DOCTOR 9020 DENTAL CLINIC- sodium fluoride liquid JANGIN PHARM CO.,LTD.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Active ingredient : Sodium fluoride

Preservatives-Sodium Benzoate

Other additives-L-Menthol, SophoraAngustifolia Root Extract, GlycyrrhizaInflata Root Extract, Glycerin, Camellia SinensisCatechins, Saccharin Sodium Hydrate, Sodium Citrate Hydrate, Citric Acid Hydrate, Xylitol, DI-Water, Caramel Colorant, Red Ginseng Extract, Polyoxyl 40 Hydrogenated Castor Oil

Effect :prevent cavities, and remove bad breath

Keep out of reach of children

Administration and doses: Once a day (mainly before bed), use it after brushing thoroughly. 6 years old to adults:10ml into your mouth, mix well for 1 minute, then spit. Be careful not to eat or drink food for 30 minutes after use to obtain sufficient effect

•Caution:1) The fluoride content of this product is 90.5ppm. 2) Do not use this product - Anyone who is hypersensitive to this product. 3) Other Precautions when using this product-Do not swallow. Do not use for children under 6 years old unless instructed by a dentist. Keep out of reach of children. If they drink lots of medicine, feed large amounts of milk and follow doctor's instructions. 4) Storage precautions and other precautions when using this product-Keep out of the reach of children. Avoid direct sunlight. To shade the light, store in a cool place. To prevent misuse and preserve quality, do not change the container

For dental use only



GS PACIFIC GROUP



로얄아이보리지 350g / 인쇄도소 : 원색 4도 / 유광라미네이팅 사이즈 : 70 X 50 X 149(mm)

71	상품기획팀 담당자	디자인팀 담당자	디자인팀장	본부장
검수				

※ 문안 내용검수 ※ 디자인 사양검수





DOCTOR 9020 DENTAL CLINIC

sodium fluoride liquid

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69255-500	
Route of Administration	DENTAL			

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
SODIUM FLUORIDE (UNII: 8ZYQ1474W7) (FLUORIDE ION - UNII:Q80 VPU408O)	FLUORIDE ION	0.02 g in 100 mL	

Inactive Ingredients			
Ingredient Name	Strength		
XYLITOL (UNII: VCQ006KQ1E)			
LEVOMENTHOL (UNII: BZ1R15MTK7)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69255-500-02	15 in 1 PACKAGE	12/16/2019	
1	NDC:69255-500-01	12 mL in 1 POUCH; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part355	12/16/2019	

Labeler - JANGIN PHARM CO.,LTD. (688733680)

Registrant - JANGIN PHARM CO.,LTD. (688733680)

Establishment				
Name	Address	ID/FEI	Business Operations	
JANGIN PHARM CO.,LTD.		688733680	manufacture(69255-500)	

Revised: 12/2019 JANGIN PHARM CO.,LTD.