

**SINUS WASH SALINE PACKETS GOOD NEIGHBOR PHARMACY- sodium bicarbonate, sodium chloride powder, for solution
AMERISOURCE BEGEN**

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.

Good Neighbor Pharmacy Sinus Wash Saline Refills 100ct

Active ingredients

(in each packet)

Sodium Bicarbonate USP (700 mg)

Sodium Chloride USP (2300 mg)

Purpose

Nasal Wash

Uses

Temporarily relieves symptoms associated with sinusitis, cold, flu or allergies

-Sneezing

-Nasal stuffiness

-Runny nose

-Post nasal drip

Removes inhaled irritants (dust, pollen)

Removes nasal and sinus drainage

Helps reduce swelling of nasal membranes

Moisturizes dry nasal passages

Keep out of reach of children

Warnings

Stop use and ask a doctor if washing is uncomfortable or symptoms are not relieved.

Do not use unfiltered tap water. see instructions inside box for proper water sources

Do not use if nasal passages are completely blocked or if you have an ear infection or blocked ears

Directions

Adults and children 4 years and over: Use 1-2 packets per 8 fl oz (240 mL) up to every 2 hours as needed.

Children under 4 years: Consult a physician

See enclosed instruction sheet for complete directions and proper use.

Inactive ingredients

None

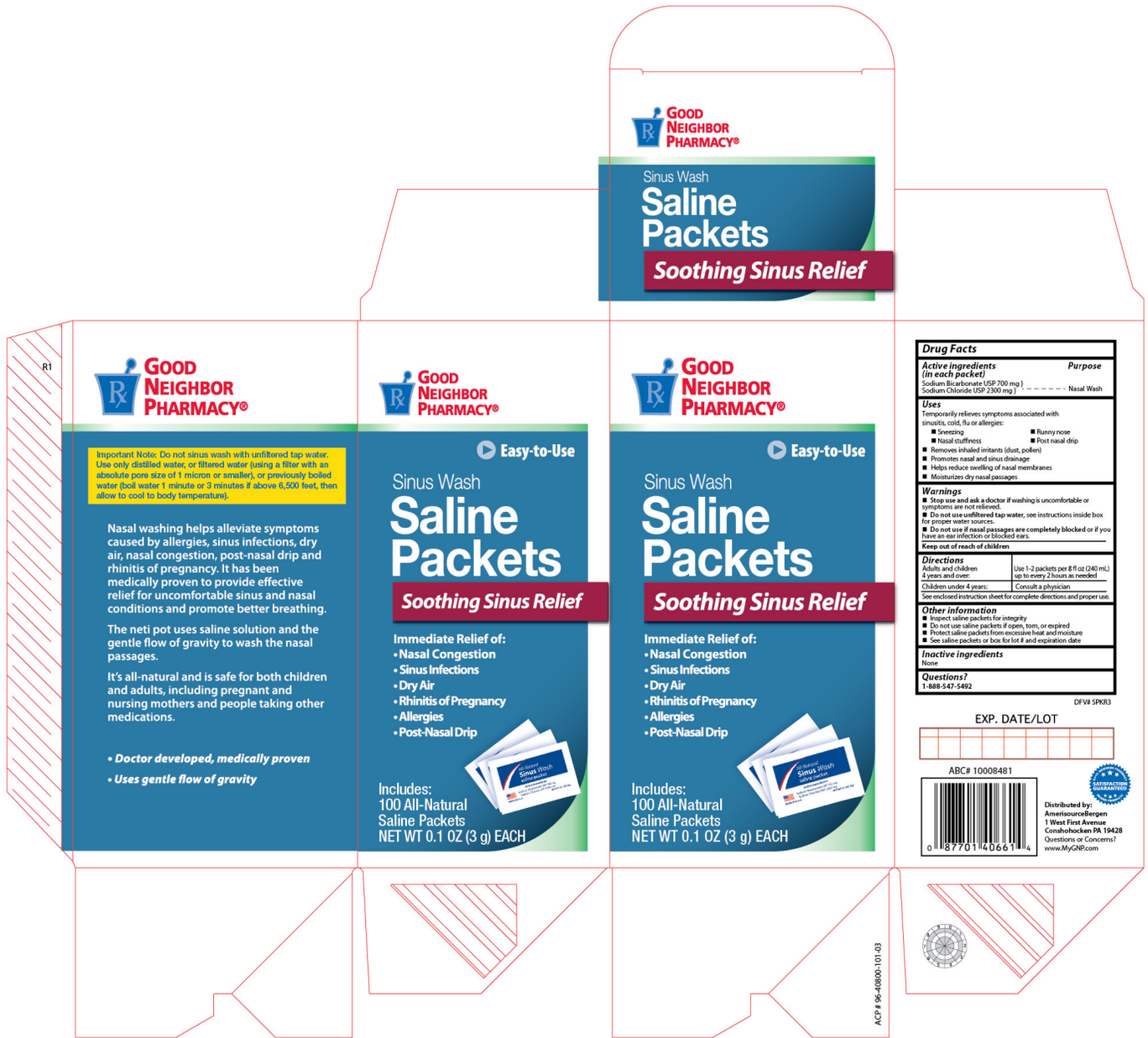
Other Information

- 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

Questions?

1-888-547-5492

- See instruction sheet for use



SINUS WASH SALINE PACKETS GOOD NEIGHBOR PHARMACY

sodium bicarbonate, sodium chloride powder, for solution

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:24385-251
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	700 mg in 3000 mg
SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37)	SODIUM CHLORIDE	2300 mg in 3000 mg

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:24385-251-10	100 in 1 CARTON	01/08/2022	
1	NDC:24385-251-01	3000 mg in 1 PACKET; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		01/08/2022	

Labeler - AMERISOURCE BEGEN (007914906)

Revised: 1/2022

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