

**NOBATON 1.5% FINE SALT AMPOULE- nobaton 1.5% fine salt ampoule
toothpaste ointment
BD TECHNOLOGY 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.

**73927-0009
Nobaton 1.5% Fine Salt ampoule toothpaste**

Active Ingredient(s)

OLAFLUR 0.0975%

SODIUM MONOFLUOROPHOSPHATE 1%

Purpose

Anticaries

Use

15% Fine Salt Ampoule Toothpaste can effectively and gently cleans teeth and mouth, and resist dental caries.

Warnings

Do not swallow; Suitable for adults: Store in a cool and dry place.

Do not use

in children less than 12 years old; People with dental fluorosis

When using this product keep out of eyes. In case of contact with eyes, rinse eyes thoroughly with water. If there is any discomfort, please stop using and consult a doctor. Keep out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

In case of contact with eyes, rinse eyes thoroughly with water. If there is any discomfort, please stop using and consult a doctor. Keep out of reach of children.

Directions

Squeeze an appropriate amount of toothpaste onto the toothbrush, and brush teeth from the inside to the outside using an up and down rubbing motion. Rinse mouth after 3 minutes. (It is recommended to brush without water.)

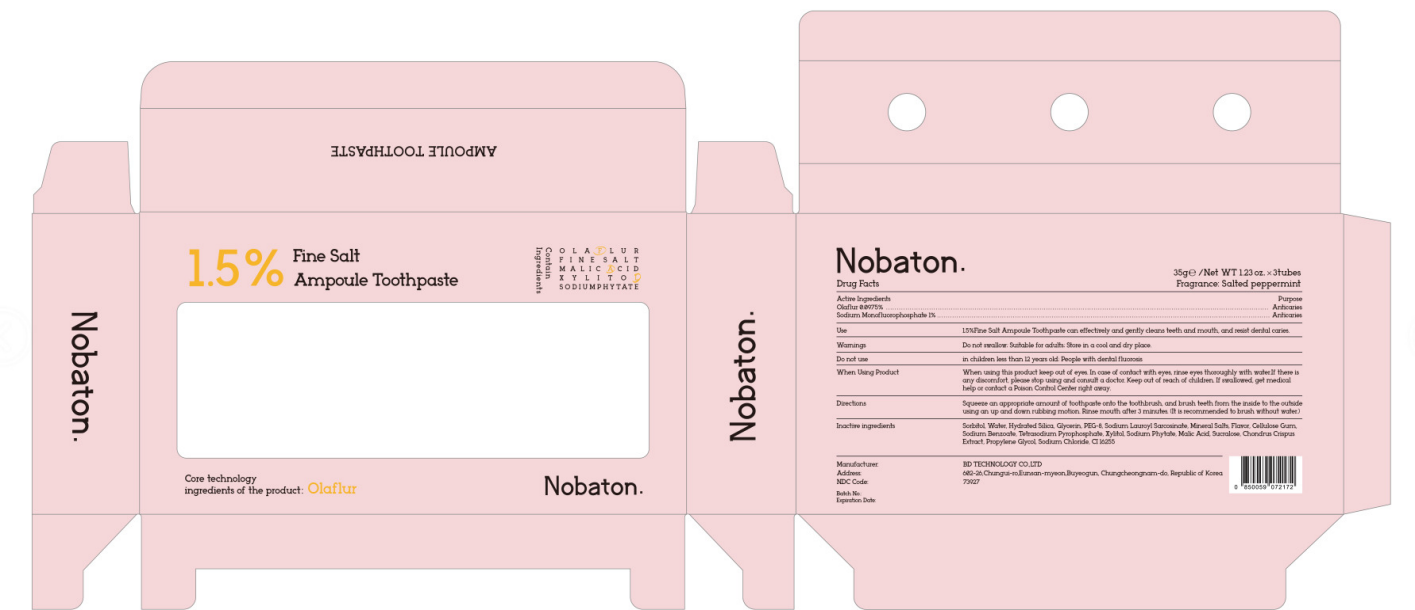
Other information

No

Inactive ingredients

Sorbitol, Water, Hydrated Silica, Glycerin, PEG-8, Sodium Lauroyl Sarcosinate, Mineral Salts, Flavor, Cellulose Gum, Sodium Benzoate, Tetrasodium Pyrophosphate, Xylitol Sodium Phytate, Malic Acid, Sucralose, Chondrus Crispus Extract, Propylene Glycol, Sodium Chloride, CI 16255

Package Label - Principal Display Panel



NOBATON 1.5%FINE SALT AMPOULE

nobaton 1.5%fine salt ampoule toothpaste ointment

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:73927-0009
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Route of Administration		DENTAL		
Active Ingredient/Active Moiety				
Ingredient Name		Basis of Strength	Strength	
SODIUM MONOFLUOROPHOSPHATE (UNII: C810JCZ56Q) (FLUORIDE ION - UNII:Q80VPU408O)		FLUORIDE ION	1 g in 100 g	
OLAFLUR (UNII: 8NY9L8837D) (OLAFLUR - UNII:8NY9L8837D)		OLAFLUR	0.0975 g in 100 g	
Inactive Ingredients				
Ingredient Name			Strength	
SODIUM BENZOATE (UNII: OJ245FE5EU)				
TETRASODIUM PYROPHOSPHATE (UNII: O352864B8Z)				
MALIC ACID (UNII: 817L1N4CKP)				
XYLITOL (UNII: VCQ006KQ1E)				
CHONDRUS CRISPUS CARRAGEENAN (UNII: UE856F2T78)				
SUCRALOSE (UNII: 96K6UQ3ZD4)				
CELLULOSE GUM (UNII: K679OBS311)				
SODIUM CHLORIDE (UNII: 451W47IQ8X)				
CI 16255 (UNII: Z525CBK9PG)				
SORBITOL (UNII: 506T60A25R)				
WATER (UNII: 059QF0KO0R)				
HYDRATED SILICA (UNII: Y6O7T4G8P9)				
PEG-8 (UNII: B697894SGQ)				
GLYCERIN (UNII: PDC6A3C0OX)				
SODIUM LAUROYL SARCOSINATE (UNII: 632GS99618)				
FLAVOR SPICE MINT N&A110589 (UNII: KG6KU384XW)				
SODIUM PHYTATE (UNII: 88496G1ERL)				
CHROMITE (UNII: W86U2Q2334)				
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:73927-0009-1	35 g in 1 TUBE; Type 0: Not a Combination Product	04/21/2025	
Marketing Information				
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other			04/21/2025	

Labeler - BD TECHNOLOGY CO.,LTD (695005161)

Establishment

Name	Address	ID/FEI	Business Operations
BD TECHNOLOGY CO.,LTD		695005161	manufacture(73927-0009)

Revised: 4/2025

BD TECHNOLOGY CO.,LTD