

SIMPLY SALINE NETI POT SINUS WASH- sinus wash powder, for solution GURUNANDA, LLC

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.

Simply Saline Neti Pot Nasal Wash

Active Ingredients

Sodium Bicarbonate USP 515 mg

Sodium Chloride USP 1685 mg

Purpose

Nasal Wash

Uses

- Restores vital moisture to provide prompt relief for dry , crusted and inflamed nasal membranes due to chronic sinusitis, colds, low humidity, overuse of nasal decongestant drops and sprays, allergies, minor nose bleeds and other minor nasal irritations.
- Temporarily relieves nasal stuffiness.
- Reduces swelling of nasal passages; shrinks swollen membranes.
- Promotes nasal and/or sinus drainage; temporarily relieves sinus congestion and pressure.

Warnings

- **Do not exceed recommended dosage.**
- This product may cause temporary discomfort such as burning, stinging, sneezing or an increase in nasal discharge.
- The use of this container by more than one person may spread infection.

Keep out of reach of children

Directions

- Adults and children 4 years and over : Use 1-2 packets per 240 ml up to every 2 hours as needed
- Children under 4 years : Consult a physician

Other Information

- Inspect saline packets for integrity
- Do not use saline packets if open or torn

- Protect saline packets from excessive heat and moisture
- See saline packets or box for lot# and expiration date

Inactive Ingredients

None



SIMPLY SALINE NETI POT SINUS WASH

sinus wash powder, for solution

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70708-269
Route of Administration	NASAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SODIUM BICARBONATE (UNII: 8MDF5V39QO) (SODIUM CATION - UNII:LYR4M0NH37, BICARBONATE ION - UNII:HN1ZRA3Q20)	SODIUM BICARBONATE	515 mg in 2200 mg
SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37)	SODIUM CHLORIDE	1685 mg in 2200 mg

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	

Product Characteristics

Color	white (WHITE CRYSTALLINE GRANULE POWDER)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:70708-269-50	50 in 1 BOX	10/01/2022	
1		2200 mg in 1 POUCH; Type 1: Convenience Kit of Co-Package		
2	NDC:70708-269-10	10 in 1 BOX	10/01/2022	
2		2200 mg in 1 POUCH; Type 1: Convenience Kit of Co-Package		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		10/01/2022	

Labeler - GURUNANDA, LLC (079671169)**Registrant** - GURUNANDA, LLC (079671169)