

BAMBOO SALT EUNGANGGO JOOK YEOM TOOTHPASTE - silicon dioxide , sodium fluoride, aminocaproic acid, glycyrrhizinate dipotassium, curcuma xanthorrhiza oil, sea salt paste

LG Household and Healthcare, 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.

Drug Facts

SILICON DIOXIDE 20 %

SODIUM FLUORIDE 0.22 %

AMINOCAPROIC ACID 0.05 %

GLYCYRRHIZINATE DIPOTASSIUM 0.04 %

URSODIOL 0.02 %

SEA SALT 3 %

Direction: Suggested brushing your teeth with proper amount for at least two times per day or following dentist / doctor's instruction.

Children between two to six should use about two to six should use about a size of a pea and should be supervised by adults while using.

This product is for brushing teeth only and uneatable. If a rash occurs, stop using this product and consult a doctor or dentist.

If it comes in contact with eyes, cleanse off immediately and consult a doctor. Store this product away from children and supervise them while using.

BAMBOO SALT

Eunganggo Toothpaste

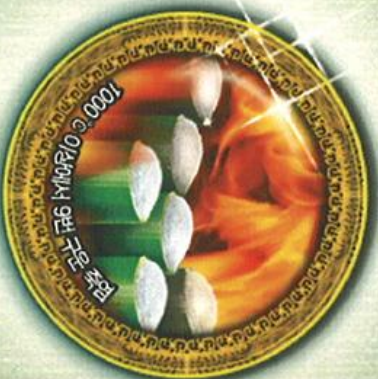


LG Household & Health Care

BAMBOO

JOOK YEOM
TOOTHPASTE

SALT



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Lifelong Gum Care

NET WT 6.0 OZ (170g)



Bamboo Salt Toothpaste

Distributed by **LG Household & Health Care America, Inc.**
17777 CENTER COURT DR. #650 CERRITOS, CA 90703, U.S.A
Manufactured by **LG Household & Health Care Ltd.**
150-32 Songjungdong Heungduk-gu Cheongju, Republic of Korea
Made in Korea
NET WT 6.0 OZ (170g)

Direction & Caution

Direction : Suggest brushing your teeth with proper amount for at least two times a day. Please follow the dentist / doctor's instructions. Children between two to six should use about a small amount and be supervised by adults while using.

Caution : This product is for brushing teeth only and uneatable. If a rash occurs or irritation, discontinue use and consult a doctor or dentist. If it comes in contact with eyes, cleanse off immediately. Store this product away from children and supervise them while using.

Ingredient

Active Ingredient : Dental-Type Silica(20%), Sodium Fluoride(0.22%), ε -Amino Caproic Acid(0.04%), Ursodesoxycholic Acid(0.02%), Curcuma Longa(3%)

Other Ingredient : Sorbitol Solution(70%), Polyethylene Glycol 300, Xanthan Gum, Cellulose, Sodium Saccharin, Methyl Parahydroxy benzoate, Poloxamer 407, Sodium Bisulfite, Titanium Dioxide, Colorant, Flavor, Purified water

nes per day or following
size of a pea and should
stop using this product
ately and consult a doc-

no-caproic Acid(0.05%),
xanthorriza oil(0.025%),
Gum, Carboxymethyl-
Sodium Lauryl Sulfate,



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silicon dioxide , sodium fluoride, aminocaproic acid, glycyrrhizinate dipotassium, curcuma xanthorrhiza oil, sea salt
paste

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:53208-459
Route of Administration	DENTAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SILICON DIOXIDE (UNII: ETJ7Z6XBU4) (SILICON DIOXIDE - UNII:ETJ7Z6XBU4)	SILICON DIOXIDE	20 g in 100 g
SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (Fluoride Ion - UNII:Q80VPU408O)	SODIUM FLUORIDE	0.22 g in 100 g
AMINOCAPROIC ACID (UNII: U6F3787206) (AMINOCAPROIC ACID - UNII:U6F3787206)	AMINOCAPROIC ACID	0.05 g in 100 g
GLYCYRRHIZINATE DIPOTASSIUM (UNII: CA2Y0FE3FX) (GLYCYRRHIZINATE DIPOTASSIUM - UNII:CA2Y0FE3FX)	GLYCYRRHIZINATE DIPOTASSIUM	0.04 g in 100 g
URSODIOL (UNII: 724L30Y2QR) (URSODIOL - UNII:724L30Y2QR)	URSODIOL	0.02 g in 100 g
Curcuma xanthorrhiza oil (UNII: F8VF0V2G7H) (Curcuma xanthorrhiza oil - UNII:F8VF0V2G7H)	Curcuma xanthorrhiza oil	0.025 g in 100 g
SEA SALT (UNII: 87GE52P74G) (SEA SALT - UNII:87GE52P74G)	SEA SALT	3 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
SORBITOL (UNII: 506T60A25R)	
Polyethylene Glycol 300 (UNII: 5655G9Y8AQ)	
XANTHAN GUM (UNII: TTV12P4NEE)	

Poloxamer 407 (UNII: TUF2IVW3M2)	
Sodium Lauryl Sulfate (UNII: 368GB5141J)	
Sodium Bisulfite (UNII: TZX5469Z6I)	
Titanium Dioxide (UNII: 15FIX9V2JP)	
WATER (UNII: 059QF0KO0R)	
Peppermint (UNII: V95R5KMY2B)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
SACCHARIN SODIUM (UNII: SB8ZUX40TY)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:53208-459-02	1 in 1 CARTON		
1	NDC:53208-459-01	170 g in 1 TUBE		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC mono graph not final	part355	05/25/2010	

Labeler - LG Household and Healthcare, Inc. (688276187)

Registrant - LG Household and Healthcare, Inc. (688276187)

Establishment

Name	Address	ID/FEI	Business Operations
LG Household and Healthcare, Inc.		688276187	manufacture

Revised: 6/2010

LG Household and Healthcare, Inc.