

**MIPASTE PLUS MINT- mipaste plus mint paste, dentifrice
GC America Inc.**

MIPaste Plus Mint

Active Ingredient

Sodium fluoride 0.20% (w/w)

Purpose

Anticavity

Warnings

Keep out of reach of children

under 6 years of age. Do not use on patients with a milk protein or hydroxybenzoates allergy. In case of allergic reaction; stop use, rinse mouth with water and seek medical advice. If you

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

Directions

- adults and children over 12:
brush teeth with your normal toothpaste at least twice a day, or as directed by your dental professional or doctor, rinse mouth after brushing, dispense a pea-sized portion of MI Paste Plus onto your finger, apply thoroughly, coating the upper teeth with an even layer, repeat for lower teeth, allow to remain on teeth for 3 minutes. Do not rinse.

- children under the age of 12:
consult your dental professional or doctor

Inactive ingredients

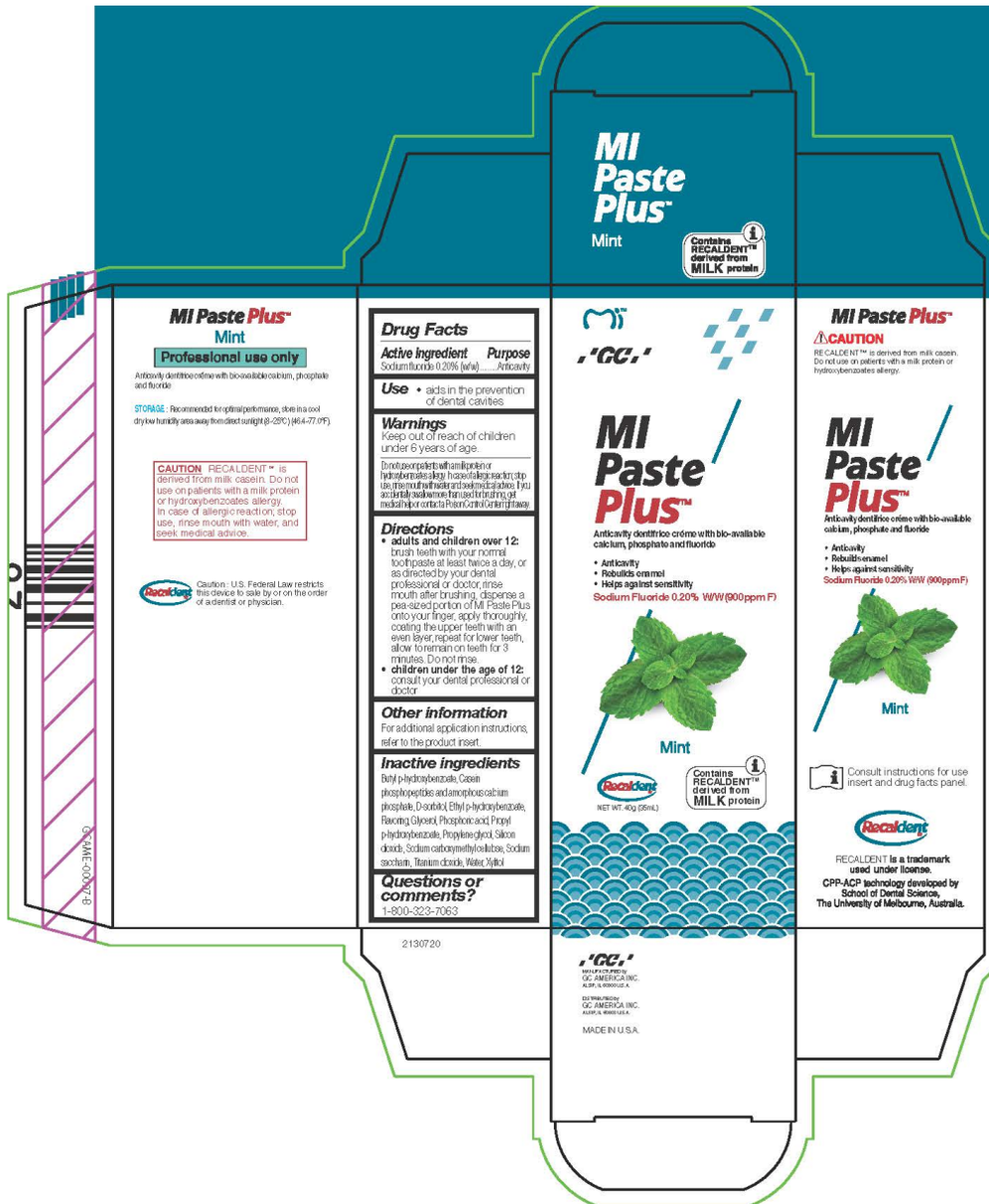
Butyl p-hydroxybenzoate, Casein phosphopeptides and amorphous calcium phosphate, D-sorbitol, Ethyl p-hydroxybenzoate, Flavoring, Glycerol, Phosphoric acid, Propyl p-hydroxybenzoate, Propylene glycol, Silicon dioxide, Sodium carboxymethyl cellulose, Sodium saccharin, Titanium dioxide, Water, Xylitol

Warnings

Keep out of reach of children under 6 years of age.

Use

- aids in the prevention of dental cavities



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MIPASTE PLUS MINT

mipaste plus mint paste, dentifrice

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:61596-801
Route of Administration	DENTAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80VPU408O)	FLUORIDE ION	0.2 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
BUTYLPARABEN (UNII: 3QPI1U3FV8)	
CALCIUM PHOSPHATE, UNSPECIFIED FORM (UNII: 97Z1W3NDX)	
ETHYLPARABEN (UNII: 14255EXE39)	
GLYCERIN (UNII: PDC6A3C0OX)	
PHOSPHORIC ACID (UNII: E4GA8884NN)	
XYLITOL (UNII: VCQ006KQ1E)	
SORBITOL (UNII: 506T60A25R)	
PROPYLPARABEN (UNII: Z8IX2SC1OH)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CARBOXYMETHYLCELLULOSE SODIUM (UNII: K679OBS311)	
SACCHARIN SODIUM (UNII: SB8ZUX40TY)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	

Product Characteristics

Color		Score	
Shape		Size	
Flavor	MINT	Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:61596-801-10	10 in 1 BOX	02/03/2022	
1	NDC:61596-801-41	1 in 1 BOX, UNIT-DOSE		
1	NDC:61596-801-40	40 g in 1 TUBE; Type 9: Other Type of Part 3 Combination Product (e.g., Drug/Device/Biological Product)		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M021	02/03/2022	

Labeler - GC America Inc. (005473608)

Registrant - GC America Inc. (005473608)

Establishment

Name	Address	ID/FEI	Business Operations
GC America Inc.		005473608	manufacture(61596-801)

Revised: 12/2025

GC America Inc.