## SINOFRESH NASAL AND SINUS CARE- eucalyptus globulus leaf, potassium dichromate liquid EMS Contract Packaging

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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#### EMS (as CMO) - SINOFRESH, NASAL (59228-103)

**ACTIVE INGREDIENTS** 

**EUCALYPTUS GLOBULUS 20X** 

KALIUM BICHROMICUM 30X

**PURPOSE** 

**ANTISEPTIC** 

RELIEVES SINUS PAIN, PRESSURE AND INFLAMMATION

**USES** 

RELIEVES NASAL AND SINUS SYMPTOMS ASSOCIATED WITH PERSISTENT SINUS CONDITIONS:

- CONGESTION
- NASAL INFLAMMATION
- SINUS PRESSURE
- FACIAL PIN
- SINUS HEADACHE
- STUFFY NOSE

#### ASK A DOCTOR BEFORE USE IF YOU HAVE

- HAD ANY MEDICAL PROCEDURES FOR YOUR NOSE OR SINUSES.
- A BLEEDING OR IRRITATED NOSE

#### KEEP OUT OF REACH OF CHILDREN.

IF MORE THAN USED FOR SPRAYING IS ACCIDENTALLY SWALLOWED, GET MEDICAL HELP OR CONTACT A POISON CONTROL CENTER RIGHT AWAY.

#### **DIRECTIONS**

 ADULTS AND CHILDREN 12 YEARS OF AGE AND OLDER (WITH ADULT SUPERVISION) MORNING AND EVENING, APPLY 1 TO 2 SPRAYS TO EACH NOSTRIL.

#### OTHER INGREDIENTS

BENZALKONIUM CHLORIDE, CETYLPYRIDINIUM CHLORIDE, DBASIC SODIUM PHOSPHATE, ESSENTIAL OIL BLEND (CONSISTING OF WINTERGREEN OIL, SPEARMINT OIL, PEPPERMINT OIL, AND EUCALYPTUS OIL), MONOBASIC SODIUM PHOSPHATE, POLYSORBATE 80, PROPYLENE GLYCOL, PURIFIED WATER, SODIUM CHLORIDE, AND SORBITOL SOLUTION



## Homeopathic Antiseptic



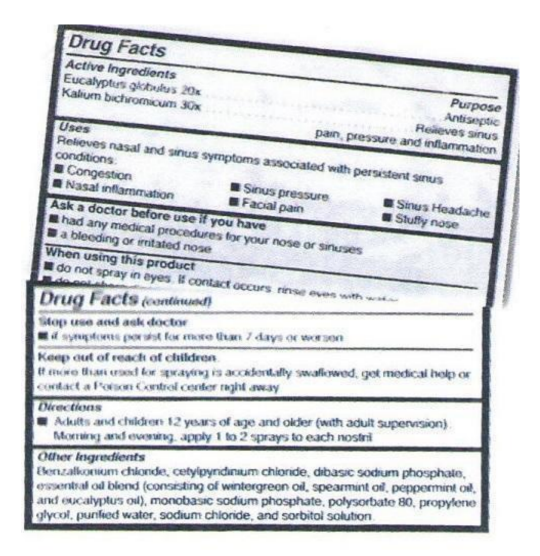
# **NASAL & SINUS CARE**

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SINO-FRESH® is a registered trademark of SinoFresh HealthCare, Inc.

Patent Numbers:

5.785,988; 6,083,525; 6,344,210 and other US and foreign patents pending



#### SINOFRESH NASAL AND SINUS CARE

eucalyptus globulus leaf, potassium dichromate liquid

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:59228-103
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
<b>EUCALYPTUS GLOBULUS LEAF</b> (UNII: S546YLW6E6) (EUCALYPTUS GLOBULUS LEAF - UNII:S546YLW6E6)	EUCALYPTUS GLOBULUS LEAF	20 [hp_X] in 100 mL	
POTASSIUM DICHROMATE (UNII: T4423S18FM) (DICHROMATE ION - UNII:9LKY4BFN2V)	POTASSIUM DICHROMATE	30 [hp_X] in 100 mL	

Inactive Ingredients		
Ingredient Name	Strength	
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7)		
CETYLPYRIDINIUM CHLORIDE (UNII: D9OM4SK49P)		

SODIUM PHOSPHATE, DIBASIC (UNII: GR686LBA74)	
METHYL SALICYLATE (UNII: LAV5U5022Y)	
SPEARMINT OIL (UNII: C3M81465G5)	
PEPPERMINT OIL (UNII: AV092KU4JH)	
SODIUM PHOSPHATE, MONOBASIC (UNII: 3980JIH2SW)	
POLYSORBATE 80 (UNII: 6OZP39ZG8H)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
WATER (UNII: 059QF0KO0R)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SORBITOL (UNII: 506T60A25R)	

l	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:59228- 103-11	29.6 mL in 1 BOTTLE; Type 0: Not a Combination Product	05/30/2014	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		05/30/2014	

### Labeler - EMS Contract Packaging (048602791)

Revised: 4/2022 EMS Contract Packaging