

SENSODYNE PRONAMEL GENTLE WHITENING ADVANCED PLUS- potassium nitrate, sodium fluoride paste
Haleon US Holdings LLC

Drug Facts

Active ingredients

Potassium nitrate 5%

Sodium fluoride 0.25% (0.15% w/v fluoride ion)

Purposes

Antihypersensitivity

Anticavity

Uses

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

Warnings

Stop use and ask a dentist if

- the problem persists or worsens. Sensitive teeth may indicate a serious problem that may need prompt care by a dentist.
- pain/sensitivity still persists after 4 weeks of use.

Keep out of reach of children.

If more than used for brushing is accidentally swallowed, get medical help or contact a Poison Control Center right away.

Directions

- **adults and children 12 years of age and older:**
 - apply at least a 1-inch strip of the product onto a soft bristle toothbrush.
 - brush teeth thoroughly for at least 1 minute twice a day (morning and evening), and not more than 3 times a day, or as recommended by a dentist or doctor. Make sure to brush all sensitive areas of the teeth. Minimize swallowing. Spit out after brushing.
- **children under 12 years of age:**Consult a dentist or doctor.

Other information

- do not store above 25°C (77°F)

Inactive ingredients

water, sorbitol, hydrated silica, glycerin, cocamidopropyl betaine, flavor, xanthan gum, titanium dioxide, sodium saccharin, sodium hydroxide

Questions or comments?

1-866-844-2797

Principal Display Panel

HALEON

4 PACK

SENSODYNE PRONAMEL

TOOTHPASTE FOR SENSITIVE TEETH AND CAVITY PREVENTION

Specialist Enamel Protection

Rebuilds, Restores, Refreshes

Alpine Breeze

GENTLE WHITENING

ADVANCED PLUS†

2 in 1 WHITENING*

+ STRENGTHENS ENAMEL

4 TUBES OF 6.5 OZ (184 g) EACH - TOTAL NET WT 26 OZ (736 g)

62000000209634 Carton Front



SENSODYNE PRONAMEL GENTLE WHITENING ADVANCED PLUS

potassium nitrate, sodium fluoride paste

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:0135-5030
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
POTASSIUM NITRATE (UNII: RU45X2JN0Z) (NITRATE ION - UNII:T93E9Y2844)	POTASSIUM NITRATE	50 mg in 1 g

SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80VPU408O) | **SODIUM FLUORIDE** | 1.11 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
SORBITOL (UNII: 506T60A25R)	
HYDRATED SILICA (UNII: Y607T4G8P9)	
GLYCERIN (UNII: PDC6A3C0OX)	
COCAMIDOPROPYL BETAINE (UNII: 5OCF3O11KX)	
XANTHAN GUM (UNII: TTV12P4NEE)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
SACCHARIN SODIUM (UNII: SB8ZUX40TY)	
SODIUM HYDROXIDE (UNII: 55X04QC32I)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0135-5030-01	4 in 1 CARTON	10/01/2024	
1		184 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
OTC Monograph Drug	M022	10/01/2024	

Labeler - Haleon US Holdings LLC (079944263)

Revised: 8/2024

Haleon US Holdings LLC