

Application Note AN-U-049

Disinfection byproducts in water

Trace bromate analysis with IC-UV/VIS according to ISO 11206

Safe drinking water is essential for human life and is also quite often a privilege. Whether the source is surface or groundwater, the presence of pathogenic bacteria, poor taste, and a detectable odor requires disinfection processes to guarantee the appropriate quality for drinking water purposes [1,2]. Chlorination was introduced at the beginning of the 20th century as a standard water treatment process. This process helped to protect human health and reduce mortality from waterborne microbial infections and diseases [3,4]. However, the chlorination process forms harmful byproducts (e.g., trihalomethanes) from the reaction of chlorine with organic water components. To avoid such reactions, modern disinfection

processes use strong oxidants like permanganate or ozone. However, if the water contains bromide, ozonation and oxidation lead to the formation of bromate, a potential carcinogen. Therefore, bromate is regulated at a maximum of 10 μ g/L in drinking water and requires regular monitoring to ensure the water quality. Ion chromatography (IC) provides a robust, efficient, and sensitive technique to monitor bromate even at trace levels in line with ISO 11206 and EPA Method 317. The specific post-column reaction (PCR) of bromate-forming triiodide enables the determination of concentrations as low as 1 μ g/L – even in carbonate- and chloride-rich matrices.

Spiked and unspiked tap water (Zurich, Switzerland) and mineral water (Evian) samples were analyzed to test the reliability and validity of this method. In addition to spiking Swiss tap water samples (0.2 and 1 μ g/L bromate), bromate traces in carbonate- and chloride-rich matrices were also investigated to show

the absence of interferences. For these tests, $0.2~\mu g/L$ bromate was added to Evian water (357 mg/L carbonate and 5 mg/L chloride) and to spiked ultrapure water (UPW containing 100–500 mg/L carbonate and 5–100 mg/L chloride).

IC separation with post-column reaction (PCR) and subsequent UV/VIS detection provides a dedicated method to determine very low concentrations of bromate in water. After separation of bromate from matrix components with an analytical column, triiodide is formed via the PCR (Reaction 1). This reaction is very specific, enabling the sensitive determination of bromate via triiodide detection at a wavelength of 352 nm. Due to the high selectivity, the influence of interferences is significantly reduced. This allows trace bromate detection even in matrices high in carbonate and chloride.

The setup is simple (930 Compact IC Flex, 947 Professional UV/VIS detector, a sample processor,

and a Dosino for precise addition of reagents) and conforms with US EPA Method 317 and DIN EN ISO 11206. The separation of bromate from other anions is achieved using the Metrosep A Supp 17 - 250/4.0 column and a sulfuric acid-molybdate eluent. The eluent, which contains molybdate as a catalyst for the PCR, is continuously pumped through the column. Before entering the reactor block, potassium iodide (Reaction 1) is added by a Dosino for triiodide formation and its subsequent UV/VIS detection. The calibration of this setup ranged from 1–20 g/L using a 1000 L injection

$$BrO_3^- + 3I^- + 3H^+ \xrightarrow{Mo(VI)} 3HOI + Br^- (a)$$

 $3HOI + 3I^- + 3H^+ \longrightarrow 3I_2 + 3H_2O (b)$
 $3I_2 + 3I^- \longrightarrow 3I_3^- (c)$
 $BrO_3^- + 9I^- + 6H^+ \xrightarrow{Mo(VI)} 3I_3^- + Br^- + 3H_2O (d)$
UV 352 nm

Reaction 1. Reaction path of bromate with iodine and molybdate as catalyst in acidic solution forming triiodide as described in the triiodide methods in US EPA 317 and ISO 11206. The reaction occurs post-column prior to the spectral detection of triiodide at 352 nm.

However, if bromate needs to be determined in very low concentrations below 1 g/L and especially next to high concentrations of chloride or carbonate, the setup can be easily modified to meet these requirements. Also in this case, the PCR and subsequent UV/VIS detection is used to guarantee a selective bromate determination. For the analytical separation of bromate in a complex matrix, the high-

capacity column **Metrosep A Supp 10 - 100/4.0** and an alkaline bicarbonate eluent is used. To provide more baseline stability and the best reaction conditions, **chemical suppression** is used prior to adding the PCR reagents (**Figure 1**). Using an injection volume of 1325 L in this case allows reliable detection of bromate from **0.05–5 g/L**.

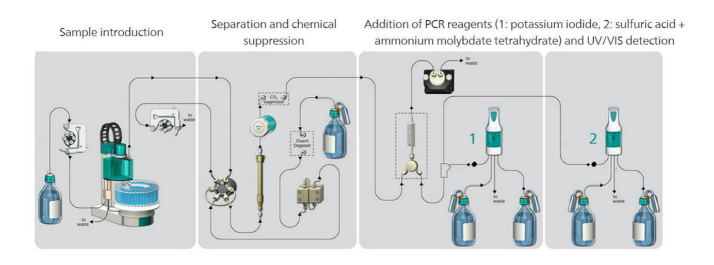


Figure 1. IC system configuration for the determination of trace bromate concentrations in water with the Metrosep A Supp 10 column, chemical suppression, PCR, and UV/VIS detection.

RESULTS AND DISCUSSION

Figure 2A shows the results of the determination of bromate with the Metrosep A Supp 17 column and acidic eluent. With the simple setup (i.e., one Dosino for reagent addition) and a 1000 L injection volume, bromate concentrations from 1–20 g/L can be determined with high accuracy.

The presence of chloride or carbonate in the sample matrix can impact the retention time and peak shape of bromate. To overcome this, the high capacity Metrosep A Supp 10 column efficiently separates bromate from these matrix components before PCR and UV/VIS detection. Thus the detection of bromate down to 0.05 g/L in matrices containing up to 200 mg/L carbonate and 100 mg/L chloride is possible (**Figure 2B**). The retention time for bromate in both

setups is comparable, eluting in less than 10 minutes, which permits analysis of at least 100 samples per day.

Sample spike recoveries for artificially prepared samples with high-matrix UPW and for Swiss tap water ranged from 85–105%. The tap water samples contained some bromate (2.3 g/L). However, no bromate was detected in the carbonate- and chloride-rich Evian samples. Trace level spikes of 0.2 g/L were determined with a recovery of 85%.

A wavelength of 352 nm was chosen for the UV/VIS detection. This decreases the baseline noise because some species from the eluent and samples do not absorb at that wavelength.

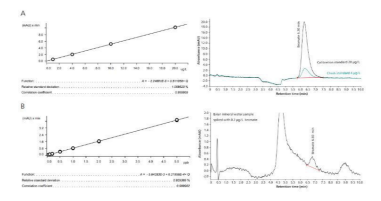


Figure 2. Bromate determination according to US EPA 317 and ISO 11206 using the Metrosep A Supp 17 column (A) and in carbonate-rich matrices with the Metrosep A Supp 10 column (B). The calibration for the simple setup with the Metrosep A Supp 17 column (A) ranged from 1–20 μ g/L bromate. The chromatogram shows the UV/VIS absorbance from bromate standards of 20 μ g/L and 4 μ g/L. The bromate calibration at trace levels using the Metrosep A Supp 10 column (B) ranged from 0.05–5.0 μ g/L (standard bracketing). The chromatogram shows a spiked Evian mineral water sample (0.2 μ g/L bromate in the carbonate- and chloride-rich matrix, spike recovery of 85%)

Water disinfection (e.g., chlorination) is a necessary process to protect us from disease. Unfortunately, it can come with disadvantages like an unpleasant chemical smell and formation of dangerous disinfection byproducts (e.g., carcinogenic trihalomethanes). Although modern technologies like ozonation impart better water flavor, carcinogenic byproducts such as bromate or haloacetic acids can be produced if bromide or other halogens are present in the source water before treatment. Therefore, monitoring drinking water for such disinfection byproducts is of great importance. EU and US EPA regulations set the maximum allowable bromate concentration in drinking water at 10 µg/L. The EPA has attempted to stipulate even lower bromate concentration limits with a maximum contaminant goal of zero for drinking water [5]. For bottled natural mineral and spring waters disinfected by ozone, the EU has reduced the limit of bromate to 3 μ g/L [6]. Regarding wastewater treatment, bromate formation

can become a critical threat for the environment, as treated effluent directly enters rivers and other water sources. Sensitive bromate detection is essential and requires flexibility to be applicable for various matrices as well as the low detection limits.

IC with PCR and UV/VIS detection offers a specific and sensitive method for bromate analysis in line with the requirements of EPA Method 317 and ISO 11206. As this technique is highly flexible, drinking water can be analyzed just as easily as water samples with a high matrix load. Only minor adjustments are necessary for the separation column and the PCR reagents. Additionally, the technique is automated, allowing efficient analysis and a high sample throughput ideal for routine operation. The complete setup can be upgraded with inline sample preparation techniques (e.g., ultrafiltration or dilution), further increasing the method efficiency and broadening the application scope to more complex sample matrices.



REFERENCES

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Internal references: AW IC CH6-1398-052020; AW IC

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CONFIGURATION



947 Professional UV/VIS Detector Vario SW
Le 947 Professional UV/VIS Detector Vario SW, un
détecteur intelligent à longueur d'onde unique,
permet une quantification sure et fiable des
substances actives dans le domaine ultraviolet ou
visible. Une longueur d'onde peut être sélectionnée.









Metrosep A Supp 17 - 250/4,0

La Metrosep A Supp 17 - 250/4,0 combine des performances de séparation élevées et un rapport performances/prix avantageux, sans nécessiter de four à colonne. Le polystyrène-divinylbenzène employé comme matériau de base garantit une longue durée de vie à la colonne. Cette colonne permet de réaliser des tâches de séparation complexes.

930 Compact IC Flex Oven/Deg

Le 930 Compact IC Flex Oven/Deg est un appareil Cl compact intelligent avec un **four à colonne**, **sans suppression** et avec un **dégazeur** intégré. L'appareil peut être utilisé avec n'importe quelles méthodes de séparation et de détection.

Domaines d'application typiques :

- Déterminations d'anions et de cations sans suppression avec détection de conductivité
- Applications simples avec UV/VIS ou détection ampérométrique

Réacteur complet pour 6.2845.100

Réacteur supplémentaire pour plaque de réacteur





Metrosep A Supp 10 - 100/4,0

La colonne de séparation Metrosep A Supp 10 - 100/4,0 est basée sur un copolymère haute capacité de polystyrène-divinylbenzène dont la dimension des particules est de seulement 4,6 µm. Cette colonne se caractérise par son nombre élevé de plateaux et sa grande sélectivité. Il est ainsi possible de séparer de manière sure les sulfites et les sulfates sans ajouter de modificateurs organiques aux éluants. Cettes propriétés sont complétées par une grande flexibilité en ce qui concerne la température de la colonne, le débit et la composition de l'éluant.

Sa construction robuste, son excellent rapport performance-prix et sa très bonne performance de séparation, tout en présentant des temps de chromatographie modérés, font de la Metrosep A Supp 10 - 100/4,0 une colonne de séparation pour anions d'utilisation universelle.

