seaFAST
Automating Seawater Analysis
Elemental Scientific
ICP | ICPMS | AA
Brief and System Schematics

Pico: REEs in Seawater (Offline)
Pico: Trace Metals in Seawater (Offline)
Pico: First Row Transition Metals in Seawater (Inline)
S2: Preconcentration Mode
S2: Direct & Preconcentration Modes
seaFAST SP3

Brief

The seaFAST is a high performance, automated sample introduction system for the determination of ultra-trace metals in undiluted seawater and other high matrix samples. It lowers procedural blanks and improves detection limits for a variety of elements through syringe-based sample preconcentration and matrix elimination.

seaFAST System

- Fully automated software control
- Seamlessly integrated with ICPMS
- Syringe controlled volumes and flow rates
- Inert fluoropolymer flow paths
- High sample capacity

Figure 1. Image and schematic of the seaFAST preconcentration column. At a pH ~ 6, transition row metals are chelated on the preconcentration column, while matrix elements (group 1 and 2) are flushed from the column.
seaFAST Pico

Seawater samples are automatically processed and concentrated fractions eluted into a destination vial for analysis.

**Offline**
- Collects samples for analysis
- Standalone operation
- Steady state signal
- User defined load and elute volumes
- 15 min/sample
- Inline mode for method development

**seaFAST S2**

Dual mode operation allows undiluted samples to be measured directly with a user-specified dilution while loading an aliquot of sample onto a column for automated, inline preconcentration and matrix removal.

**Inline**
- Elutes sample directly to ICPMS
- Real time data
- High preconcentration factors
- Dual mode operation
- < 9 min/sample
- Offline Mode

Figure 2. seaFAST Pico system with integrated mobile stand.

Figure 3. seaFAST S2 system with integrated mobile stand.
Offline Application

Accurate and precise quantification of Rare Earth Elements (REEs) in open ocean seawater remains a difficult task. This is due to low dissolved analyte concentration (10s of pg L\(^{-1}\)), BaO\(^+\) interferences, and a complex matrix (3.5% TDS). The seaFAST-Pico™ is an ultra-clean, automated, low-pressure ion chromatography system capable of single digit picogram L\(^{-1}\) detection limits. In an offline configuration, the automated system buffers user defined volumes (10 to 100mLs) of acidified seawater before loading it onto a column. The matrix is then removed and the concentrated sample is eluted in user defined volumes (250 to 1000 µLs) of acid. Rapid sample loading rates combined with low elution volumes provides high concentration factors and unprecedented detection limits (pg L\(^{-1}\)).
Figure 5. Similar to Figure 1, REEs determined and consensus concentrations for a deep sample (2000 m) from the Bermuda Atlantic Time-series Station (BATS) are plotted. The error bars (2 SD) illustrate the precision of the method, whereas accuracy is demonstrated through agreement with consensus values. Data courtesy of Katharine Pahnke, Melanie Behrens and Ronja Paffrath, Max Planck Research Group for Marine Isotope Geochemistry.

Figure 6. A comparison of a seaFAST-Pico inline elution profile with a seaFAST-Pico offline injection is shown above. Offline fraction collection of programmable elution volume is injected to the ICPMS with a microFAST system. A profile collected from a sample eluted in 200 µL and introduced at 100 µL min⁻¹ illustrates two minutes of steady state signal.
Figure 7. By adjusting elution volume (1000, 500, to 250 µL), 10 mLs of Indian Ocean surface water (with 10 ppt REE spike) is concentrated 10, 20, and 40x. A linear correlation between concentration and intensity indicates that even 250 µLs is sufficient for quantitative recovery.

Figure 8. By adjusting sample volume (10 - 60 mL), Indian Ocean surface water (with 10 ppt REE spike) eluted in 1 mL is concentrated 10 - 60x. A linear correlation between concentration and intensity indicates the columns capacity to process large volume samples with quantitative recovery.
Figure 9. Limits of detection are compared to SAFe surface water concentrations. Matrix removal combined with a 20-fold preconcentration factor is sufficient to determine the lowest level of REEs in open ocean seawater.

Figure 10. The interpretation of dissolved REEs requires high quality concentration values from which very precise patterns and elemental ratios can be determined. Three NASS-6 replicates, run on five different analytical dates, indicate good precision for concentration data and better than ± 3.5% (2-sigma) for LREE/HREE (Nd/Er) ratios.
**Offline Application**

Accurate and precise quantification of transition row metals in seawater remains a difficult task due to low dissolved analyte concentration (10s of ng L\(^{-1}\)), a complex matrix (3.5% TDS) and difficulty obtaining low procedural blanks. The seaFAST-Pico™ is an ultra-clean, fully automated system with a fluoropolymer flow path that reduces blanks by minimizing sample exposure and handling. Rapid sample loading rates combined with small elution volumes provide high concentration factors (~20-100 fold) and unprecedented detection limits (pg L\(^{-1}\)).

**Figure 11.** Determined (two separate dates) and certified values for seawater reference material (CRM) NASS-6 are plotted (on a log scale). Excellent agreement between determined and certified values for both low (Pb, 6 ng L\(^{-1}\)) and high (Fe and Mn ~500 ng L\(^{-1}\)) concentration elements illustrates the accuracy of the method. No internal standard was used.
Figure 12. Indium is reproducibly and quantitatively recovered from seawater samples making it an excellent Internal Standard (IS) candidate.

Figure 13. Three replicates of a three step elution are plotted. The first step, after loading NASS-6, recovers trace metals from the column. Two subsequent elution steps illustrate greater than a 1000-fold washout, indicating minimal sample to sample carryover. Low carryover is sufficient to measure trace metals in a wide variety of oceanic environments.
Figure 14. Limits of detection are compared to NASS-6 certified values. Matrix removal combined with a 20-fold preconcentration factor is appropriate for many seawater applications. If lower detection limits are required, the system can easily be programmed for larger preconcentration factors.
Inline Application

The seaFAST Pico can be operated inline. The automated method loads 10 mL of seawater and elutes the transition row metals directly to an ICPMS. Rapid sample loading rates combined with low elution volumes provide high preconcentration factors (~200 fold) and unprecedented detection limits (pg L⁻¹). The whole process is complete in 9 minutes providing real-time data with invaluable insight into blank reduction and other aspects of method development.

**Figure 15.** Elution profiles for a selection of first row transition metals (2 point running average).

**In collaboration with:**

The Institute of Marine and Coastal Sciences Rutgers the State University of New Jersey

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Figure 16. Limits of detection for contamination-prone elements (Fe and Zn) are in the high pg L⁻¹ range and are reproducible on a day-to-day basis. These limits of detection are appropriate for determination of ultra-trace metal concentrations in open ocean surface seawater.

Figure 17. An internal laboratory standard of Ross seawater collected from the Southern Ocean is run several times on each of 9 discrete analytical dates. Excellent day-to-day reproducibility is illustrated for Fe, a difficult contamination-prone element.
**Figure 18.** Accuracy of Manganese, Nickel and Copper at very low concentrations typical of the open ocean is demonstrated for SAFe and GEOTRACES surface water.

**Figure 19.** Accuracy of Iron and Zinc at very low concentrations typical of the open ocean is demonstrated for SAFe and GEOTRACES surface water.
Abstract

A seaFAST S2 system is used in Preconcentration mode to determine the Cd and Pb concentration in arctic seawater. Validation of the method is illustrated by obtaining Cd and Pb concentrations that agree with the consensus values for SAFe (S and D1) and GEOTRACES (GS and GD) samples. Advantages of the system include the small sample volumes required (12 mL for duplicate analysis) for low level multi element analysis, fast throughput (6 samples per hour) and low procedural blanks.

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Figure 21. Concentrations of dissolved Cd determined by isotope dilution for SAFe and GEOTRACES reference standards.

Figure 22. Concentrations of dissolved Pb determined by isotope dilution for SAFe and GEOTRACES reference standards.
Abstract

The seaFAST S2 is a high performance, automated sample introduction system for the determination of trace elements in seawater and other high matrix samples by ICPMS. Undiluted samples can be measured directly with a user-specified dilution while simultaneously loading an aliquot of sample onto a column for automated, online preconcentration and matrix removal. Both Direct and Preconcentration modes reduce procedural blanks associated with offline sample preparation; Preconcentration mode dramatically improves detection limits by eliminating matrix effects and increasing sensitivity. The seaFAST S2 offers complete flexibility and can run in Direct, Preconcentration, or both modes by simply selecting the desired mode(s) in the ESI software.

![Figure 23. Periodic table showing the possible modes for each element.](image)

Modes

**Preconcentration mode:**

A chelation column binds transition metals and rare earth elements but allows matrix Na⁺, Cl⁻, Ca²⁺ and Mg²⁺ ions to be rinsed out. After the preconcentration step, analytes are eluted and detected by ICPMS.

**Direct mode:**

Sample is automatically diluted inline by high-precision syringe pumps. Dilution reduces matrix effects and allows the determination of elements whose chemistry is not compatible with the preconcentration column.
Figure 24. Direct mode calibration of Tl in 3.5% NaCl. Calibration spikes are 1, 10, 50, 250 and 500 ppt.

Figure 25. Preconcentration mode calibration of Fe in 3.5% NaCl. Calibration spikes are 1, 10, 50 and 250 ppt.
Figure 26. Results for NASS-6 and CASS-5 Seawater Reference Materials are shown. Elements determined with the seaFAST S2 are accurate and precise for elements with certified concentrations from < 10 ppt to > 10 ppb.
Figure 27. Washout in Preconcentration mode is excellent with both 1% nitric acid calibration standards and matrix-matched standards. Matrix matching the standards with seaBlank is not required for Preconcentration mode, but it enhances washout.

Figure 28. Detection limits are near or better than 1 ppt on a quadrupole ICPMS (n=10, 3σ).
A new, automated analysis system for ultratrace determination of low and sub-ppt metals in high matrix samples is described. Simple external autocalibration from a single multielement stock standard eliminates common sources of contamination for ultratrace metals determination. Inline, automatic matrix matching provides exceptional accuracy. Metals may be measured in up to three modes: preconcentration and matrix removal, direct analysis with inline dilution, and hydride generation.

Figure 29. Periodic table showing the possible modes for each element.
Introduction

Matrix effects from samples with high total dissolved solids (TDS) can cause severe problems for accurate determination of many elements by ICPMS. High dilution factors attenuate the matrix effects but are undesirable when excellent detection limits are required. Historically, a variety of advanced techniques have been used to accurately determine trace elements in high TDS samples. One common approach is preconcentration and matrix removal, which offers superb results for elements whose chemistry is compatible with the chosen chelation resin. For other elements, manually matching the matrix of calibration blanks and standards to that of the diluted samples can achieve very accurate results in high TDS samples. Lastly, hydride generation improves quantification for elements such as As and Se that are otherwise difficult to determine.

The new seaFAST SP3 analyzes samples using one or more advanced analytical techniques: preconcentration and matrix removal, direct analysis with inline dilution, and hydride generation. For improved accuracy the seaFAST performs automatic inline matrix matching for calibration blanks, standard, and QC samples, as well as autodiluted samples.

Figure 30. Autocalibration from a single 1000 ppt calibration standard. Calibration curve was matrix matched to ~3% TDS.

Figure 31. Direct mode with 10x inline dilution: Autocalibration from a single 1000 ppt calibration standard. Calibration curve was matrix matched to ~3% TDS.

Figure 32. Autocalibration from a single 1000 ppt calibration standard. Calibration curve was matrix matched to ~3% TDS.
Direct Mode: Long Term Stability

Figure 34. Long term stability of Cr in synthetic seawater shows >14 hours of repeated determinations with internal standard correction. Direct mode automatically diluted the sample inline by 10x. The determined concentration in the synthetic seawater was 33 ± 3 ppt.

Preconcentration Mode: Elution of 6 Elements at 1 ppb

Figure 33. Simultaneous elution of 6 elements in Preconcentration mode at 1 ppb.
Figure 36. NASS-6 was analyzed in all three modes, Preconcentration (P), Direct with fixed 10x inline dilution (D) and Hydride (H). NASS-6 was run undiluted and with a 2x autodilution using automatic, inline matrix matching with seaBlank ultrapure sodium chloride solution. The seaFAST achieves excellent accuracy in all three modes, with and without autodilution.