ODYSSEY ROLL-ON ANTI-PERSPIRANT DEODORANT- aluminum chlorohydrate gel New Avon LLC

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.

Odyssey Anti-Perspirant Roll-On Deodorant

Active ingredient

Aluminum Chlorohydrate 14.5%, anhydrous...

Purpose

..Antiperspirant

Uses

•reduces underarm perspiration

□*Warnings*

IFor 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

□ **Keep out of reach of children.** □ If swallowed, get medical help or contact a poison control center right away.

Directions

•apply to underarms only

Inactive ingredients

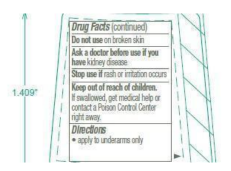
WATER/EAU, STEARETH-2, PPG-15 STEARYL ETHER, DICAPRYL ADIPATE, PARFUM/FRAGRANCE, STEARETH-20, ISOPROPYL PALMITATE

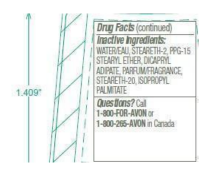
Questions?

Call **1-800-FOR-AVON** or 1 **-800-265-AVON** in Canada









ODYSSEY ROLL-ON ANTI-PERSPIRANT DEODORANT

aluminum chlorohydrate gel

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:10096-9360

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name

ALUMINUM CHLO RO HYDRATE (UNII: HPN8 MZW13M) (ALUMINUM CHLO RO HYDRATE - UNII: HPN8 MZW13M)

Basis of Strength

ALUMINUM
CHLORO HYDRATE - ALUMINUM
CHLORO HYDRATE in 1 mL

Inactive Ingredients

Ingredient Name Strength

WATER (UNII: 059QF0KO0R)

Packaging

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	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:10096- 9360-1	50 mL in 1 BOTTLE, DISPENSING; Type 0: Not a Combination Product	11/12/2012	
	2 NDC:10096- 9360-2	75 mL in 1 BOTTLE, DISPENSING; Type 0: Not a Combination Product	11/12/2012	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part350	11/12/2012	

Labeler - New Avon LLC (080143520)

Establishment						
Name	Address	ID/FEI	Business Operations			
Fareva Morton Grove Inc.		116752326	manufacture(10096-9360)			

Revised: 3/2019 New Avon LLC