



Application Note AN-CIC-033

Monitoring PFASs in water sources

Determining adsorbable organically bound fluorine (AOF) in aqueous matrices according to U.S. EPA Method 1621

Per- and polyfluorinated alkyl substances (PFASs), a group of thousands of organic molecules, are widely used in different industries (e.g., as surfactants for film-forming foams or as impregnating agents for packaging) [1,2]. Due to their extreme persistence, they are called «forever chemicals», accumulating in the environment and biomagnifying [3]. Negative health impacts have forced governmental and standardization bodies to take action against the most harmful PFASs. The determination of the non-targeted sum parameter AOF (adsorbable organic fluorine, also called adsorbable organically bound fluorine) is an easy, straightforward way to screen for

PFASs. AOF is a sum parameter covering a broad spectrum of organofluorines. Based on the same principle, i.e., adsorption of organic fluorines on a carbon cartridge, pyrohydrolytic combustion, and determination of fluorine by ion chromatography, the standardization bodies U.S. EPA (U.S. EPA Method 1621), DIN (DIN 38409-59), and ISO (ISO 18127) developed appropriate analytical approaches to estimate a screening level for «total» PFASs in aqueous matrices. This Application Note focuses on the described analytical approach for AOF analysis by combining **pyrohydrolytic combustion** and **ion chromatography (CIC)**.

SAMPLE AND SAMPLE PREPARATION

Three different aqueous environmental samples—one surface water and two waste waters—were analyzed for their AOF content, as described below.

In contrast to other adsorbable organically bound halogens (i.e., chlorine – AOCl, bromine – AOBr, and iodine – AOI), it is crucial for the determination of AOF that the samples have a neutral pH to prevent absorption of any inorganic fluorine. Therefore, each 100 mL sample was pretreated with 0.5 mL of a 2 mol/L sodium nitrate solution.

Organofluorine adsorption was achieved on activated carbon as an automated sample preparation step

(APU sim, Analytik Jena). Automation standardizes the preparation method resulting in excellent repeatability and allows for a high sample throughput. In this step, two carbon cartridges connected in series are flushed with 100 mL sample with a flow rate of 3 mL/min. After adsorption, the two carbon cartridges are washed with 25 mL of a 0.01 mol/L sodium nitrate solution at a flow rate of 3 mL/min. After finishing the sample preparation, the complete content of the two cartridges is transferred into two separate ceramic boats for analysis by CIC.

EXPERIMENTAL

The activated carbon containing all adsorbable organically bound fluorine is analyzed by pyrohydrolytic combustion. The CIC system consists of an autosampler for solid samples, a combustion module, an absorber module, and an ion chromatograph (IC) (Figure 1).

The autosampler automatically transfers the ceramic sample boats into the combustion module, where they are combusted at 1050 °C. With the gas stream,

volatilized fluorine (next to other halogens and sulfur) is transferred into the 920 Absorber Module and absorbed into the aqueous phase. Precise, automated liquid handling is done using Dosinos to transfer the aqueous sample into the IC (930 Compact IC flex) for analysis. To keep the background and limits of detection of fluorine low, it is essential to use chemical reagents which are at least of the purity grade «per analysis».



Figure 1. Combustion IC setup consisting of a 930 Compact IC flex (2.930.2560), a 920 Absorber Module (2.920.0010), a Combustion Module (Oven + ABD, 2.136.0700), and an MMS 5000 Autosampler (2.136.0800) configured for solid samples (6.7302.000).

The separation of fluoride (retention time 6.2 minutes) from other halogens is achieved on a Metrosep A Supp 5 - 250/4.0 separation column in combination with the A Supp 5 Guard/4.0 (**Figure 2**). Automated eluent production with the 941 Eluent Production Module enables continuous and almost unattended operation of the CIC, increasing the overall performance and analysis efficiency. The calibration (0.01–0.5 mg/L) was performed automatically from one standard solution (sodium

fluoride, 0.5 mg/L) by applying the Metrohm intelligent Partial-Loop Injection Technique (MiPT). By injecting one standard with different injection volumes (4–200 µL), a calibration range of 0.01–0.5 mg/L was achieved.

The method detection limit and the method performance were checked with standardized reference materials (4-fluorobenzoic acid) and blanks (ultrapure water), prepared in the same way as the samples and analyzed for their AOF content.

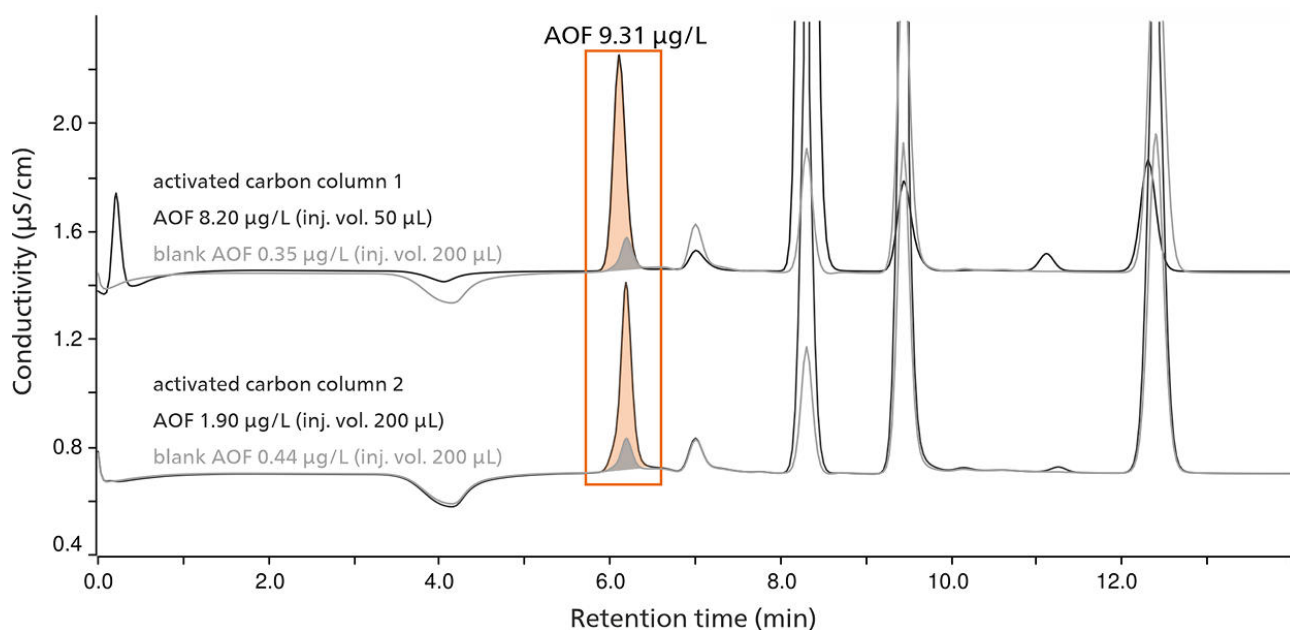


Figure 2. Chromatograms for a wastewater sample. An AOF concentration of 7.85 µg/L was found on the first carbon column and 1.46 µg/L on the second carbon column. This adds up to a total AOF concentration of 9.31 µg/L for this sample. This is the result after blank subtraction. The respective AOF blanks are also shown in grey.

The final sample concentrations are calculated according to the formula below. Thereby the final AOF concentration is the sum of the content

measured for the two subsequent cartridges after blank subtraction (Figure 2).

$$c(AOF) = \left(c(F^-)_{IC} * \frac{V_{Abs}}{V_{Smpl}} \right) - \left(c(F_{BW}^-)_{IC} * \frac{V_{AbsBW}}{V_{SmplBW}} \right)$$

$c(AOF)$	Mass concentration of AOF in µg/L
$c(F^-)_{IC}$	Fluoride concentration in the sample's absorption solution in µg/L
V_{Abs}	Final volume of the absorption solution in L
V_{Smpl}	Volume of the sample that was used for adsorption in L
$c(F_{BW}^-)_{IC}$	Fluoride concentration in the absorption solution of the blank in µg/L
V_{AbsBW}	Final volume of the absorption solution of the blank in L
V_{SmplBW}	Volume of the blank solution that was used for adsorption in L

RESULTS

All samples were analyzed in replicates ($n = 4$). All waters contained trace concentrations of AOF ranging from an average of 6.52 µg/L to 9.70 µg/L, with lower concentrations found in surface water compared to wastewater (Table 1). Although concentrations of

AOF are generally low and sample preparation can be complex, the automation of sample processing and the analysis guarantees excellent repeatability. For the replicates, RSDs of 3.6–5.3% were achieved ($n = 4$). For routine analysis, the method blank was

determined to be 1.1 µg/L for AOF (based on ultrapure water and including all sample preparation

and combustion steps).

Table 1. Results of the AOF analyses for surface water and wastewater samples. The table shows AOF results for the four measured replicates of each sample, the average and standard deviation (SD), and the relative standard deviation (RSD) as determined with the formula shown above. The AOF concentrations are corrected for the blank content.

Sample	AOF #1 (µg/L)	AOF #2 (µg/L)	AOF #3 (µg/L)	AOF #4 (µg/L)	Average ± SD (µg/L)	RSD (%)
Surface water	6.26	6.27	6.79	6.77	6.52±0.30	4.6
Wastewater 1	10.23	4.56	9.31	9.21	9.70±0.51	5.3
Wastewater 2	7.36	6.99	7.61	7.21	7.29±0.26	3.6

CONCLUSION

Determination of the sum parameter **AOF** by adsorption of organic fluorine, pyrohydrolytic combustion, and subsequent fluorine determination by ion chromatography as described in **U.S. EPA 1621**, **DIN 38409-59**, and **ISO 18127** enables a fast and reliable way for **screening PFASs** in various water samples. Ideal for monitoring, this approach can serve as a supplementary method to the comprehensive,

time-consuming, and expensive targeted analysis of PFASs by e.g., LC-MS/MS [4]. With the possibility of automated sample preparation in combination with fully automated analysis by CIC, this is an easy, reliable, economic, time-saving, and straightforward technique for routine AOF analysis and estimation of «total» PFASs.

REFERENCES

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CONFIGURATION



930 Combustion IC PP (AJ)

The 930 Combustion IC PP (AJ) enables the analysis of halogens and sulfur in flammable samples of all types using inline combustion digestion (pyrohydrolysis) with subsequent ion chromatography determination (Combustion IC). It comprises all required components, e.g., the Combustion Module from Analytik Jena (2.136.0700), the 920 Absorber Module, the 930 Compact IC Flex Oven/SeS/PP/Deg, and the MagIC Net software. If necessary, the 930 Metrohm Combustion IC package can be supplemented with an Autosampler for solid or liquid samples (MMS 5000 Autosampler). The entire analysis, including sample input and sample digestion, is completely automated and is completely controlled by MagIC Net.



Metrosep A Supp 5 Guard/4.0

The Metrosep A Supp 5 Guard/4.0 reliably protects the Metrosep A Supp 5 and 7 IC anion columns against contamination from the sample or the eluent.

It contains the same separation material as the Metrosep A Supp 5, is also made of PEEK, and is screwed directly onto the respective separation column with virtually no dead volume ("On Column Guard System"). The guard column prolongs the service life of the analytical column, with practically no influence on its chromatographic separating efficiency. The economical price and simple handling make using the A Supp 5 Guard/4.0 highly recommended.



Metrosep A Supp 5 - 250/4.0

The high-performance separation column from Metrohm with an extremely high number of plates for the most demanding separation tasks. Even complex separation problems can be solved easily and reproducibly with the Metrosep A Supp 5 - 250/4.0. The high capacity of the column allows, for example, the detection of 1 µg/L bromate along with 150 mg/L chloride without sample preparation. The range of applications possible with this column far exceeds the detection of standard anions. The Metrosep A Supp 5 - 250/4.0 is the column of choice when it comes to reliable monitoring of the high purity standards in the semiconductor industry or of the boiler feed water of power plants.



930 Compact IC Flex Oven/SeS/PP/Deg

The 930 Compact IC Flex Oven/SeS/PP/Deg is the intelligent Compact IC instrument with **column oven**, **sequential suppression**, **peristaltic pump** for suppressor regeneration and built-in **degasser**. The instrument can be used with any separation and detection methods.

Typical areas of application:

- Anion or cation determinations with sequential suppression and conductivity detection



920 Absorber Module

The 920 Absorber Module combines the Combustion Module with the ion chromatograph. The 920 Absorber Module ensures that the gaseous compounds of the analytes are dissolved and channeled to the IC. It is responsible for the entire Liquid Handling. In addition to Combustion IC, it can also be used for gas analysis.



MMS 5000 Autosampler (AJ)

Autosampler MMS 5000 (AJ) made by Analytik Jena for use with the Metrohm Combustion IC for fully automatic analysis of liquid and solid samples. To adapt the modular Multi-Matrix sampler to the correct sample type, either the liquids kit (6.7303.000) or the solids kit (6.7302.000) must be used.