

LISTERINE GUM THERAPY MOUTHWASH GLACIER MINT- eucalyptol, menthol, methyl salicylate, thymol mouthwash
Kenvue Brands LLC

Listerine Gum Therapy Mouthwash Glacier Mint

Drug Facts

<i>Active ingredients</i>	<i>Purposes</i>
Eucalyptol (0.092%)	Antiplaque/antigingivitis
Menthol (0.042%)	Antiplaque/antigingivitis
Methyl Salicylate (0.060%)	Antiplaque/antigingivitis
Thymol (0.064%)	Antiplaque/antigingivitis

Use

helps prevent and reduce:

- plaque
- gingivitis

Warnings

Do not use in children under 12 years of age

Ask a dentist if symptoms persist, new symptoms appear, or conditions worsen after regular use

Keep out of reach of children. If more than used for rinsing is accidentally swallowed, get medical help or contact a Poison Control Center right away.

Directions

- rinse full strength for 30 seconds with 20 mL (2/3 fluid ounce or 4 teaspoonfuls) morning and night
- do not swallow

Other information

- this rinse is not intended to replace brushing or flossing
- store at room temperature
- cold weather may cloud this product. Its antiseptic properties are not affected.

Inactive ingredients

Water, Alcohol (21.6% v/v), Sorbitol, Poloxamer 407, Benzoic Acid, Zinc Chloride, Sodium Benzoate, Sucralose, Flavor, Sodium Saccharin, Blue 1

Questions?

call toll-free **888-222-0182** or **215-273-8755** (collect)

Distributed by:

JOHNSON & JOHNSON CONSUMER INC.

Skillman, NJ 08558

PRINCIPAL DISPLAY PANEL - 1.0 L Bottle Label

ANTINGIVITIS / ANTIPLAQUE MOUTHWASH

LISTERINE®

GUM THERAPY

4X

HEALTHIER*

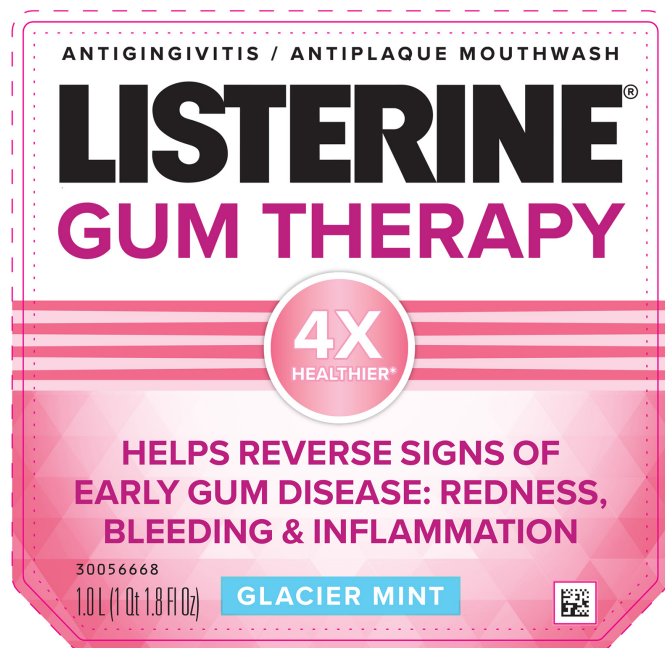
HELPS REVERSE SIGNS OF

EARLY GUM DISEASE: REDNESS,

BLEEDING & INFLAMMATION

GLACIER MINT

1.0 L (1 Qt 1.8 Fl Oz)



LISTERINE GUM THERAPY MOUTHWASH GLACIER MINT

eucalyptol, menthol, methyl salicylate, thymol mouthwash

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:69968-0948	
Route of Administration	ORAL			
Active Ingredient/Active Moiety				
Ingredient Name		Basis of Strength	Strength	
EUCALYPTOL (UNII: RV6J6604TK) (EUCALYPTOL - UNII:RV6J6604TK)		EUCALYPTOL	0.92 mg in 1 mL	
MENTHOL, UNSPECIFIED FORM (UNII: L7T10EIP3A) (MENTHOL, UNSPECIFIED FORM - UNII:L7T10EIP3A)		MENTHOL, UNSPECIFIED FORM	0.42 mg in 1 mL	
METHYL SALICYLATE (UNII: LAV5U5022Y) (SALICYLIC ACID - UNII:O414PZ4LPZ)		METHYL SALICYLATE	0.6 mg in 1 mL	
THYMOL (UNII: 3J50XA376E) (THYMOL - UNII:3J50XA376E)		THYMOL	0.64 mg in 1 mL	
Inactive Ingredients				
Ingredient Name			Strength	
WATER (UNII: 059QF0KO0R)				
ALCOHOL (UNII: 3K9958V90M)				
SORBITOL (UNII: 506T60A25R)				
POLOXAMER 407 (UNII: TUF2IVW3M2)				
BENZOIC ACID (UNII: 8SKN0B0MIM)				
ZINC CHLORIDE (UNII: 86Q357L16B)				
SODIUM BENZOATE (UNII: OJ245FE5EU)				
SUCRALOSE (UNII: 96K6UQ3ZD4)				
SACCHARIN SODIUM (UNII: SB8ZUX40TY)				
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69968-0948-5	500 mL in 1 BOTTLE; Type 0: Not a Combination Product	10/15/2025	
2	NDC:69968-0948-1	1000 mL in 1 BOTTLE; Type 0: Not a Combination Product	10/15/2025	
Marketing Information				
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug		M022	10/15/2025	

Labeler - Kenvue Brands LLC (118772437)