

1. Overview

An accurate mass, single quadrupole liquid chromatography mass spectrometry (SQ-LCMS) method was developed to facilitate screening environmentally relevant Per- and polyfluoroalkyl substances (PFAS)

2. Introduction

PFAS are ubiquitous and enduring contaminants that have harmful impacts on the environment and public health. Analysis of non-volatile PFAS is commonly performed by triple quadrupole liquid chromatography tandem mass spectrometry (TQ-LCMS); the ability to screen PFAS using SQ-LCMS provides a novel and economical alternative for these laboratories. SQ-LCMS is less selective compared to TQ-LCMS; therefore, accurate mass determination through line-shape calibration is a powerful technique for improving compound confirmation.

3. Method

The 40 PFAS targets and 31 internal standards were diluted to match EPA Method 1633A (EPAM1633A) analysis¹. Neat standards were analyzed from 0.2-1560 ppb in-vial with an injection volume of 5 μ L. Cerno Bioscience MassWorks was used to improve mass accuracy by testing differing calibration criteria and evaluating efficacy using the new Target Analysis function.

Table 1. HPLC and MS acquisition parameters

Nexera XR		LCMS-2050	
Analytical Column:	Shim-pack Scepter C18-120, 3 μ m; 2.1x50 mm	Ionization Type:	ESI
Delay Column:	Shim-pack Scepter C18-120, 3 μ m; 2.1x100 mm	Desolvation Temp:	250 °C
Mobile Phase	A: 2 mM Ammonium acetate in water B: Acetonitrile	DL Temp:	200 °C
Flow rate:	0.4 mL/min	Nebulizing Gas:	2.0 L/min
Oven temp (°C):	40 °C	Heating Gas:	7.0 L/min
Injection volume:	5 μ L	Probe Position:	+2 mm
		Interface Voltage:	-0.5 kV

4. Results and Discussion

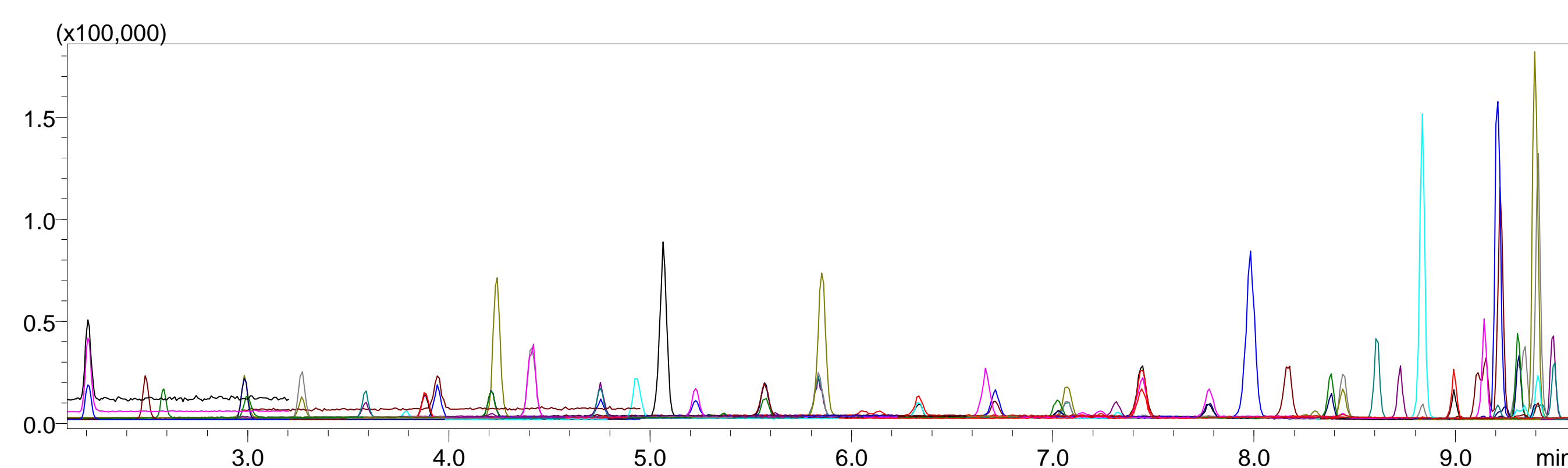


Figure 1. Chromatogram for all analytes at the mid-range calibration standard (1.25-62.4 ppb).

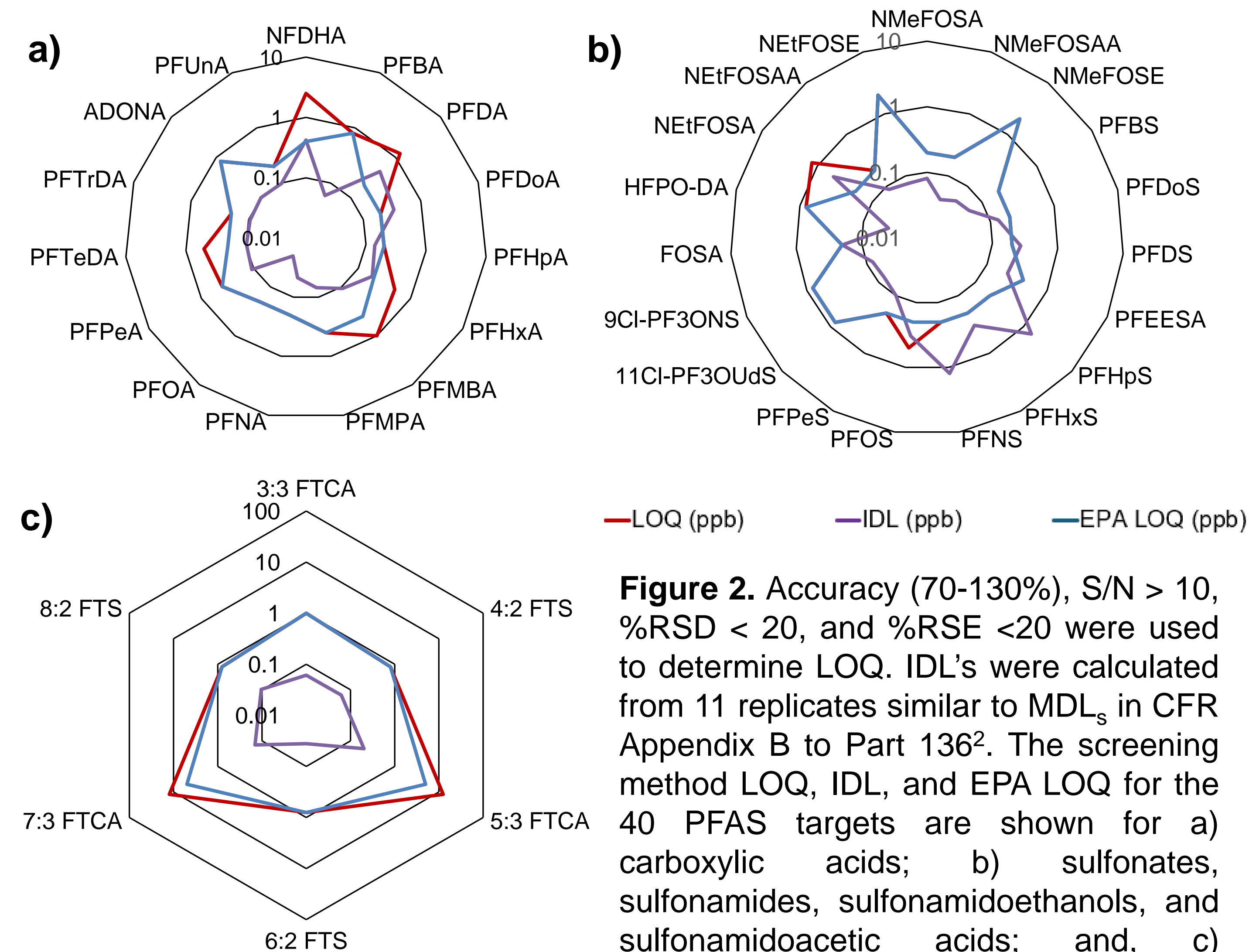


Figure 2. Accuracy (70-130%), S/N > 10, %RSD < 20, and %RSE < 20 were used to determine LOQ. IDL's were calculated from 11 replicates similar to MDLs in CFR Appendix B to Part 136². The screening method LOQ, IDL, and EPA LOQ for the 40 PFAS targets are shown for a) carboxylic acids; b) sulfonates, sulfonamides, sulfonamidoethanols, and sulfonamidoacetic acids; and, c) fluorinated telomer acids and sulfonates.

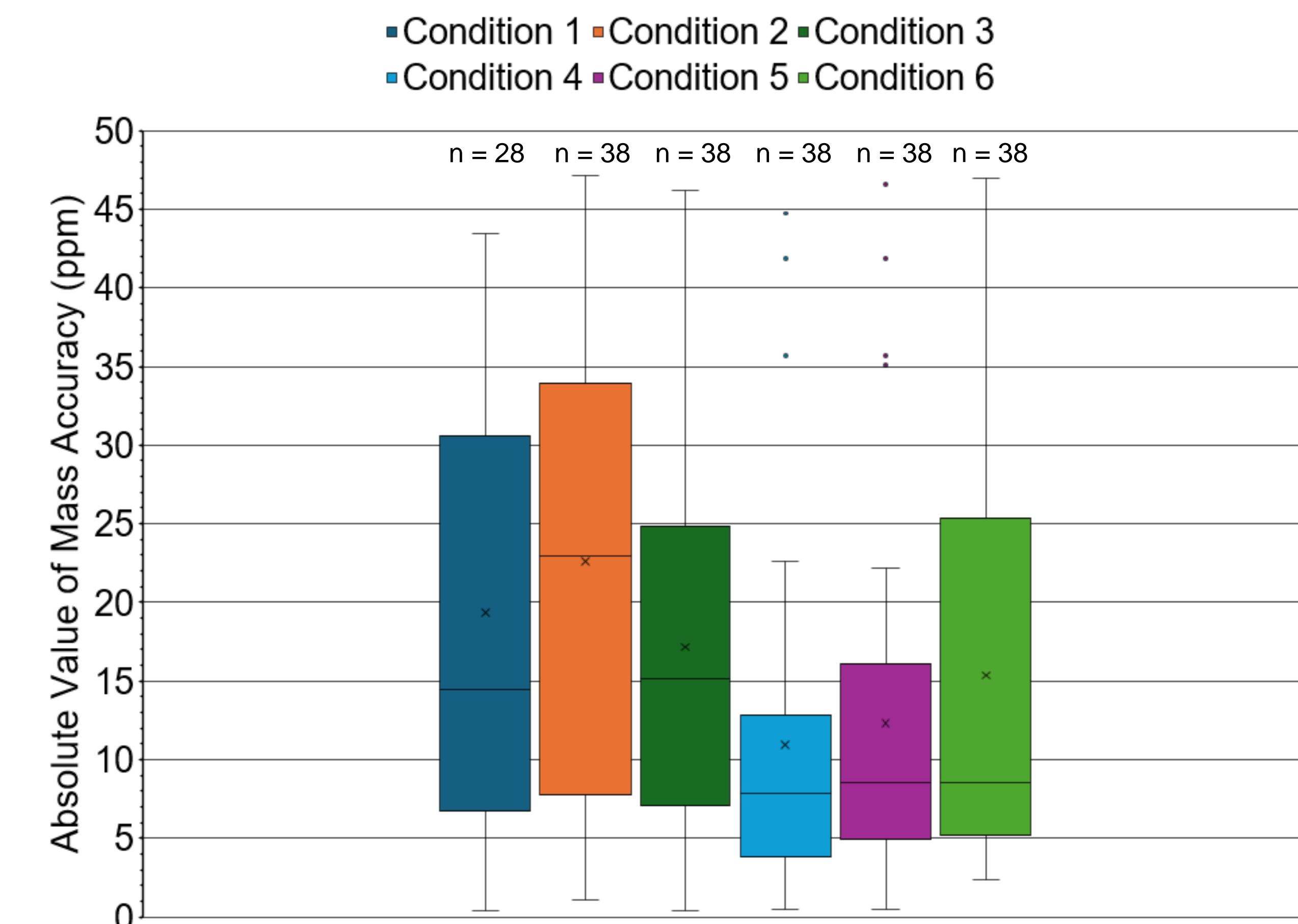


Figure 3. Mass accuracy distribution for the highest calibration standard. Compounds with low S/N and mass accuracy \leq 50 ppm were not identified. The differing criteria and compounds used for analysis were as follows: **Condition 1** – Sigma ESI Tune mix; **Condition 2** – Condition 1 & benzenesulfonamide; **Condition 3** - 7 internal standards with mass accuracy \leq 20 mDa ; **Condition 4** – Target PFAS with sufficient S/N; **Condition 5** – Target PFAS with mass accuracy \leq 20 mDa and spectral accuracy \geq 90%; and, **Condition 6** – Target PFAS with mass accuracy \leq 10 mDa and spectral accuracy \geq 95%.

5. Conclusion

- Effective screen for regulated PFAS for SQ-LCMS analysis with low LOQs comparable to EPA.
- Utilizing a calibration with more compounds (Condition 4) resulted in the best overall mass accuracies.
- Target Analysis in MassWorks enabled accurate mass determination for most PFAS by SQ-LCMS.

6. Reference

1. U.S. Environmental Protection Agency, Draft Method 1633, Determination of 6PPD-quinone in aqueous matrices using liquid chromatography with tandem mass spectrometry (LC MS/MS), December 2023.
2. CFR Appendix B to Part 136, Title 40 -- Definition and Procedure for the Determination of the Method Detection Limit-- Revision 2

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