## IONITE APF- sodium fluoride gel Dharma Research, 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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- Store at 59° 86°F (15° 30°C)
- Protect from freezing
- Ionite APF Gel is free of gluten, soymilk, egg peanut and free of nut products

Citric Acid, FD & C Red No. 40 (C.I. 16035), Flavor, Magnesium Aluminum Silicate, Phosphoric Acid, Polysorbate 20, Purified Water, Sodium Benzoate, Sodium Saccharin, Sweetness Enhancer, Titanium Dioxide, Tocopheryl Acetate, Xanthan Gum, Xylitol

- Keep out of the 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.
- Do not use on patients with an allergy Fluoride.
- Ionite APF Gel contains artificial color, confirm that no kno.wn patient allergies exist.
- For professional use only.
- Shake well before use.
- This is a four minute or one minute Fluoride gel for in-office patient use.
- For best results, use at least twice a year.
- After thorough prophylaxis, fill two single or one dual tray one third full with gel. Air dry teeth and insert trays into the mouth.
- Instruct patient to bite down lightly but firmly for one minute (80% effectiveness) or four minutes (100% effectiveness)
- Remove trays, instruct patient to expectorate any excess gel and not to eat or drink for at least 30 minutes

	Drug Facts	Ľ,	
	Active Ingredient Purpose Sodium Fluoride 2.72% Anticaries	$\langle \rangle$	
Valencia Orange	Warnings ■ Keep out of the 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. ■ Do not use on patients with an allergy to fluoride. ■ Ionite APF Gel contains artificial color, confirm that no known patient allergies exist. ■ For professional use only.		
APF THIXOTROPIC GEL with Xylitol & Vitamin E Acidulated Phosphate Fluoride	Directions Shake well before use. This is a four minute or one minute fluoride gel for in-office patient use. For best results, use at least twice a year. After thorough prophylaxis, fill two sigle or one dual tray one third full with gel. Air dry teeth and insert trays into the mouth. Instruct patient to bite down lightly but firmly for one minute (80% effectiveness) or four minutes (100% effectiveness). Remove trays, instruct patient to expectorate any excess gel and not to eat or drink for at least 30 minutes.		
Treatment Gel 1.23% Fluoride Ion	Other information   ■ Store at 59° - 80°F (15°-30° C),   ■ Protect from freezing,   ■ Ionite APF Gel is free of gluten, soy milk, egg, peanut and tree nut products.		
GLUTEN FREE	Inactive ingredients Citric Acid, FD&C Red No. 40 (C.I. 16035), Flavor, Magnesium Aluminum Silicate. Phosphoric Acid, Polysorbate 20, Purified Water, Sodium Benzoate, Sodium Saccharin, Sweetness Enhancer, Titanium Dioxide, Tocopheryl Acetate, Xantham Gum, Xyllol.	te,	
1 minute or 4 minute Treatment Note: (500ml)	Manufactured by: Charles Besarce INC WWW.DHARMARESTARCH.INC 5220 N.W 72 Avenue Unit 15, Miami, FI, 33166 1-877-833-3725 3 53045 00017/188		

MADE IN USA

NDC#53045-204-17

## **IONITE APF** sodium fluoride gel **Product Information** Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:53045-204 **Route of Administration** DENTAL, ORAL **Active Ingredient/Active Moiety Ingredient Name** Basis of Strength Strength **SODIUM FLUORIDE** (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80VPU408O) FLUORIDE ION 6.027 g in 490 g **Inactive Ingredients Ingredient Name** Strength CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP) FD&C RED NO. 40 (UNII: WZB9127XOA)

MAGNESIUM ALUMINUM SILICATE (UNII: 6M3P64V0NC)

Pł	HOSPHORIC ACID (U	JNII: E4GA8884NN)						
PO	DLYSORBATE 20 (U	JNII: 7T1F30V5YH)						
W	ATER (UNII: 059QF0	KO0R)						
so	DIUM BENZOATE	(UNII: OJ245FE5EU)						
SA	SACCHARIN SO DIUM (UNII: SB8ZUX40TY)							
TI	TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)							
.A	.ALPHATO COPHEROL ACETATE, DL- (UNII: WR1WPI7EW8)							
X	XANTHAN GUM (UNII: TTV12P4NEE)							
X	YLITOL (UNII: VCQ0	06KQ1E)						
_								
Product Characteristics								
C	olor	or Score						
SI	ıape			Size				
Fl	avor	ORANGE (Valencia Orange)		Imprint Code				
Contains								
Packaging								
#	Item Code	Package Description	Marke	ting Start Date	Marketing End	Date		
1	NDC:53045-204-17	490 g in 1 BOTTLE; Type 0: Not a Combination Product	0 1/0 1/20 13					
Marketing Information								
	0		Marka	ting Start Data	Maxima End I	) a ta		
	Iarketing Category approved drug other	Application Number or Monograph Citation	0 1/0 1/20	ting Start Date	Marketing End I	Jale		
un	approved drug other		01/01/20	15				

Labeler - Dharma Research, inc. (078444642)

Registrant - Dharma Research, inc. (078444642)

Establishment								
Name	Address	ID/FEI	<b>Business Operations</b>					
Dharma Research, inc.		078444642	manufacture(53045-204)					

Revised: 11/2020

Dharma Research, inc.