PRIMO APF- sodium fluoride gel Tri State Dental

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.

Active Ingredient:

Fluoride Ion 1.23%

Available from 2.09% Sodium Fluoride and Hydrofluoric Acid

Purpose:

Fluoride Treatment Gel

Indications and Usage:

- A stable thixotropic fluoride treatment gel used to help prevent dental decay.
- For Professional Use Only. This product is not intended for home or unsupervised 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).

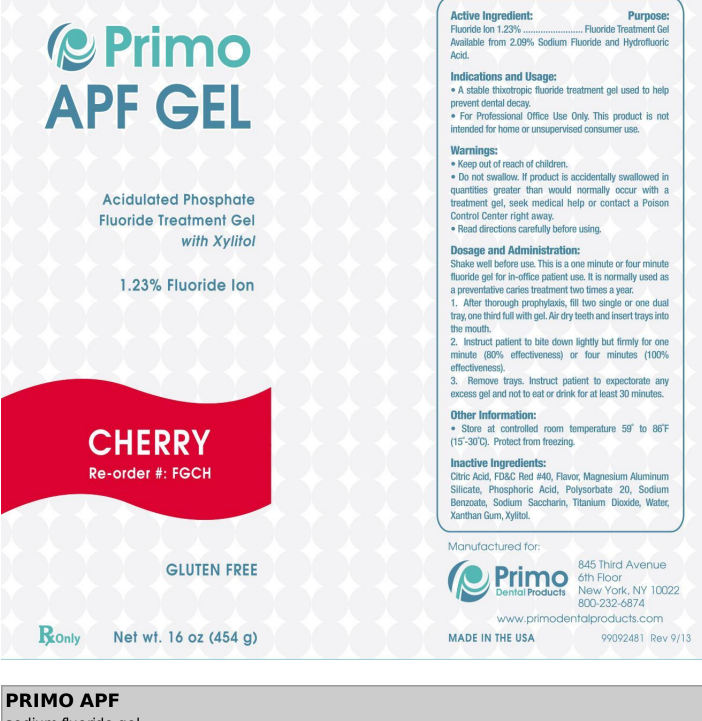
3. Remove trays. Instruct patient to expectorate any excess gel and not to eat or drink for at least 30 minutes.

Other Information:

Store at controlled room temperature 59° to 86°F (15°-30°C).

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.



sodium fluoride gel

Product Information

Product Type

HUMAN PRESCRIPTION DRUG

Item Code (Source)

NDC:69509-051

Route of Administ	ration DEN
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Route of Administration	DENTAL			
Active Ingredient/Active	Moiety			
Ingre	edient Name		Basis of Strength	n Strength
SODIUM FLUORIDE (UNII: 8ZYQ1	474W7) (FLUORIDE ION - U	JNII:Q80VPU408O)	FLUORIDE ION	5.6 g in 454 g
Inactive Ingredients				
	Ingredient Name			Strength
CITRIC ACID MONOHYDRATE (U	- NII: 2968PHW8QP)			
MAGNESIUM ALUMINUM SILICA	TE (UNII: 6M3P64V0NC)			
PHOSPHORIC ACID (UNII: E4GA8884NN)				
POLYSORBATE 20 (UNII: 7T1F30V5YH)				
SODIUM BENZOATE (UNII: OJ245	SODIUM BENZOATE (UNII: OJ245FE5EU)			
SACCHARIN SODIUM (UNII: SB8ZUX40TY)				
WATER (UNII: 059QF0KO0R)				
XANTHAN GUM (UNII: TTV12P4NE	E)			
XYLITOL (UNII: VCQ006KQ1E)				
FD&C RED NO. 40 (UNII: WZ B912	•			
TITANIUM DIOXIDE (UNII: 15FIX9)	/2JP)			
Product Characteristics				
Color	red	Score		
Shape		Size		
Flavor	CHERRY	Imprint Code		
Contains				

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Par	K - C I I C I
1 4 4	NAMINA

-	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:69509- 051-15	454 g in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	09/01/2013	

Marketing Information

unapproved drug 09/01/2013	Marketing	Application Number or Monograph	Marketing Start	Marketing End
	Category	Citation	Date	Date
	unapproved drug other		09/01/2013	

Labeler - Tri State Dental (176048478)