

Application Note AN-S-376

Le fluorure dans le gel de fluorure de sodium à usage pharmaceutique

Method validation according to the U.S. Pharmacopoeia

Fluoride is a mineral that is naturally found in water and some foods. It has been proven to strengthen the enamel of the teeth and protect them from decay [1]. However, exposure to too much fluoride can cause dental fluorosis, a condition that affects the appearance of the teeth. Therefore, it is important to monitor the amount of fluoride in dental care products such as gels and toothpastes.

Sodium fluoride gel is a beneficial product that effectively helps prevent cavities (caries). As specified by the authoritative United States Pharmacopeia – National Formulary (USP-NF) Monograph «Sodium Fluoride Gel» [2], ion chromatography (IC) with suppressed conductivity detection is a reliable method to measure fluoride and impurities in sodium fluoride gel.

This study validates an IC method using a Metrosep A Supp 16 - 250/4.0 column and a hydroxide eluent, which meets the USP-NF criteria. Fluoride is separated from chloride and other contaminants in gel toothpaste with high accuracy and precision. The IC method has been validated according to USP General Chapters <621> Chromatography [**3**] and <1225> Validation of Compendial Procedures [**4**].



SAMPLE AND STANDARD PREPARATION

Commercial gel toothpaste was diluted to a known concentration of approximately 2 μ g/mL sodium fluoride (NaF). Here, 1.585 g of a gel toothpaste containing 331.5 mg NaF/100 g was diluted in 500 mL ultrapure water (UPW). The solution was sonicated for 10 minutes and further diluted 1:8.8 with UPW. Afterwards, the diluted solution was filtered using 0.2 μ m pore size filters. The nominal sodium fluoride concentration for these samples was 1.19 μ g/mL.

No additional sample preparation is required. The standard solutions and the system suitability solutions are prepared from the respective 1000 μ g/mL certified standards by dilution with UPW.

For the assay, the standard solution is obtained by diluting a sodium fluoride solution to 2 μ g/mL. The system suitability solution contains 2 μ g/mL sodium fluoride and 1 μ g/mL sodium acetate. For the impurity test, the standard solution consists of 0.2 μ g/mL sodium chloride in UPW. The system suitability solution for the impurity test contains 1 mg/mL sodium fluoride and 1 μ g/mL sodium chloride in UPW.

EXPERIMENTAL

Samples and standard solutions were directly injected

into the IC using a 919 IC Autosampler plus (Figure 1).



Figure 1. Instrumental setup including a 930 Compact IC Flex, 919 IC Autosampler plus, and an 800 Dosino for automatic regeneration of the Metrohm Suppressor Module (MSM).



Fluoride was separated from acetate and chloride by using a potassium hydroxide eluent and the Metrosep A Supp 16 column (column material <u>L91</u>, **Table 1**). The analytes were quantified by evaluating their conductivity signal after chemical suppression. The calibration was performed using a single 2.0 μ g/mL sodium fluoride standard injected six times. The sample was analyzed in duplicate.

Table 1. Requirements for IC method as per USP Monograph «Sodium Fluoride Gel» [2].

| Column with L91 packing | Metrosep A Supp 16 - 250/4.0 |
|-------------------------|-------------------------------|
| Eluent | 15 mmol/L potassium hydroxide |
| Flow rate | 1.0 mL/min |
| Temperature | 40 °C |
| Injection volume | 20 μL |
| Detection | Conductivity with suppression |

RESULTS

The IC assay for fluoride content was validated according to the USP Monograph «Sodium Fluoride Gel» [2]. Suitability requirements for resolution, tailing

factor, and relative standard deviation were fulfilled (Table 2).

Table 2. Suitability requirements for the fluoride assay.

| Parameter (assay) | Actual | USP requirement | Status |
|------------------------------------|--------|-----------------|--------|
| Resolution F ⁻ /acetate | 5.9 | NLT 1.5 | Pass |
| Tailing factor | 1.1 | NMT 2.0 | Pass |
| RSD fluoride (%, n=5) | 0.52 | NMT 0.73 | Pass |

Commercial gel toothpaste samples were analyzed for their sodium fluoride content and the results showed the concentration at 104% of the label claim (Figure 2). The recovery of fluoride for the sample analysis was within the USP acceptance criteria of 90–110%.



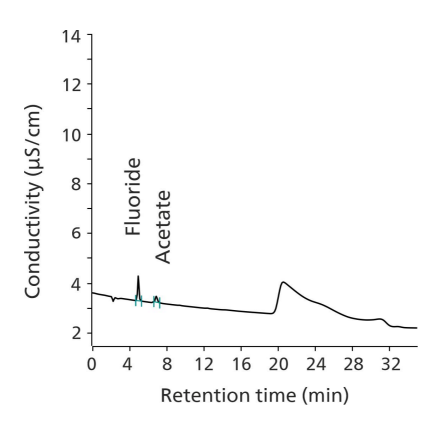


Figure 2. Chromatogram of a commercial toothpaste sample containing 1.24 µg/mL sodium fluoride (104% of the label claim).

When performing the impurity tests for potential contamination with chloride, the IC method showed

excellent compliance with the USP requirements (Table 3).

Table 3. Suitability requirements for the chloride impurity in sodium fluoride gel.

| Parameter (impurity) | Actual | USP requirement | Status |
|--------------------------------------------|--------|-----------------|--------|
| Resolution F ⁻ /Cl ⁻ | 7.7 | NLT 4 | Pass |
| RSD fluoride (%, n=5) | 4.2 | NMT 5 | Pass |
| S/N ratio Cl ⁻ | >740 | NLT 20 | Pass |

RÉSUMÉ

The presented IC method complies with the USP General Chapters <621> and <1225> [**3,4**]. It is suitable to determine sodium fluoride in gels

containing sodium fluoride according to the USP Monograph «Sodium Fluoride Gel» [**2**].



REFERENCES

- Yeung, C. A. A Systematic Review of the Efficacy and Safety of Fluoridation. *Evid Based Dent* 2008, 9 (2), 39–43. <u>https://doi.org/10.1038/sj.ebd.6400578</u>.
- Sodium Fluoride Gel; Monograph; U.S. Pharmacopeia/National Formulary: Rockville, MD.

https://doi.org/10.31003/USPNF_M3947_02_01.

CONTACT

Metrohm France 13, avenue du Québec - CS 90038 91978 VILLEBON COURTABOEUF CEDEX

CONFIGURATION



 621 Chromatography, General Chapter; U.S. Pharmacopeia/National Formulary: Rockville, MD.

https://doi.org/10.31003/USPNF_M99380_01 _01.

 1225 Validation of Compendial Procedures; General Chapter; U.S. Pharmacopeia/National Formulary: Rockville, MD. <u>https://doi.org/10.31003/USPNF_M99945_04</u> <u>01</u>.

info@metrohm.fr

930 Compact IC Flex Oven/ChS/PP/Deg

Le 930 Compact IC Flex Oven/ChS/PP/Deg est un appareil CI compact intelligent avec un **four à colonne**, **suppression chimique** et une **pompe péristaltique** pour la régénération du suppresseur et un **dégazeur** intégré. L'appareil peut etre utilisé avec n'importe quelles méthodes de séparation et de détection.

Domaines d'application typiques :

- Déterminations d'anions avec suppression chimique et détection de conductivité
- Acides organiques avec chromatographie d'exclusion d'ions et suppression inversée





919 IC Autosampler plus

Le 919 IC Autosampler plus satisfait les exigences des laboratoires ayant un nombre moyen d'échantillons à traiter. Il permet l'automatisation de tous les systèmes de chromatographie ionique de Metrohm.



800 Dosino

Moteur de burette avec système de lecture/écriture pour les Unités de distribution intelligentes. Avec cable attenant (150 cm).

