### OHTRUST MOUTH AND NASAL RELIEF- hydroxide ions liquid Nanoplus Life Biomedical Technology Co. Ltd.

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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#### Nanoplus 007-01

## Active Ingredient(s)

Hydroxide ions (544ppm)

#### Purpose

Anti-bacterial

#### Use

• For mouth:

Kills bacteria and virus Balance pH Instant minty breath

#### • For nasal:

Moisturizes and flushes irritants in nostrils

Relieves stuffy nose

#### Warnings

- Avoid contact with eyes. In case of contact, rinse eyes thoroughly with water immediately.
- Use of this container by more than one person may spread infection.
- Temporary discomforts such as burning, stinging, sneezing or an increase in nasal discharge may occur.
- Stop use if rash or irritation develops. Do not drink it!

Keep out of reach of children except under adult supervision.

#### Directions

Spray products 3-4 times directly into the back of the mouth and teeth or into nostril. Allow product to flow down the throat. Repeat when necessary.

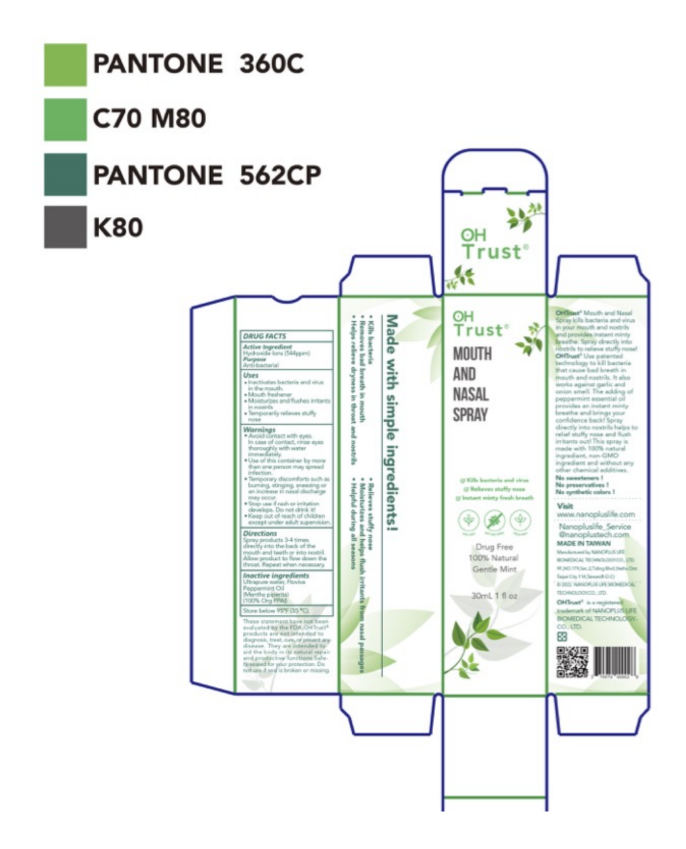
# Other information

Store below 95°F (35°C).

# **Inactive ingredients**

Ultrapure water, Floviva Peppermint Oil (Mentha piperita) (100% Org PPAI)

# Package Label - Principal Display Panel







8X7.5cm

# 30 mL NDC: 70970-007-01

OHTRUST MOUTH AND NASAL RELIEF hydroxide ions liquid											
Ρ	roduct Info	rmation									
Pı	roduct Type		HUMAN OTC DRUG	ltem Code (Source)			NDC:70970-007				
Ro	oute of Admir	nistration	TOPICAL								
Active Ingredient/Active Moiety											
Ingredient Name B						Basis of Strength St					
HYDROXIDE ION (UNII: 9159UV381P) (HYDROXIDE ION - UNII:9159UV381P)					) HYDROXIDE I	HYDROXIDE ION					
In	active Ingr	edients									
		I	ngredient Name			Strength					
PEPPERMINT OIL (UNII: AV092KU4JH)											
W	<b>ATER</b> (UNII: 059	QF0KO0R)									
Pa	ackaging										
#	ltem Code	Pa	ckage Description		Marketing S Date	tart l	Marketing End Date				
1	NDC:70970- 007-01	30 mL in 1 BOT Combination Pr	TLE, SPRAY; Type 0: Not a oduct		04/28/2022						

Marketing Information									
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date						
unapproved drug other		04/28/2022							

Labeler - Nanoplus Life Biomedical Technology Co. Ltd. (657425219)

# Establishment

Name	Address	ID/FEI	<b>Business Operations</b>
Nanoplus Life Biomedical Technology Co. Ltd.		657425219	manufacture(70970-007)

Revised: 12/2023

Nanoplus Life Biomedical Technology Co. Ltd.