

Status: Currently Official on 14-Feb-2025
 Official Date: Official as of 01-Nov-2020
 Document Type: USP Monographs
 DocId: GUID-C7BA6F7D-3D05-47E5-96CC-7EE91B7C9AB3_2_en-US
 DOI: https://doi.org/10.31003/USPNF_M20544_02_01
 DOI Ref: xo4wp

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Cromolyn Sodium Nasal Solution

DEFINITION

Change to read:

Cromolyn Sodium Nasal Solution is an aqueous solution of Cromolyn Sodium. It contains NLT 90.0% and NMT 110.0% of the labeled amount of cromolyn sodium ($C_{23}H_{14}Na_2O_{11}$). It may contain suitable \blacktriangle antimicrobial and stabilizing agents. \blacktriangle (USP 1-Aug-2020)

IDENTIFICATION

• **A.** The retention time of the major peak of the *Sample solution* corresponds to that of the *Standard solution*, as obtained in the Assay.

Add the following:

\blacktriangle • **B.** The UV absorption spectrum of the major peak of the *Sample solution* exhibits maxima and minima at the same wavelengths as those of the *Standard solution*, as obtained in the Assay. \blacktriangle (USP 1-Aug-2020)

ASSAY

Change to read:

• PROCEDURE

Buffer: 5.6 g/L of [monobasic potassium phosphate](#) and 22.2 g/L of [myristyltrimethylammonium bromide](#) in [water](#). Adjust with \blacktriangle 50% sodium hydroxide TS \blacktriangle (USP 1-Aug-2020) to a pH of 6.5.

Mobile phase: [Methanol](#) and *Buffer* (55:45)

Diluent: [Acetonitrile](#) and [water](#) (30:70)

System suitability solution: 0.5 mg/mL of [USP Cromolyn Sodium RS](#) and 0.02 mg/mL each of [USP Cromolyn Related Compound A RS](#) and [USP Cromolyn Related Compound B RS](#) in *Diluent*

Standard solution: 0.5 mg/mL of [USP Cromolyn Sodium RS](#) in *Diluent*. Sonication may be needed to aid dissolution.

Sample solution: Nominally equivalent to 0.5 mg/mL of cromolyn sodium from a volume of Nasal Solution in *Diluent*

Chromatographic system

(See [Chromatography \(621\), System Suitability](#).)

Mode: LC

Detector: UV 326 nm. \blacktriangle For *Identification B*, use a diode array detector in the range of 220–400 nm. \blacktriangle (USP 1-Aug-2020)

Column: 4.6-mm \times 10-cm; 3.5- μ m packing [L7](#)

Column temperature: 40°

Flow rate: 1.5 mL/min

Injection volume: 20 μ L

\blacktriangle **Run time:** NLT 2 times the retention time of cromolyn \blacktriangle (USP 1-Aug-2020)

System suitability

Samples: *System suitability solution* and *Standard solution*

[NOTE—See [Table 1](#) for the relative retention times.]

Suitability requirements

Resolution: NLT 2.0 between cromolyn related compound B and cromolyn related compound A; NLT 2.0 between cromolyn related compound A and cromolyn, *System suitability solution*

Tailing factor: NMT 2.0, *Standard solution*

Relative standard deviation: NMT 0.73%, *Standard solution*

Analysis

Samples: *Standard solution* and *Sample solution*

Calculate the percentage of the labeled amount of cromolyn sodium ($C_{23}H_{14}Na_2O_{11}$) in the portion of Nasal Solution taken:

$$\text{Result} = (r_U/r_S) \times (C_S/C_U) \times 100$$

r_U = peak response of cromolyn from the *Sample solution*

r_S = peak response of cromolyn from the *Standard solution*

C_S = concentration of [USP Cromolyn Sodium RS](#) in the *Standard solution* (mg/mL)

C_U = nominal concentration of cromolyn sodium in the *Sample solution* (mg/mL)

Acceptance criteria: 90.0%–110.0%

IMPURITIES

Change to read:

• ORGANIC IMPURITIES

Mobile phase, Diluent, and System suitability solution: Prepare as directed in the Assay.

Standard solution: 0.002 mg/mL each of [USP Cromolyn Related Compound A RS](#), [USP Cromolyn Related Compound B RS](#), and [USP Cromolyn Sodium RS](#) in *Diluent*

Sample solution: Nominally equivalent to 2 mg/mL of cromolyn sodium from a volume of Nasal Solution in *Diluent*

Chromatographic system

(See [Chromatography \(621\), System Suitability](#).)

Mode: LC

Detector: UV 326 nm

Column: 4.6-mm × 10-cm; 3.5-μm packing [L7](#)

Temperatures

Autosampler: 4°

Column: 40°

Flow rate: 1.5 mL/min

Injection volume: 20 μL

Run time: ▲NLT▲ (USP 1-Aug-2020) 2 times the retention time of cromolyn

System suitability

Samples: *System suitability solution* and *Standard solution*

▲[NOTE—See [Table 1](#) for the relative retention times.]▲ (USP 1-Aug-2020)

Suitability requirements

Resolution: NLT 2.0 between cromolyn related compound B and cromolyn related compound A; NLT 2.0 between cromolyn related compound A and cromolyn, *System suitability solution*

Relative standard deviation: NMT 3% for 6 replicate injections for each peak, *Standard solution*

Analysis

Samples: *Standard solution* and *Sample solution*

Calculate the percentage of cromolyn related compound A or cromolyn related compound B in the portion of Nasal Solution taken:

$$\text{Result} = (r_U/r_S) \times (C_S/C_U) \times 100$$

r_U = peak response of cromolyn related compound A or cromolyn related compound B from the *Sample solution*

r_S = peak response of cromolyn related compound A or cromolyn related compound B from the *Standard solution*

C_S = concentration of ▲[USP Cromolyn Related Compound A RS](#) or [USP Cromolyn Related Compound B RS](#)▲ (USP 1-Aug-2020) in the *Standard solution* (mg/mL)

C_U = nominal concentration of cromolyn sodium in the *Sample solution* (mg/mL)

Calculate the percentage of any individual unspecified degradation product in the portion of Nasal Solution taken:

$$\text{Result} = (r_U/r_S) \times (C_S/C_U) \times 100$$

r_U = peak response of each ▲individual▲ (USP 1-Aug-2020) unspecified degradation product from the *Sample solution*

r_S = peak response of cromolyn from the *Standard solution*

C_s = concentration of [USP Cromolyn Sodium RS](#) in the *Standard solution* (mg/mL)

C_u = nominal concentration of cromolyn sodium in the *Sample solution* (mg/mL)

Acceptance criteria: See [Table 1](#).

Table 1

Name	Relative Retention Time	Acceptance Criteria, NMT (%)
Cromolyn related compound B	0.4	1.0
Cromolyn related compound A	0.5	1.0
Cromolyn	1.0	—
Any individual unspecified degradation product	—	1.0
Total impurities	—	2.0

SPECIFIC TESTS

- [pH \(791\)](#): 4.0–7.0

Add the following:

- ▲ • [MICROBIAL ENUMERATION TESTS \(61\)](#) and [TESTS FOR SPECIFIED MICROORGANISMS \(62\)](#): The total aerobic microbial count is NMT 10^2 cfu/mL, and the total combined molds and yeasts count is NMT 10^1 cfu/mL. It meets the requirements of the tests for absence of *Staphylococcus aureus* and *Pseudomonas aeruginosa*. ▲ (USP 1-Aug-2020)

ADDITIONAL REQUIREMENTS

- **PACKAGING AND STORAGE:** Preserve in tight, light-resistant containers. Store at controlled room temperature.

- [USP REFERENCE STANDARDS \(11\)](#).

[USP Cromolyn Related Compound A RS](#)

1,3-Bis(2-acetyl-3-hydroxyphenoxy) propan-2-ol.

$C_{19}H_{20}O_7$ 360.36

[USP Cromolyn Related Compound B RS](#)

Diethyl 5,5'-[(2-hydroxypropane-1,3-diyl)bis(oxy)]bis(4-oxo-4H-chromene-2-carboxylate).

$C_{27}H_{24}O_{11}$ 524.48

[USP Cromolyn Sodium RS](#)

Auxiliary Information - Please [check for your question in the FAQs](#) before contacting USP.

Topic/Question	Contact	Expert Committee
CROMOLYN SODIUM NASAL SOLUTION	Documentary Standards Support	SM52020 Small Molecules 5

Chromatographic Database Information: [Chromatographic Database](#)

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Pharmacopeial Forum: Volume No. 45(3)

Current DocID: GUID-C7BA6F7D-3D05-47E5-96CC-7EE91B7C9AB3_2_en-US

DOI: https://doi.org/10.31003/USPNF_M20544_02_01

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