**RESEARCH PAPER** 



# Detection of nanoparticles in edible plant tissues exposed to nano-copper using single-particle ICP-MS

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Abstract The increasing use of nanopesticides has raised concerns about their effects on crop plants and the impact of human health as well as ecological effects. While increased uptake of metal ions has been observed before, to date, very few studies have demonstrated the presence of nanoparticles in edible tissues. Singleparticle inductively coupled plasma–mass spectrometry (sp-ICP-MS) has been suggested as a powerful tool to detect inorganic nanoparticles (NPs) in environmental samples. Here, we exposed edible plant tissues from lettuce, kale, and collard green to nano-CuO, simulating its use as a nanopesticide. We applied sp-ICP-MS to demonstrate the presence of nanoparticles, both in the water used to rinse crop leaf surfaces exposed to nano-

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J. Nelson Agilent Technologies Inc., Santa Clara, CA, USA CuO and within the leaf tissues. Lettuces retained the highest amounts of nCuO NPs on the leaf surface, followed by collard green and then kale. Surface hydrophilicity and roughness of the leaf surfaces played an important role in retaining nano-CuO. The results indicate that most of the nanoparticles are removed via washing, but that a certain fraction is taken up by the leaves and can result in human exposure, albeit at low levels.

Keywords sp-ICP-MS  $\cdot$  Copper  $\cdot$  Vegetable  $\cdot$  Inorganic nanoparticles extraction  $\cdot$  Retention  $\cdot$  Sensors

#### Introduction

Determining the environmental concentrations of engineered nanomaterials (ENMs) is a major challenge for assessing their potential implications. Without this information, it is not possible to establish whether they pose a human or ecological risk. To overcome this challenge, predictive tools (Nowack 2017) have been developed to generate estimates using material flow analysis at global (Keller et al. 2013; Song et al. 2017), continental (Lazareva and Keller 2014; Liu et al. 2014; Sun et al. 2017), regional (Bornhoft et al. 2016; Gottschalk and Nowack 2011; Gottschalk et al. 2013; Keller and Lazareva 2014; Sun et al. 2014), and local levels (Keller and Lazareva 2014; Lazareva and Keller 2014). Detailed fate and transport models have recently been developed to go a step further and predict the residence time of ENMs, their concentrations, and their accumulation in environmental compartments (e.g., soils, sediments) (Dale et al. 2015; Garner et al. 2017; Liu et al. 2015; Praetorius et al. 2012; Meesters et al. 2014). While these estimates can serve to bridge the gap, they rely on a large number of assumptions and generalizations and have a wide range of uncertainty. Accurate detection of nanoparticles (NPs) in different environmental media is needed, to evaluate the predictions and risk assessments. There are a number of methods available for detecting the presence of inorganic ENMs in different media, with various limitations. In particular, detecting ENMs at the low concentrations predicted (parts per billion or less) to be present in most media is a major challenge. In addition, for ENMs that transform, there are additional challenges in determining whether the NP detected is the original one or a transformed one.

The most common methods for detecting NPs in an aqueous media are dynamic light scattering (DLS) and UV/Vis spectrophotometry. While these approaches have been used extensively to study the behavior of NPs in aqueous media (e.g., aggregation, sedimentation, stability of the suspension) (Adeleye et al. 2014; Burns et al. 1997; Chen and Elimelech 2006; Chowdhury et al. 2012; Conway et al. 2015b; Domingos et al. 2009; Keller et al. 2010; Thio et al. 2011; Tiraferri et al. 2008) and can be used to track NP concentrations over time, they have several limitations. Concentrations have to be sufficiently high (typically > 10 mg/L) for accurate detection; there is no information on NP composition, and in the case of UV/Vis, the wavelength needs to be tailored to the particle in question, bi- or multimodal size fractions cannot be handled well; and the presence of other (e.g., natural or incidental) NPs would seriously affect the measurement. Thus, these approaches cannot be used for more realistic samples (e.g., multimodal size, other non-target NPs). The nanoparticle tracking analysis system (Gallego-Urrea et al. 2011; Mehrabi et al. 2017; Mitrano et al. 2017; Shang and Gao 2014) offers some improvements, since it can track particles with more complex particle-size distributions, but also requires concentrations in the ppm range, provides no information on composition, and would be challenging to use with more realistic samples containing natural and incidental NPs. These approaches will continue to be extremely useful for understanding the fundamental behavior of NPs in different media, but there is a need to be aware of the limitations.

Transmission electron microscopy (TEM) is useful for visually detecting NPs in a sample and may be used to determine the number of particles in a subset of the sample, as well as their shape and size distribution (Hassellöv and Kaegi 2009; Mitrano et al. 2017; Plathe et al. 2013; Shang and Gao 2014; Su et al. 2015; Zhou et al. 2013); a large number of subsamples may be needed to ensure reproducibility. However, the probability of detecting NPs in a sample via TEM decreases rapidly as the NP concentration decreases. Therefore, high to very high initial concentrations are needed for best visualization. Feature composition can be determined using X-ray diffraction and energydispersive X-ray spectroscopy (Adeleye et al. 2016; Mitrano et al. 2017; Plathe et al. 2013; Su et al. 2015; Zhou et al. 2013), although confirmation that these indeed are the introduced NPs may be challenging. Xray absorption near edge structure and extended X-Ray absorption fine structure can also be used to confirm the presence of NPs within a sample, with exquisite precision as to their location within a particular cell or tissue (Gardea-Torresdey et al. 2014; Hernandez-Viezcas et al. 2016; Rui et al. 2015), but also requires high ENM exposure concentrations, as well as access to the beam lines. This method cannot by itself provide information on the concentration of the particles, although it does provide speciation, which can be very useful for confirming the presence of the target ENM.

Single-particle inductively coupled plasma-mass spectrometry (sp-ICP-MS) has the potential for addressing many of these limitations for inorganic NPs. It can provide information on the number of particles in a sample and their mass concentration, particle size, and elemental composition. Laborda et al. present a very detailed explanation of the principles and key considerations for successful sp-ICP-MS (Laborda et al. 2013, 2016). Earlier studies discussed a number of challenges even for the measurement of very well-defined NPs in simple aqueous matrices (Liu et al. 2017; Mitrano et al. 2012, 2014; Reed et al. 2012), but more recent work indicates that this technique is becoming more common for aqueous samples (Montano et al. 2016). As the technique has matured, it has been applied to a wider range of elemental compositions (Frechette-Viens et al. 2017; Lee et al. 2014; Praetorius et al. 2017) and samples (Azodi et al. 2016; Hadioui et al. 2015; Schwertfeger et al. 2016). A few studies have reported the use of sp-ICP-MS in food products, for example to determine the presence of nanoTiO<sub>2</sub> in different processed foods (Peters et al. 2014) or nanoAg in food stimulants (Linsinger et al. 2014). sp-ICP-MS was also used to track the release of copper-based NPs from an external marine paint (Adeleye et al. 2016). One significant advantage of sp-ICP-MS is the ability to detect NPs at very low concentrations (a few parts per billion) and in fact requires dilution of concentrated samples, overcoming a major limitation in having realistic exposure concentrations.

Copper has been used extensively in agriculture to protect against diseases (Keller et al. 2017), particularly fungal infections and other microbial pathogens (El-Habbaa et al. 2016; Salma et al. 2015). Recently a number of copper-based nanopesticides have been introduced in the market. Early studies indicate that in general, these nanopesticides cause little or no negative physiological effects at the recommended doses, and that in fact, plants may gain biomass when protected with these materials (Elmer and White 2016; Ma et al. 2015; Stampoulis et al. 2009; Trujillo-Reyes et al. 2014; Zuverza-Mena et al. 2015). Metabolomics studies do indicate that several crop plants respond to the exposure to these copper-based nanopesticides by using some of their antioxidants (Zhao et al. 2016a, b, c, d, 2017a, b, c, d, e), as well as upregulating metal chelators and downregulating some organic acids in root exudates (Huang et al. 2017), to minimize the impact of excess copper.

The objective of this study was to detect and quantify the presence of nanoparticles in edible plant tissues exposed to nano-CuO using sp-ICP-MS. Three different types of leaf tissues were evaluated, to determine to what extent leaf surface roughness influences the retention of NPs. In addition, we sought to determine whether NPs would enter the leaf tissues and remain within them even after washing.

# Materials and methods

*Reagents* Ultrapure water (DI) was obtained from a Barnstead NANOpure water purification system (Thermo Scientific, Boston, MA, USA) and used throughout the work. Nitric acid (67–70%) of ultra high purity for quantitative trace metal analysis at the partsper-trillion (ppt) level (BDH Aristar® Ultra grade) was used to prepare calibration standards. A certified copper reference standard solution of 1000  $\mu$ g/mL in 2% nitric acid was obtained from High-Purity Standards, Inc. (Charleston, SC, USA).

*Nanomaterials* Gold NPs with nominal diameters of  $60 \pm 4$  nm were purchased from nanoComposix (San Diego, CA) as bare, dispersible, gold nanospheres in aqueous suspension (2 mM sodium citrate) and are hereafter referenced by their nominal diameters.

Uncoated nano-CuO was purchased from U.S. Research Nanomaterials (Houston, TX, USA); a detailed characterization was presented in a previous study (Adeleye et al. 2014). Briefly, the primary particle size is 20–100 nm (as provided by the manufacturer) and the hydrodynamic diameter (HDD) is  $280 \pm 15$  nm in DI water at pH 7 (0.5 mM phosphate buffer). The surface charge in 0.5 mM phosphate buffer solution is  $-34.4 \pm$ 0.5 mV at pH 7, with the isoelectric point (IEP) at 6.3. The helium density of nano-CuO is 6.349 g/cm<sup>3</sup> with 74.3 ± 1.2 wt% copper content and the balance is oxygen.

*Leaf tissues* Three USDA organic vegetables were purchased on the day of the experiment to ensure freshness. The vegetables, kale (*Brassica oleracea*, var. *Acephala Lacinato*), lettuce (*Lactuca sativa* var. *green leaf cultivar*), and collard green (*Brassica oleracea*, var. *Acephala*) were produced by Cal-Organic Farms (Bakersfield, CA, USA). The surface roughness of the leaf tissues and microscale details were visualized using an environmental scanning electron microscope (ESEM, Philips Electron Optics, Eindoven, The Netherlands).

Exposure and retention of nanoparticles To quantitatively analyze the exposure and retention of nano-Cu onto leaf tissues, vegetable leaves from each type were cut into a  $75 \times 75$  mm square (Fig. S1). Nano-CuO was suspended in DI water and sonicated for 30 min before being applied to the leaf tissues. Ten drops of 20 µL nano-CuO suspension (1 mg/L particle concentration) were evenly spread onto each leaf square with an analytical pipette to ensure that all the vegetable tissues were exposed to equal amounts of nano-CuO (Fig. S2). The final amount of nano-CuO applied onto leaf tissues was  $34.4 \text{ g/m}^2$ , based on the recommendation for copper-based nanopesticides (Conway et al. 2015a). The sprayed-on nano-CuO suspension was air-dried on the surface of the leaf tissues for 2 h (Fig. S3). Then, the nano-CuO-exposed leaves were rinsed three times with 10 mL DI water, and the drained water of each rinse cycle was collected in metal-free polypropylene tubes (VWR International, West Chester, PA, USA). The collected rinse water samples were analyzed via sp-ICP-

MS to detect NPs and determine the size and particle concentration of nano-CuO. Triplicates from each type of vegetable leaves were treated; control samples were also rinsed following the same procedure. The morphology and particle size of nano-CuO before and after application were studied via electron microscopy using a FEI Tecnai G2 Sphera 200 kV TEM.

Enzymatic digestion To investigate if nano-CuO remained on the rinsed leaf surfaces or entered the leaf tissues via the stomata, Macerozyme R-10 (bioWORLD, Dublin, OH, USA) was used to digest the plant tissues for NP extraction at neutral pH. Macerozyme R-10 is a multi-component (e.g., cellulase, hemicellulase, and pectinase) enzyme mixture, which has been shown to digest plant tissues without changing the NP size (Dan et al. 2015). Circular pieces of leaf tissue (diameter = 6.35 mm) were cut near the area where nano-CuO was applied and also downstream of rinse water runoff, after each rinse (Fig. S3). Each sample was digested in 2 mL of the enzyme solution (containing 100 mg R-10 powder). The samples were shaken with the enzyme solution at 25 °C for 24 h to ensure complete digestion of leaf tissues. A final centrifugation of the digested samples at 20,000×g was done for 20 min, and then the supernatant (0.1 mL) was diluted 100 times for sp-ICP-MS analysis.

sp-ICP-MS measurement An Agilent 7900 ICP-MS (Santa Clara, CA, USA) equipped with standard nickel sampling and skimmer cones, standard glass concentric nebulizer, quartz spray chamber, and small internal diameter (1.0 mm) quartz torch was used to perform sp-ICP-MS analysis to determine the particle size and concentration in all collected samples. Samples were introduced directly into the ICP-MS with the standard peristaltic pump and tubing (i.d. 1.02 mm). Analyses were performed in time-resolved analysis (TRA) mode using an integration time (dwell time) of 100 µs per point with no settling time between measurements. The instrumental settings used for the sp-ICP-MS analysis are summarized in Table 1. The sp-ICP-MS method setup, data collection, and analysis were conducted via the Single Nanoparticle Application Module (method wizard) in the Agilent ICP-MS MassHunter software (Version C.01.03 Build 505.16 Patch 3).

The 60 nm gold NP standard was diluted to 100 ng/L Au (99%) with DI in metal-free polypropylene tubes, to evaluate the nebulization efficiency to be used in the

 Table 1
 Optimized instrumental setting for the nano-CuO analysis in sp-ICP-MS (Agilent 7900)

Parameter	Value		
RF power	1550 W		
Carrier gas	0.67 L/min		
Spray chamber temperature	2 °C		
Nebulizer pump	0.1 rps		
Sample depth	8.0 mm		
Integration time	100 µs		
Acquisition time	60 s		
Mass monitored	<sup>63</sup> Cu		

data conversion from raw signal to NP size. A Cu ionic standard of 1  $\mu$ g/L was prepared with 1% HNO<sub>3</sub> and was used to determine the elemental response factor. The samples were diluted with DI water to ensure the nano-CuO particle concentration was between 10 and 100 ng/L, and a sample inlet flow of 0.346 mL/min was used. Analyte mass fraction was set to 0.75 while CuO particle density was set to 6 g/cm<sup>3</sup>. NP sample preparation and dilution were performed on the day of the analysis to avoid sample degradation and minimize nano-CuO dissolution after processing. Before dilution of the samples and again prior to their analysis, all solutions were placed in an ultrasonic bath for 10 min to ensure that the samples were fully homogenized.

# **Results and discussion**

*Characteristics of nano-CuO* Nano-CuO as received forms agglomerates of 30–100 nm-size particles, shown in TEM images (Fig. 1a), which agrees with the size information provided by the manufacturer (20–100 nm). Individual 30–50 nm nano-CuO primary particles can be identified in Fig. 1b. In DI, agglomerates exhibited a larger size of 500–1000 nm (Fig. 1c). The TEM images indicate most of the primary nano-CuO particles are nearly spherical, while the aggregates appeared as rods and flakes (Fig. 1c). NP morphology is important for the accuracy of particle size measurements since NP sizes are calculated based on the assumption that the NPs are spherical (Sannac 2015). Thus, the measured size of nano-CuO agglomerates by sp-ICP-MS may show some difference from other techniques, for example, TEM. In

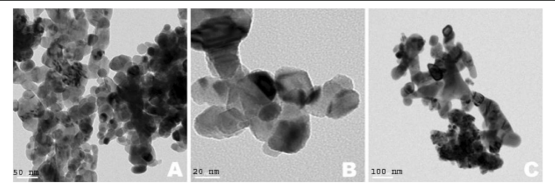


Fig. 1 TEM images of nano-CuO with a scale bar of a 50, b 20, and c 100 nm

a previous study (Adeleye et al. 2014), the same nano-CuO were characterized with dynamic light scattering (Malvern Zetasizer Nano-ZS90) to obtain the HDD, which was reported as  $280 \pm 15$  nm in DI water at pH 7 (0.5 mM phosphate buffer).

The calibration of nano-CuO concentration in DI water was obtained by analyzing from 1 to 500 ng/L (Fig. 2a), with very good linearity ( $R^2 = 0.993$ ) across the entire concentration range. When the nominal nano-CuO concentration was higher than 100 µg/L (e.g., 250 and 500 µg/L), the measured nano-CuO concentration exhibited a larger root square difference (RSD ~ 12%), indicating decreased precision. Similar RSD values were obtained when the nominal nano-CuO concentration was lower than 10 µg/L (e.g., at 1 µg/L, RSD = 9.1%), which suggests the optimal particle concentration for analysis is between

10 and 100  $\mu$ g/L (RSD < 5%). In the following experiments, all the samples were diluted to the concentration range of 10–100  $\mu$ g/L as nano-CuO.

Table 2 summarizes the ionic Cu concentration, median size, and mean size of nano-CuO aqueous samples as measured by sp-ICP-MS. The ionic Cu concentration reflects the dissolved fraction within the initial nano-CuO suspension. As the nominal nano-CuO concentration increased, the detected/ calculated ionic Cu concentration increased more markedly, especially at higher nano-CuO concentrations (e.g., 250 and 500  $\mu$ g/L nano-CuO). Since the nano-CuO suspension was prepared fresh on the day of analysis, minimal dissolution of CuO was expected (Adeleye et al. 2014). At NP concentrations higher than 100  $\mu$ g/L, there may be more than one NP per monitored event, or there

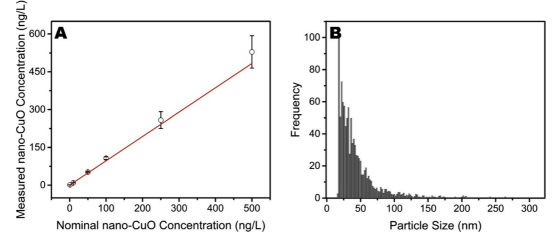


Fig. 2 a Calibration curve of nano-CuO concentration in DI water; symbols represent experimental data, and red line represents a linear regression. b Size distribution of nano-CuO (at 50  $\mu$ g/L) measured by sp-ICP-MS.

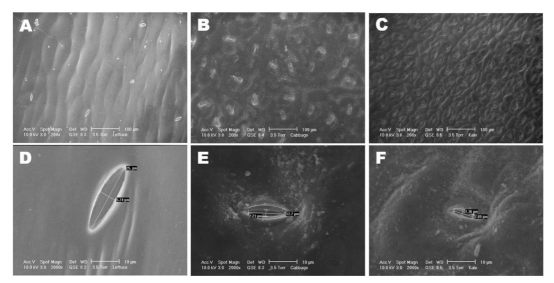
Nominal nano-CuO concentration (ng/L)	Ionic Cu concentration (ng/L)			Median size (nm)			Mean size (nm)		
	Avg.	Std.	RSD (%)	Avg.	Std.	RSD (%)	Avg.	Std.	RSD (%)
1	0.11	0.01	9.3	16.48	0.81	4.9	18.51	1.56	8.4
10	0.39	0.04	10.8	15.99	0.00	0.0	23.62	0.54	2.3
50	2.33	0.06	2.5	24.84	4.40	17.7	36.73	3.62	9.9
100	2.78	0.20	7.3	25.26	2.36	9.3	37.15	2.24	6.0
250	32.12	3.36	10.4	31.76	0.11	0.3	34.01	0.60	1.8
500	35.10	2.47	7.0	31.63	0.10	0.3	34.01	0.72	2.1

Table 2 Ionic Cu concentration, median, and mean size of nano-CuO aqueous samples at different nominal nano-CuO concentrations, as measured by sp-ICP-MS

could be "carry-over" from the previous event. Thus, at higher NP concentrations, a fraction of the signal may be recorded as dissolved Cu ion instead of CuO NPs, as observed by Montano et al. (2016). Thus, it is best to dilute the nano-CuO samples to <100  $\mu$ g/L. The reported median and mean size of nano-CuO ranged from 15 to 37 nm at different nominal nano-CuO concentrations (Table 2). However, the primary particle mean size was in the 24–37 nm range at concentrations >10  $\mu$ g/L, with a size distribution from 20 to 210 nm (Fig. 2b), which agrees well with the TEM results. Although the majority of the CuO NPs in solution are agglomerates (Fig. 1), some

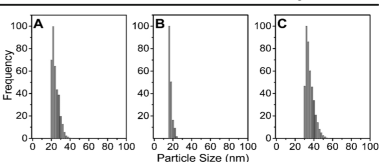
may be soft agglomerates, releasing primary particles or smaller agglomerates as they pass through the sp-ICP-MS (Fig. 2b).

*Characteristics of leaf tissues* The surface morphology of selected editable vegetable leaf tissues was characterized via ESEM, as shown in Fig. 3. The lettuce surface had the smoothest texture (Fig. 3a) and was the most hydrophilic with good wettability (Fig. S2A). Kale leaf tissues exhibited the roughest surface with dense texture (Fig. 3c), and very good hydrophobicity with a small contact angle observed when an aqueous droplet was added (Fig. S2C). Collard green also showed good hydrophobicity when interacting with aqueous droplets



**Fig. 3** Surface morphology of **a** green leaf lettuce, **b** collard green, and **c** kale leaf tissues and stomatal openings of **d** green leaf lettuce, **e** collard green, and **f** kale leaf tissues. ESEM images were obtained under a magnification of  $\times 200$  and  $\times 2000$ , respectively

Fig. 4 Particle size distribution of Cu-based nanoparticles from the initial rinse of as-purchased leaf tissues of a green leaf lettuce, b collard green, and c kale. Data obtained via sp-ICP-MS measurement



(Fig. S2B) with some roughness (Fig. 3b). Under higher magnification, the stomatal openings of the various leaves could be observed (Fig. 3d–f). All three species had stomatal openings from 1 to 25  $\mu$ m; lettuce had the largest stomatal openings while kale had the smallest.

Before applying the nano-CuO suspension onto leaf tissues,  $75 \times 75$  mm samples of as-received leaves were rinsed with 10 mL DI water three times. The collected rinse water was analyzed by sp-ICP-MS to determine whether the unexposed leaves contained nanosized Cubased particles. As no information was available regarding the shape, density, and composition of these particles, an assumption was made that any initial particles had the same physicochemical properties as nano-CuO (e.g., density, Cu content, and density). In all three leaf types, Cubased NPs were detected, mostly in the 20-50 nm size range (Fig. 4). However, these NPs were at very low concentration, from 30 to 90 ng/L (Fig. 5) in the first rinse. Additional rinses removed most of these Cu-based NPs. For example, the NP concentrations in the third rinse were 95% less than the first rinse for lettuce and collard green (Fig. 5). The higher roughness of kale (Furmidge 1962) may have shielded some of the NPs from the rinse water, since the third rinse still had 43% of the concentration of the first rinse. These NPs may be from dust in the fields or steps along the distribution chain.

*Nano-CuO in leaf tissue rinse water* Nano-CuO was detected via sp-ICP-MS in the rinse water of all three leaf tissues after exposure to nano-CuO (Fig. 6). In the first rinse, most of the detected nano-CuO exhibited sizes in the range between 30 and 50 nm (as shown in Fig. 6a, d, g), which agree with the original primary size distribution. Aggregates with particle sizes up to 400 nm were also detected in the first rinse water, although at much lower frequency compared to primary size NPs (Fig. 6a. d. g). In the second rinse cycle, the sizes of

nano-CuO particles in the rinse water were larger (Fig. 6b, e, h), especially in the exposed lettuce leaf tissues, with a mean size of 230 nm (Table 3). By the third rinse, the remaining Cu-based NPs have a wide size distribution (Figs. 6c, f, i). These larger NPs may have been trapped in surface crevasses or even stomata and would take more effort to remove via rinsing. The same effect is seen in all three types of vegetables.

Nano-CuO concentrations in the first rinse water were around 500–750  $\mu$ g/L (Fig. 7, note the logarithmic scale), with the highest concentration detected in collard green leaf rinse water (752.5  $\mu$ g/L), followed closely by kale, and the lowest concentration in lettuce leaves (508.6  $\mu$ g/L). There is a strong relationship between first rinse water concentration and leaf surface characteristics. Kale leaf has the highest surface hydrophobicity as well as surface roughness (Fig. 3), which likely channels the applied NP suspension and rinse water through a limited region; lettuce leaves are smoother and much more hydrophilic, spreading the

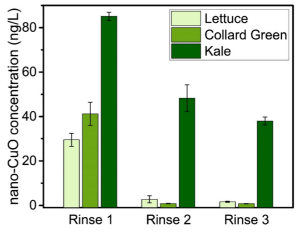


Fig. 5 Cu-based NP concentration in rinse water from aspurchased leaf tissues. Data obtained via sp-ICP-MS measurement

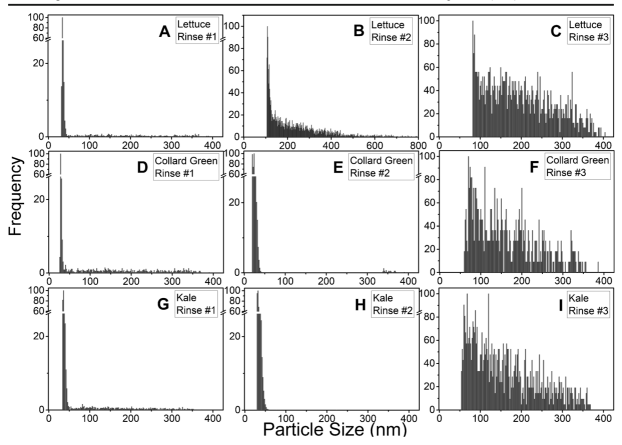


Fig. 6 Particle size distribution in rinse water of nano-CuO exposed lettuce leaves after a first rinse, b second rinse, and c third rinse; collard green leaves after d first rinse, e second rinse, and f

third rinse; and kale leaves after  ${\bf g}$  first rinse,  ${\bf h}$  second rinse, and  ${\bf i}$ third rinse. Data obtained via sp-ICP-MS

applied nano-CuO suspension. Thus, nano-CuO were more easily flushed out by the rinse water from the surface of kale and collard green than from lettuce; these results may vary depending on nanopesticide

Species	Rinse cycle	Median siz	e (nm)		Mean size (nm)			
		Avg.	Std. Dev.	RSD (%)	Avg.	Std. Dev.	RSD (%)	
Lettuce	1	34.80	1.88	5.4	45.61	5.21	11.4	
Lettuce	2	233.68	10.51	4.5	187.09	14.46	6.3	
Lettuce	3	236.17	12.79	5.4	221.87	17.88	8.1	
Collard green	1	30.36	2.53	8.3	78.32	6.09	7.8	
Collard green	2	22.92	1.68	7.3	23.76	1.96	8.2	
Collard green	3	135.48	13.01	9.6	153.04	11.77	7.7	
Kale	1	35.14	2.63	7.5	76.00	3.40	4.5	
Kale	2	32.63	0.74	2.3	36.37	3.16	8.7	
Kale	3	135.48	13.01	9.6	163.91	11.98	7.3	

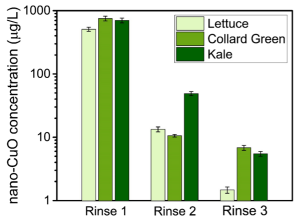


Fig. 7 Particle concentration in rinse water from leaf tissues exposed to nano-CuO. Data obtained via sp-ICP-MS.

application method. The subsequent rinses decreased the nano-CuO concentrations significantly in all three leaf types (Fig. 7), although more pronouncedly in lettuce leaves.

*Nano-CuO in leaf tissues* Since the stomatal openings of the leaf tissues are in the microscale (Fig. 3), it is possible that nano-CuO enter leaf tissues during the exposure (Zhao et al. 2016b). In addition, some nano-CuO may remain on the leaf surfaces even after several rinses. Figure S4 presents images of the positions where the samples were taken from controls leaves and leaf squares exposed to nano-CuO. Nano-CuO particles were detected in the enzymatically digested leaf tissue samples (Fig. 8). The size distribution of released nano-CuO from the enzymatic digestion ranges from 50 to

150 nm (Fig. 8) and is generally consistent for all three leaf types.

Compared to unexposed controls, nano-CuO concentrations in the exposed crops were 2-3 orders of magnitude greater near the location where nano-CuO was applied (Fig. 9a, note the logarithmic scale). Lettuce retained the highest amount of nano-CuO, followed by collard green and then kale, which was highly related to the stomatal opening sizes on the leaf surfaces (Fig. 3). Lettuce had the largest stomatal openings, making it easier for nano-CuO to enter its tissue (Larue et al. 2014; Zhao et al. 2016b). After the DI water rinse, the residual nano-CuO on or within lettuce tissues decreased significantly from 430 to  $\sim$ 180  $\mu$ g/L, while residual nano-CuO levels on or within collard green (~100  $\mu$ g/L) and kale (~ 45  $\mu$ g/L) leaf tissues were approximately the same rinse after rinse. This suggests that while lettuce retains the most nano-CuO, it is easiest to remove them via rinsing, in part due to its higher surface hydrophilicity and larger stomatal openings.

For the samples taken further downstream from the position where nano-CuO was applied (see Fig. S4), there were no significant differences in nano-CuO levels between unexposed controls and exposed collard green and kale (Fig. 9b), suggesting that nano-CuO transported by the rinse water is not deposited further downstream for these leaf surfaces. However, nano-CuO concentrations in these downstream locations from exposed lettuce tissues were 10–30-fold higher than control lettuce leaves. The hydrophilic surface on the lettuce leaves may enhance the mobility of nano-CuO.

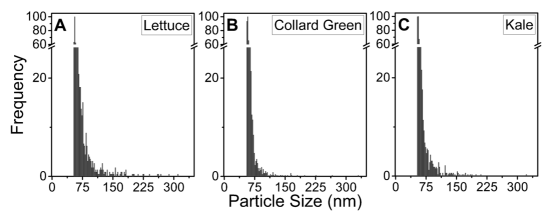


Fig. 8 Nano-CuO particle size distribution in leaves exposed to nano-CuO, rinsed three times and then digested enzymatically: a lettuce, b collard green, and c kale. Data obtained via sp-ICP-MS

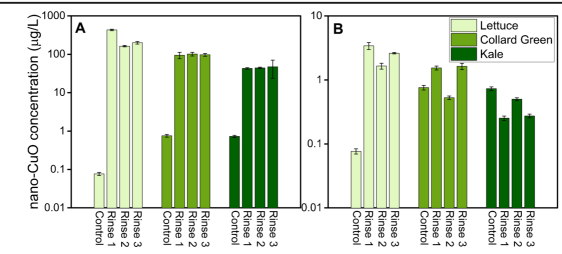


Fig. 9 Particle concentration in enzymatically digested leaf tissues after exposure to nano-CuO spot sampled from positions:  $\mathbf{a}$  close to where the nano-CuO suspension was applied and  $\mathbf{b}$  further downstream of rinse water runoff. Data obtained via sp-ICP-MS.

Thus, nano-CuO may enter lettuce tissues via the stomata during the rinse process as well. Note that the concentrations in these locations are orders of magnitude smaller than near the point of nano-CuO application. In any case, the residual levels of Cu are well below any toxicity concern for humans, based on FDA guidance for daily Cu intake (Zhao et al. 2016a).

### Conclusions

We developed a sp-ICP-MS method for detecting NPs in environmental samples, including the water used to rinse exposed crop tissues and plant tissue that were enzymatically digested. sp-ICP-MS can accurately measure nanoparticle size, particle concentration, and dissolved Cu concentration in aqueous samples with low RSD values and very good particle concentration calibration linearity. NPs were detected in the ppb range, and the best conditions for analysis were within 10–100 ppb.

Three different species of edible vegetable leaf tissues, including lettuce, collard green, and kale, were exposed to nano-CuO. Particles were found in all rinse water samples, as individual nanoparticles as well as aggregates. The concentration of nano-CuO in rinse water was highly related to leaf surface characteristics. The highly hydrophilic and smooth lettuce leaves retained fewer nanoparticles after washing. Kale and collard green leaf tissues, with higher surface roughness and somewhat lower hydrophobic affinity, retained more nanoparticles even after three washes. From a practical perspective, washing does remove a significant amount of the nano-CuO, but rougher leaf surfaces will retain more.

Digestion of controls and exposed leaf tissues with macerozyme R-10, followed by sp-ICP-MS analysis, indicates that a substantial fraction of the nano-CuO can remain on the leaf surface or perhaps even enter via the stomata. Lettuce leaves retained the highest amount of nano-CuO within leaf tissues, followed by collard green and then kale. Levels in exposed crops were 2–3 orders of magnitude greater than in unexposed controls. After three cycles of rinse, the residual Cu concentrations were below any toxicity concern for humans (Trumbo et al. 2001).

sp-ICP-MS is a very useful tool which can serve to better understand the interactions between ENMs and plants, including uptake, accumulation, translocation, and biological implications. It can serve to reveal the distribution of nanopesticides in agriculture, whether in runoff water, plant tissues, and wash water. This understanding can be used to design more effective nanopesticides and reduce risk of exposure to humans.

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#### Compliance with ethical standard

**Conflict of interest** The authors declare that they have no conflict of interest.

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