Quantification and Clustering of Inorganic Nanoparticles in Wastewater Treatment Plants across Switzerland*

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Abstract: Single particle Inductively Coupled Plasma Time-of-Flight Mass Spectrometry (sp-ICP-TOFMS), in combination with online microdroplet calibration, allows the determination of particle number concentrations (PNCs) and the masses of elements in individual particles. Because sp-ICP-TOFMS analyses of environmental samples produce rich datasets composed of both single-metal nanoparticles (smNPs) and many types of multi-metal NPs (mmNPs), interpretation of these data is well suited to automated analysis schemes. Here, we present a data analysis approach that includes automatic particle detection and elemental mass determinations based on online microdroplet calibration, and unsupervised clustering analysis of mmNPs to identify unique classes of NPs based on their element compositions. To demonstrate the potential of our approach, we analyzed wastewater samples collected from the influent and effluent of five wastewater treatment plants (WWTPs) across Switzerland. We determined elemental masses in individual NPs, as well as PNCs, to estimate the NP removal efficiencies of the individual WWTPs. Through hierarchical clustering, we identified NP classes conserved across all WWTPs; the most abundant particle types were those rich in Ce-La, Fe-Al, Ti-Zr, and Zn-Cu. In addition, we found particle types that are unique to one or a few WWTPs, which could indicate point sources of anthropogenic NPs.

Keywords: Clustering · ICP-TOFMS · Microdroplet calibration · Nanoparticle · Wastewater

Kamyar Mehrabi obtained his Bachelor’s degree from Sharif University of Technology, Tehran, Iran and his Master’s degree from EPF Lausanne in material science and engineering. During his Master’s studies he was a visiting researcher at the Swiss Federal Laboratories for Materials Science and Technology (Empa) in St. Gallen and Brigham and Women’s Hospital (BWH) in Boston, USA. In 2018, he started his doctoral studies in the group of Prof. Detlef Günther at ETH Zürich on multiplexed nanoparticle analyses. He was awarded the METAS prize 2020 for his work on an online microdroplet calibration method to compensate for matrix effects in single-particle ICP-TOFMS.

1. Introduction

In ICP-TOFMS, the complete elemental mass spectrum (from Li to U) is recorded continuously at time resolutions down to 1 ms. At this time resolution, ICP-TOFMS can be used to record NP-derived signals, which are short transients typically between 300–500 µs in duration.[1] Single particle-ICP-TOFMS can be used to quantitatively determine the amounts of several elements and isotopes in single NPs. Through multi-element and multi-isotope detection, sp-ICP-TOFMS provides a more comprehensive analysis of the composition of individual NPs, which is ideal for clustering, sorting, and identifying unique NP types from particle mixtures that occur in environmental samples. However, some elements commonly present in NPs, such as carbon, nitrogen, oxygen, sulphur, and fluorine, are not readily detectable at the single-particle level by ICP-TOFMS.[3] Throughout this article, we use the terms ‘single-metal’ and ‘multi-metal’ NPs (smNP and mmNP) to refer to particles measured with just one and with two or more ICP-TOFMS-detectable elements, respectively.

Distinguishing between naturally occurring NPs (NNPs) and engineered NPs (ENPs) has been the topic of several studies.[3] Although there is ambiguity on the best method of investigation of these particle types, the most prominent NP characteristics used to categorize a NP as either engineered or natural include composition, crystal structure, particle morphology, and shape.[4] With sp-ICP-TOFMS, the simultaneous measurement of most metal and metalloid elements allows for the detection of variable and unique elemental fingerprints from different particle types. Due to manufacturing processes, it is assumed that ENPs will have fairly pure and controlled elemental fingerprints, which will differ from those of NNPs.[5] However, human-based evaluation of elemental associations through visual inspection of data and user-based pattern recognition is extremely time consuming and
is prone to user error and bias. For example, a user might limit the analyte element list to simplify multi-metal patterns in the data and so miss unanticipated multi-metal associations. Similarly, low-abundance multi-metal associations are more likely to be overlooked in non-automated data analysis schemes.

In this article, we show a high-throughput data evaluation approach for non-targeted analysis of NPs measured via sp-ICP-TOFMS. We apply these techniques to extract NP information from sp-ICP-TOFMS measurements of influent and effluent from five wastewater treatment plants (WWTPs) across Switzerland. Through the analysis of mmNP compositions, we report a means – via hierarchical clustering – to discover both conserved and unique mmNP types across the wastewater samples. This mmNP clustering approach provides insights into the origins of various nanoparticle types present in the wastewater samples.

2. Experimental Approach

2.1 Sample Selection and Preparation

We collected influent (I) samples after primary clarification and effluent (E) samples in 500 mL glass media bottles from five WWTPs (Fig. 1) and measured the samples by sp-ICP-TOFMS on the day of collection. Prior to sp-ICP-TOFMS analysis, we sonicated the samples. More details on sample preparation can be found in the original article.[7]

Fig. 1 WWTP sampling sites.[8] For each site, the respective locality as well as the abbreviation for it in the main text are highlighted. Influent samples are indicated by ‘I’ and effluent by ‘E’. This figure is reproduced from Environ. Sci.: Nano, 2021, 8, 1211–1225 with permission from the Royal Society of Chemistry.

2.2 Sample Introduction and MS Measurement

Measurements were performed on an icpTOF-2R (TOFWERK AG, Thun, Switzerland) instrument, which has a native mass resolving power (m/Δm at FWHM) of ~6000 and an extraction frequency of 21.739 kHz. Element-mass and PNC determinations were performed according to our previously published online microdroplet calibration method.[9] As shown in Fig. 2, we used a dual-sample introduction setup, which was composed of nebulizer/spray chamber for wastewater sample introduction and a falling tube and microdroplet generator (50-µm diameter Autodrop Pipette, AD-KH-501-L6, Microdrop Technologies GmbH, Germany) for introduction of microdroplets calibration standards. Droplets were introduced into a vertical falling tube filled with a helium/argon gas[10] mixture and then introduced into the plasma concomitant with sample aerosols from the conventional pneumatic nebulizer/cyclonic spray chamber. A known amount of a plasma-uptake standard (i.e., cesium, Cs) was spiked into the WWTP samples and the microdroplet calibration solution. Through detection of the nebulized and microdroplet-contained Cs, we determined a sample plasma uptake rate for every sample, which allowed us to directly calculate PNCs.

2.3 Data Processing

We developed an automated data analysis procedure to detect, quantify, and cluster smNPs and mmNPs for sp-ICP-TOFMS data. With online microdroplet calibration, we determined the mass of detectable elements in individual particles and the PNCs for each type of found NP.[7,9,11] In addition, our sp-ICP-TOFMS particles detection and clustering software, ‘TOF_NanoFinder’ is available under an open-source license (http://doi.org/10.5905/ethz-1007-363) and is hosted at https://github.com/ethz-tema/TEMAsingleParticle.

We only clustered the multi-metal particles (mmNPs). For smNPs, we obtain mass distributions and PNCs; however, this information is not sufficient to develop meaningful clusters of the smNPs. Therefore, smNPs can be thought of as each representing their own particle cluster. If we had additional information about smNPs, such as shape or crystal structure, we could consider developing clustering approaches to identify the most similar smNP families.

3. Automated Clustering to Discover Multi-metal Nanoparticle Types

The possibility to differentiate between natural and anthropogenic NPs based on multi-metal fingerprints has received considerable interest.[12] However, as an inventory of natural and
Anthropogenic particle types is currently lacking. NP identification based on database searching or spectral matching approaches is not possible. To identify different clusters of mmNPs – which may be assigned to specific sources – we developed a two-stage hierarchical clustering analyses. Clustering was accomplished with a custom-written MATLAB (verR2020b Mathworks, MA, USA) script, accessible by TOF_NanoFinder software; a schematic of our clustering approach is provided in Fig. 3. In the first step of HC, we performed intra-sample clustering of elemental masses present in each mmNP using the correlation distance and averaging method for HC. After calculating the average correlation distance between all the particle-events, we generated an agglomerative hierarchical cluster tree, in which each particle is connected to the cluster tree by a single linkage and linkages are merged together to minimize average distance between all mmNPs. We used an empirically determined cut-off distance of 0.85 to identify major clusters in each wastewater sample, and then created representative mmNP proxies for each of the major clusters found. The mmNP proxies were composed of elements that occurred in more than 10% of the particles part of given cluster and the amplitude of elements in mmNP proxies was the median of the ratio of element mass to the mass of the most-frequent occurring element in the cluster.

In order to compare the similarities between the mmNP clusters developed for the different WWTPs, we performed a second inter-sample HC analysis. In this analysis, we clustered a pooled sample of the representative mmNP proxies from the first HC. For inter-sample clustering of the mmNP proxies, we again used the correlation distance metric with average unweighted linkages, but used a lower cluster cut-off distance of 0.5 to define the major clusters. Subgroups of the major intra-sample clusters, which break up the major clusters into contributions from different WWTPs, are apparent below correlation distances of 0.5 on the resultant dendrogram (see Fig. 5).

5. Characterization of Multi-metal NP Clusters

In Fig. 5, we present results from two-stage HC analysis of the mmNP data from the influent of WWTPs. The major clusters from each WWTP are ‘leaves’ in the cluster tree; names of each leaf correspond to the wastewater samples (i.e. I1–I5) and the most frequent elements in the mmNP cluster. Branches of the cluster tree show the relation of mmNP clusters across the different WWTPs. Overall, we identify 23 mmNP clusters across the five WWTP samples. The dendrogram reveals that some mmNP clusters are conserved across all WWTPs and other branches result from one or a few WWTPs. While HC analysis reduces mmNP data into 23 clusters and 77 leaves; the original data from each of these mmNPs is still accessible. Cluster tree representation offers the possibility to select branches or leaves of interest, and then extract and examine mmNP data from that subset. For example, one can evaluate the PNCs or element masses per particle for any given branch or leaf. For the remainder of this manuscript, we explore selected aspects of the mmNP dataset that become apparent through clustering analysis.

The dendrogram in Fig. 5a shows that some clusters of mmNPs are more conserved than others: mmNP types that exist in all samples include those rich in Ti-Zr, Zn-Cu, Au-Ag, Ce-La, and Zr-Y. The ubiquity of these mmNP types suggests a natural
Fig. 5. a) Dendrogram of hierarchical clustering analysis of samples I1–I5. In this cluster tree, we identify 23 major mmNP types. Each unique mmNP type is plotted in a different color. Six of the unique mmNP clusters are highlighted in colored rectangles. For these mmNP clusters, the representative mmNP proxies with elements present at an occurrence rate limit of 1% are plotted in b–g. Note that not all mmNPs types are found in all WWTP influent samples. While the Ti-Zr, Zn-Cu, Au-Ag, Ce-La, and Zr-Y mmNP clusters are found in all influent samples, the Bi-V-type mmNPs are only found in I1 and I3. Error bars mean in b–g are standard deviation of normalized masses. This figure is reproduced from Environ. Sci.: Nano 2021, 8, 1211–1225 with permission from the Royal Society of Chemistry.

origin, or at least common anthropogenic sources; if mmNPs were from specific point sources of anthropogenic NP pollution, they would likely be found only in selected WWTPs. In addition, we know from previous research that our mmNP types match, at least in part, results from studies focused on Ce, Ti, Zr, and Zn natural NPs.[3,5,12,14] In Fig. 5b–g, we extract data from selected mmNP clusters and plot the normalized mass of elements that are measured in at least 1% of the particles that make up each clusters. Normalization was done at the particle level: the quantified masses of each element were divided by the mass of the most frequently occurring element from the cluster. For example, in Fig. 3h–g, the normalizing elements were Ti, Zn, Ag, V, Ce, and Zr, respectively. The rather well conserved element ratios in these mmNP clusters further emphasizes that these mmNP clusters likely originate from comparable sources and are likely naturally occurring.

In addition to ubiquitous NNP, hierarchical clustering also identifies uncommon particle types including NPs rich in Au-Ag, Rh-Pd, Bi-V-(Mo), Sb-W, Cr-Ni, Zn-Cu, Mn-Cu, and Ba-La. We find a Bi-V-rich mmNP cluster in two of the WWTP influent samples. As shown in Fig. 5e, the mean mass ratios of Bi:V in this cluster are 5:1 in I3 and 6.8:1 in I1. Because 209Bi is about four-times heavier than 51V, the measured element ratios are reasonably consistent with a one-to-one atomic ratio (4.1:1, Bi:V mass ratio) of bismuth vanadate (BiVO₄), especially considering the low numbers of particles recorded. Bismuth vanadate – which is sometimes doped with Mo (as also found in the Bi-V-Mo cluster from I3) – is used as yellow pigment in industry[15] and as a catalyst,[16] and is very likely anthropogenic in origin. Another unexpected class is the Au-Ag cluster (Fig. 3d) that contains 5–10 times more Au than Ag on a mass basis. We find the Au-Ag cluster in all WWTP samples from both of our collection days. These NPs may originate from gold alloys used in the jewellery manufacturing industry, and thus are likely anthropogenic in origin.
6. Conclusions

We presented a strategy for the analysis of NPs in environmental waters or other water samples. Wastewater samples are complex and likely have a mix of different types of natural and anthropogenic NPs. Here, we created rich sp-ICP-TOFMS datasets composed of a range of smNPs and mmNP types. To break down the inherent complexity of sp-ICP-TOFMS datasets and reduce them to an interpretable form, we combined online microdroplet calibration with an automated data analysis system. With this approach, we are able to extract NP data from raw ICP-TOFMS time traces, determine the element masses in single NPs, record PNCs, correct for both high-throughput data collection and data analysis, we are now able to rapidly deliver multi-element data – for example, through the identification of unique mmNP types.

Interpretation of our sp-ICP-TOFMS data set in terms of NP origin is an ongoing process and we expect continued refinement of our understandings. In fact, the continued refinement of our knowledge about mmNP fingerprints (both of natural and anthropogenic NP types) and the expected types of mmNPs in various environmental compartments is a substantial benefit of our approach. Because our approach allows for both high-throughput data collection and data analysis, we are now able to expand our inventory of NPs – which will, in turn, improve our classification models. An eventual aim of such studies is to establish a database approach for the identification of different NP types.

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