

PFAS analysis on the SCIEX 7500 system: 15 months of robustness data

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Introduction

This technical note demonstrates the robustness of the SCIEX 7500 system over 15 months of routine PFAS analysis with various water samples. Quality control (OC) samples, spikL-PFHxS, L-PFOS, L-PFOA and L-PFNA, showed accuracies generally within ±1 standard deviation of the mean and all OC samples were with 30% of the mean values (Figure 1). Only 1 preventative maintenance (PM) service was performed during the 15-month ed at 10 ng/L with timeframe. During the analysis, approximately 100–200 injections were performed per week and other non-PFAS applications were also run. These results highlight the strong robustness of both the analysis method and the SCIEX 7500 system.

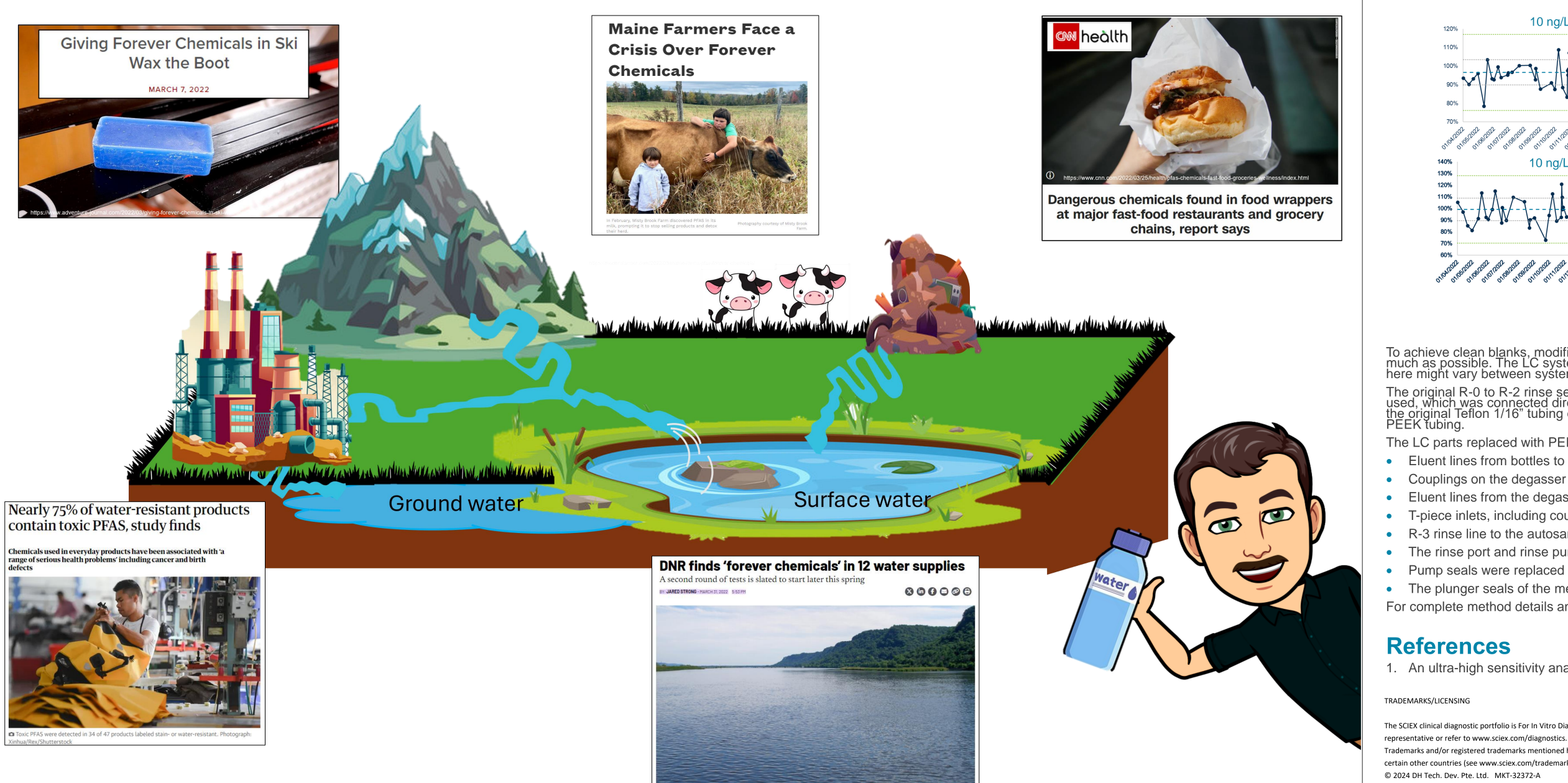
A robust analysis is paramount to the long-term viability of routine PEAS analysis. In this technical note, the developed method was validated for 26 PEAS compounds of interest in LC-MS-grade, drinking, ground and surface water samples.¹ Modifications were made to the LC system to reduce background contamination, including replacing or bypassing any components of the system that contribute to PEAS contamination. Specifically, system components containing fluorinated ethylene propylene (FEP) and Teflon were bypassed or replaced with polyether ether ketone (PEEK), when possible.



Analysis details

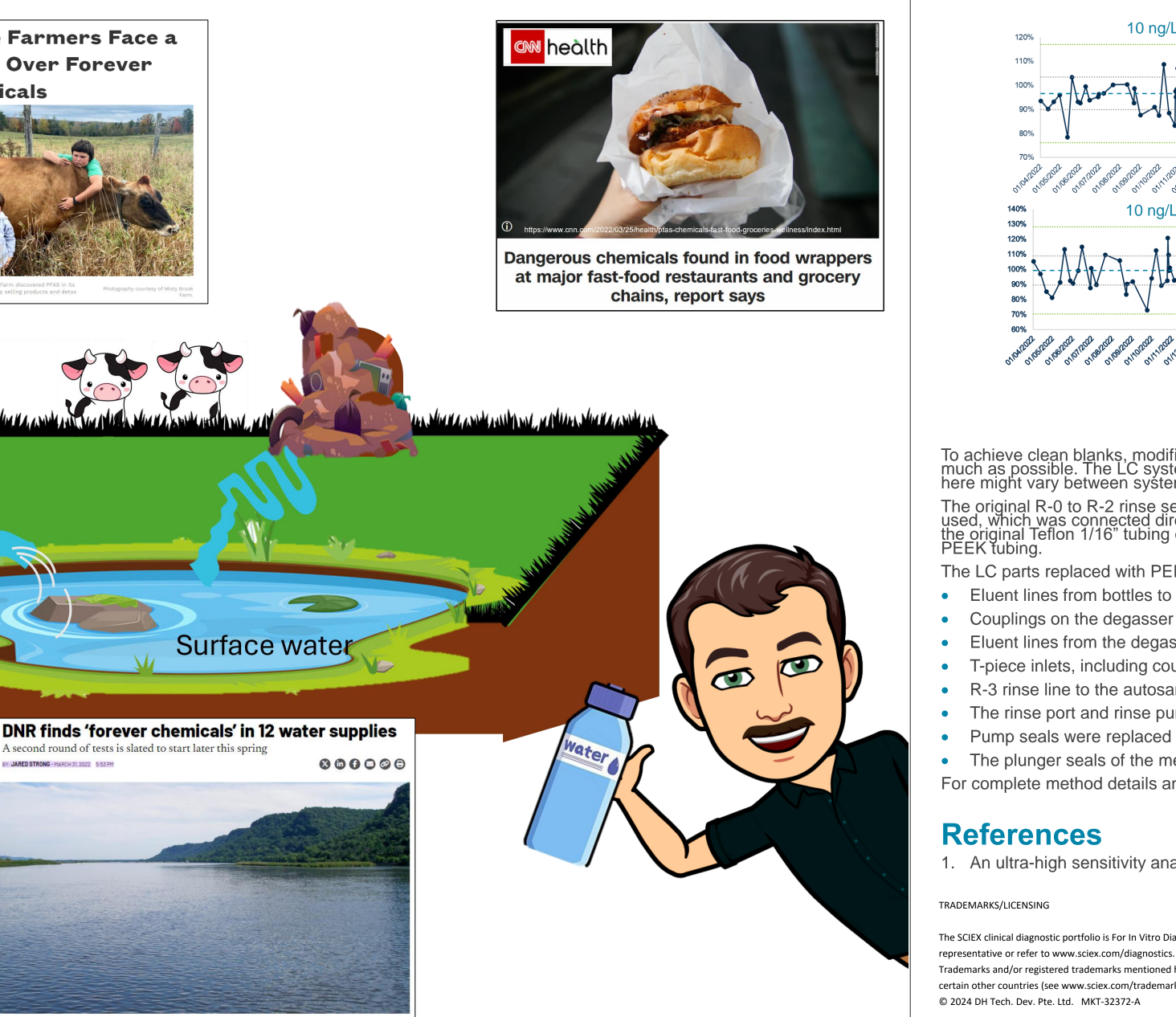
To further clarify how the QC data were collected, the analysis batch is included below. Step 8 introduces the QC samples analyzed in matrix. Step 14 outlines the flushing procedure used following sample analysis.

- 1. 10 ng/L standard: Injected in triplicate to check the instrument performance
- 2. Diluent blank: Used as the 0 ng/L calibration curve standard
- 3. Calibration standards and sample diluent: A calibration curve was constructed across concentrations ranging from 0 to 50 ng/L in ultrapure water
- 4. Diluent blank
- 5. Procedure blank: Blank sample that underwent all sample preparation steps
- 6. QC sample in ultrapure water
- 7. Blank QC sample in matrix
- 8. QC sample at 10 ng/L in matrix
- 9. Diluent blank
- 10.10 water samples (various sources)
- 11. QC in ultrapure water and sample diluent: Repeated every 10 samples to check for drift
- 12. Diluent blank: Repeated every 10 samples
- 13. Calibration curve
- 14. End of method: The column was flushed with 50:50 (v/v), mobile phase A/mobile phase B for 15 minutes. The column was flushed with 99% mobile phase B for 5 minutes. The flow stopped and the instrument was placed into standby.



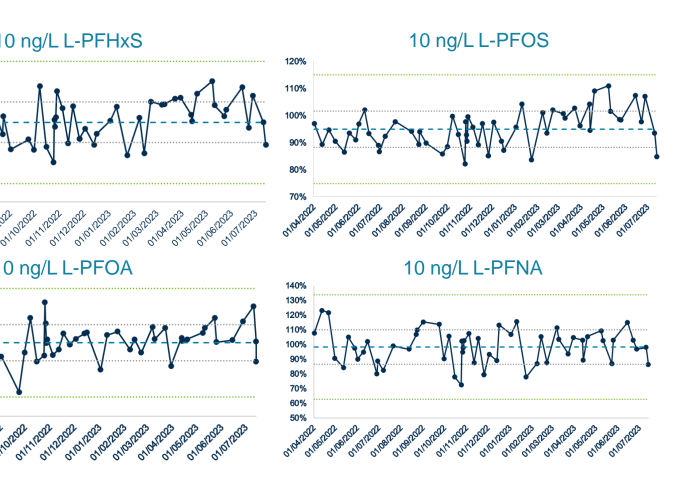












Modifications to LC system

To achieve clean blanks, modifications were made to the LC system to mitigate PFAS contamination as much as possible. The LC system used here was a Shimadzu LC-40 system. The changes described here might vary between systems, depending on configuration and flow path.

The original R-0 to R-2 rinse selection block in the autosampler was bypassed. Only 1 rinse liquid was used, which was connected directly to the low pressure valve (LPV) using a PEEK tube. Subsequently, the original Teflon 1/16" tubing connecting the LPV to the bottom of the rinse port was replaced by 1/16" PEEK tubing.

- The rinse port and rinse pump tubing from the autosampler to the ports, including couplings
- For complete method details and more information please refer to reference #1.

1. An ultra-high sensitivity analysis of PFAS compounds in multiple water sources,

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