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-0.5

Introduction

Water naturally contains differing levels of chloride, bromide, and iodide ions. These ions are typically not harmful to human health at reasonable levels; however, these halides can be incorporated into organic structures when water containing these ions is treated using oxidative processes such as chlorination, ozonation, and chloramination[1, 2]. The oxidative conditions employed in the treatment process can convert these halides into "activated" forms, which can then react with organics present in wastewater to form halogenated disinfection byproducts (DBPs)[3], where many species are largely unidentified[4]. While some DBPs are regulated in drinking water by the United States EPA, the regulated DBPs are a small list of identifiable compounds[5]. Given that iodinated and brominated DBPs are more toxic than their chlorinated analogs[6-8], it is prudent that occurrence data is acquired quickly in order to better understand the magnitude and prevalence of these emerging DBPs.

Most analytical methods such as EPA method 551.1 employ GC with electron capture detection (GC-ECD) for determining the concentrations of volatile halogenated DBPs in extracts prepared from waters. These methods are limited in their ability to differentiate halogenated species from each other, or from other interfering nonhalogenated species. Indeed, methods do exist for determining these species by GC-MS or GC-MS/MS. Unfortunately these forms of molecular MS are limited in their ability to screen a wide array of unidentified molecules for halogen content in a single analysis given due to ionization issues. Chemical Ionization (CI) does not ionize all types of organic molecules, and electron impact (EI) can lead to excessive, undesirable insource fragmentation. In addition, GC-MS/MS analysis is limited by the inefficient fragmentation of halogenated precursor ions into monatomic halogen product ions (i.e., Br⁺ and Br⁻). In this poster, we describe our work in using GC-ICP-MS to examine and quantitate the presence of these halogenated DBPs.

Instrumentation

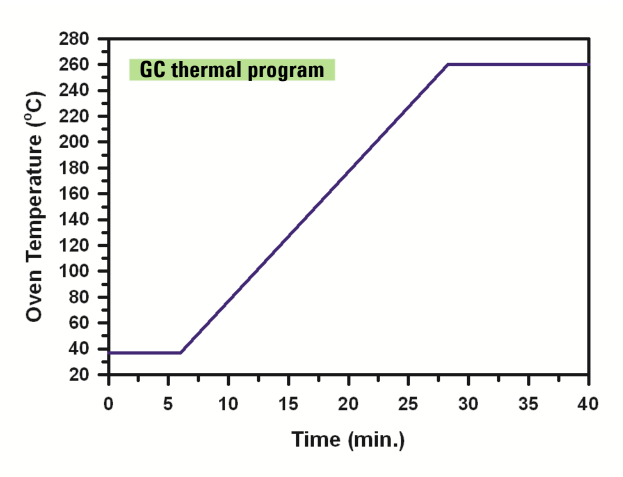
Sample preparation

Municipal wastewater samples were collected from geographically separated areas. Samples were split in two, with one half untreated and the other half treated with aqueous monochloramine. For extraction, 35 mL of these wastewater samples were extracted using 5 mL of MTBE in a modified version of EPA method 551.1. The organic layers were carefully separated and then placed into 2.0 mL amber Agilent GC vials.

Instrument Configuration and Conditions

Gas Chromatograph:

- Agilent 7890A w/ heated ICP-MS transfer line & injector
- Agilent 30 m HP-5 column (320 μm x 0.25 μm)
- 200 °C inlet & 260 °C transfer line/injector temperatures
- Pulsed splitless inection (10 psi until 0.75 min, 5.8 psi afterwards)
- Oven 37 °C for 6 min, then 10 °C/min rise to 260 °C, then hold for 11 min.



ICP-MS:

- Agilent 7700x ICP-MS operated in No gas mode
- 3.0 mm sample depth, RF power 700 W
- 0.4 L/min dilution gas (Ar) delivered to transfer line
- 0.15 second integrations of m/z 79, 81, and 127
- Calibration standards prepared in MTBE using 1-bromo-4iodobenzene (0, 1, 2, 5, 10, 25, 100 ng/mL)

Element-specific examination of volatile halogenated organics in wastewater extracts using GC-ICP-MS. Armando Durazo and Shane A. Snyder

Results and Discussion Analysis of Calibration Standards & Samples Using the method described previously, 1-bromo-4-iodobenzene elutes from the column at 20.5 minutes. We have been able to detect iodine in all non-zero calibration standards and bromine in all standards with (compound) concentrations above 5 ng/mL Conc. (ppb) Calc. Conc. Counts Conc. (ppb) Carc. Conc. Counts 1 1.1 190.6 2 1.7 303.8 5 5.6 971.0 10 9.9 1717.0 25 26.5 4605.0 100 99.6 17341.0

The chloramination of these samples causes clear and profound changes to occur in the concentration and speciation of halogenated organics. The total volatile organohalogen levels increase upon chloramination(as evidenced below) and there is an increased presence of more volatile I-DBPs.

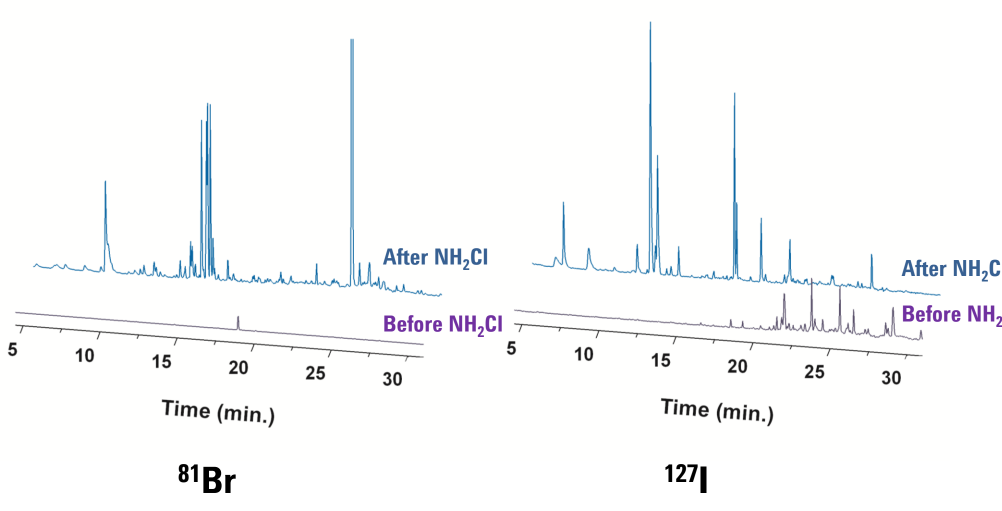
Bromine lodine (pp

We have succeeded in using an Agilent 7890A coupled to an Agilent 7700x ICP-MS to determine the presence, transformation, and formation of halogenated DBPs in wastewaters that are treated by chloramination. Our instrument configuration allows for easy tuning, chromatographic separation of an array of diverse compounds, and interference-free, high sensitivity analysis After NH₂Cl for halogens covalently bound within organic molecules. The use of CICs allows us to quantitate the halogen contents of Before NH₂CI these compounds based on the responses obtained from a commercially available dihaloaromatic. An added benefit to using GC-ICP-MS for the analysis of these complex mixtures of halogenated organics is the element-specificity that our **ICP-MS** platform provides over other halogen detection methodologies like GC-ECD. In the near future, we aim to identify these DBPs using GC-QToF and to investigate water treatment technologies that will minimize the formation of iodinated and brominated DBPs.

Wastewater samples were then examined before and after treatment with monochloramine. The reaction with monochloramine leads to an increase in the concentration of brominated and iodinated species in the extracts. These data reveals several interesting facts. First, there are volatile halogenated organics present in wastewaters prior to chloramination and some of these species are resistant to transformation upon monochloramine treatment while others are consumed (and likely transformed into new halogenated DBPs). Thus it is likely that many nonhalogenated organics in untreated wastewaters are converted into new halogenated DBPs, as well.

0.5 1.0 1.5 2.0

Log₁₀ ([Brl-benzene], ppb)





Agilent Technologies

WP 340

Summary of Results

The results of our study indicate that the monochloramination of wastewater samples does indeed dramatically change the concentration and speciation of halogenated volatile organics in these waters.. The effects of monochloramination are seen most profoundly in terms of the differences between chromatograms for brominated and iodinated DBPs. There are two reinforcing explanations for this, one dealing with the reactivity of bromide and iodide during oxidative treatments, and the higher sensitivity for detection for I and Br in our assays due to their elemental ionization potentials.

	Blank Extract	Before CINH ₂	After CINH ₂
(ppb)	103	1134	98229
pb)	93	490	777

Conclusions

References

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- 4. Jeong, C.H., et al., Occurrence and Toxicity of **Disinfection Byproducts in European Drinking** Waters in Relation with the HIWATE Epidemiology Study. Environmental Science & Technology, 2012. 46(21): p. 12120-12128.
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- 6. Krasner, S.W., et al., Occurrence of a New Generation of Disinfection Byproducts. Environmental Science & Technology, 2006. 40(23): p. 7175-7185.
- 7. Smith, E.M., et al., Comparison of Byproduct Formation in Waters Treated with Chlorine and Iodine: Relevance to Point-of-Use Treatment. **Environmental Science & Technology, 2010. 44(22):** p. 8446-8452.
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For More Information

Please see Agilent publication 5991-4398EN, which is available at www.agilent.com/chem