

**GD11 RX SCM C5- mannitol liquid**  
**Coson 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](#).*

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**ACTIVE INGREDIENT**

Active ingredients: MANNITOL 8.0%

**INACTIVE INGREDIENT**

Inactive ingredients: Water, Human Cord Blood Cell Conditioned Media, Sodium Hyaluronate, Betaine, Trehalose

**PURPOSE**

Purpose: Skin Moisturizing

**WARNINGS**

Warnings: For external use only

Caution in usage 1. In case of having problems such as red rash, swollenness, itching, you need to consult a dermatologist. 2. You are banned to use it on the part where you have a scar, eczema, or dermatitis 3. Caution for treatment and keeping. - Keep it out of infants or children's reach. - Don't keep it exposed to the direct sunlight

**KEEP OUT OF REACH OF CHILDREN**

Keep it out of infants or children's reach.

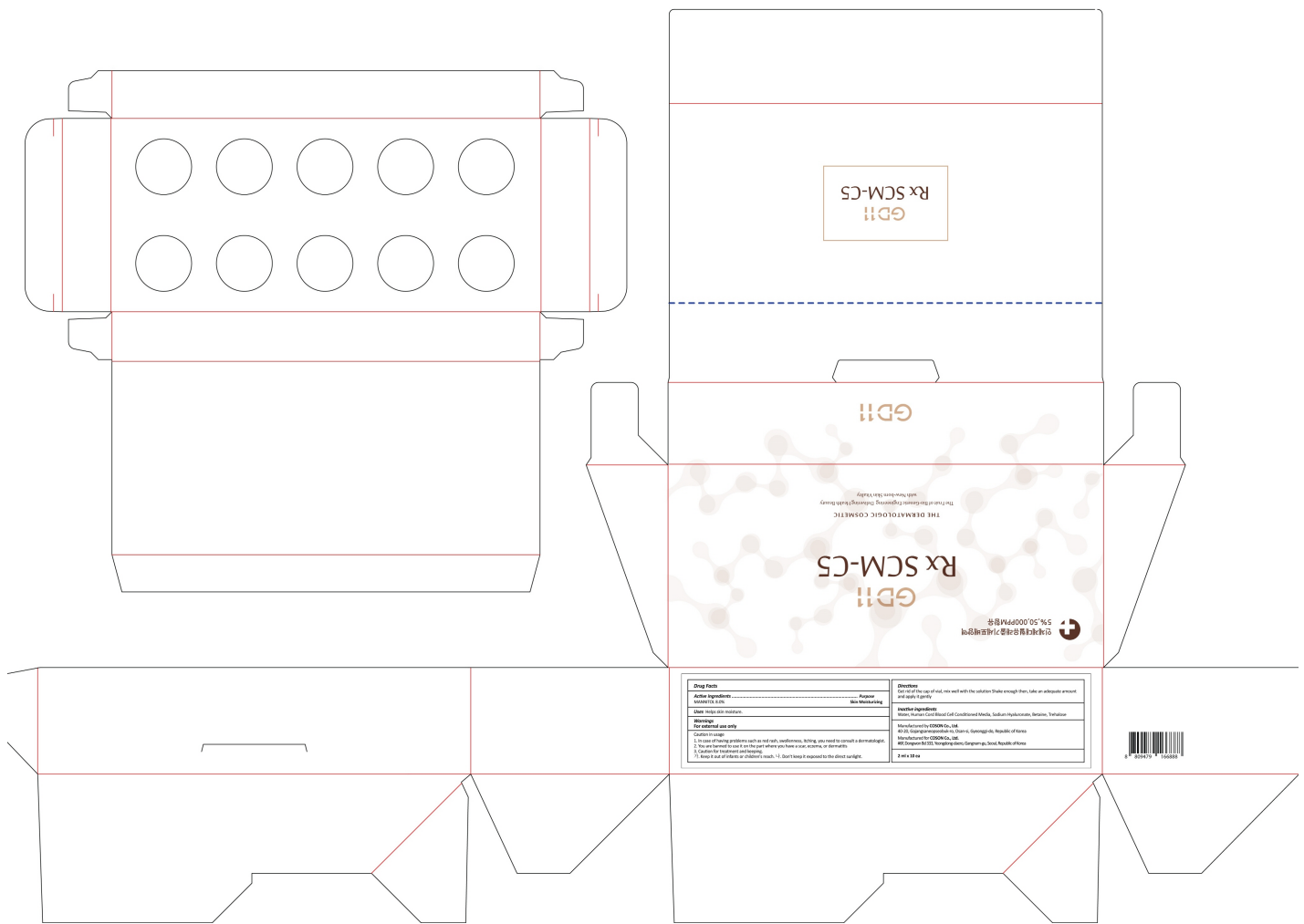
**Uses**

Uses: Helps skin moisture.

**Directions**

Directions: Get rid of the cap, replace with the enclosed white plastic cap, take adequate amount apply it gently

**PACKAGE LABEL.PRINCIPAL DISPLAY PANEL**



## GD11 RX SCM C5

mannitol liquid

### Product Information

<b>Product Type</b>	HUMAN OTC DRUG	<b>Item Code (Source)</b>	NDC:62171-100
<b>Route of Administration</b>	TOPICAL		

### Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
MANNITOL (UNII: 3OWL53L36A) (MANNITOL - UNII:3OWL53L36A)	MANNITOL	0.10 g in 2 mL

### Inactive Ingredients

Ingredient Name	Strength
<b>Water</b> (UNII: 059QF0KO0R)	
<b>Betaine</b> (UNII: 3SCV180C9W)	

### Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:62171-100-02	10 in 1 CARTON	07/01/2017	
1	NDC:62171-100-01	2 mL in 1 CONTAINER; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		07/01/2017	

**Labeler** - Coson Co., Ltd. (689835593)

**Registrant** - Coson Co., Ltd. (689835593)

**Establishment**

Name	Address	ID/FEI	Business Operations
COSON Co., Ltd._Osan Plant		689847210	manufacture(62171-100)

Revised: 8/2017

Coson Co., Ltd.