

HI AND DRI ROLL ON- aluminum chlorohydrate liquid
Revlon Consumer Products Corp

Hi & Dri Antiperspirant Roll-On - Powder Fresh

Aluminum chlorohydrate 18%

Purpose

Antiperspirant

Warnings:

For external use only.

Do not use on broken skin

Ask a doctor before use if you have kidney disease

Stop use if rash or irritation occurs

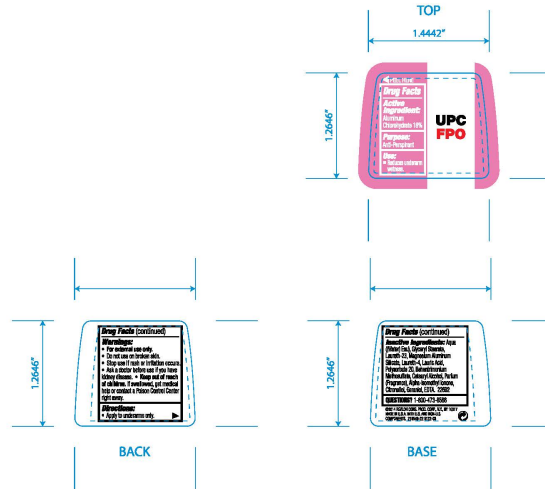
If swallowed, get medical help or contact a Poison Control Center right away.

Inactive Ingredients

aqua((water) eau), glyceryl stearate, laureth-23, magnesium aluminum silicate, laureth-4, lauric acid, cetearyl alcohol, behentrimonium methosulfate, edta

Reduces underarm wetness

**HI & DRI ROLL ON 1.7OZ POWDER FRESH
BACK LABEL
TR05967 / 5122-03
1.4442"W x 1.2646"H**



ENLARGED 200% FOR PROOFREADING

◀ PEEL HERE

Drug Facts

Active Ingredient:
Aluminum Chlorohydrate 18%

Purpose:
Anti-Perspirant

Use:
• Reduces underarm wetness.

Drug Facts (continued)

Warnings:

- For external use only.
- Do not use on broken skin.
- Stop use if rash or irritation occurs.
- Ask a doctor before use if you have kidney disease.
- Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions:

- Apply to underarms only.

Drug Facts (continued)

Inactive Ingredients: Aqua (Water), Glycerol Stearate, Laureth-23, Magnesium Aluminum Silicate, Laureth-4, Lauric Acid, Polysorbate 20, Behentrimonium Methosulfate, Cetearyl Alcohol, Parfum (Fragrance), Alpha-Isomethyl Ionone, Citronellol, Geraniol, EDTA, Z2692.

QUESTIONS? 1-800-473-8566

©2014 REVLOM COSM. PROD. CORP., N.Y., NY 10017
MADE IN U.S.A. WITH U.S. AND FOREIGN COMPONENTS. 216549-22 5122-03

UPC: 309975122039

Mechanical Approval Disk Release: 03-11-2014

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<p>REVISIONS</p> <table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th>Rev</th> <th>Reason</th> <th>Date</th> </tr> </thead> <tbody> <tr> <td> </td> <td> </td> <td> </td> </tr> <tr> <td> </td> <td> </td> <td> </td> </tr> <tr> <td> </td> <td> </td> <td> </td> </tr> <tr> <td> </td> <td> </td> <td> </td> </tr> </tbody> </table>		Rev	Reason	Date													<p>Reasons for change:</p> <p>1. Release: _____ Date: 03-11-14</p> <p>2. Change Description: _____ Date: 03-03-09</p> <p>3. Change Description: _____ Date: 03-03-09</p> <p>4. Change Description: _____ Date: 03-03-09</p>	
Rev	Reason	Date																

SUBSTRATE IS WHITE

BACKGROUND PRINTS:
100% PMS 211

BLACK TEXT, RULES & UPC PRINT:
100% BLACK

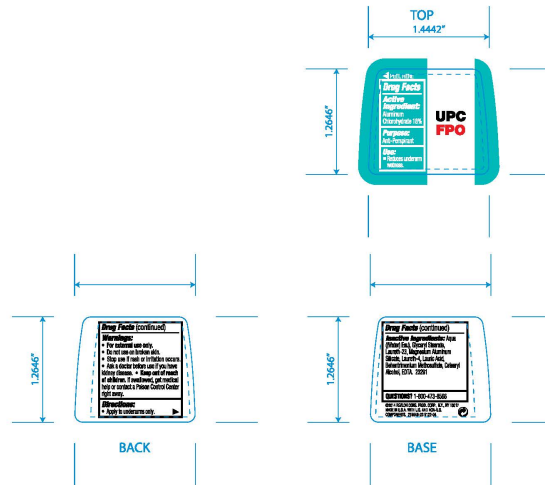
WHITE TEXT & RULES:
K/O WHITE

CYAN RULES REPRESENT DIE AND MEASUREMENTS:
DO NOT PRINT

CYAN DASHED RULES REPRESENT LIVE AREA:
DO NOT PRINT

Hi & Dri Roll-On Antiperspirant

**HI & DRI ROLL ON 1.7OZ UNSCENTED
BACK LABEL
TR05968 / 5122-04
1.4442" W x 1.2646" H**



ENLARGED 200% FOR PROOFREADING

◀ PEEL HERE

Drug Facts

Active Ingredient:
Aluminum Chlorohydrate 18%

Purpose:
Anti-Perspirant

Use:
• Reduces underarm wetness.

Drug Facts (continued)

Warnings:

- For external use only.
- Do not use on broken skin.
- Stop use if rash or irritation occurs.
- Ask a doctor before use if you have kidney disease.
- **Keep out of reach of children.** If swallowed, get medical help or contact a Poison Control Center right away.

Directions:

- Apply to underarms only.

Drug Facts (continued)

Inactive Ingredients: Aqua (Water), Ecol, Glycerol Stearate, Laureth-23, Magnesium Aluminum Silicate, Laureth-4, Lauric Acid, Behentrimonium Methosulfate, Cetaryl Alcohol, EDTA, Z3291

QUESTIONS? 1-800-473-8566

©2014 REVLOW CORP. PRSD, CORP., N.Y., NY 10017
MADE IN U.S.A. METALS AND METALS COMPONENTS: 215549-23 5122-04

UPC: 309975122046

Mechanical Approval Disk Release: 03-11-2014

SCHAWKE

1600 East Stearns Avenue
Des Plaines, IL 60015
Phone: 847-296-6000
800-621-1929
Fax: 847-296-6486

For support please contact the following:
Level 1: Customer Service: Susan Gwynn
susan.gwynn@schawke.com - 847-296-7045
Level 2: Customer Service: Tim Dawson
tim.dawson@schawke.com - 847-759-7070
Package Developer: Donna Lisowski
donna.lisowski@schawke.com - 755-292-7644

100%	75%	50%	25%	15%	10%	5%	1%
Color 319							
							Black

Client: Revlon **Brand:** HI & Dri **Job #:** 5253359 **Authem #:** 40261
Description: HI & Dri 1.7oz Roll On Unscented Back Label
Component: Back Label **Program #:** **Project #:** TR05968
Stock #: 216549-23 **MLI #:** 23291 **Product #:** 5122-04
Size: 1.4442" W x 1.2646" H **Die #:**

DC1	DC2	DC3
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File Name: REV_525335902_TR05968.ai **Software:** 15
Op/Date: HY 03-9-2014, ZCL 03-11-2014

DATE	DESIGN	PROOF	PRINT	INSTR	CHK	APP	REV	BY	DATE

Reasons for change:
1. Release Date: 03-11-14
2. Change description Date: 03-09-14
3. Change description Date: 03-09-14
4. Change description Date: 03-09-14

SUBSTRATE IS WHITE

BACKGROUND PRINTS:
100% PMS 319

BLACK TEXT, RULES & UPC PRINT:
100% BLACK

WHITE TEXT & RULES:
K/O WHITE

CYAN RULES REPRESENT DIE AND MEASUREMENTS:
DO NOT PRINT

CYAN DASHED RULES REPRESENT LIVE AREA:
DO NOT PRINT

HI AND DRI ROLL ON

aluminum chlorohydrate liquid

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:10967-786

Route of Administration

TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
ALUMINUM CHLOROHYDRATE (UNII: HPN8MZ W13M) (ALUMINUM CHLOROHYDRATE - UNII:HPN8MZ W13M)	ALUMINUM CHLOROHYDRATE	0.18 g in 1 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
MAGNESIUM ALUMINUM SILICATE (UNII: 6M3P64V0NC)	
LAURETH-4 (UNII: 6HQ855798J)	
LAURIC ACID (UNII: 1160N9NU9U)	
CETEARYL ALCOHOL (UNII: 2DMT128M1S)	
BEHENTRIMONIUM METHOSULFATE (UNII: 5SHP745C61)	
EDTA (UNII: 9G34HU7RV0)	
GLYCERYL MONOSTEARATE (UNII: 230OU9XXE4)	
LAURETH-23 (UNII: N72LMW566G)	
POLYSORBATE 20 (UNII: 7T1F30V5YH)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:10967-786-17	50 mL in 1 BOTTLE, WITH APPLICATOR; Type 0: Not a Combination Product	01/01/2022	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M019	01/01/2022	

Labeler - Revlon Consumer Products Corp (788820165)

Registrant - REVLON, INC. (188442578)

Establishment

Name	Address	ID/FEI	Business Operations
Revlon, Inc		809725570	manufacture(10967-786)