

ANTISEPTIC- eucalyptol, menthol, methyl salicylate, thymol mouthwash
Vi Jon, LLC

Swan 072.003/072AN
Spring Mint Antiseptic Mouthrinse

Active ingredients

Eucalyptol 0.092%

Menthol 0.042%

Methyl salicylate 0.060%

Thymol 0.064%

Purpose

Antigingivitis, Antiplaque

Use

helps control plaque that leads to gingivitis

Warnings

for this product

Do not use

if you have painful or swollen gums, pus from the gum line, loose teeth or increased spacing between the teeth. See your dentist immediately. These may be signs of periodontitis, a serious form of gum disease.

Stop use and ask a dentist if

gingivitis, bleeding, or redness persists for more than 2 weeks

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

adults and children 12 years of age and older vigorously swish 20 mL (2/3 FL OZ or 4 teaspoonfuls) between teeth for 30 seconds then spit out; do not swallow

children under 12 years of age consult a dentist or doctor

- this rinse is not intended to replace brushing or flossing

Other information

cold weather may cloud this product. Its antiseptic properties are not affected. Store at room temperature (59°-77°F).

inactive ingredients

water, alcohol 21.6%, sorbitol solution, flavor, poloxamer 407, benzoic acid, sodium saccharin, sodium citrate, D&C yellow no. 10, FD&C green no. 3

Claims

"The ADA Council on Scientific Affairs Acceptance of Swan Spring Mint Antiseptic Mouth Rinse is based on its finding that the product is effective in helping to prevent and reduce gingivitis and plaque about the gumline, when used as directed."

Disclaimer

*This product is not manufactured or distributed by Johnson & Johnson Healthcare Products, distributor FreshBurst Listerine Antiseptic Mouthwash.

Adverse reaction

Distributed By:

Vi-Jon, LLC

One Swan Drive

Smyrna, TN 37167

DSP-TN-15000

DSP-MO-34

SDS-TN-15012

Principal display panel

Sealed With Printed Neckband For Your Protection

NDC 0869-0072-77

swan ®

ANTISEPTIC

Mouth Rinse

spring mint [®]

Kills Germs that Cause Bad Breath, Plaque & the Gum Disease Gingivitis.

Compare to the active ingredients in Listerine ^{®*}

ADA ACCEPTED

AMERICAN DENTAL ASSOCIATION

500 mL (16.9 FL OZ)



| ANTISEPTIC | | | |
|--|-------------------|---------------------------|---------------|
| eucalyptol, menthol, methyl salicylate, thymol mouthwash | | | |
| Product Information | | | |
| Product Type | HUMAN OTC DRUG | Item Code (Source) | NDC:0869-0072 |
| 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 (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A) | MENTHOL | 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) | |
| SACCHARIN SODIUM (UNII: SB8ZUX40TY) | |
| SODIUM CITRATE (UNII: 1Q73Q2JULR) | |
| D&C YELLOW NO. 10 (UNII: 35SW5USQ3G) | |
| FD&C GREEN NO. 3 (UNII: 3P3ONR6O1S) | |

Packaging

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:0869-0072-21 | 89 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 03/13/2024 |
| 2 | NDC:0869-0072-88 | 2000 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 03/13/2024 |
| 3 | NDC:0869-0072-69 | 250 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 03/17/2020 |
| 4 | NDC:0869-0072-77 | 500 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | |
| 5 | NDC:0869-0072-86 | 1000 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 10/19/2025 |
| 6 | NDC:0869-0072-19 | 94 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 03/31/2017 |
| 7 | NDC:0869-0072-50 | 710 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 03/13/2024 |
| 8 | NDC:0869-0072-13 | 1250 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 03/13/2024 |
| 9 | NDC:0869-0072-12 | 1500 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product | 07/20/1988 | 06/06/2021 |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|--------------------|--|----------------------|--------------------|
| OTC Monograph Drug | 505G(a)(3) | 07/20/1988 | |

Labeler - Vi Jon, LLC (088520668)

Registrant - Consumer Product Partners, LLC (119091520)

Establishment

| Name | Address | ID/FEI | Business Operations |
|--------------------------------|---------|-----------|------------------------|
| Consumer Product Partners, LLC | | 119091514 | manufacture(0869-0072) |

