (ALLANTO IN - UNII:344S277G0Z)	ALLANTOIN	0.0005 g in 1 mL

# Inactive Ingredients Ingredient Name Strength WATER (UNII: 059QF0KO0R) GLYCERIN (UNII: PDC6A3C0OX) CYCLOMETHICONE (UNII: NMQ347994Z) PROPYLENE GLYCOL (UNII: 6DC9Q167V3) ARBUTIN (UNII: C5INA23HXF)

P	Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:62695-1001-1	40 mL in 1 TUBE					

Marketing Information							
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date				
OTC monograph final	part347	03/15/2014					

### **Labeler -** K.N.Life Co., Ltd. (688270562)

#### **Registrant -** K.N.Life Co., Ltd. (688270562)

Establishment								
Name	Address	ID/FEI	Business Operations					
K.N.Life Co., Ltd.		688270562	manufacture(62695-1001)					

Revised: 3/2014 K.N.Life Co., Ltd.