

BAN ROLL-ON ANTIPERSPIRANT DEODORANT REGULAR- aluminum chlorohydrate liquid
Kao USA Inc.

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.

Ban Roll-On Antiperspirant Deodorant Regular

Drug Facts

Active ingredient

Aluminum chlorohydrate 17.60%

Purpose

Antiperspirant

Use

reduces underarm perspiration

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

Other information

store at room temperature

Inactive ingredients

water, PPG-11 stearyl ether, steareth-2, steareth-20, fragrance, disodium EDTA, helianthus annuus (sunflower) seed oil, phellodendron amurense bark extract, hordeum distichon (barley) extract, santalum album (sandalwood) extract, amyl cinnamal, benzyl alcohol, benzyl benzoate, cinnamyl alcohol, citronellol, eugenol, geraniol, hexyl cinnamal, linalool

Questions? 1-866-226-3363

www.bandeodorant.com

BAN is a trademark of Kao Corp.

Dist. by Kao USA Inc. Cincinnati, OH 45214 ©2023

Made in Canada

44 mL PACKAGE

ban

regular

INVISIBLE ROLL-ON

CRUELTY FREE • PARABEN FREE • DYE FREE

antiperspirant deodorant

1.5 FL OZ (44 mL)



103 mL PACKAGE

ban

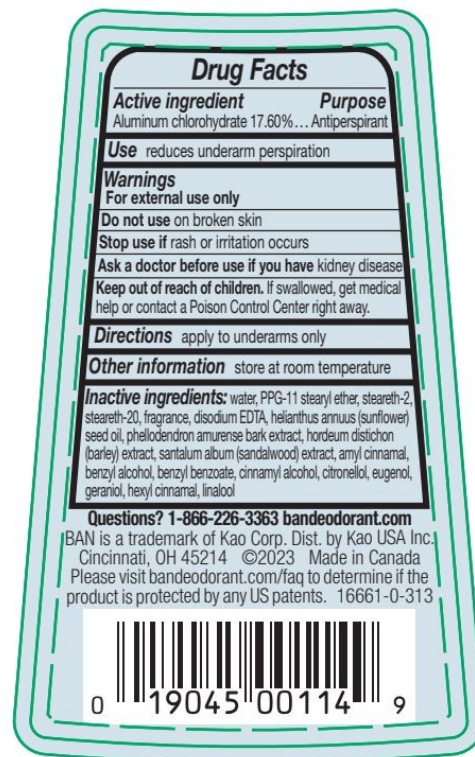
regular

INVISIBLE ROLL-ON

CRUELTY FREE • PARABEN FREE • DYE FREE

antiperspirant deodorant

3.5 FL OZ (103 mL)



VALUE PACK

2X THE BAN PROTECTION

ANTIPERSPIRANT DEODORANT

2- NET WT 3.5 FL OZ (103 mL)



BAN ROLL-ON ANTIPERSPIRANT DEODORANT REGULAR

aluminum chlorohydrate liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:10596-336
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	20 g in 103 mL

Inactive Ingredients

Ingredient Name	Strength
PPG-11 STEARYL ETHER (UNII: S4G2J0Y0LG)	
WATER (UNII: 059QF0K00R)	
STEARETH-2 (UNII: V56DFE46J5)	
STEARETH-20 (UNII: L0Q8IK9E08)	
SANDALWOOD (UNII: 3641YW25N2)	
PHELLODENDRON AMURENSE BARK (UNII: PBG27B754G)	
BARLEY (UNII: 5PWM7YLI7R)	
EDETATE DISODIUM (UNII: 7FLD91C86K)	
BUTYLATED HYDROXYTOLUENE (UNII: 1P9D0Z 171K)	
SUNFLOWER OIL (UNII: 3W1JG795YI)	
.ALPHA.-AMYLCINNAMALDEHYDE (UNII: WC51CA3418)	
BENZYL ALCOHOL (UNII: LKG8494WBH)	
CINNAMYL ALCOHOL (UNII: SS8YOP444F)	
.BETA.-CITRONELLOL, (R)- (UNII: P01OUT964K)	
GERANIOL (UNII: L837108USY)	
.ALPHA.-HEXYLCINNAMALDEHYDE (UNII: 7X6O37OK2I)	
LINALOOL, (+/-)- (UNII: D81QY6I88E)	
BENZYL BENZOATE (UNII: N863NB338G)	
EUGENOL (UNII: 3T8H1794QW)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:10596-336-35	103 mL in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product	01/01/2012	
2	NDC:10596-336-70	2 in 1 PACKAGE	09/12/2016	
2		103 mL in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product		
3	NDC:10596-336-15	44 mL in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product	11/04/2016	
4	NDC:10596-336-14	4 in 1 PACKAGE	12/12/2019	
4		103 mL in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part350	01/01/2012	

Labeler - Kao USA Inc. (004251617)

Revised: 8/2023

Kao USA Inc.