## ADVANCED WHITENING ANTI CAVITY FLUORIDE- sodium monofluorophosphate paste Universal Distribution Center LLC

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#### UNIVERSAL ADVANCED WHITENING ANTI-CAVITY FLUORIDE TOOTHPASTE

#### **Active Ingredient**

Sodium Monofluorophosphate 0.76%

(1000 ppm)

#### **Purpose**

Anticavity toothpaste

#### Uses

- builds increasing protection against painful sensitivity of teeth due to cold, heat, acids, sweets or contact.
- aids in the prevention of dental cavities.

#### Warning

#### When using this product

• if pain\ sensitivity still persists after 4 weeks of use, please visit your dentist.

#### Stop and ask a dentist

• if the problem persists or worsens.

Sensitivity teeth may indicate a serious problem that may need prompt care by a dentist.

## Keep out of reach of children

• If accidentally swallowed more than used for brushing, seek professional help or contact a Poison Control Center immediately.

#### **Directions**

- adults and children of 2 years and older: brush teeth thoroughly after meals or at least twice a day, or as directed by a dentist.
- do not swallow.
- to minimize swallowing, use a pea-sized amount in children under 6 years old.
- supervise children's brushing until good habits are established.
- children under 2 years: ask a dentist before use.

### Children under 12 years of age: Consult a dentist or doctor

#### Other information

• store in a cool, dry place.

## **Inactive Ingredients**

Calcium carbonate, Water, Glycerine, Silica, Sodium Lauryl sulphate, Sorbitol, Xanthan gum, Sodium saccharine, Sodium benzoate, Sodium carboxy methyl cellulose, Flavor and FD&C blue #1

#### PRINCIPAL DISPLAY PANEL

#### ADVANCED WHITENING ANTI-CAVITY FLUORIDE TOOTHPASTE



#### **ADVANCED WHITENING ANTI CAVITY FLUORIDE**

sodium monofluorophosphate paste

# Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:52000-108 Route of Administration DENTAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
<b>SODIUM MONOFLUOROPHOSPHATE</b> (UNII: C810JCZ56Q) (FLUORIDE ION - UNII:Q80VPU408O)	FLUORIDE ION	0.76 g in 100 g		

Inactive Ingredients		
Ingredient Name	Strength	
CALCIUM CARBONATE (UNII: H0G9379FGK)		
WATER (UNII: 059QF0KO0R)		

GLYCERIN (UNII: PDC6A3C0OX)	
SILICON DIOXIDE (UNII: ETJ7Z 6XBU4)	
SODIUM LAURYL SULFATE (UNII: 368GB5141J)	
SORBITOL (UNII: 506T60A25R)	
XANTHAN GUM (UNII: TTV12P4NEE)	
SACCHARIN SODIUM (UNII: SB8ZUX40TY)	
SODIUM BENZOATE (UNII: OJ245FE5EU)	
CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM (UNII: K6790BS311)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	

	Packaging				
7	tem Code	Package Description	Marketing Start Date	Marketing End Date	
:	NDC:52000-108- 01	181 g in 1 TUBE; Type 0: Not a Combination Product	05/18/2020		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M021	05/18/2020		

## **Labeler -** Universal Distribution Center LLC (019180459)

Revised: 11/2023 Universal Distribution Center LLC