# ZAP APF- sodium fluoride gel Crosstex International Inc.

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.

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# **Active Ingredient:**

Fluoride Ion 1.23%. A stable thixotropic gel providing 1.23% fluoride ion. Available from 2.09% Sodium Fluoride and Hydrofluoric Acid.

# **Purpose:**

Anticaries Treatment.

# Indications and Usage:

This is a prescription fluoride treatment gel used to help prevent dental decay.

• For Professional Use Only. This product is not intended for home or unsupervised consumer use.

# **Warnings:**

- Keep out of reach of children.
- Do not swallow. If product is accidentally swallowed in quantities greater than would normally occur with a treatment gel, seek medical help or contact a Poison Control Center right away.
- Read directions carefully before using.

# **Dosage and Administration:**

Shake well before use. This is a one minute or four minute fluoride gel for in-office patient use. It is normally used as a preventative caries treatment two times a year.

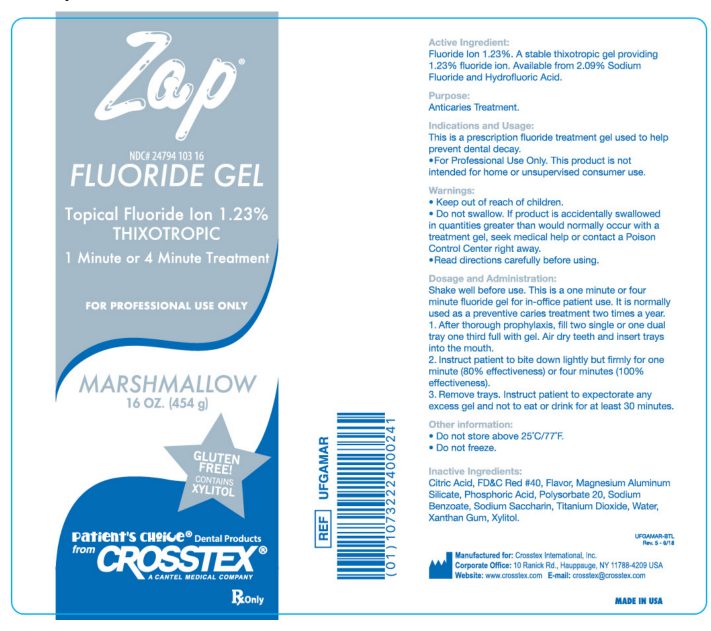
- 1. After thorough prophylaxsis, fill two single or one dual tray, one third full with gel. Air dry teeth and insert trays into the mouth.
- 2. Instruct patient to bite down lightly but firmly for one minute (80% effectiveness) or four minutes (100% effectiveness).
- 3. Remove trays. Instruct patient to expectorate any excess gel and not to eat or drink for at least 30 minutes.

### Other Information:

Do not store above 25°C/77°F. Do not freeze.

# **Inactive Ingredients:**

Citric Acid, FD&C Red #40, Flavor, Magnesium Aluminum Silicate, Phosphoric Acid, Polysorbate 20, Sodium Benzoate, Sodium Saccharin, Titanium Dioxide, Water, Xanthan Gum, Xylitol.



# ZAP APF sodium fluoride gel Product Information Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:24794-103 Route of Administration DENTAL Active Ingredient/Active Moiety

Ingredient Name	<b>Basis of Strength</b>	Strength
SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80VPU4080)	FLUORIDE ION	5.6 g in 454 g

Inactive Ingredients	
Ingredient Name	Strength
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)	
MAGNESIUM ALUMINUM SILICATE (UNII: 6M3P64V0NC)	
PHOSPHORIC ACID (UNII: E4GA8884NN)	
POLYSORBATE 20 (UNII: 7T1F30V5YH)	
SODIUM BENZOATE (UNII: OJ245FE5EU)	
SACCHARIN SODIUM (UNII: SB8ZUX40TY)	
WATER (UNII: 059QF0KO0R)	
XANTHAN GUM (UNII: TTV12P4NEE)	
XYLITOL (UNII: VCQ006KQ1E)	
FD&C RED NO. 40 (UNII: WZB9127XOA)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	

Product Characteristics			
Color pink Score			
Shape		Size	
Flavor	MARSHMALLOW	Imprint Code	
Contains			

l	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:24794- 103-16	454 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/01/2018	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		12/14/2012	
outer			

# **Labeler -** Crosstex International Inc. (057728685)

Revised: 1/2022 Crosstex International Inc.