

EPA Method 533:

Determination of Per- and Polyfluoroalkyl Substances in Drinking Water via Anion-Exchange SPE and LC-MS/MS

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INTRODUCTION

Per- and polyfluoroalkyl substances (PFAS) are a diverse group of synthetic organofluorine compounds that are widely used in industrial applications and consumer products. PFAS are persistent in the environment, are resistant to degradation, and are known to bioaccumulate in humans and wildlife. EPA 533 measures PFAS by isotope dilution anion-exchange solid-phase extraction (SPE) and liquid chromatography/tandem mass spectrometry (LC-MS/MS). In total, EPA 533 covers 25 PFAS. Furthermore, it includes the use of 16 isotopically labeled Isotope Dilution Standards and 3 Isotope Performance Standards to ensure optimum method performance. In light of the EPA's recent announcement on the fifth Unregulated Contaminant Monitoring Rule (UCMR5), which includes 29 PFAS, the use

SPE METHOD

a) Rinse SPE cartridge (ECWAX156-P) with 10mL methanol.

of EPA 533 is set to become increasingly important to water testing labs [4].

- b) Rinse the cartridge with 10 mL of 0.1M pH 7 phosphate buffer, being sure to not allow the water to drop below the top edge of the packing.
- c) Close the valve and add 2–3 mL of phosphate buffer to the cartridge reservoir and fill the remaining volume with reagent water.
- Note: Do not allow cartridge packing material to go dry during any of the conditioning steps. If the cartridge goes dry during the conditioning phase, the conditioning must be repeated.

Sample Extraction/Drying

- a) Attach a large volume sample transfer tube (VMFSTFR06-PFC) to the top of each SPE cartridge and place the stainless-steel end of the transfer tube directly into the sample
- b) Adjust the vacuum so that the flow rate is approximately 5 mL/min. Flow rates above 5 mL/min during loading may cause low analyte recovery.
- c) After the entire sample has passed through the cartridge, rinse the sample bottle with 10 mL of 1 g/L ammonium acetate in reagent water. Draw the rinsate through the sample transfer tubes and the cartridges.
- d) Add 1 mL of methanol to the sample bottle and draw through the transfer tubes and SPE cartridges. This step is designed to remove most of the water from the transfer line and cartridge resulting in the reduction of the salt and water present in the eluate. e) Dry the cartridge under high vacuum (15-20 in.Hg) for 5 minutes.

- a) Insert a collection rack containing 15 mL polypropylene collection tubes into the extraction manifold (VMF016GL-PFAS).
- b) Add 5 mL of methanol containing 2% NH4OH (v/v) to the sample container, cap and thoroughly rinse the sides with the elution solvent.
- Note: Due to the volatility of NH4OH, it is highly recommended to use fresh elution solvent. Rinsing sides of container is important for obtaining good recovery of the long-chain hydrophobic PFAS.
- c) Elute the analytes from the cartridges by pulling the elution solvent through the sample transfer tubes and the cartridges. Use a low vacuum such that the solvent exits the cartridge in a dropwise fashion.
- d) Repeat sample bottle rinse and cartridge elution with a second 5 mL aliquot of elution

Concentration

- a) Concentrate the extract to dryness under a gentle stream of nitrogen in a heated water bath (55–60 °C).
- b) Reconstitute the extract with 1.0 mL of 20% reagent water in methanol (v/v). c) Add the isotope performance standards to the extract and vortex.
- d) Transfer an aliquot of the final extract to a polypropylene autosampler vial (PTFE free).

Figure 3: Chromatogram of a LRB sample containing Isotope Dilution & Isotope Performance standards

LC-MS/MS PARAMETERS Shimadzu Nexara LC-30AD UCT Selectra[®] C18, 50 × 4.6 mm, 5 μm (p/n: SLC-1850ID46-5UM) UCT Selectra® C18, 100 × 2.1 mm, 3 μm (p/n: SLC-18100ID21-3UM) UCT Selectra® C18, 10 × 2.0 mm, 3 μm (p/n: SLC-18GDC20-3UM) 300 μL/min

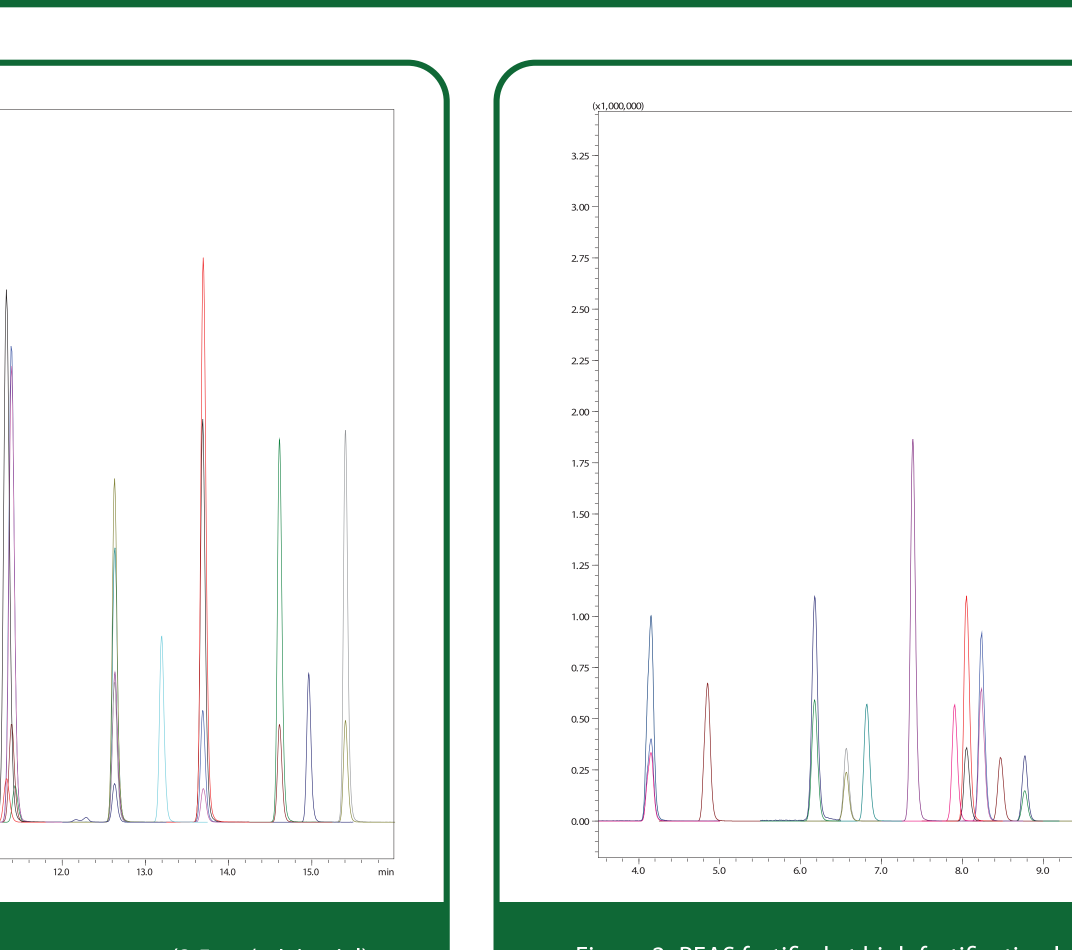


Figure 1: PFAS fortified at low fortification level 10 ng/L in reagent water (2.5 ng/mL in vial). Figure 2: PFAS fortified at high fortification level 80 ng/L in reagent water (20 ng/mL in vial).



Figure 4. Chromatogram showing baseline separation of PFOS isomers (branched vs linear).

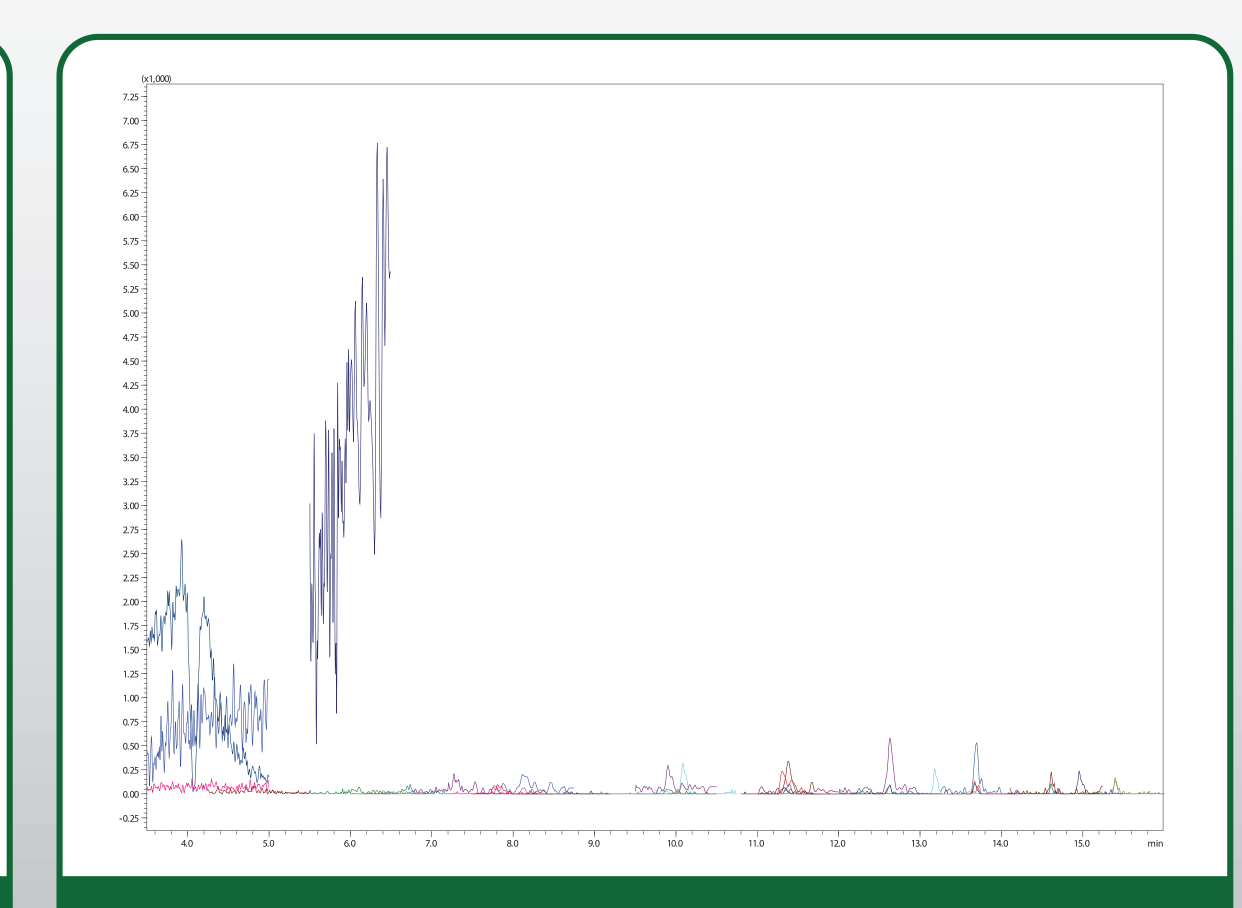


Figure 4: Chromatogram of a blank solvent injection demonstrating low system background levels.

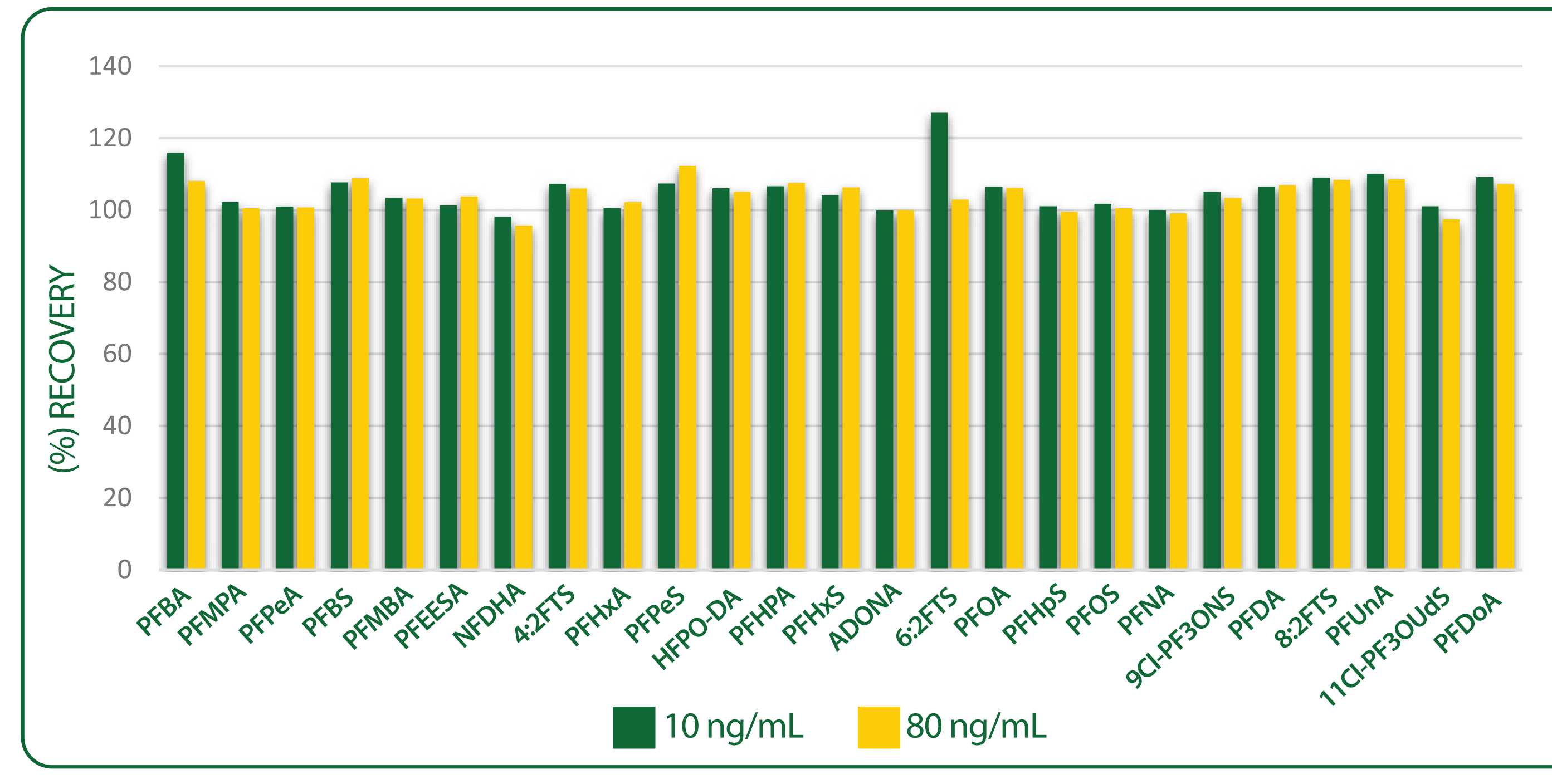
Results in Reagent Water

Analyte	Low Fortification (10 ng/L; n=6)		High Fortification (80 ng/L; n=6)	
	Recovery (%)	RSD (%)	Recovery (%)	RSD (%)
PFBA	115.89	9.22	108.04	10.71
PFMPA	102.20	4.71	100.47	1.24
PFPeA	100.93	5.10	100.70	2.33
PFBS	107.67	5.92	108.84	2.09
PFMBA	103.33	5,45	103.18	0.45
PFEESA	101.27	5.28	103.78	1.90
NFDHA	98.07	5.46	95.73	2.29
4:2FTS	107.27	5.89	105.98	1.90
PFHxA	100.47	5.73	102.22	2.34
PFPeS	107.40	5.97	112.27	2.69
HFPO-DA	106.07	7.09	105.08	2.50
PFHPA	106.60	5.01	107.54	1.96
PFHxS	104.13	5.32	106.32	1.35
ADONA	99.87	5.16	99.94	1.76
6:2FTS	127.00	13.24	102.90	1.54
PFOA	106.47	5.74	106.16	1.91
PFHpS	101.07	5.84	99.50	3.68
PFOS	101.73	4.96	100.48	2.05
PFNA	99.93	5.61	99.12	2.60
9CI-PF3ONS	105.07	5.10	103.32	5.22
PFDA	106.47	4.84	106.89	2.38
8:2FTS	108.93	4.42	108.40	1.94
PFUnA	110.00	5.21	108.53	2.46
11Cl-PF3OUdS	101.07	5.28	97.37	10.74
PFDoA	109.13	4.85	107.22	1.99

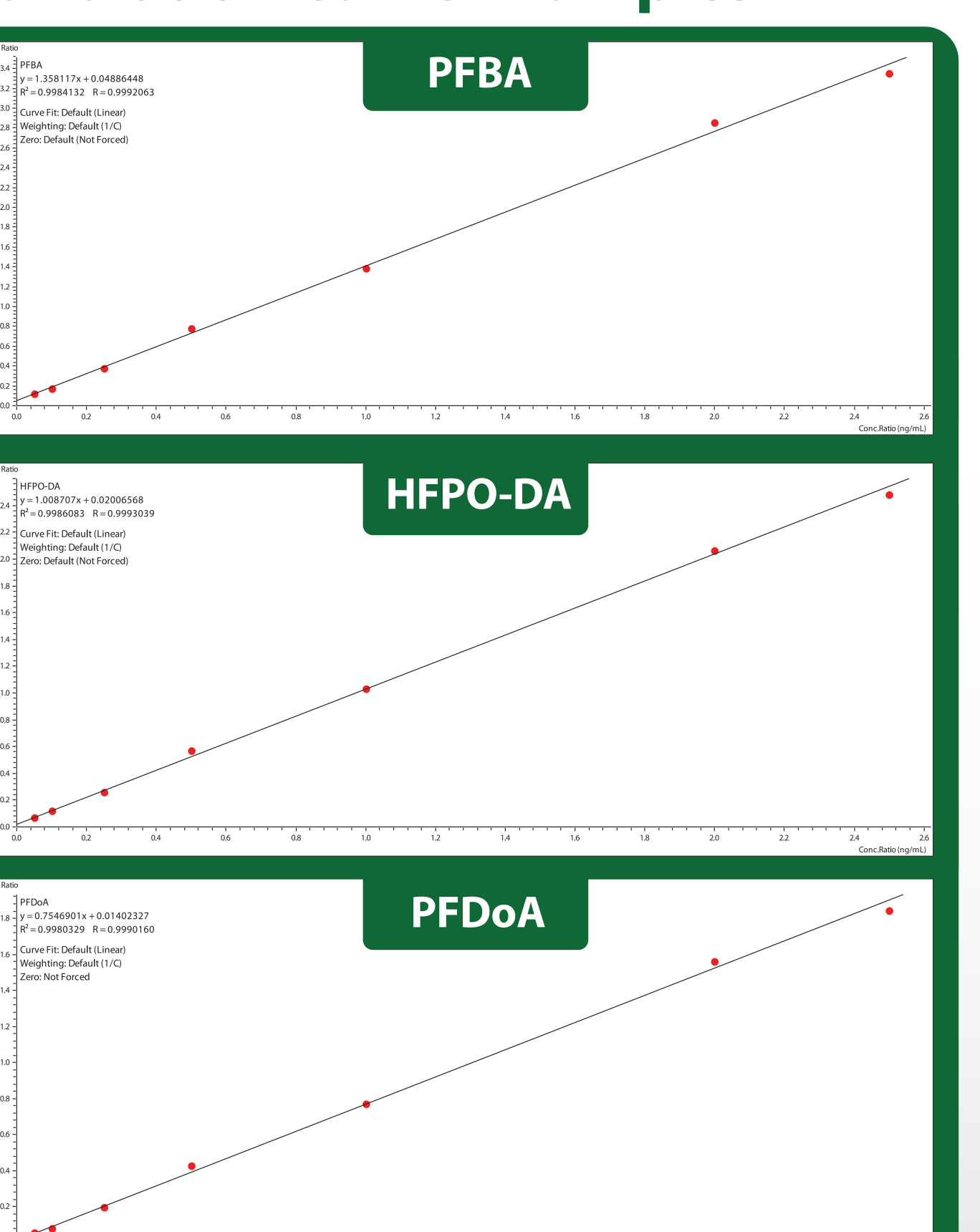
SPE RESULTS



EPA 533 TARGET ANALYTES



Calibration Curve Examples



CONCLUSION

This poster outlines the analysis of PFAS in drinking water according to EPA 533 utilizing UCT's Enviro-Clean®polymeric weak-anion exchange (WAX) SPE cartridges (ECWAX156-P). LC-MS/MS analysis was carried out using a Selectra® C18 HPLC analytical column (SLC- 18100ID21-3UM), while a short (5cm) C18 delay column (SLC-1850ID46-5UM) was used to reduce potential PFAS contamination from the HPLC system.

The chromatography was optimized to obtain maximum resolution with minimum co-elution of isomers, including the critical linear and branched isomers of PFHxS and PFOS. For quantitation, a seven-point calibration (0.5-25 ng/mL) was performed, and all compounds were found to be linear with R2 values > 0.99.

The extraction method was evaluated by spiking reagent water samples with PFAS at 10 and 80 ng/L. Recoveries of all analytes were within a range of 70-130% and RSD values <20%. Due to the prevalence of fluorochemicals used in lab equipment, excluding the use of any PTFE labware throughout the sampling and analytical processes (including HPLC solvent inlet tubing) is essential for accurate analysis of PFAS.

The use of UCT's linear low-density polyethylene (LLDPE) large volume sample transfer tubes (VMFSTFR06-PFC) in conjunction with our complete Glass Block Manifold kit (VMF016GL-PFAS) geared towards PFAS analysis allows for simplified sample preparation and prevent any further introduction of contaminants

References

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- 1. EPA Method 537: Determination of Selected Perfluorinated Alkyl Acids in Drinking Water by Solid Phase Extraction and Liquid Chromatography/Tandem Mass Spectrometry (LC/MS/MS). https://cfpub.epa.gov/si/si_public_file_download.cfm?p_dowload_id=525468&Lab=NERL.
- N. Method 537.1: Determination of Selected Per- and Polyfluorinated Alkyl Substances in Drinking Water by Solid Phase Extraction and Liquid Chromatography/Tandem Mass Spectrometry (LC/MS/MS). https://cfpub.epa.gov/si/si_public_file_download.cfm?p_dowload_id=537290&Lab=NERL.
- 3. Method 533: Determination of Per- and Polyfluoroalkyl Substances in Drinking Water by Isotope Dilution Anion Exchange Solid Phase Extraction and Liquid Chromatography/Tandem Mass Spectrometry. https://www.epa.gov/sites/production/files/2019-12/documents/method-533-815b19020.pdf
- . Unregulated Contaminant Monitoring Rule 5 (UCMR5), accessed online March 2021, https://www.epa.gov/dwucmr/fifth-unregulated-contaminant-monitoring-rule; https://www.govinfo.gov/content/pkg/FR-2021-03-11/pdf/2021-03920.pdf.



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