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Monodisperse microdroplets: a tool that advances single-particle ICP-MS measurements

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In single-particle inductively coupled plasma mass spectrometry (sp-ICP-MS), individual nanoparticles are detected by measuring ICP-MS signals with high temporal resolution. At high time resolution, particlebased signals—which are around 200-500 us in duration—make up a larger fraction of the signal measured. If the dissolved background is low enough and the mass of element(s) of interest in the particles is high enough, then nanoparticle (NP) signals are recognized as signal spikes on the time trace. With modern instrumentation, sp-ICP-MS can be used to quantify NPs with element mass down to single-digit attogram levels. Monodisperse microdroplets composed of elemental solutions are vehicles for the transport of discrete amounts of elemental species into the ICP, and signals produced from microdroplets closely match those produced by NPs. Temporal durations and element sensitivities obtained from microdroplets and NPs are so similar that microdroplets may be used as NP proxies. Unlike NPs, microdroplets offer a flexible platform for user-designed sp-ICP-MS measurements because—with microdroplets—we can precisely control the absolute mass of elements injected into the plasma in each droplet. Controlled introduction of analyte mass into the ICP enables us to use microdroplets to generate sensitivity calibration factors and also to use microdroplets as NP proxies to study sp-ICP-MS measurement accuracy. Here, were report several instances of how measuring microdroplet-based element signals with ICP-TOFMS has allowed us to expand the versatility, as well as the general understanding, of sp-ICP-TOFMS measurements. First, we summarize how microdroplets can be used for online calibration of analyte NP element mass and particle-number concentration. Second, we describe how the measurement of microdroplets with tailored analyte masses helps us to refine, test, and validate sp-ICP-TOFMS data analysis strategies. Specifically, we use microdroplets to study the accuracy and robustness of split-event correction and signal-thresholding approaches for NP detection. Together, these experiments describe how the use of monodisperse microdroplets allows us to design better sp-ICP-MS experiments.

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Introduction

Monodisperse microdroplets are not a new addition to the atomic spectroscopist's toolbox. In the past sixty years, many researchers have employed monodisperse microdroplets to study the interaction of aerosolized analyte with various atomization, excitation, and ionization sources for both optical and mass spectrometry. In principle, the use of uniform microdroplets is advantageous for fundamental studies because microdroplets replicate typical sample introduction approaches, i.e. nebulized aerosol, but are a well-controlled system which allows for the isolation of flame/plasma interactions from phenomena arising from the size heterogeneity of an ensemble of droplets.

The use of monodisperse microdroplets in atomic spectrometry was pioneered in the late 1960's by Hieftje and Malmstadt for the study of droplet behavior in air-acetylene laminar flames.^{1,2} From the onset, microdroplet sample introduction was identified as an approach well-suited to probe sample-source interactions, with Hieftje's 1968 paper in Analytical Chemistry entitled, "A unique system for studying flame spectrometric processes." Following this initial work, the Hieftje research group employed microdroplet measurements to study various fundamental aspects of sample-laminar flame interaction including processes of liquid-aerosol desolvation,3 particle evaporation,4 and atomic diffusion,5 and determinations of aerosol velocity.^{6,7} Several research groups have used monodisperse microdroplets to study fundamental properties of the inductively coupled plasma combined with either optical emission or mass spectrometric detection.8 Of specific note, most contemporary microdroplet introduction systems trace their roots back the monodisperse dried microparticulate injector (MDMI) system first reported by French et al.9 Over the

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years, variations of this microdroplet injection system have

been used to study many processes in the ICP, including droplet desolvation,10,11 particle evaporation,12,13 local heating or cooling of the plasma,14,15 plasma-mass spectrometer interface characteristics, 16 space-charge effects, 17 and other matrix effects.18-21

Researchers have also employed monodisperse microdroplets as a sample carrier. Dissolved analyte can be introduced into the ICP as monodisperse microdroplets (or dried microdroplet particles) in order to have better control of sample-introduction related phenomena and thus better control of matrix effects and optimization for best absolute sensitivities. 9,22-25 Discrete samples, such as single nanoparticles²⁶⁻²⁹ or cells³⁰⁻³² can also be introduced within microdroplets for a more controlled sample introduction strategy compared to conventional pneumatic nebulization. For both single-nanoparticle and single-cell ICP-MS measurements, introduction of a sample with monodisperse microdroplets presents an attractive approach for quantitative transfer of analyte particles/cells into the plasma with the possibility of direct counting experiments for number concentration determinations. Quantitative transfer of individual microdroplets into the ICP allows for direct determination of absolute detection efficiencies (counts per g) of analyte elements in the microdroplet solution,26 which can be used to quantify the mass of elements in discrete entities injected into the plasma, such as single nanoparticles, 28,29,33-38 or cells.

In 1994, Olesik and Hobbs wrote an article in Analytical Chemistry entitled "Monodisperse Dried Microparticulate Injector: A New Tool for Studying Fundamental Processes in Inductively Coupled Plasmas."39 In that article, they discussed how time-resolved measurements of monodisperse microdroplets allow researchers to study previously inaccessible characteristics of ICP-OES and -MS. Likewise, here we explore how monodisperse microdroplets are a "tool" that we can use to help us understand and develop new measurement approaches for time-resolved ICP-MS measurements, such as with singleparticle and single-cell ICP-MS. For this purpose, we use microdroplets in two general ways: as a calibration standard for absolute mass quantification and as a nanoparticle proxy to study acquisition and data processing strategies for accurate sp-ICP-MS quantification. Microdroplets are an excellent option for studying fundamentals of sp-ICP-MS because they overcome limitations of studies based solely on nanoparticle materials as a reference: namely, microdroplets can be delivered at known frequency, have a narrow analyte-mass distribution, and their elemental composition is easily adjusted.

Experimental

All experiments and experiment concepts presented here make use of a dual sample introduction setup, as depicted in Fig. 1. This setup has been described elsewhere. 33,35-38,40,41 In this setup, microdroplets are produced by a microdroplet generator (50 µm diameter Autodrop Pipette, AD-KH-501-L6, MD-E-3000 dispensing system, Microdrop Technologies, GmbH, Germany) and introduced via a falling tube that is filled with a He/

Ar gas mixture to accelerate solvent evaporation so that microdroplets are reduced to dried droplet residues by the end of the falling tube.26,42 Continuously nebulized samples are introduced via a pneumatic nebulizer and spray chamber, and then mixed with microdroplet-aerosols via a t-piece just in front of the injector base of the ICP torch. A video camera and triggered LED are used to stroboscopically image droplets to measure diameter—and, by calculation, volume—of spherical monodisperse microdroplets. 43 RSDs for dropet volume are \sim 1%. For ICP-MS measurements presented here, we used an ICP-TOFMS instrument (icpTOF 2R, TOFWERK AG, Switzerland), as already described elsewhere.34,44 Because the TOF mass analyzer provides simultaneous full mass spectrum detection at a high speed (21.739 kSpectra per s) and continuous readout of averaged spectra at ~ 1000 Hz, the full analyte composition of each microdroplet can be quantitatively measured. Solutions for microdroplets were prepared by dilution of commercially available single-element standard solutions into either 1% subboiled HNO₃ or a mixture of 1% HNO₃ and 1% HCl. All dilutions were performed gravimetrically. TOFMS data are stored in the HDF5 file format, and all post-acquisition data processing was performed in LabVIEW (ver. 18.0, National Instruments, TX, USA). Data were plotted in OriginPro (ver. 2019b, OriginLab Corp., MA, USA) and final figures were assembled in Adobe Illustrator (ver. 24.1, Adobe Systems Inc., USA).

Results and discussion

sp-ICP-MS is often recognized as a premier analytical approach for the high-throughput determination of nanoparticle size (via element-mass quantification) and particle-number concentration (PNC). One reason sp-ICP-MS has been adapted broadly is because it makes use of widely available ICP-MS instrumentation. In addition, sp-ICP-MS remains the most viable approach for quantitative detection of inorganic nanomaterials at environmentally relevant concentrations. 45,46 Since the first description of sp-ICP-MS for colloid measurements,47 hundreds of papers have been published about sp-ICP-MS, as well as several review papers. 48-50 In those years, researchers have developed novel instrumentation^{26,51} and measurement strategies in order to improve the approach.52-55 Further, ICP-MS instrumentation companies have released updated instruments and software that enable more-or-less "plug-and-play" sp-ICP-MS measurements. Today, sp-ICP-MS is used as a routine approach.

When nanoparticles (or micron-sized particles) are introduced into the ICP at low enough number concentrations, the ion clouds formed by the vaporization, atomization, and ionization of these particles are measurably discrete in both time and space. In sp-ICP-MS, these discrete ion clouds are registered as individual events by discrimination of their signals across time. The temporal duration of an individual particle in the ICP is usually 200–500 μs, and so by recording MS signals with millisecond (or sub-millisecond) time resolution, responses from individual particles can be measured as signal spikes. If analyte particles are quantitatively vaporized and ionized in the plasma, then sp-ICP-MS can-in principle-be

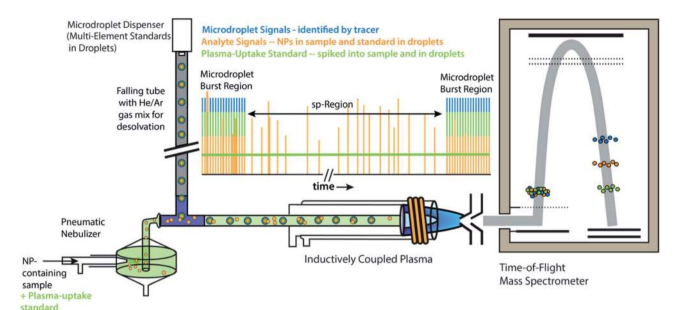


Fig. 1 Schematic diagram online microdroplet calibration. In this setup, signals from analyte standards in microdroplets from the "Burst Regions" are used to determine absolute detection efficiencies (counts per g) for each analyte, which are then used to quantify the mass of elements in NPs from the "sp-Region." The ratio of sensitivities from a plasma uptake standard in both microdroplets and spiked into NP-containing samples is used to determine volumetric flux of sample into the plasma (q_{plasma}), which is necessary for particle-number concentration (PNC) determinations. This figure is adapted from Mehrabi *et al.*³⁷ and is used under the creative common license.

used determine particle number concentrations (PNCs) and the amount of element(s) in individual particles. Very simply, quantification in sp-ICP-MS involves three steps, all of which affect the measurement's accuracy: first, NP signals must be accurately found, *i.e.* distinguished from background signals; second, found particle signals must be calibrated to reflect the amount of analyte elements in the particles; third, sample flux into the plasma must be determined to relate the number of found particles to number concentration.

It is an easy task itemize the fundamental requirements of a quantitative sp-ICP-MS experiment; however, validation of these measurements for a range of NP types and in various samples matrices is difficult. Basic understanding of the accuracy of sp-ICP-MS measurement and data analysis approaches is hampered by a lack of NP reference materials certified for both element mass and number concentration. This lack of standards has led to inconsistent measurement practices among researchers, which may cause confusion for new practitioners of sp-ICP-MS. For example, there is little consensus on significant measurement conditions such as optimal dwell time, most accurate method to determine transport efficiency and absolute elemental sensitivities, 55-58 or robust signal-thresholding criteria for NP identification.59 There are, indeed, many researchers working to harmonize and standardize sp-ICP-MS measurements,60-63 but, without a suite of well-characterized reference NPs, this process remains a challenge.

In this paper, we present some recent efforts we've made to develop accurate methods for NP quantification for sp-ICP-TOFMS. Unlike ICP-MS with a quadrupole or sector-field mass spectrometer, with TOFMS, we have the ability to measure full elemental mass spectra at a time resolution suitable for single-

particle analysis. Full-mass-spectrum detection with TOFMS allows us to perform multiplexed sp-ICP-MS experiments in which we can simultaneously measure signals from singleelement NPs, multi-element NPs, microdroplets, and dissolved elemental species. Recently, sp-ICP-TOFMS has gained heightened interest for high-throughput screening of NPs and for measurement of multi-element signatures in NPs as a means to source single particles, i.e. as natural or engineered. 37,64-68 In our sp-ICP-TOFMS measurements, we use microdroplet sample introduction in two distinct ways: (1) for online calibration of NP mass in individual NPs, and (2) as a tool to study data analysis approaches for the quantitative detection of fast transient signals, which is critical for reliable PNC determination in sp-ICP-TOFMS. In both of these applications, microdroplets are a flexible platform for controlled introduction of discrete masses of analytes which allow us to better probe sp-ICP-TOFMS measurements.

Calibration with microdroplets

We have described online microdroplet calibration in several recent publications. There, we provide a brief overview of how microdroplet-generated signals provide a route for calibration of element mass in *individual* analyte nanomaterials. Unlike all other approaches for sp-ICP-MS quantification, online microdroplet calibration does not rely on the use of external reference nanoparticles or the stability of analyte sensitivities and sample transmission efficiencies into the plasma between measurements. Instead, analyte sensitivities and plasma uptake rate are calibrated in *every* measurement for *each* sample. A schematic diagram of our sample introduction

setup is provided as Fig. 1, and a flow chart of the overall online microdroplet calibration procedure is provided in Fig. 2.

In online microdroplet calibration, we use a dual sample introduction approach in which microdroplets composed of solutions with known multielement concentrations are introduced into the ICP along with sample nebulized *via* conventional pneumatic nebulizer and spray chamber. Because the

volume of each microdroplet can be measured, we know the mass of analyte injected with each microdroplet. Based on the average intensities recorded from microdroplets, we obtain absolute detection efficiencies for each analyte, i, $(S_{\rm drop,i}, {\rm counts} {\rm per} {\rm g})$, which are then used to calibrate the mass(es) of element(s) in NPs. In addition to analyte standards, we add a known amount of a plasma-uptake standard into both

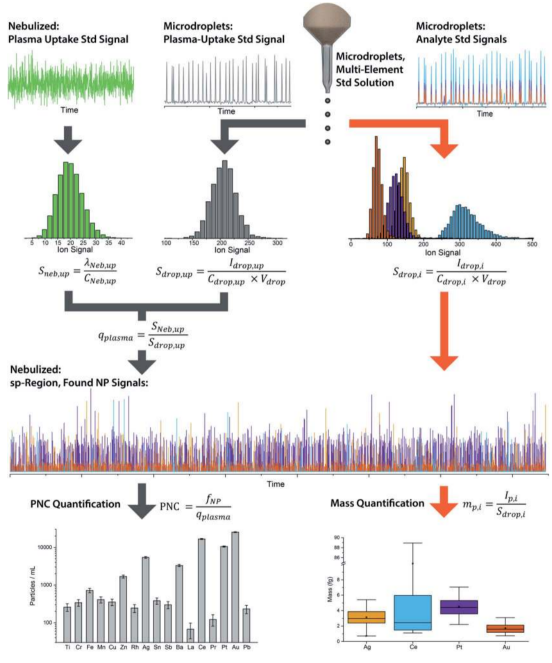


Fig. 2 Flow chart of the basic steps and equations in online microdroplet calibration. In this calibration approach, no external NP reference materials are required. For each measurement, sample-dependent absolute sensitivities ($S_{drop,i}$) of each element, i, and volumetric flux into the plasma (q_{plasma}) are determined in order to correct for any potential matrix-effects. Mass calibration of analytes in NPs is done based on the average absolute sensitivities of analytes in microdroplets ($S_{drop,up}$, counts per g) divided by the concentration-based sensitivity for the plasma uptake standard spiked into the nebulized sample ($S_{neb,up}$, count per s per g per mL) is used to determine q_{plasma} . Data used here is for explanatory purposes only and is taken from our study of the quantification of Ag, Pt, and Au NPs spiked into waste-water treatment plant effluent.³⁷

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microdroplets and nebulized samples in order to determine the sample-specific flow rate (mL s⁻¹) into the plasma (q_{plasma}). This plasma uptake rate is equal to the ratio of the relative sensitivity of the plasma-uptake standard in the nebulized sample ($S_{neb,up}$) cps g mL⁻¹) to the absolute sensitivity from microdroplet signal $(S_{\text{drop,up}}, \text{ counts per g}).$

With online microdroplet calibration, microdroplet standards, analyte particles, and sample matrix all are introduced concurrently, and so experience the same plasma conditions, which leads to inherit matrix-matched calibration.36 Because sensitivities from the plasma-uptake standard in the sample and in microdroplets are matrix-matched, online determination of q_{plasma} corrects for sample-introduction related matrix effects.37 To date, we have tested our system for direct quantification of NPs in beverages,35 solutions with variable acid concentrations,36 heavy-element-rich matrices,36 surfactantcontaining matrices,37 waste-water,37 and biologically-relevant matrices.36,37 In all of these cases, online microdroplet calibration has provided accurate element mass quantification, and in those studies where we studied PNC-accurate PNCs. While we have demonstrated online microdroplet calibration with ICP-TOFMS analysis, the approach is also applicable for analysis with quadrupole or sector-field MS instruments; however, signals from multiple element standards will likely need to be obtained in separate runs.38

Microdroplets as NP proxies

Monodisperse microdroplets are an excellent tool to study the effects of instrument parameters and data analysis procedures on the results produced by sp-ICP-MS. Here, microdroplets can be thought of as a nanoparticle proxy: microdroplet-derived ICP-MS signals have comparable temporal durations and sensitivities as their NP counterparts. However, unlike NPs or other particle reference materials, such as element-doped polymer beads,69 researchers can easily adjust the composition of microdroplets by controlling analyte concentration and the size of produced microdroplets. This allows researchers to quickly probe an instrument's analytical figures of merit such as absolute mass detection limits or linear dynamic range for single-particle analysis.23,34 It also allows for the design of unique experiments to quantitatively investigate critical parameters of sp-ICP-MS, such as the ability to accurately count NP signals.

In Fig. 3, we provide a general outline of microdroplet traceranalyte experiments that can be used to assess the accuracy of NP counting by sp-ICP-MS. In these experiments, microdroplets are composed of a solution with at least two elements: one of which serves as the "tracer" and is at high concentration so that it is easily detectable, and the other of which is the "analyte" at a lower concentration to simulate a challenging sp-ICP-MS measurement. Analyte element(s) may also be introduced via pneumatic nebulizer/spray chamber in order to simulate dissolved background. With this setup, we can probe the ability to quantify signals from analyte microdroplets at controlled particle-to-background signal ratios.

Accurate single-particle finding

A limiting factor in the determination of PNCs is the accurate counting of particles. To detect particle signals against a dissolved-ionic background, one must be able to statistically determine the likelihood that a given signal comes from a steady-state background. If that signal is considered "unlikely" to come from the background, i.e. it is an "outlier," then it is counted as a particle. The threshold above which signals can be considered to reside outside the signal distribution from dissolved background is called the "critical value,"70 and is calculated based on the shape of the background distribution and the acceptable false-positive rate as defined by the experimenter. In typical sp-ICP-MS analyses, the background distribution is considered normally distributed and the critical value for particle detection is usually defined as $\mu + n\sigma$, where μ is the average background signal, σ is the background standard deviation, and n is a user-defined factor that controls the number of standard deviations away from the mean to set the single-particle threshold. 53,71 In most cases, n is selected as an integer, and for a Normal Distribution, critical values of 3σ or 5σ have defined false-positive rates of 0.135% or 0.0032%, respectively. For low-count-rate backgrounds, several researchers have proposed defining critical values based on the Poisson distribution, such that the critical value is $\lambda_{\rm bkgd} + n \sqrt{\lambda_{\rm bkgd+1}}$, where $\lambda_{\rm bkgd}$ is the average background count rate and n is an integer. ^{59,72}

For ICP-TOFMS, we have shown that ion signals are neither Gaussian nor Poisson distributed; instead, signals follow a compound-Poisson distribution that is the result of Poissondistributed ion arrival and the response function of the electron-multiplier detection system. 40,41 Because distributions of ICP-TOFMS signals have a unique compound-Poisson shape, accurate separation of background signals from particle signals requires a distinct critical value, $L_{C(ADC)}$, that is calculated via Monte Carlo methods. 40 $L_{C(ADC)}$ is the net-signal critical value, i.e. critical value for background-subtracted-signal, and can be calculated for any false-positive rate, which is defined as the alpha (α) value. For particle identification, we typically set $\alpha =$ 0.01%, and term this the single-particle critical value, $L_{C,sp}$. Every element has a unique $L_{C,sp}$ based on their average background count rates (λ_{bkgd}) in each measurement. Detailed explanation of how element-specific background rates and critical values are determined is beyond the scope of this paper. What is essential for discussion here is that the measurement of monodisperse microdroplets offers a unique possibility to experimentally examine and validate sp-ICP-TOFMS data analysis strategies.

While various signal-thresholding approaches have been proposed to discriminate particle signals from background in sp-ICP-MS, there is no direct way to study the accuracy of singleparticle finding using NPs. Even with well-defined NP reference materials, the possibilities of particle dissolution, aggregation, or adsorption to sample-introduction surfaces always cause uncertainty in the true PNC of sample reaching the ICP-MS. Here, the detection of multi-elemental microdroplets in a tracer-analyte experiment offers an absolute method for

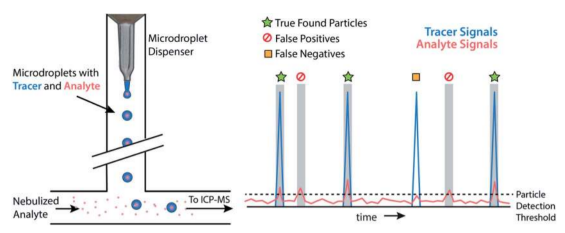


Fig. 3 Schematic diagram of a generic tracer-analyte experiment with microdroplets. Low-abundance "analyte" species in the microdroplets are treated as analyte particles and measured via sp-ICP-TOFMS. The accuracy of analyte particle detection is assessed based on the presence or absence of tracer signal concurrent with the found analyte particle signals.

assessing the accuracy of single-particle detection thresholds. For example, in Fig. 4, we present results in which microdroplets were composed of a solution with 100 ng mL-1 In (tracer) and 10 ng mL⁻¹ Ce (analyte). The microdroplets were 66.7 µm in diameter, so each droplet contained 15 fg 115 In and 1.4 fg ¹⁴⁰Ce, and microdroplets were introduced in a background created by nebulization of a solution containing 1 ng mL⁻¹ Ce. Microdroplets were introduced at 20 Hz for 202.4 s, which resulted in 4048 true microdroplet events. Analyte Ce particle events are then "detected" based on either 3σ or $L_{C.sp.}$ In both cases, data were background-subtracted and split-event corrected prior to particle-event thresholding. After Ce single particles are identified, the amount of tracer signal concurrent with each detected particle event is determined. If 115 In signal is present concurrent with a found Ce-particle signal, then that particle is a true detection event. However, if no ¹¹⁵In signal is present with the found Ce particle event, then the Ce signal

actually originates from the dissolved background and is classified as a false positive.

As can be seen in Fig. 4a, with a 3σ single-particle detection threshold, many false positive events are measured: of the 5782 Ce particles "found," 1805 of them are false positives, which corresponds to a false-positive rate of $\sim 0.9\%$ —well above the 0.135% false-positive rate predicted for a 3σ critical value. Because the number of measured particle-events is always low compared to background events, small discrepancies in falsepositive rates can lead to large errors in particle-number determination. The 3σ particle-event detection threshold is unsatisfactory for sp-ICP-TOFMS because background distributions are not normally distributed, and so the standard deviation does not accurately describe the spread of the background signals. On the other hand, thresholding data with $L_{C,sp}$ results in just 108 false positives, i.e. a 0.05% false-positive rate. However, even with $L_{C,sp}$, we overshoot the number of predicted

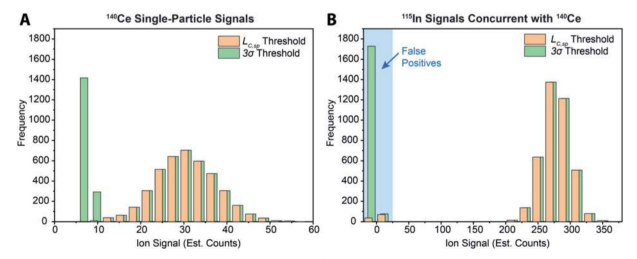


Fig. 4 (A) Histogram of microdroplet signals from analyte cerium (i.e. 140 Ce) found using a 3σ -based threshold and the critical value ($L_{C,sp}$) with the background treated as compound-Poisson distributed. (B) Tracer indium (i.e. 115 In) signals recorded in the same measurements bin as found Ce-particle events. Absence of 115 In signal indicates that the found Ce-particle event did not originate from a microdroplet, i.e. it is a false positive. Data for both Ce and In were background-subtracted and split-event corrected before particle-detection thresholds were applied.

false positives: at an α value of 0.01%, we expect to measure just 20 false-positive events. As will be shown in the next section, about 70% of the false-positive Ce signals are caused by misregistration of data following split-event correction; the remaining false-positive Ce signals are in-line with that predicted by thresholding with $L_{\rm C,sp}$.

In Table 1, we provide NP-detection thresholds and falsepositive rates found for thresholding our Ce-microdroplet data with normal statistics using standard-deviation-based critical values $(3\sigma \text{ and } 5\sigma)$ and Poisson statistics using lambda-based critical values $(3\sqrt{\lambda_{bkgd}+1})$ and $5\sqrt{\lambda_{bkgd}+1}$, ⁵⁹ as well as compound-Poisson based critical values $(L_{C(ADC)})$ for false-positive rates of 0.1% and 0.01%. We find that neither the sigma-based method or the solely Poisson method for critical value assignment is able to accurately predict the number of false positives measured. Reasonable discrimination of singleparticle signals does seem to be possible with both 5σ and $5\sqrt{\lambda_{\rm bkgd}+1}$; however, we still measure a much higher false positive rate than we would expect for signals that were truly Gaussian or Poisson distributed. In the case of unknown particles analysis, it is crucial to be able to predict the number of false positive detection events in order to evaluate their contribution to measured PNCs and particle-mass distributions. Because we know the TOFMS background signal follows a compound-Poisson distribution, it is best to assign NPdetection thresholds according to this distribution. As shown in Table 1, the thresholding particle signals based on $L_{\rm C(ADC)}$ is more robust: the false-positive rates found nearly match the numbers predicted.

Split-event correction

In sp-ICP-MS measurements in which dwell times are larger than the temporal duration of particle events, particle-signals that span across data measurement bins are called "split events."⁷³ If not corrected, these split-event signals will lead to the truncation of signal intensities, and can lead to errors in particle counting because single NP events may be registered as multiple particles. A distinction should be made for between split events with millisecond sp-ICP-MS and sp-ICP-MS in which microsecond dwell times are used. $^{51,74-76}$ In the latter, all NP signal durations are longer than measurement bins and so signals from every NP are "split." Microsecond data acquisition can be used to improve S/N ratio to some extent because less noise is integrated in each measurement bin. 77 We will not discuss microsecond sp-ICP-MS here; instead, we study the necessity of finding and correcting split events for millisecond sp-ICP-MS.

In the beginning of sp-ICP-MS development, the quadrupole-based ICP-MS instruments employed had a settling time between each measurement bin so that data from split events was not recoverable and dwell times were chosen to be long (ca. 10 ms) to reduce the chance of split events.^{78,79} However, most ICP-MS instruments now offer continuous data acquisition, so that single-particle data spread across two or more measurement bins can be put back together.⁸⁰ This is the case for our ICP-TOFMS measurements: with our setup, mass spectra are measured continuously at a rate of \sim 1000 Hz, which is comparable to a "dwell time" of 1 ms, though all isotopes are measured in this "dwell." With a measurement time resolution of 1 ms or higher, signals from single particles are most often contained within a single spectral acquisition, but are sometimes split across two acquisitions.

To correct for split events with sp-ICP-TOFMS, we again must consider the shape of the background-signal distribution. In the case of ICP-TOFMS, the background distribution has a compound-Poisson shape that we can model by Monte Carlo methods. 40 For split-event correction, we'd like to be able to find as many true split events as possible, but also avoid adding noise to the particle distribution through false identification of

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	Threshold ion signal	Number particles found	Number false positives ^a	False- positive rate (%)	Predicted false- positive rate (%)	Error in particle counting (%)
Sigma-based thresholding						
$\sigma_{ m bkgd}$	2.03					
$3\sigma_{ m bkgd}$	6.08	5782	1734	0.867	0.135	42.8
$5\sigma_{ m bkgd}$	10.13	4079	34	0.017	0.003	0.8
Poisson-distribution based thr	resholding					
$\lambda_{ m bkgd}$	3.76					
$3\sqrt{\lambda_{\text{bkgd}}+1}$	6.55	5203	1155	0.578	\sim 0.135	28.5
$5\sqrt{\lambda_{\text{bkgd}}+1}$	10.91	4074	30	0.015	\sim 0.003	0.6
Compound-Poisson based thro	$\mathbf{e}\mathbf{s}\mathbf{holding}^{b}$					
$L_{\rm C(ADC)}$ ($\alpha = 0.1\%$), 3.45 $\sqrt{\lambda_{\rm bkgd}}$	•	4272	225	0.113	0.100	5.5
$L_{\rm C(ADC)}$ ($\alpha = 0.01\%$), $4.22\sqrt{\lambda_{\rm bkgd}} + 2.27$	10.45	4082	37	0.019	0.010	0.8

^a Number of false positives is corrected for mis-registered Ce–In multi-element particle events. $^bL_{C(ADC)}$ expressions are determined empirically from Monte-Carlo simulations of the Poisson-distributed arrival of ions compounded with the detector response function of the TOF detector.

split events. To find split events, we use an outlier approach similar to that used for identifying single NPs, but use a lower split-event (S-E) threshold in order to find split events for low-intensity NPs that might not have a signal above the single-particle threshold (*i.e.* $L_{\rm C,sp}$). In particular, we set the S-E threshold to $L_{\rm C(ADC)}$ with $\alpha=1\%$: split events are found if two adjacent signals are both above the S-E threshold ($L_{\rm C,S-E}$). With the $L_{\rm C,S-E}$ set at a false positive rate of 1%, the chances of having two adjacent events above this critical value is $(0.01)^2$ or 0.01%. Following identification of a split event, the signals are summed and this summed signal is registered into the measurement bin that has the largest ion signal of the pair; a zero is placed in the other measurement bin. For adjacent signals, I_i and I_{i+1} , located at times t_i and t_{i+1} , the split event correction algorithm can be written:

if
$$I_i$$
 and $I_{i+1} > L_{\text{C,S-E}}$
then if $\frac{I_i}{I_i + I_{i+1}} > 0.5$, $I_i = I_i + I_{i+1}$ and $I_{i+1} = 0$,
else $I_i = 0$ and $I_{i+1} = I_i + I_{i+1}$

For sp-ICP-TOFMS, the signals from all m/z-channels of interest are processed individually for split-event correction because multi-elemental composition of NPs cannot be assumed.

In Fig. 5, we provide a subsection of the background-subtracted time traces before and after split-event correction for the same tracer-analyte microdroplet experiment described in the previous section. In these time traces, split-event signals are marked with an asterisk, and it can be seen that split events found for analyte (¹⁴⁰Ce) match those from the tracer (¹¹⁵In). In

Fig. 5c, we plot the number of split-event signals found for the tracer and analyte. Many more split events are found for $^{115}{\rm In}$ because the signal:background ratio (SBR) is over 50 000, whereas the SBR for Ce is $\sim\!\!10$. For $^{115}{\rm In}$ signals, tails of the split-particle events are well above the background and so are easily found based on $L_{\rm C,S-E}$. Because $^{115}{\rm In}$ signals have such high SBR, it is extremely important to find all split events; without S-E correction, In signals would be counted 43% higher than the true number, and the average signal intensity would be 11% too low. For the Ce signals, which have lower SBR, S-E correction is less important; however, no S-E correction would still result in a 5% overshoot in particle number and 5% undershoot in terms of average signal intensity.

After split-event correction, the particle number for both the tracer (115In) matches the number of microdroplets introduced into the ICP: 4048 droplets. The number of found Ce-particle events is slightly more: 4082 events. However, as seen in Fig. 5c, the number of concurrent In-Ce is not quantitatively determined. This occurs because 115 In and 140 Ce signals from split events are summed and then registered into the first or second bin of the split event. For low Ce-signals, ion statistics are a dominate factor in whether the first or second bin of the split event is more intense, and so the ratio of signal from the first to the second bin of a S-E for 140Ce does not necessarily match the ratio of these signals for 115 In. Additionally, in the icpTOF instrument, m/z-dependent travel time through ion optics upstream of the TOF mass analyzer can cause slight m/zdependent time shifts in measured droplet or NP signals.44 Overall, 74 of the 4048 microdroplet signals, or 1.8%, had misregistered 115In-140Ce signals. This multi-element misregistration rate is a limitation of current ICP-TOFMS instrumentation and split-event correction strategies. However, it only

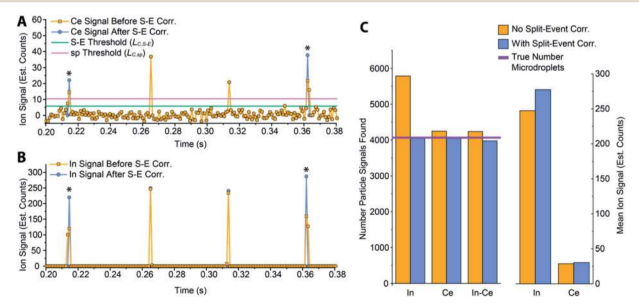


Fig. 5 (A and B) A 180 ms subset of the In–Ce, tracer-analyte experiment. The time traces of the analyte (A) and tracer (B) are presented before and after split-event correction. For split-event correction, a critical value, $L_{C,S-E}$, with a moderate false-positive rate of 1% can be used because identification of a split event requires two adjacent signals to be above $L_{C,S-E}$, which lowers the overall probability. (C) Number of particles found and median signal intensities for both the tracer and analyte elements with and without split-event correction. Tracer and analyte particle signals are found independently.

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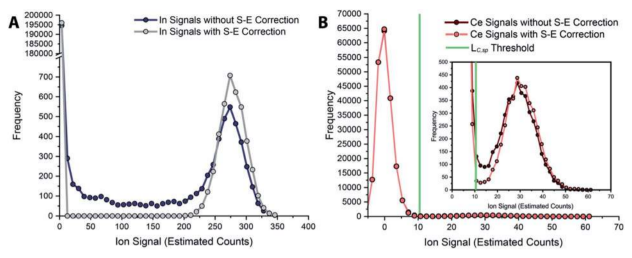


Fig. 6 Histograms of tracer (A) and analyte (B) signals without and with split-event correction. Signals from both datasets are background subtracted.

through the development of analyte-tracer microdroplet experiments that we are able to quantify and begin to understand the possibility of such sources of error in multi-elemental sp-ICP-TOFMS analysis.

In Fig. 6, we report uncorrected and split-event-corrected histograms for 115 In and 140 Ce. As shown, without split-event correction, a broad tail of signal intensities due to partial microdroplet detection events stretches across the ion signal range. Through using $L_{C,S-E}$ —which is controlled by the average dissolved background signal—as a decision threshold for the presence of split-events, there is no defined ratio of high-to-low signals that must be present to flag a split event. 80 In turn, this means that our split-event correction method functions equally well for high-intensity and low-intensity particle signals and will capture as many split-event signals as possible. Independence of split-event correction on particle signal intensity is essential for characterization of unknown particles in real samples, where the particle intensity distributions will likely be broad and varied for different elements present in particle form. For our split-event correction method to function appropriately, the complete signal duration from a single-particle event needs to be less than the duration of one spectral acquisition, e.g. 1 ms for the given experiment, so that a given particle can have signal split across a maximum of two measurement bins. In addition, PNC needs to be low enough to have a low likelihood of single particles in adjacent measurement bins.80 Here, we demonstrate capabilities of split-event correction for a two-element microdroplet system; however, in the future this could be extended to explore split-event correction and particle-signal registration for many-element systems.

Conclusions

For highly time-resolved ICP-MS measurements, microdroplets composed of multi-elemental acidic solutions behave very similarly to nanoparticles with similar total analyte masses (attograms to 100s of femtograms). This similar response allows

us to substitute microdroplets for NP reference materials for the quantification of NP analytes, as well as for the characterization of sp-ICP-MS data collection and analysis strategies. Microdroplets are an attractive tool for use with sp-ICP-TOFMS due to a number of special characteristics:

- (1) Tunable droplet size with highly repeatable volume.
- (2) Controllable generation frequency.
- (3) Flexible elemental composition.
- (4) Quantitative analyte transfer; known injection of discrete element masses into plasma with no sample carryover.
- (5) Well-defined temporal response; comparable to that of NPs.
- (6) Similar detection efficiencies (counts per g) as NPs with ICP-MS.
- (7) Possibility of concurrent microdroplet and nebulized sample introduction.

Here, we have explored how microdroplets can be used to calibrate sp-ICP-MS, both in terms of NP mass and particle number concentration. When combined with sp-ICP-TOFMS, online microdroplet calibration is a powerful method for untargeted quantitative single-particle analysis. This methodology enables detection of NPs *in situ* in varied matrices and at relevant PNCs. We anticipate expanded application and validation of online microdroplet calibration as an alternative to NP-based calibration approaches for sp-ICP-MS.

In addition to the benefits of microdroplets for particle calibration, microdroplets are an ideal medium through which to test performance of sp-ICP-MS measurements. Through the development of tracer-analyte microdroplet experiments, we can—for the first time—provide a truly quantitative assessment of the accuracy of sp-ICP-MS data acquisition and processing procedures. Here, we demonstrate microdroplets as a tool to study approaches for split-event correction and single-particle detection with ICP-TOFMS. In the future, microdroplets as proxies for multi-elemental NPs may serve as tools to improve robustness of multi-elemental fingerprinting approaches.

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Finally, while we focused on use of microdroplets for calibration and as NP proxies, microdroplets can also be carriers of analyte particles. With even the most sophisticated pneumatic nebulization/spray chamber designs, sample transport is nonquantitative and particle-size fractionation in polydisperse aerosols from the nebulizer can lead to sampling biases, which could affect the particles measured by ICP-MS. With microdroplet-based sample introduction, total consumption and quantitative transport of analyte in each microdroplet could reduce sampling bias and thus improve quantification of element-mass distributions in analyte particles or cells. Low volumetric throughput for microdroplet sample introduction remains a challenge for detection of anthropogenic NPs at environmentally relevant PNCs (10² to 10⁶). Nonetheless, we anticipate continued development of high-frequency microdroplet sample introduction systems for improved low-volume sample introduction and also for single-NP and single-cell analyses.

Conflicts of interest

There are no conflicts to declare.

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