

DENTI BUONGIORNO PLAQUE CARE LEMON MINT TOOTH- silica, sodium fluoride, tocopheryl acetate, tetrasodium pyrophosphate paste, dentifrice DONG IL PHARMS 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.

Active ingredient(s)

Silica 17.0%
Sodium Fluoride 0.22%
Tocopheryl Acetate 0.20%
Tetrasodium Pyrophosphate 0.50%

Purpose

Antiplaque, Anticavity, Antigingivitis, Antitartar

Warnings

- Keep out of reach of children under 6 years of age.
- If more than used for brushing is accidentally swallowed, get medical help or contact a Poison Control Center right away.

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Use(s)

- Helps remove plaque and tartar buildup
- Helps prevent cavities
- Helps reduce gum inflammation and promote healthy teeth

Directions

- Adults and children 2 years of age and older : Brush teeth thoroughly, preferably after each meal or at least twice a day, or as directed by a dentist or physician
- Children 2 to 6 years : Use only a pea sized amount and supervise child's brushing and rinsing (to minimize swallowing)
- Children under 2 years : Ask a dentist or physician

Other Information

- Store at temperatures between 1°C and 30°C
- Use within 36 months from the date of manufacture
- Do not store in high/low temperatures or direct sunlight

Questions

+82-1588-9709

Inactive ingredients

D-Sorbitol Solution, Water(Aqua), Silicon Dioxide, Concentrated Glycerin, Sodium Cocoyl Glutamate, Polyethylene Glycol 1500, Carboxymethylcellulose Sodium, Sodium Fluoride, Tocopherol Acetate, Sodium Pyrophosphate, Sodium Bicarbonate, Titanium Oxide,

Xylitol, Enzymatically Modified Stevia, Propolis Extract, Aloe Extract, Matricaria Extract, Eucalyptus Extract, Sage Extract, Green Tea Extract, Rosemary Extract, Calendula Extract, Centella Extract, Lemon Extract, Leontopodium Alpinum Extract, Lactic Acid Bacteria Fermented Solution, Grapefruit Seed Extract, L-Menthol, Lemon Oil, Herb Mint Flavor, Herb



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Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:73242-1126
Route of Administration	ORAL		

Active Ingredient/Active Moiety				
Ingredient Name			Basis of Strength	Strength
.ALPHA.-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0) (.ALPHA.-TOCOPHEROL - UNII:H4N855PNZ1)			.ALPHA.-TOCOPHEROL ACETATE	0.2 g in 100 g
SODIUM PYROPHOSPHATE (UNII: O352864B8Z) (PYROPHOSPHORIC ACID - UNII:4E862E7GRQ)			SODIUM PYROPHOSPHATE	0.5 g in 100 g
SILICON DIOXIDE (UNII: ETJ7Z6XBU4) (SILICON DIOXIDE - UNII:ETJ7Z6XBU4)			SILICON DIOXIDE	17 g in 100 g
SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80VPU408O)			FLUORIDE ION	0.22 g in 100 g

Inactive Ingredients	
Ingredient Name	Strength
SORBITOL (UNII: 506T60A25R)	
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6A3C0OX)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:73242-1126-2	1 in 1 CARTON	01/02/2026	
1	NDC:73242-1126-1	100 g in 1 TUBE; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		01/02/2026	

Labeler - DONG IL PHARMS CO.,LTD (557810721)

Establishment			
Name	Address	ID/FEI	Business Operations
DONG IL PHARMS CO., LTD.		557810721	manufacture(73242-1126)

Revised: 1/2026

DONG IL PHARMS CO.,LTD