SUPER VANILLA PRESCRIPTION- anticavity toothpaste paste, dentifrice SuperMouth, LLC.

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.

SuperMouth Super Vanilla Prescription Toothpaste

ACTIVE INGREDIENT Sodium Fluoride 1.1% (5000 ppm)

PURPOSE Sodium Fluoride Anticavity

USE: This medication is used as part of a professional program for the prevention and control of dental caries. Use as directed by your health professional.

WARNINGS: Do not swallow. Do not use in children under 6 years of age unless recommended by a dentist or physician. 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.

See your dentist if the problem persists or worsens. Do not use this product longer than 4 weeks unless recommended by a dentist or physician.

DIRECTIONS: Use as directed by a dentist or physician.

INACTIVE INGREDIENTS

Glycerin, Water, Hydrated Silica, *Hydroxamin® (Nano-Hydroxyapatite, Menquinone-y (Vit K2), Cholecalciferol (Vit D3)), Xylitol, Inulin, Sodium Gluconate, Methylsulfonylmethane, Xanthan Gum, Quillaja Saponaria Extract, Natural Flavor, Natural Benzoic Acid, Sodium Ascorbate (Vit C), Stevia Leaf Extract, Cranberry Seed Oil. *Patent-Pending Formulation

PRINCIPAL DISPLAY PANEL - SuperMouth Super Vanilla Prescription Toothpaste

Supermouth Pro

1.1% sodium fluoride

5000 ppm

Anticavity Toothpaste

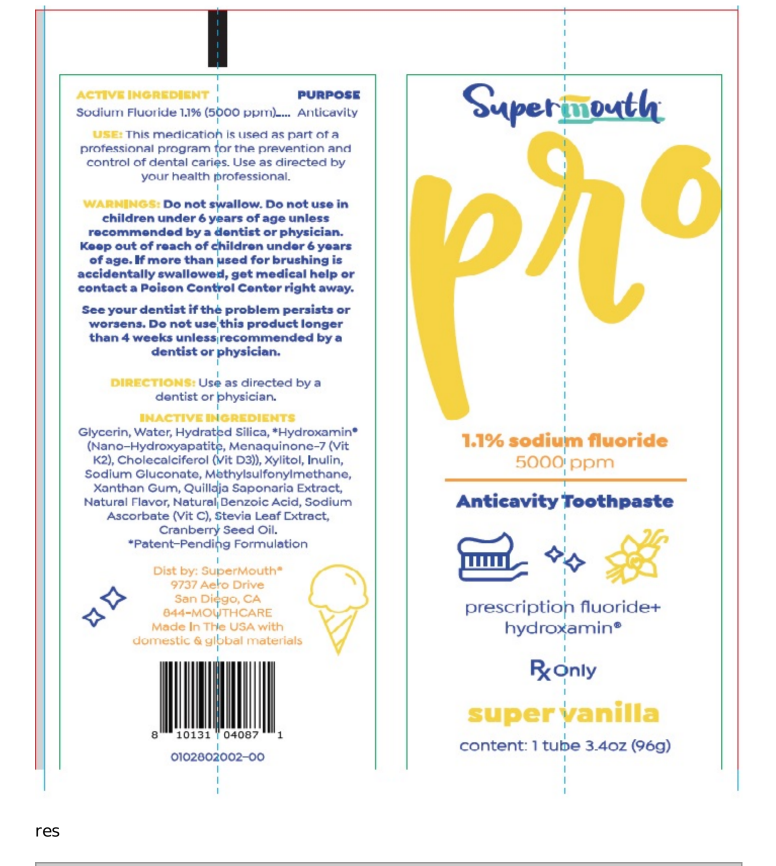
Rx Only

super vanilla

content: 1 tube 3.4oz (96g)

Dist by: SuperMouth[®] 9737 Aero Drive San Diego, CA 844-MOUTHCARE Made In The USA with Domestic & global materials





SUPER VANILLA PRESCRIPTION

anticavity toothpaste paste, dentifrice

Product Information

Product Type

HUMAN PRESCRIPTION DRUG

Item Code (Source)

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80VPU4080)	FLUORIDE ION	1.1 g in 100 g
Inactive Ingredients		
Ingredient Name		Strength
GLYCERIN (UNII: PDC6A3C0OX)		onengen
WATER (UNII: 059QF0KO0R)		
HYDRATED SILICA (UNII: Y607T4G8P9)		
TRIBASIC CALCIUM PHOSPHATE (UNII: 91D9GV0Z28)		
MENAQUINONE 7 (UNII: 8427BML8NY)		
CHOLECALCIFEROL (UNII: 1C6V77QF41)		
XYLITOL (UNII: VCQ006KQ1E)		
INULIN (UNII: JOS53KRJ01)		
SODIUM GLUCONATE (UNII: R6Q3791S76)		
DIMETHYL SULFONE (UNII: 9H4PO4Z4FT)		
XANTHAN GUM (UNII: TTV12P4NEE)		
QUILLAJA SAPONARIA WHOLE (UNII: HIU9R169Y7)		
BENZOIC ACID (UNII: 85KN0B0MIM)		
SODIUM ASCORBATE (UNII: S033EH8359)		
STEVIA LEAF (UNII: 6TC6NN0876)		
CRANBERRY SEED OIL (UNII: 73KDS3BW5E)		

Product Characteristics

Color	brown	Score	
Shape		Size	
Flavor	VANILLA	Imprint Code	
Contains			

Packaging

	#	ltem Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:83729-100- 10	1 in 1 CARTON	10/13/2023	
	1		96 g in 1 TUBE; Type 0: Not a Combination Product		
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Marketing In	larketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		10/13/2023		

Labeler - SuperMouth, LLC. (049384038)

Registrant - SuperMouth, LLC. (049384038)

Revised: 10/2023

SuperMouth, LLC.