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Author manuscript

Environ Sci Technol Lett. Author manuscript; available in PMC 2020 January 06.

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Published in final edited form as:

Environ Sci Technol Lett. 2019 ; 6(11): 662–668. doi:10.1021/acs.estlett.9b00525.

Fate of Per- and Polyfluoroalkyl Ether Acids in the Total Oxidizable Precursor Assay and Implications for the Analysis of Impacted Water

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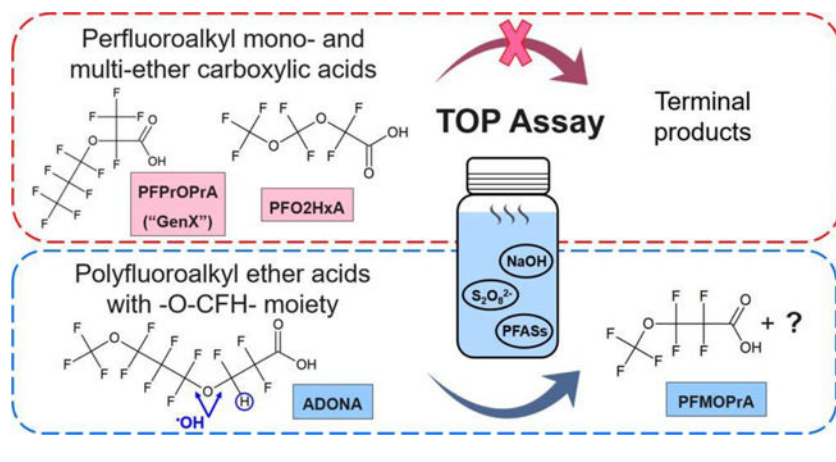
Abstract

Per- and polyfluoroalkyl substances (PFASs) are widely used anthropogenic chemicals. The PFAS class includes almost 5000 registered compounds, but analytical methods are lacking for most PFASs. The total oxidizable precursor (TOP) assay was developed to indirectly quantify unknown PFASs that are precursors to commonly measured perfluoroalkyl acids. To understand the behavior of recently identified per- and polyfluoroalkyl ether acids (PFEAs), including fluorinated replacements and manufacturing byproducts, we determined the fate of 15 PFEAs in the TOP assay. Ten perfluoroalkyl ether acids and a chlorinated polyfluoroalkyl ether acid (F-53B) were stable in the TOP assay and represent terminal products that are likely as persistent as historically used PFASs. Adding perfluoroalkyl ether acids and F-53B to the target analyte list for the TOP assay is recommended to capture a higher percentage of the total PFAS concentration in environmental samples. In contrast, polyfluoroalkyl ether acids with a -O-CFH- moiety were oxidized, typically to products that could not be identified by liquid chromatography and high-resolution mass spectrometry. Application of the TOP assay in its proposed enhanced form revealed high levels of PFEAs, the presence of precursors that form perfluoroalkyl carboxylic acids, and the absence of precursors that form PFEAs in surface water impacted by PFAS-containing wastewater discharges.

Graphical Abstract

Supporting Information

Tables, figures, experimental details about the TOP assay, SPE and LC-MS analysis, and PFAS quantification



INTRODUCTION

Per- and polyfluoroalkyl substances (PFASs) are extensively used anthropogenic chemicals. The unique properties of PFASs have led to a variety of applications, including their use in firefighting foams, stain repellents, paper coatings, and fluoropolymer production.^{1–7} To date, almost 5000 PFASs have been registered.⁸ Two long-chain PFASs, perfluorooctanesulfonic acid (PFOS) and perfluorooctanoic acid (PFOA), were historically produced in large quantities, but a manufacturing shift toward short-chain PFASs and fluorinated alternatives started in the early 2000s as a result of links between long-chain PFAS exposure and adverse health outcomes.^{4,9–12}

Several per- and polyfluoroalkyl ether acids (PFEAs) have been produced to replace PFOS and PFOA. For example, ammonium 4,8-dioxo-3H-perfluorononanoic acid [ADONA (Table 1)] was developed to replace the ammonium salt of PFOA as an emulsifier in the manufacture of fluoropolymers.¹³ It was detected downstream of a fluoropolymer manufacturing plant in Germany in 2008.^{14,15} As an alternative to PFOS, 6:2 chlorinated polyfluoroalkyl ether sulfonic acid [6:2 Cl-PFESA, trade name F-53B (Table 1)] is widely used in Chinese chromium-plating facilities.¹⁶ After decades of F-53B usage in China, it has been detected in Chinese surface water, sewage sludge, wildlife, and humans.^{17–21} The ammonium salt of hexafluoropropylene oxide-dimer acid [HFPO-DA (Table 1)] with the trade name “GenX” has been used as a processing aid to replace the ammonium salt of PFOA in fluoropolymer manufacturing processes since 2010.¹⁴ The presence of GenX was reported in river water downstream of fluorochemical plants in China,²² Germany,²² The Netherlands,²³ and the United States (U.S.).^{4,24} The U.S. study further showed that manufacturing byproducts, which are generated during the production of fluoropolymer building blocks, were discharged into an important drinking water supply. These byproducts can be broadly classified as (1) perfluoroalkyl monoether carboxylic acids (monoether PFECAs), (2) perfluoroalkyl multiether carboxylic acids (multiether PFECAs), and (3) polyfluoroalkyl ether acids (Table 1, Table S1, and Figure S1).^{4,24–26}

Analytical methods and reference standards are lacking for the vast majority of PFASs.²⁷ For example, U.S. EPA Method 537.1 targets only 18 PFASs.²⁸ Approaches for quantifying total PFAS concentrations in environmental samples are therefore needed.²⁷ The total

oxidizable precursor (TOP) assay was developed to indirectly quantify the concentration of perfluoroalkyl acid (PFAA) precursors, i.e., PFASs, that may degrade in the environment to terminal products such as perfluoroalkyl carboxylic and sulfonic acids (PFCAs and PFSAs, respectively).²⁹ To date, the TOP assay has been applied to characterize wastewater,^{30,31} urban runoff,^{29,32} precipitation,³³ groundwater,^{34–36} aqueous film-forming foams,³⁷ paper, and textiles.^{38,39} However, no information about the fate of PFEAs in the TOP assay is available. It is unclear whether PFEAs are PFAA precursors in the TOP assay or whether some PFEAs are terminal products that need to be added to the analyte list for the TOP assay. Similarly, it is unknown whether the TOP assay can uncover the presence of PFEAs and/or PFEA precursors in environmental samples. The aims of this research were, therefore, (1) to determine the fate of 15 PFEAs in the TOP assay, (2) to identify possible new terminal products of the TOP assay, and (3) to apply the TOP assay to surface water impacted by a wastewater discharge containing PFEAs.

MATERIALS AND METHODS

Materials

Twenty-five PFASs in three classes (i.e., 7 PFCAs, 3 PFSAs, and 15 PFEAs) were targeted on the basis of the availability of analytical standards and the significance to the Cape Fear River (CFR) watershed in North Carolina (Table S1 and Figure S1). Analytical standards and isotopically labeled internal standards were obtained from Wellington Laboratories (Guelph, ON), Fluoryx Laboratories (Carson City, NV), The Chemours Company (Wilmington, DE), and SynQuest Laboratories (Alachua, FL) (see Table S1 for details). All other chemicals and solvents were purchased from Fisher Scientific (Hampton, NH) and Sigma-Aldrich (St. Louis, MO).

Sample Collection

CFR water samples were collected at the William O. Huske Lock and Dam (HLD) immediately downstream from a fluorochemical manufacturing site in August 2014 and April 2018, as well as at a drinking water intake (Lock and Dam #1 or LD1) located approximately 140 km downstream from the manufacturer in July 2015 (Figure S2). River discharges on the sampling dates are depicted in Figure S3. At each site, samples were collected in 20 L high density polyethylene (HDPE) carboys and stored at 4 °C until use. Details of water sample collection have been described previously.⁴

Total Oxidizable Precursor (TOP) Assay

TOP assays were conducted according to a previously developed protocol.^{29,35} For experiments using individual PFEAs, an aliquot of a 5 ng/μL individual PFEA stock solution in methanol was added to each HDPE bottle and evaporated to dryness under nitrogen. The PFEA was subsequently dissolved in 125 mL of deionized water to yield a starting concentration of ~1000 ng/L, and the solution was preheated for 30 min at 60 °C. For experiments involving CFR water, 125 mL aliquots were added directly to HDPE bottles. To initiate precursor oxidation, potassium persulfate (5 mM) and sodium hydroxide (150 mM) were added. Samples were heated in a temperature controlled water shaker bath for 6 h at 85 °C. The pH of each sample was measured to confirm a pH of >12 was maintained during

the TOP assay. At elevated temperatures, persulfate is converted to sulfate radicals ($\text{SO}_4^{\bullet-}$), which, at elevated pH values, rapidly react to form hydroxyl radicals ($^{\bullet}\text{OH}$).^{29,35} Upon reaction, samples were cooled, neutralized to pH 5–9²⁹ with nitric acid,⁴⁰ and stored at 4 °C until offline solid phase extraction (SPE) and liquid chromatography–high-resolution mass spectrometry (LC–HRMS) analysis. For each sample, PFAS concentrations were measured before and after the TOP assay. TOP assay method validation and control experiments are described in the Supporting Information (Texts S1 and S2).

Analytical Methods

SPE using Oasis WAX Plus cartridges and LC–MS analysis were carried out as previously described.^{24,41} While sample preparation by SPE may have led to low bias for or the loss of some oxidation products (e.g., neutral products), SPE was used to avoid the injection of large quantities of salt from the TOP assay into the LC instrument. All PFAS concentrations were determined using authentic standards. When possible, an isotope dilution approach was used, in which the analyte response was normalized to that of an isotopically labeled analogue.²⁵ For other PFASs, the analyte response was normalized to that of an isotopically labeled PFAS with an LC retention time similar to that of the analyte (Table S1). Details of the analytical methods, including approaches to identifying reaction products by LC–HRMS, are provided in the Supporting Information (Texts S3–S7).

RESULTS AND DISCUSSION

Fate of Perfluoroalkyl Ether Carboxylic Acids

The fate of 10 perfluoroalkyl ether carboxylic acids [PFECAs, including six monoether PFECAs and four multiether PFECAs (Table S1)] in the TOP assay was evaluated. For the six monoether PFECAs, concentrations before and after oxidation showed no statistically significant difference (t test; $n = 2$; $p < 0.05$); i.e., monoether PFECAs were stable in the TOP assay and represent new terminal products (Figure 1 and Table S2). For monoether PFECAs, both linear and branched isomers (PFMOPrA and PMPA; PFMOBA and PEPA) were persistent during the TOP assay, implying that branching does not change the stability of monoether PFECAs in the TOP assay. Similarly, the four multiether PFECAs with repeating $-\text{CF}_2\text{O}$ subunits were stable during the TOP assay ($p < 0.05$, except for PFO2HxA, for which the duplicate was lost), illustrating that they also represent new terminal products that are likely persistent in the environment.

TOP assay results indicate that PFECAs are resistant to attack by $^{\bullet}\text{OH}$, in agreement with the finding that PFECA levels did not change measurably after ozonation in a full-scale surface water treatment plant.⁴ Ozonation produced $^{\bullet}\text{OH}$ at this plant, as evidenced by oxidation of 1,4-dioxane,⁴² but the oxidation conditions were less severe than those employed here. To date, no study has evaluated the reaction between $^{\bullet}\text{OH}$ and PFPrOPrA (“GenX”), but a recent study indicated that GenX was not oxidized in systems involving both $^{\bullet}\text{OH}$ and $\text{SO}_4^{\bullet-}$.⁴³ Although one rationale behind replacing PFOA and PFOS with PFEAs was to make the replacements more labile by inserting ether linkage(s),²⁴ the TOP assay results highlight that PFECAs are highly resistant to chemical degradation even under extreme oxidative conditions. Because the TOP assay currently employed by most research

and commercial laboratories does not include PFECAs in the analyte list, it would significantly underestimate the total PFAS concentrations in samples containing PFECAs or PFECA precursors. Thus, we propose the addition of PFECAs to the analyte list for an expanded TOP assay, as they represent new terminal products (Table S4).

Fate of Polyfluoroalkyl Ether Acids

To assess the fate of polyfluoroalkyl ether acids in the TOP assay, we conducted experiments with two polyfluoroalkyl ether sulfonic acids (Nafion byproduct 2, NVHOS), two polyfluoroalkyl ether carboxylic acids (ADONA, HydroEVE), and one chlorofluoroalkyl ether sulfonic acid (F-53B) in deionized water (Table S1, Figure S1). These compounds were selected because they have been detected in the environment,^{14,24,26} and they allowed us to assess the effects of different structural features, such as the introduction of a C–H bond adjacent to an ether oxygen, different headgroups (sulfonic and carboxylic acids), and the presence of a C–Cl bond, on the fate of PFASs during the TOP assay.

An alternative to PFOS, F-53B, has two structural modifications, i.e., an ether linkage between C2 and C3 and a fluorine-to-chlorine substitution on C8 (Table 1).⁴⁴ In the TOP assay, F-53B was persistent and represents a new terminal product that should be added to the TOP assay analyte list (Figure 2, Table S2, and Table S4). The TOP assay result is consistent with the reported stability of F-53B in advanced oxidation processes, including ultraviolet light/H₂O₂, ozonation, and O³/H₂O₂.^{44,45}

The PFOA replacement ADONA has two ether linkages and a fluorine-to-hydrogen substitution on C3, next to the first ether bond (Table 1). In the TOP assay, ADONA was oxidized to PFMOPrA (Figure S1), a linear four-carbon PFECA and isomer of PMPA. The molar yield of PFMOPrA was 98 ± 20%, indicating complete conversion of ADONA (Figure 2 and Table S2). Formation of PFMOPrA suggests that the site of hydroxyl radical attack on the ADONA anion is the -O-CFH- moiety. This experimental observation is supported by results of quantum mechanical models, which indicate that H-abstraction of the aliphatic proton of ADONA by a hydroxyl radical is the thermodynamically favored reaction path.⁴⁶ PFMOPrA represents a new terminal product of the TOP assay, further highlighting the need to add PFECAs to the TOP assay analyte list. No additional oxidation product(s) could be identified by LC–HRMS. It is unclear whether additional low-molecular weight fluorinated oxidation products were formed and/or whether a fraction of ADONA was mineralized to fluoride.⁴⁷ Fluoride can be generated from some polyfluoroalkyl substances (e.g., 6:2 fluorotelomer sulfonate) in advanced oxidation processes involving hydroxyl radicals.⁴⁸ In our study, fluoride production could not be measured because our starting ADONA concentration was ~1000 ng/L. We lacked access to sufficient ADONA to complete TOP assays with ADONA concentrations that would have allowed us to quantify fluoride.

The three other polyfluoroalkyl ether acids containing the -O-CFH- moiety [Nafion byproduct 2, NVHOS, and HydroEVE (Figure S1)] were readily oxidized in the TOP assay, implying the acid headgroup (sulfonic or carboxylic acid) does not affect the fate of polyfluoroalkyl ether acids in the TOP assay (Figure 2 and Table S2). It should be noted that transformation of these compounds in the TOP assay does not suggest they are readily

transformed in the environment or in (waste)water treatment, where oxidative conditions are less severe than in the TOP assay. For example, we have observed no measurable change in Nafion byproduct 2 concentrations in a full-scale surface water treatment plant that employs ozonation (unpublished data). No oxidation product(s) could be identified by LC–HRMS, suggesting transformation of these compounds to low-molecular weight organofluorine species, volatile organofluorine species, and/or fluoride.⁴⁷ Because the longest perfluoroalkyl group in the studied polyfluoroalkyl ether acids with -O-CFH- moieties contains three carbon atoms (Figure S1), the longest possible PFAA formed is shorter than PFBA, the shortest PFAA that is typically quantified in the TOP assay. Hence, the results demonstrate an important weakness of the TOP assay; i.e., some compounds are oxidized to products that are not usually identified. The presence of PFASs like Nafion byproduct 2, NVHOS, and HydroEVE would be missed if analytical methods were not to target these compounds prior to oxidation.

Application of the Expanded TOP Assay to Cape Fear River Water

Direct measurement of legacy PFAS (PFCA and PFSA) concentrations in the water sample collected at HLD in August 2014 indicated the presence of five PFCAs and one PFSA, with their total concentrations being 89 and 17 ng/L, respectively (Figure 3A and Table S5). The dominant PFCAs were PFBA (32 ng/L) and PFPeA (22 ng/L), consistent with a previous study.⁴ Upon application of the TOP assay with only the traditionally analyzed PFCAs and PFSAs, the total PFCA concentration increased by 38% on a mass basis, largely as a result of the presence of precursors that form PFPeA (increased by 13 ng/L or 60%) and PFBA (increased by 10 ng/L or 31%). Precursor sources are not known; precursors could be introduced by the fluorochemical manufacturer and/or point or nonpoint sources located further upstream in the CFR watershed. When PFEAs were included in the analyte list (before TOP assay oxidation), the sum concentration of PFEAs (Σ PFEA \sim 990000 ng/L) was nearly 4 orders of magnitude larger than that of the traditionally analyzed PFCAs and PFSAs (Figure 3B and Table S5). The dominant PFEAs were PFMOAA, PFO2HxA, and PFO3OA, consistent with a previous publication.²⁵ The astoundingly high levels of PFEAs were attributed, in part, to a lack of complete mixing of the industrial wastewater with the bulk river water because HLD is located immediately downstream of the fluorochemical manufacturing site. By expanding the TOP assay analyte list to include PFEAs, we observed that three polyfluoroalkyl ether acids (i.e., Nafion byproduct 2, NVHOS, and HydroEVE) were oxidized during the TOP assay while the eight detected PFECAs were stable, consistent with our results for individual PFEAs. Furthermore, the expanded TOP assay results suggest that PFEA precursors were not present at significant levels in our sample. The environmental levels of PFEAs in CFR highlight the importance of adding PFEAs to the analyte list of the TOP assay and targeted LC-MS methods to capture a larger fraction of total PFASs.

In the water sample collected at a drinking water intake (i.e., LD1) located approximately 140 km downstream from the manufacturer, concentrations of PFCAs and PFSAs were 120 and 45 ng/L, respectively (Figure S8 and Table S6). Higher legacy PFAS levels compared to those in the HLD sample were attributed, in part, to a lower river discharge (930 ft³/s at LD1 in July 2015 and 1600 ft³/s at HLD in August 2014). Upon application of the TOP assay, the

PFCA concentration increased by 21% on a mass basis, which was mainly attributed to increases in PFPeA (33%) and PFHxA (24%). When PFEAs were included in the PFAS analyte list (before TOP assay oxidation), PFEAs dominated (Σ PFEA \sim 130,00 ng/L), accounting for >99% of the total quantified PFAS mass concentration. Lower PFEA concentrations compared to those in HLD samples were primarily explained by the complete mixing of wastewater discharge from the fluorochemical manufacturer. As was the case for the HLD sample, PFEA precursors were not found in the LD1 sample.

Only PFMOAA (92 ng/L) was detected in the water sample collected at HLD in April 2018, and its concentration changed little in the TOP assay (Figure S9 and Table S7). Low PFAS levels were explained by (1) high streamflow (14000 ft³/s) and (2) the fluorochemical manufacturer beginning to capture PFEA-containing process wastewater starting in mid-June 2017.²⁵

To the best of our knowledge, this is the first paper describing (1) the fate of PFEAs in the TOP assay and (2) the application of the TOP assay with 15 PFEAs added to the analyte list to impacted surface water. Results highlight that current targeted approaches (e.g., EPA Method 537.1) and a TOP assay that considers only PFCAs and PFSAs as terminal products can miss a large fraction of the total PFAS concentration in environmental samples. To more comprehensively characterize the occurrence of PFASs, we recommend a suite of complementary analytical approaches, including LC–HRMS and gas chromatography (GC)–HRMS, supplemented by “total organofluorine” methods such as the TOP assay with an expanded analyte list to include PFEAs, extractable organic fluorine (EOF), and/or adsorbable organic fluorine (AOF).^{27,38,49} Such approaches are needed to assess, for example, whether PFEAs and/or precursors to PFEAs occur primarily near manufacturing sites or whether they occur more widely.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

This work was supported in part by the North Carolina Policy Collaborative. This document has been reviewed by the U.S. Environmental Protection Agency, Office of Research and Development, and approved for publication. The views expressed in this article are those of the authors and do not necessarily represent the views or policies of the U.S. Environmental Protection Agency.

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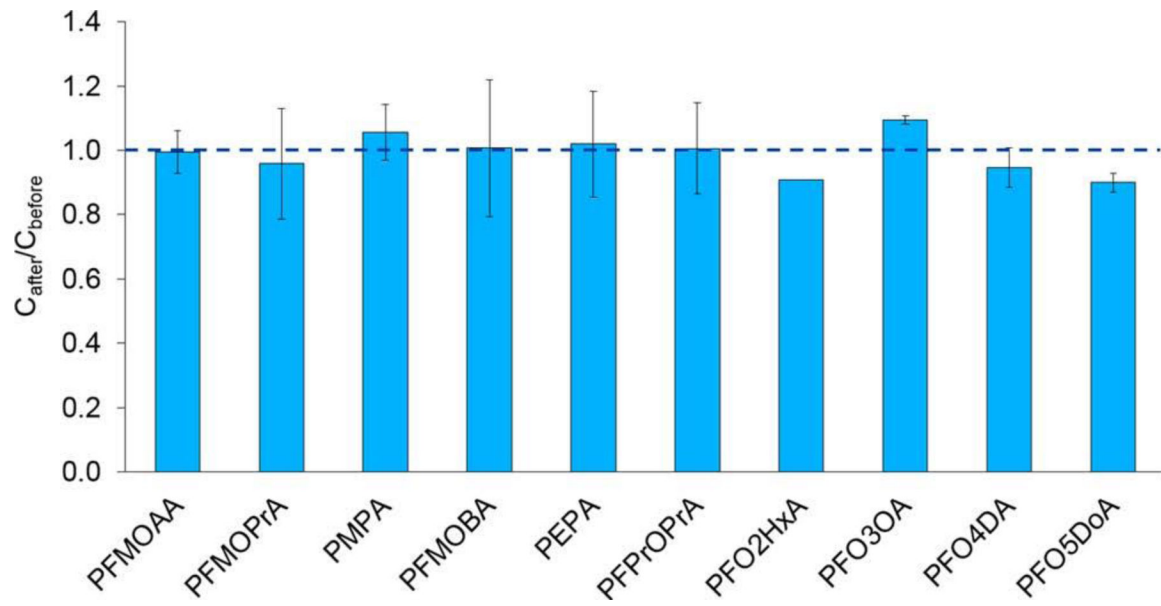


Figure 1. Fractions of perfluoroalkyl ether carboxylic acid (PFECA) concentrations remaining after oxidation relative to that measured before oxidation in the TOP assay. Averages and standard deviations of duplicate experiments are shown. See Figure S1 for compound structures.

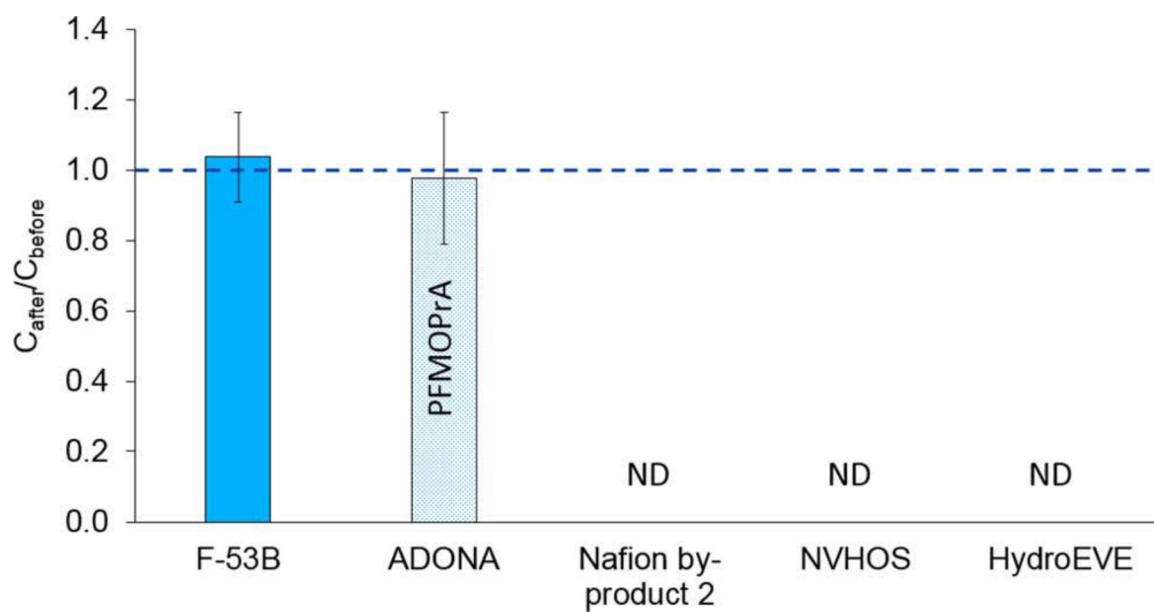


Figure 2.

Fractions of polyfluoroalkyl ether acid concentrations remaining after oxidation relative to that measured before oxidation in the TOP assay. Averages and standard deviations of duplicate experiments are shown. In the TOP assay, ADONA was oxidized to PFMOPrA. ND means no product or intermediate was detected by LC–HRMS in the sample after oxidation.

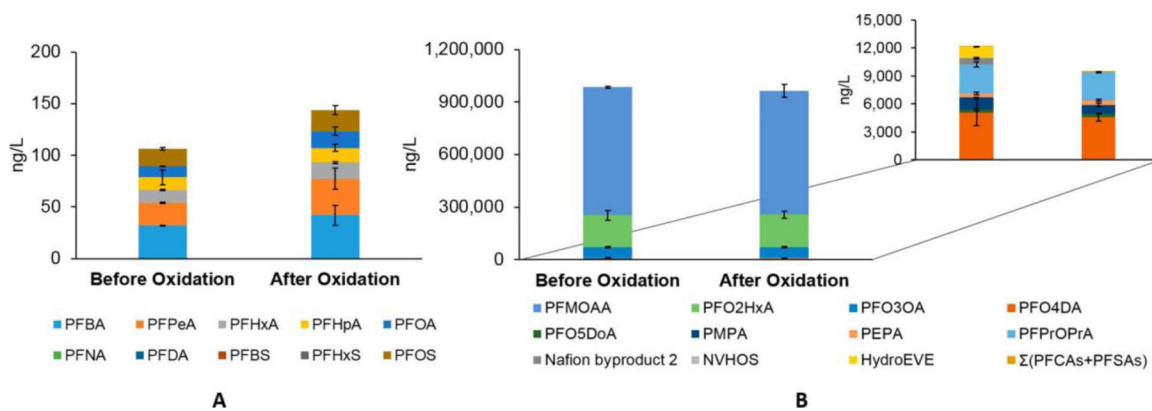
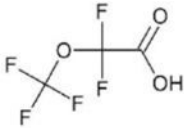
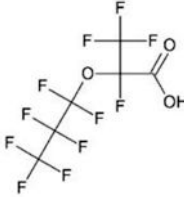

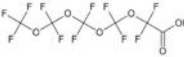
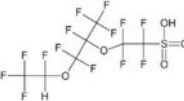



Figure 3. Application of the TOP assay (A) with traditionally analyzed PFCAs and PFSA (ng/L) and (B) with an expanded PFEA analyte list to a water sample collected at the William O. Huske Lock and Dam (HLD) in August 2014. The inset in panel B highlights the concentrations of PFASs other than PFMOAA, PFO2HxA, and PFO3OA. Averages and standard deviations of duplicate experiments are shown. The dissolved organic carbon (DOC) concentration was 9.5 mg/L.

Table 1.

Examples of Per- and Polyfluoroalkyl Ether Acids (PFEAs) Targeted in This Study

Compound	Formula	CAS#	Molecular structure
Perfluoroalkyl mono-ether carboxylic acids (mono-ether PFECAs)			
Perfluoro-2-methoxyacetic acid (PFMOAA)	$C_3HF_5O_3$	674-13-5	
Perfluoro-2-propoxypropanoic acid (PFPrOPrA) = Hexafluoropropylene oxide-dimer acid (HFPO-DA) = parent acid of "GenX"	$C_6HF_{11}O_3$	13252-13-6	
Perfluoroalkyl multi-ether carboxylic acids (multi-ether PFECAs)			
Perfluoro(3,5-dioxahexanoic) acid (PF02HxA)	$C_4HF_7O_4$	39492-88-1	
Perfluoro(3,5,7,9-tetraoxadecanoic) acid (PF04DA)	$C_6HF_{11}O_6$	39492-90-5	
Polyfluoroalkyl ether acids			
Ethanesulfonic acid, 2-[1-(difluoro(1,2,2,2-tetrafluoroethoxy)methyl)-1,2,2,2-tetrafluoroethoxy]-1,1,2,2-tetrafluoro- = "Nation byproduct 2"	$C_7H_2F_{14}SO_5$	749836-20-2	
4,8-dioxa-3H-perfluorononanoic acid = parent acid of "ADONA"	$C_7H_2F_{12}O_4$	919005-14-4	
9-chlorohexadecafluoro-3-oxanone-1-sulfonic acid (9C1-PF30NS, main component of F-53B)	$C_8HF_{16}SO_4Cl$	756426-58-1	