

Fate, accumulation and impact of metallic nanomaterials in the terrestrial environment

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Chapter 3

Quantifying the relative contribution of particulate versus dissolved silver to toxicity and uptake kinetics of silver nanowires in lettuce: impact of size and coating

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Abstract: Functionalized high-aspect-ratio silver nanowires (AgNWs) have been recognized as one of the most promising alternatives for fabricating products, with their use ranging from electronic devices to biomedical fields. Given concerns on the safety of AgNWs, there is an urgent need to investigate the relation between intrinsic properties of AgNWs and their toxicity. In this study, lettuce was exposed for either 6 or 18 days to different AgNWs to determine how the size/aspect ratio and coating of AgNWs affect the contributions of the dissolved and particulate Ag to the overall phytotoxicity and uptake kinetics. We found that the uncoated AgNW (39 nm diameter×8.4 µm length) dissolved fastest of all AgNWs investigated. The phytotoxicity, uptake rate constants and bioaccumulation factors of the PVP-coated AgNW (43 nm diameter×1.8 µm length) and the uncoated AgNW (39 nm diameter \times 8.4 µm length) were similar, and both were higher than that of the PVPcoated AgNW with the larger diameter(65 nm diameter×4.4 µm length). These results showed that the diameter of the AgNWs predominantly affected toxicity and Ag accumulation in plants. Particulate Ag was found to be the predominant driver/descriptor of overall toxicity and Ag accumulation in the plants rather than dissolved Ag for all AgNWs tested. The relative contribution of dissolved versus particulate Ag to the overall effects was influenced by the exposure concentration and the extent of dissolution of AgNWs. This work highlights inherent particulatedependent effects of AgNWs in plants, and suggests that toxicokinetics should explicitly be considered for more nanoparticles and organisms, consequently providing more realistic input information for their environmental risk assessment.

3.1. Introduction

The widespread use of silver nanoparticles in products and applications has raised concerns about their potential side-effects on environmental and human health^{41,57,62,202,203}. Numerous investigations have reported adverse impacts on a range of endpoints of spherical-shaped silver nanoparticles to both aquatic and terrestrial organisms, including impacts on growth or reproductive inhibition, generation of ROS, alteration of enzyme expression, DNA damage and genotoxicity^{35,69, 204}. High aspect ratio silver nanowires and silver nanotubes are also an active area for commercialization and nanotechnology research, given their superior electrical,

plasmonic, optical and antibacterial properties^{205,206,207,208}.

To date, a very limited number of studies have discussed the biological effects of wireshaped metallic nanoparticles on aquatic and terrestrial organisms. From those papers, morphological comparison studies of nanoparticles showed that the dissolution behavior and the biological effects of wire-shaped nanoparticles are distinctly different from the dissolution behavior and adverse effects of nanosphere or nanoplate analogues^{87,89,209,210}. Intrinsic properties such as size, surface chemistry and charge of nanowires could play a key role in controlling their adverse effects to organisms, as has been widely confirmed by studies on spherical nanoparticles^{81,89,211}. To the best of our knowledge, less than 15 studies have been published in which the impact of physicochemical properties of nanowires on their toxic effects were examined. Based on these studies, aspect ratio and length seem to play a dominant role in regulating the adverse effects of nanowires. However, four contradictory patterns of toxicity have been reported;

- 1) higher aspect ratio and longer nanowires induced increased cytotoxic^{212,213,214,215} or;
- lower aspect ratio and shorter nanowires induced higher toxicity to daphnia^{208,216} or;
- shorter nanowires presented more cytotoxicity than the longer nanowires without a consistent aspect ratio dependent pattern ²¹⁷ or;
- 4) the length and aspect ratio do not affect toxicity 218,219,220 .

Moreover, surface coating can affect the dissolution and the stability or aggregation of nanoparticles, which in turn may modulate the adverse effects of nanoparticles to organisms ²⁹. So far, most studies on nanowires concentrated on *in vitro* cytotoxicity and no studies on the effects of size/aspect ratio and coating of nanowires on plants are available. In addition, numerous studies suggested that the toxicity of silver nanospheres to aquatic and terrestrial organisms can be attributed to the release of ions from NPs^{29,57}. However, our previous results showed that particulate Ag dominates phytotoxicity²²¹. Given the importance of shape, it is therefore key to better understand whether the toxicity of silver nanowires suspension is exerted by nano-particulate Ag or by the released ionic Ag.

Extensive research on Ag nanospheres suggests that bioaccumulation of AgNPs is a determinant for their toxicological effects³⁸. Importantly, previous studies showed that the cellular uptake of gold nanowires by mammalian cells^{220,222} and silver nanowires by daphnids^{216,223} were affected by their aspect ratio and length. To the best of our knowledge, there is currently no data is available on the bioaccumulation and phytotoxicity of AgNWs in terrestrial plants. Moreover, toxicokinetic and toxicodynamic models for nanoparticles have placed considerable emphasis on tracing metal accumulation in organisms in order to interpret their toxicological effects. Some studies have reported on parameters associated with uptake biokinetics of metal nanoparticles (e.g. uptake and elimination rate constant, as well as bioaccumulation factor)224,225,226. However, in most cases the measurements performed within these studies focused on total metal concentrations of metallic nanoparticles (particulate plus dissolved form), which prevents differentiating between the relative contributions of dissolved and particulate Ag to the phytotoxicity and toxicokinetics of AgNWs. It is critical to collect this data, as distinguishing the dissolved metal concentration and particulate concentration could increase the accuracy of quantification of kinetics parameters of soluble metallic nanoparticles²²⁷. This would help to provide a more accurate understanding of the particle-specific accumulation of nanoparticles in plants. However, currently it remains unclear whether the inherent properties of AgNWs not only influence dissolution, but also the uptake kinetics of AgNWs in plants. Plants, which play a vital role in providing ecosystems services and form the base of most food webs, are likely exposed to silver nanoparticles via air, soil and water²²¹. Therefore, much more efforts in exploring the adverse impacts of silver nanowires on plants are necessary.

This study therefore aims to i) investigate the importance of the physicochemical properties of AgNWs on their toxicity and on the uptake of Ag in higher plants, and ii) to distinguish the corresponding relative contribution to suspension toxicity and Ag accumulation of ionic and particulate Ag present in suspensions. To this end, the widely cultivated vegetable lettuce (*Lactuca sativa*) was exposed to a series of concentrations of different types of silver nanowires and ionic Ag in hydroponic systems for 6 days to generate dose-toxic response curves. Subsequently, the EC₂₅ level for each of the AgNWs was selected as the exposure concentration to quantify

the biokinetics parameters of Ag uptake in plants after 18 days of exposure. Since there are limited data available on the uptake and toxicity of Ag nanowires in higher plants, this study facilitates the establishment of toxicokinetic models to describe the accumulation of silver nanowires in higher plants.

3.2. Materials and methods

3.2.1 AgNW suspensions: preparation and characterization

AgNO₃ was obtained from Sigma-Aldrich (Zwijndrecht, the Netherlands). Original stock suspensions of polyvinylpyrrolidone (PVP) coated low aspect ratio AgNW (LAR-AgNW:43 nm diameter×1.8 µm length) and medium aspect ratio AgNW (MAR-AgNW: 65 nm diameter×4.4 µm length) were purchased from Nanogap (Milladoiro, Spain). Noteworthily, the PVP used in the two PVP-coated AgNWs was of the same composition and the same method and the same materials were used to coat the two nanowires. This means that there are no differences in the nature and the thickness of the PVP-coating between the two tested PVP-coated AgNWs. The uncoated high aspect ratio AgNWs (HAR-AgNW: 39 nm diameter×8.4 µm length) were purchased from Ras-Ag (Regensburg, Germany). All original stock suspensions of AgNWs were stored at 4 °C in the dark. The actual silver concentrations of the original stock AgNWs suspensions were measured in triplicate by means of Atomic Absorption Spectroscopy (AAS, PerkinElmer 1100 B, Waltham, MA, USA). Taking aliquots from the original stock AgNWs suspensions was always performed under a nitrogen atmosphere. Suspensions of AgNWs dispersed in 1/4 Hoagland solution $(pH 6.0 \pm 0.1)$ with a concentration of 50 mg/L were used to determine the actual size and shape by Transmission Electron Microscopy (TEM, JEOL 1010, JEOL Ltd., Tokyo, Japan) after sonication for 10 min. The composition of the Hoagland solution is described in Table S2.1, Supplementary material. The surface area and volume of the AgNWs were calculated assuming that the AgNWs possessed a perfect cylindrical structure. The basic physico-chemical data of the three AgNWs are presented in Table S3.1. The results discussed in this study are based on actual measured data and all data obtained from the TEM excluded the PVP coating.

3.2.2 AgNWs dissolution testing

The dissolution kinetics of suspensions of the AgNWs were monitored over 72 hours at the concentrations equaling the EC_{25} level of each AgNW. The dissolution experiments were conducted under the same conditions as used in the toxicity and uptake experiments but without plants to avoid underestimation due to the uptake of dissolved Ag by plants. For each AgNW suspension, 6 mL aliquots dispersed in 1/4 Hoagland solution for 1, 6, 12, 24, 48 and 72 h were collected in duplicate from the top 8 cm of the tubes. One of these samples was digested for at least 3 d at room temperature by addition of a few drops of concentrated nitric acid (65%) to determine the total concentration of AgNWs in the suspension (defined as AgNWs_(total) hereafter). The other aliquot was centrifuged at 30,392 g for 30 min at 4 °C (Sorvall RC5Bplus centrifuge, Bleiswijk, Netherlands) and subsequently the supernatants were filtered via a syringe filter of 0.02 µm pore diameter (Anotop 25, Whatman, Eindhoven, The Netherlands). The obtained solutions represented the concentration of dissolved Ag in the suspensions (defined as AgNWs_{(dissolved})), as determined using AAS after addition of a few drops of concentrated HNO₃. The particulate Ag concentrations were obtained as a function of exposure time by calculating the difference between AgNWs(total) and AgNWs(dissolved) at each timepoint. The assessments of the dissolution of each AgNW at each timepoint were performed in triplicate.

3.2.3 Plant pre-culture and toxicity assay

Lactuca sativa seeds obtained from Floveg GmbH (Kall, Germany) were sterilized with NaClO (0.5% w/v) for 15 min and cleaned with deionized tap water. After immersion in deionized water for 24 h, 15 seeds were germinated and grown in a Petri-dish containing 1/8 Hoagland solution for 2 weeks. The young seedlings were then transferred to tubes (one seedling per tube) with 22 mL of 1/4 Hoagland solution, and left to grow 7 d before being exposed to AgNWs. All experiments were performed in a climate room with 16 /8 h of light/darkness and 60 % relative humidity at a 20 /16 °C day/night temperature regime.

The toxicity tests in this study were carried out to establish the dose-response curves

of the AgNWs. The EC_{50} and EC_{25} values were calculated to compare the toxicity of the three AgNWs. Briefly, a series of actual exposure concentrations ranging from 0.006 to 3.94 (LAR-AgNW_(total)), 3.92 (MAR-AgNW_(total)) or 4.48 mg/L (HAR-AgNW_(total)) were prepared in 1/4 Hoagland solution of exposure groups containing uniform pre-grown seedlings. Exposure concentrations of Ag-ions (ranging from 0.005 to 0.80 mg/L) were used as a reference to obtain the dose-response curve of the ions (AgNWs_(dissolved)) released from the AgNWs in suspension. After exposure for 6 days, the biomass of the plants was recorded after the plants were washed thoroughly with deionized tap water and air-dried for 4 hours. The tubes used in this study were covered with aluminum foil and the medium was refreshed every 3 d. All treatments, including the control, were exposed for 6 days in quadruplicate at the same conditions of plants pre-growth.

3.2.4 Uptake experiments of AgNWs and dissolved Ag in plants

The EC₂₅ level of each AgNW_(total) based on the toxicity experiments described above, was selected to perform the uptake experiments of AgNWs in plants over 18 d. Dispersing the AgNWs into 1/4 Hoagland solution results in a mixture of particulate Ag and dissolved Ag ions which changes over time due to sedimentation of particle agglomerates and continuous dissolution of Ag-ions. Thus, in addition to exposure experiments with AgNWs suspensions, dissolved Ag exposure experiments using AgNO₃ were performed separately to obtain the relevant toxicity and uptake information about the dissolved ionic Ag. To study this, time weighted average (TWA) concentrations of ionic Ag at corresponding exposure time points obtained from the dissolution kinetic experiments were performed using the same protocol as in the above mentioned toxicity experiments. One tube contained one plant, with three replicates per treatment. Ag-spiked 1/4 Hoagland solutions were refreshed every 3 d for a total of 18 d. At each exposure time point of 0.25, 0.5 1, 2, 3, 6, 9, 12 and 18 d, plants were collected to determine the Ag content in plant tissues.

After harvesting the plants from each exposure period, the whole plants were immersed in 10 mM HNO_3 for 30 min, transferred into 10 mM EDTA (Ethylenediaminetetraacetic acid) for 30 min, and finally thoroughly rinsed with

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Milli-Q water to remove the strongly attached AgNWs/Ag-ions. Subsequently, the plants were divided into the root and shoot and oven-dried for at least 72 h at 70 °C. After determining the dry biomass of roots and shoots, the samples were digested by adding 6 mL of concentrated HNO₃ for 1 h at 120 °C, followed by adding 2 mL of H_2O_2 at 120 °C until the solutions were clear. Finally, the digest solutions were diluted by adding deionized water to a final volume of 3 mL and the concentrations of Ag were subsequently analysed using AAS. For quality control, blanks containing a Ag standard solution were digested with the same digestion procedure as used for the plants. Standard solutions of Ag at 0.5 mg/L were monitored every 20 samples to examine the stability of the machine. The exposure concentrations of each refreshed batch were determined and the standard deviations were less than 5%.

3.2.5 Data analysis

As Ag-ions are continuously released from AgNWs and the particulate Ag will sediment into the bottom of tubes, the exposure concentration of each AgNW and of the corresponding dissolved ions changes over time. The TWA method can catch the dynamic changes of NPs associated with exposure conditions and offers a more accurate and naturally relevant expression of the actual effective exposure of organisms to NPs than when using the static initial exposure concentrations ²²⁸. Therefore, the expression of the exposure concentrations of AgNWs_(total), AgNWs_(particulate) and AgNWs_(dissolved) was based on the time weighted average (TWA) concentration for each exposure period instead of expressing the exposure concentrations by means of the initially measured concentrations. The TWA

$$C_{TWA} = \frac{\sum_{n=0}^{N} \left(\Delta t_n \frac{C_{n-1}+C_n}{2} \right)}{\sum_{n=1}^{N} \Delta t_n} \tag{1}$$

Where n is the time interval number, N is the total number of intervals, Δt is the time interval, C is the concentration at the end of the time interval.

A first order kinetics equation was used to model the sedimentation process of AgNWs:

$$[Ag_{NW}]_t = ([Ag_{NP}]_0 - P_2) \times exp^{-K_S t} + P_1 \qquad (2)$$

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The increase of the concentration of dissolved Ag released from each AgNW in 1/4 Hoagland solution followed first order kinetics:

$$[Ag^+]_t = (P_1 - [Ag^+]_0) \times exp^{-K_d t} + P_2 \qquad (3)$$

Incorporation of the size and length of AgNWs, the dissolution of each AgNW in 1/4 Hoagland solution can be described as follows:

$$\frac{d[Ag^+]_t}{dt} = N_0 \rho \frac{dV}{dt} = N_0 KA = N_0 K_D (2\pi R^2 + 2\pi RL)$$
(4)

Where $[Ag^+]_t$ and $[Ag_{NW}]_t$ are the concentration of dissolved Ag-ions and particulate Ag at a given point in time with the units of mg/L, P₁ and P₂ are the concentration of particulate Ag and dissolved Ag-ions at equilibrium; K_d and K_s is the first-order rate constants of dissolution and sedimentation (d⁻¹), respectively. N₀ is the number of AgNWs present in one liter of suspension at t = 0 (1/L, N₀= $\frac{C_{NP(0)}}{\rho V(0)}$). V is the volume of AgNWs (µm³). A is the area of the AgNWs (µm²). R is the mean radius of AgNWs (nm). L is the mean length of the AgNWs (µm). K_D is the dissolution rate constant of AgNWs (ng/cm²/h). $[Ag^+]_t$, $[Ag_{NW}]_t$, $[Ag^+]_0$, $[Ag_{NW}]_0$ and time t were measured experimentally. Thus the rate constants can be calculated by fitting equations (2), (3) and (4).

The decrease of biomass of plants exposed to each AgNW suspension is induced by particulate and dissolved Ag together, with the modes of action widely believed to be independent. The response addition model is therefore used to calculate the relative contributions of particulate Ag and dissolved Ag in AgNWs suspensions to the observed toxicity^{136,180}:

$$E_{(total)} = 1 - \left(\left(1 - E_{(particulate)} \right) \left(1 - E_{(dissolved)} \right) \right) \quad (5)$$

where $E_{(total)}$ represents the biomass decrease caused by the suspension of AgNWs as quantified experimentally. $E_{(dissolved)}$ is the decrease of biomass induced by ionic Ag present in the AgNWs suspension as calculated based on the dose-response curve of AgNO₃ towards lettuce biomass decrease. The biomass decrease caused by the particulate Ag in AgNWs can be calculated directly as $E_{(particulate)}$ is the only unknown in the equation.

The concentrations of EC_{25} and EC_{50} for biomass decrease of plants exposed to three kinds of $AgNWs_{(total)}$, $AgNWs_{(particulate)}$ and ionic Ag were calculated with the dose-response-inhibition model in GraphPad using TWA and initial exposure concentrations, respectively.

The mass of Ag accumulated in plants tissues exposed to each AgNW and the corresponding dissolved Ag-ions over time can be described by equation (6):

$$\frac{d[Ag]_{plant}}{dt} = K_u[Ag]_{exposure} - K_e[Ag]_{plant} \quad (6)$$

Where K_u and K_e are the uptake rate constant (d⁻¹) and elimination rate constant (d⁻¹) of Ag from AgNWs_(total), AgNWs_(particulate) and the corresponding dissolved Ag, respectively.

Since at the initial phase of the accumulation process the uptake rate is expected to be much faster than the elimination rate, the elimination of Ag was assumed to be negligible, and equation (6) was simplified into

$$\frac{d[Ag]_{plant}}{dt} = K_u[Ag]_{exposure} \quad (7)$$

Integration of this differential equation leads to equation (8):

$$[Ag]_{plant}(t) = K_u[Ag]_{exposure}(t) \times t \quad (8)$$

Where $[Ag]_{plant}(t)$ is the mass of Ag accumulated in plant tissues (µg/plant), $[Ag]_{exposure}(t)$ is the exposure content of AgNWs_(total), AgNWs_(particulate) and the corresponding dissolved Ag-ions (µg), respectively. K_u is the uptake rate constant. The dynamic changes of exposure concentrations of AgNWs_(total), AgNWs_(particulate) and AgNWs_(dissolved) should be considered when fitting this model. Thus, the uptake rate constant of AgNWs_(total), AgNWs_(particulate) and AgNWs_(dissolved) from each AgNW were obtained by fitting equation (8) to the measured Ag content in plants and TWA exposure content of AgNWs_(total), AgNWs_(particulate) and AgNWs_(dissolved) at each time point ranging from 1 h to 3 d.

The Ag bioaccumulation factor (BAF) was calculated using the following equation:

$$BAF = \frac{[Ag]_{plant}}{[Ag]_{medium}} \quad (9)$$

Where $[Ag]_{plant}$, the content of Ag in the plants ($\mu g/plant$), and $[Ag]_{medium}$, the content of Ag in the exposure medium (μg), were obtained experimentally.

The Ag translocation factor (TF), defined to evaluate the capacity of plants to transfer Ag from roots to the shoots, was calculated as follows:

$$TF = \frac{[Ag]_{shoots}}{[Ag]_{roots}} \quad (10)$$

Where $[Ag]_{shoots}$ and $[Ag]_{root}$ represent the Ag content in plant shoot tissues and root tissues (µg/plant), respectively.

The differences for EC₂₅, EC₅₀ and toxicokinetic parameters of accumulation of Ag among the different AgNWs were analysed for significance using one-way ANOVA followed by Turkey's honestly significant difference tests at $\alpha < 0.05$ using IBM SPSS Statistics 25. The Shapiro-Wilk test was used to check for normality and the Bartlett test for homogeneity of the variance of the data, with no deviations found for both. All results are expressed as mean \pm standard error, based on 4 replicates for biomass decrease and 3 replicates for Ag bioaccumulation

3.3.Results

3.3.1 TEM characterization and dissolution behavior of AgNWs

The TEM pictures confirmed that the shapes of the three types of Ag nanoparticles used in this study were wire-like and that the diameters were in agreement with the information provided by the producer (**Table S3.1**, **Figure 3.1**). However, the length of MAR-AgNW and HAR-AgNW, as determined by TEM, was much shorter than indicated by the producer. Based on these values, the surface area of three AgNWs followed the order of HAR-AgNW > MAR-AgNW > LAR-AgNW (**Table S3.1**). No obvious aggregation was observed for the three types of freshly prepared AgNWs in the exposure medium based on at least 10 TEM images for each AgNW.

As shown in **Figure 3.2**, the dissolution behavior of the NWs over time was different across the different types of AgNW. The percentage of AgNW_(dissolved) (calculated by

using [Ag]_{dissolved}/[Ag]_{total}) increased by 4.2%, 3.9% and 8.5% for LAR-AgNW, MAR-AgNW and HAR-AgNW respectively, after incubation in 1/4 Hoagland solution from 1 h to 72 h. The dissolution extent of HAR-AgNW was 8% higher than that of LAR-AgNW and MAR-AgNW (**Figure 3.2B**). Similarly, the dissolution rate constant, K_d, of uncoated HAR-AgNW was slightly higher than the dissolution rate constant of PVP-coated LAR-AgNW and MAR-AgNW (**Table S3.2**). However, once incorporated the surface area of AgNWs to model the dissolution rate constant, the K_D of HAR-AgNW was the lowest (**Table S3.2**). Even though the initial exposure concentration of MAR-AgNW was a bit higher than that of LAR-AgNW, their sedimentation rate constants were similar (**Table S3.2**).



Figure 3.1. TEM images of AgNWs after 1 h of incubation in the exposure medium (scale bar for A, B and C: 200 μm; scale bar for D, E and F: 500 nm). LAR-AgNW: 43 nm diameter×1.8 μm, PVP-coated, MAR-AgNW: 65 nm diameter×4.4 μm, PVP coated, HAR-AgNW: 39 nm diameter×8.4 μm, uncoated.



Figure 3.2. Ion release profiles of AgNWs suspensions at the EC₂₅ level in the exposure medium over time. (A) Total Ag concentrations in the AgNW suspensions. (B) Percentage of dissolved Ag released in the AgNW suspensions, (C) particulate Ag concentrations in AgNWs suspensions. Data are mean \pm SE (N=3). (LAR-AgNW: 43 nm diameter×1.8 µm, PVP-coated, MAR-AgNW: 65 nm diameter×4.4 µm, PVP coated, HAR-AgNW: 39 nm diameter×8.4 µm, uncoated).

3.3.2 Acute toxicity of AgNWs and AgNO3 to Lactuca sativa

The dose-response curves of plants exposed to the different kinds of AgNWs and ionic Ag for 6 d show that both AgNWs and ionic Ag can induce significant toxicity to plants, decreasing the biomass of plants as a function of increasing exposure concentrations of suspensions of Ag-species (**Figure 3.3**, **Figure S3.1**). Based on the dose-response curves, the effect concentrations (EC) of AgNWs_(total) and their corresponding AgNWs_(particulate) causing 25% (EC₂₅) and 50% (EC₅₀) inhibition of plant growth were calculated. The EC₂₅ (0.07 mg/L) and EC₅₀ (0.15 mg/L) values based on TWA exposure concentrations of Ag-ions were about 1.3-2.4 fold and 2.1-3.3 fold lower than the corresponding effect levels of the AgNWs tested, respectively, indicating that Ag-ions were the most toxic to lettuce.

We also investigated differences in toxicity among different kinds of AgNWs to plants on the basis of total Ag. The relative EC₅₀ values based on AgNWs_(total) was significantly higher for MAR-AgNW compared to LAR-AgNW and HAR-AgNW, which had a comparable EC₅₀ value to each other (LAR-AgNW \approx HAR-AgNW<MAR-AgNW), regardless of the way in which EC₅₀ was expressed (ANOVA, C_{TWA} *vs* C_I, P=0.008 *vs* 0.016, **Table 3.1**). This suggests that the suspensions of LAR-AgNW and HAR-AgNW and HAR-AgNW induced higher toxicity than suspensions of MAR-AgNW.

when expressed as $AgNWs_{(particulate)}$, no significant differences were observed for the EC_{50} values among the three AgNWs tested (ANOVA, P=0.135 for C_{TWA} and P=0.287 for C_I , **Table 3.1**). In addition, the EC_{25} values of $AgNW_{(total)}$ (ANOVA, P=0.003) and $AgNW_{(particulate)}$ (ANOVA, P=0.022) for both LAR-AgNW and HAR-AgNW were significantly lower than the corresponding values of MAR-AgNW regardless of the way in which EC_{25} was expressed. This indicates that the effects of aspect ratio and coating of AgNWs on the phytotoxicity were more obvious at low effective concentration levels.



Figure 3.3. Dose–response curves of biomass decrease of *Lactuca sativa* exposed to different concentrations of (A)AgNWs_(total) and AgNO₃, and (B) AgNWs_(particulate) expressed as time weighted concentrations. Data are mean \pm SE (N=3).(LAR-AgNW: 43 nm diameter×1.8 µm, PVP-coated, MAR-AgNW: 65 nm diameter×4.4 µm, PVP coated, HAR-AgNW: 39 nm diameter×8.4 µm, uncoated).

3.3.3 Bioaccumulation kinetics and translocation of AgNWs and dissolved Ag in plant tissues

After 18 d the Ag content in control plants was below the detection limit. The mass of silver accumulated in plants was positively correlated with exposure time regardless of AgNW type or Ag-form. After exposure to the suspensions of LAR-AgNW, MAR-AgNW and HAR-AgNW at the EC_{25} level for 18 d, the content of Ag in the plants increased to 7.5, 11.1 and 6.5 µg/plant, which was respectively 50, 33 and 11 times higher compared to the corresponding amount of dissolved Ag released from the AgNWs at the EC_{25} value (**Figure 3.4**A and 4B). This suggests particulatespecific uptake of Ag for all three AgNWs tested. According to **Table S3.3**, exposure time had no significant effect on the BAFs values of AgNWs_(total), AgNWs_(particulate) and AgNWs_(dissolved) for all AgNWs after exposure for more than 9 d (ANOVA, P>0.25, **Table S3.3**).

Meanwhile, the type of AgNW was found to have a significant effect on the bioaccumulation process as the K_u and BAF values of MAR-AgNW_(particulate) were much lower compared to LAR-AgNW_(particulate) and HAR-AgNW_(particulate) (ANOVA, P<0.05). Furthermore, the Ag content in plant shoots was only detectable for the AgNWs_(total) groups exposed for more than 9 d. However, the transfer factors for all treatments were less than 0.02, suggesting that a major portion of the Ag remained in the plant roots. In addition, all type of AgNWs (ANOVA, P=0.055, **Table 3.2**) and exposure time (ANOVA, P=0.117, **Table 3.2**) did not significantly influence the Ag translocation from roots to shoots.



Figure 3.4. Measured Ag uptake in *Lactuca sativa* exposed to different (A) AgNWs suspension, (B) the particulate Ag and (C) the corresponding concentrations of dissolved Ag at EC₂₅ level over 18 days. Data are mean \pm SE (N=3). (LAR-AgNW: 43 nm diameter×1.8 µm, PVP-coated, MAR-AgNW: 65 nm diameter×4.4 µm, PVP coated, HAR-AgNW: 39 nm diameter×8.4 µm, uncoated)

3.3.4 Relative contribution of dissolved Ag and particulate Ag to toxicity and Ag uptake in plants

In general, the dominant role of AgNWs_(particulate) to the overall toxicity relative to the corresponding dissolved Ag was observed in all exposure cases (Figure 3.5). In addition, exposure concentration was found to have a significant impact on the relative contribution of AgNWs_(particulate) to the overall toxicity regardless of the type of AgNWs exposed. For instance, the relative contributions of AgNWs_(dissolved) to the overall suspension toxicity were found to increase by 11.1%, 17.2% and 41.5% for LAR-AgNW, MAR-AgNW and HAR-AgNW when the exposure concentrations increased from 0.04, 0.02 and 0.03 to 2.42, 2.71 and 3.39 mg/L, respectively. Therefore the relative contribution of HAR-AgNW_(dissolved) to toxicity at high concentrations was much higher than that of LAR-AgNW(dissolved) and MAR-AgNW_(dissolved) (ANOVA, P<0.001, Figure 3.5). Similarly, uptake of AgNWs_(particulate) dominated the contribution to overall Ag accumulation in plants compared to the uptake of AgNWs(dissolved) for all AgNWs tested as the contribution of the NWs accounted for more than 85% of the Ag accumulation in the plants. Additionally, the relative contributions of AgNWs_(dissolved) increased significantly after exposure from 0.25 d to 3 d and then tended to be stable for LAR-AgNW and MAR-AgNW (P<0.001, Figure 3.5D). Among the three types of AgNWs, the HAR-AgNW_(dissolved) contributed most to the overall Ag accumulation in plants with a contribution of about 10% at the EC₂₅ exposure level for 18 d, which was much higher than that of LAR-AgNW_(dissolved) (2%) and HAR-AgNW_(dissolved) (3%) (ANOVA, P<0.001, Figure 3.5D).



Figure 3.5. Relative contribution (%) of different AgNWs_(particle) and AgNWs_(ion) to toxicity at different concentrations (A, B and C), and to Ag accumulation at EC25 concentrations for different exposure time (D). The data represent the mean \pm SE (n = 3). The different letters in the same group indicate statistically significant differences between treatments at p < 0.05. (LAR-AgNW: 43 nm diameter×1.8 µm, PVP-coated, MAR-AgNW: 65 nm diameter×4.4 µm, PVP coated, HAR-AgNW: 39 nm diameter×8.4 µm, uncoated)

		$EC_{25}(mg/L)$		$EC_{s0}(mg/L)$	
		C _I (95% confidence Interval)	C _{TWA} (95% confidence Interval)	C _I (95% confidence Interval)	C _{TWA} (95% confidence Interval)
	LAR-AgNW	0.137a (0.098-0.175)	0.084a ($0.060-0.107$)	0.513a (0.387-0.800)	0.315a (0.238-0.491)
$AgNWS_{\rm (total)}$	MAR-AgNW	0.237b (0.179-0.290)	0.164b ($0.124-0.200$)	0.718b (0.549-1.045)	0.497b ($0.380-0.724$)
	HAR-AgNW	0.116a (0.071-0.164)	0.088a ($0.054-0.124$)	0.528ab (0.366-1.174)	0.417ab (0.286-0.965)
	Ag+	0.057c ($0.047-0.088$)	1	0.150c	
	LAR-AgNW	0.144a ($0.102-0.191$)	0.086a ($0.060-0.115$)	0.542a (0.385-0.866)	0.318a (0.226-0.503)
${ m AgNW}_{({ m particulate})}$	MAR-AgNW	0.246b (0.184-0.300)	0.157b (0.114-0.197)	0.701a (0.520-0.910)	0.422a ($0.317-0.546$)
	HAR-AgNW	0.105a (0.057-0.178)	0.061a ($0.022-0.096$)	0.616a (0.351-1.117)	0.382a (0.201-0.902)

as time weighted average concentrations. The data represent the mean \pm SE (n = $\overline{4}$). The different letters of the same Ag form in **Table 3.1.** The EC₂₅ and EC₅₀ values of AgNWs(total) and AgNWs(particulate) expressed as initial concentrations and expressed the same column indicate statistically significant differences between treatments among three AgNWs at p < 0.05.

Table 3.2. Uptake rate constants, BAFs and TFs for Lactuca sativa exposed to different AgNWs at EC25 level over 18 d. The data
represent the mean \pm SE (n = 3). The different letters in the same column indicate statistically significant differences between
treatments at $p < 0.05$. Asterisks indicate statistically differences between different Ag forms of each AgNW for the same row at p
< 0.05.

TFs	Total Ag	0.010±0.001a	0.010±0.002a	0.015±0.001a	nm dia meter X8 4
	Dissolved ions	$0.758\pm0.018a^{*}$	0.745±0.009a*	0.759±0.014a*	$HAR_A \sigma NW 30$
BAFs	Particulate Ag	0.621±0.006a	0.507±0.024b	$0.684\pm0.011c$	IIII DVP coated
	Total Ag	0.630±0.006a	0.531±0.026b	0.700±0.011c	m diameter×4.4
	Dissolved ions	0.245±0.021a*	0.244±0.014a*	0.229±0.009a*	MAR-AdNIM-65 n
${ m K}_{ m uptake}\left({ m d}^{-1} ight)$	Particulate Ag	$0.136\pm 0.006a$	0.098±0.008b	0.149±0.015a	8 IIII DVD-coated
	Total Ag	0.139±0.007a	0.105±0.008b	0.153±0.013a	nm diameter×1
Type of AgNIWs	0 4 4 TQ 1	LAR-AgNW	MAR-AgNW	HAR-AgNW	$I A R_A \sigma NIM \cdot 43$

The state of the s coated, HAK-AGN W: 39 nm 4 4.4 Jum, F annanne CUALCU, INIAN-AGIN W.: UJ ц > LAK-AgNW: 43 nm diameter×1.8 μm, P[·] μm, uncoated

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3.4. Discussion

3.4.1 Dissolution behavior of AgNWs

Our results showed that the dissolution of uncoated HAR-AgNW was much higher compared to PVP-coated AgNWs with lower aspect ratio values (LAR-AgNW and MAR-AgNW). The dissolution rate constant (h^{-1}) of AgNWs was observed to follow the order of HAR-AgNW > MAR-AgNW > LAR-AgNW. Scanlan et al.²¹⁶ made the generic statement that dissolution increases for short AgNWs with a low aspect ratio compared to longer AgNWs with a higher aspect ratio in EPA media. The discrepancy with our results suggests that restricting or placing too much focus on aspect ratio for toxicity assessment is not giving the full picture. A larger surface area of nanoparticles can increase the contact between NPs and the oxidant, thereby accelerating the dissolution⁸⁷. Although the surface area of HAR-AgNW was larger compared to LAR-AgNW and MAR-AgNW, the dissolution rate constant (ng/cm²/h) which incorporated the surface area of AgNWs was lowest for HAR-AgNW (LAR-AgNW≈MAR-AgNW > HAR-AgNW, Table S3.2). This indicates that the dissolution behaviors of AgNWs in this study cannot be explained by the larger surface area only. Therefore, we postulate that the differences in dissolution kinetics of HAR-AgNW compared to LAR-AgNW and MAR-AgNW were due to the absence of PVP-coating on HAR-AgNW. This suggests that a PVP-coating largely played a role with regard to the dissolution behaviors in the current study, and was more important than the dimensions of AgNWs. In addition, the dissolution of AgNWs in the present study was found to be less than 15%, which was much lower than the extent of Ag released from Ag nanospheres (about 30%) obtained by our previous study under similar exposure conditions²²¹. This was consistent with the findings that spherical AgNPs displayed the highest dissolution, followed by Ag nanocubes and AgNWs⁸⁵, confirming that the dissolution of AgNPs is shape-dependent.

3.4.2 Coating related factors cannot fully explain toxicity differences among AgNWs

All three AgNWs suspensions tested in this study inhibited the growth of plants significantly with the EC_{50} values ranging from 0.32 to 0.49 mg/L expressed on TWA

basis (compared to 0.52 to 0.71 mg/L expressed as the initial concentrations). Comparison of the toxicity levels deduced for the three AgNWs tested suggests that a low aspect ratio and a lack of coating of AgNWs are slightly more toxic than high aspect ratio and PVP-coated AgNWs. This finding is consistent with the results of Scanlan et al²¹⁶, who concluded that short AgNWs with low aspect ratio exhibited higher toxicity compared to long AgNWs with high aspect ratio. However, Chae et al.²²³ reported that larger and longer PVP-coated AgNWs dispersed in TAP medium exhibited greater toxicity to algae than the smaller and shorter ones. These apparent contradictions indicate that the impacts of AgNWs characteristics on their toxicity can also depend on the test species and/or the chemistry of the exposure medium. An important goal of PVP-coating is to promote the dispersion and to reduce the agglomeration of nanoparticles^{82,229,230}, which in turn affects their toxicity as sedimentation and aggregation can contribute to reduced toxicity^{231,232}. However, in the current study we found that the sedimentation of PVP-coated LAR-AgNW and MAR-AgNW was similar, and both sedimentation rate constants were higher than the sedimentation rate constants of uncoated HAR-AgNW (Figure 3.2 and Table \$3.2), whereas the toxicity of MAR-AgNW was lowest. In addition, the higher toxicity of uncoated nanoparticles has been commonly attributed to their faster dissolution^{231,233,234}. However, the toxicity of LAR-AgNW and HAR-AgNW was comparable, while the dissolution of HAR-AgNW was significantly increased compared to LAR-AgNW. Our results indicate that the sedimentation and dissolution of AgNWs associated with the lack of coating may not be indicative of their phytotoxicity.

3.4.3 Accumulation kinetics of AgNWs and dissolved Ag

All three AgNWs suspensions induced Ag uptake in plants roots. Contaminants can be taken up by plant roots via apoplastic transport through the intracellular spaces of adjacent cells along cell walls and via symplastic/transmembrane pathway through plasmodesmata/cell membranes between cells^{235,236}. In this study, Ag uptake via the plant roots was rapid and equilibrium was reached quickly. In addition, the data of AgNWs accumulation in plant roots fitted the one-compartment kinetic model well (R²>0.9). Taken together, these results indicate that in our study apoplastic transport was likely the major pathway for the uptake of Ag by plant roots after AgNWs adhere to the root epidermis. The limited translocation of Ag from roots to shoots in all AgNWs exposure treatments further confirmed this, as it is difficult for materials taken up by the apoplastic pathway to cross the Casparian strip (a barrier limits the entrance of substances to xylem or phloem) and thus cannot be transported into the above-ground parts. Wang et al.¹⁹² also confirmed that CuO NPs pass through the epidermis into root tissues via the apoplastic route. All toxicokinetic parameters of dissolved Ag were much higher than those of AgNWs_(particulate), demonstrating that the accumulation of Ag-ions in plants proceeds much faster than the accumulation of particulate Ag. This implies that dissolved Ag is more bioavailable than particulate Ag.

Among the three types of AgNWs tested, the uptake rate constants and BAFs of both AgNWs_(total) and AgNWs_(particulate) followed the order of LAR-AgNW (PVP-coated) \approx HAR-AgNW (uncoated) > MAR-AgNW (PVP-coated). As the LAR-AgNW and HAR-AgNW have a similar diameter but different length and coating, this result suggests that the dimension, more specifically, the diameter of AgNWs is the dominant factor related to their cellular uptake. The AgNPs with smaller diameter might easier pass through the pores in/between cell walls due to the size exclusion limit of cell walls and/or apoplast⁶⁸. Torrent et al.⁸² showed that silver nanospheres accumulated in lettuces, and that accumulation was diameter (size) dependent, but coating independent, which is in line with our findings. Previous studies of wireshaped nanoparticles are shown to behave totally different compared to the results found in our study - but it should be noted that those were all conducted on nonflora species. For example, the cellular uptake of Au nanowires in MCF-7²²⁰ and HeLa²²² cells, of Ag nanowires in daphnids²¹⁶, and of Fe nanowires in HeLa cell²³⁷ and NIH/3T3 fibroblast cells²¹⁸ was length dependent: shorter NWs are more likely to be taken up compared with longer ones in general. However, from our results, no definite dependency on length-specific uptake of AgNWs in plants can be concluded. Our results highlight that interaction between metal-based NPs and plants should consider the combined effects of diameter, length, shape and surface chemistry.

3.4.4 Particulate Ag-dominant effects

Ionic Ag was more toxic in comparison to all AgNWs (see EC_{50} values) when plants were exposed to an equal dose of total Ag. A great number of studies (based on silver, zinc, copper, gold, iron and nickel-based nanoparticles) suggested that the toxicity of metal-based nanoparticles is mainly driven by the dissolved ions shed from the particulate form. However, following exposure to the specific amount of Ag-ions which corresponded to the amount released from metallic NPs, toxicity originating from the nanoform rather than from the ionic Ag present in suspension has also been confirmed⁸⁵. The dominant contribution of particulate Ag to the overall toxicity of AgNWs was also found in this study on the basis of assessing the relative contributions of particulate Ag and the corresponding dissolved Ag to the reduction of plants biomass. The almost similar EC₅₀ values of AgNWs(total) and AgNWs(particulate) also revealed that the particulate Ag was the major source of toxicity of AgNWs. Similar particulate-originated toxicity was observed following exposure of algae to PVP-AgNWs as the dissolved concentration of Ag, as released from the AgNWs, was below 0.05 ppm ²²³. In addition, our previous study in which we exposed lettuce to spherical AgNPs at similar conditions as employed in the current study, also verified the particle-ruling toxicity ²²¹. Gorka and Liu⁸⁵ also suggested that the toxicity observed for Lolium multiflorum cannot be explained by ionic silver solely when the plants are exposed to AgNWs, AgNPs and Ag nanocubes. The similar contribution of particulate Ag among different shapes of Ag nanoparticles confirms that the dominant role of particles in suspension toxicity of Ag materials to higher plants is shape-independent.

For the accumulation experiments, the exposure concentrations of dissolved Ag at each time point were comparable to the TWA concentrations of dissolved Ag released from each AgNWs at the EC_{25} level at the same exposure time. The dominant contribution patterns of AgNWs_(particulate) to the total Ag accumulation in plants were similar to the contribution of AgNWs_(particulate) to the overall toxicity. The similar patterns strongly indicate that a relationship between the phytotoxicity of AgNWs and the Ag accumulation in plants exists. This potentially indicates that the accumulation originates from particulate Ag, which is also responsible for the phytotoxicity of AgNWs. Once the AgNWs are taken up by plants, the *in-vivo* dissolution of AgNWs and in-place biological transformation of new particles can occur to some extent^{85,90}. These processes could also affect the plant growth partly^{90,190}. Dang et al.²²⁷ also reported that particulate Ag dominates the overall Ag accumulation in wheat (*Triticum aestivum*) exposed to suspensions of AgNPs. In addition, the relative importance of ions versus particles of AgNPs are highly dependent on the dissolution extent of AgNPs. This observation is consistent with our findings that the particulate Ag contribution to the overall Ag accumulation of HAR-AgNW (which displayed the highest dissolution ability) was higher than that of LAR-AgNW and MAR-AgNW. Our results showed furthermore the relative contribution of ion versus particulate to the overall adverse effects depends on the exposure concentration (**Figure 3.5**).

3.5. Conclusions

To the best of our knowledge, this is the first investigation that explored the effects of PVP-coated and uncoated AgNWs with different dimensions on their phytotoxicity and toxicokinetics of the dissolved and particulate Ag in terrestrial plants. Our results showed that PVP-coating affected the dissolution behaviors of AgNWs. This coating associated dissolution behavior was found not to affect the phytotoxicity of the AgNWs studied. The toxicity and the toxicokinetics parameters of the three AgNWs followed the order of LAR-AgNW≈HAR-AgNW>MAR-AgNW. This indicates that the diameter of the AgNWs plays a crucial role in determining the toxicity and Ag accumulation of AgNWs. In addition, the particulate Ag dominates the overall toxicity and Ag accumulation in plants of AgNWs suspensions compared to dissolved Ag. Overall, the results of this study highlighted that the toxicokinetics and toxicodynamics of AgNWs associated with different dimension and coating should be taken into account when studying the interactions between NPs and biological systems. Our findings provide in depth understanding of nanosafety to plants as well as can contribute to further developing knowledge regarding their safe design to mitigate the side effects of nanowires. Furthermore, the established EC₅₀values and the toxicokinetic parameters of AgNWs can be incorporated into predictive models for assessing risk NWs for environmental health and safety

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	Volume coating	7 0.003 PVP	§ 0.144 PVP	1 0.010 uncoated	30 nm diametery8 4
Mean values	Aspect SA ratio (µm ²)	42 0.247	68 0.898	218 1.021	HAP AGNIM
Actual	r Length / (µm)	1.8	4.4	8.4	1 A IIIII DVD COat
	Diameter (nm)	43.1	64.5	38.6	n diametery/
	Volume (µm³)	0.004	0.115	0.031	A a NTM/ 65 nn
values	$SA (\mu m^2)$	3.80	6.61	3.14	oated MAR-
ominal Mean	Aspect ratio	75	428	625	8 um DV/D_C
Ň	Length (µm)	3	30	25	diametery1
	Diameter (nm)	40	70	40	*NW/· 13 nm
	AgNWs type	LAR-AgNW	MAR-AgNW	HAR-AgNW	I AR-AG

Table S3.1. The nominal and actual values of the characteristics of the AgNWs used in this study.

þ um, uncoated; All data obtained from the TEM excluded the PVP coating Table S3.2. Dissolution rate constants of AgNWs_(dissolved) and sedimentation rate constant of AgNWs_(particulate) for each AgNWs at EC25 level over 72 h

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Type of AgNWs	K _D (ng.c	$2m^{-2}.h^{-1}$	K.	$_{\rm d}({\rm h}^{-1})$	Ks((h^{-1})
	K value	\mathbb{R}^2	K value	\mathbb{R}^2	K value	\mathbb{R}^2
LAR-AgNW	0.031	0.99	0.011	0.95	0.039	0.92
MAR-AgNW	0.039	0.99	0.021	0.95	0.039	0.98
HAR-AgNW	0.016	0.97	0.027	0.94	0.031	0.90
LAR-AgNW: 43 nm di uncoated	ameter×1.8 μm,	PVP-coated, MAR	-AgNW: 65 nm diam	ieter×4.4 μm, PVP coat	ted, HARAgNW: 39 nn	ı diameter×8.4 μm,

Type of AgNW	Exposure duration	Applied dose (µg)		BAFs		TFs
C	(d)	total	Total Ag	Particulate	Dissolved ions	Total Ag
	3	1.98	0.70±0.05a	0.69±0.05a	0.95±0.08a*	/
LAR-	9	5.93	0.47±0.03b	0.46±0.03b	0.81±0.04a*	0.015±0.003a
AgNW	15	9.90	0.56±0.02ab	0.56±0.02ab	0.77±0.02a	/
	18	11.85	0.63±0.01a	0.62±0.01a	0.76±0.02a*	0.010±0.001a
	3	3.50	0.54±0.09a	0.52±0.09a	0.96±0.04a*	/
MAR-	9	10.50	0.49±0.03a	0.46±0.03a	0.83±0.01b*	0.012±0.0002a
AgNW	15	17.5	0.44±0.03a	0.42±0.03a	0.75±0.01b	/
	18	21.00	0.53±0.03a	0.51±0.02a	0.75±0.01b*	0.010±0.002a
	3	1.55	0.67±0.10a	0.64±0.11a	0.96±0.03a*	/
HAR-	9	4.64	0.55±0.05a	0.51±0.06a	0.85±0.01b*	0.018±0.002 a
AgNW	15	7.75	0.56±0.02a	0.54±0.03a	0.78±0.03b	/
	18	9.29	0.70±0.01a	0.68±0.01a	0.76±0.01b*	0.015±0.001 a

Table S3.3. Bioaccumulation (BAFs) and transfer (TF) factors of Ag for *Lactuca sativa* exposed at the EC25 concentrations of AgNWs over 3, 9, 15 and 18 d.

The data represent the mean \pm SE (n = 3). The different lower-case letters in the same column of each AgNW indicate statistically significant differences among exposure time at p < 0.05. (LAR-AgNW: 43 nm diameter×1.8 µm, PVP-coated; MAR-AgNW: 65 nm diameter×4.4 µm, PVP coated; HARAgNW: 39 nm diameter×8.4 µm, uncoated.)

Type of		$K_{uptake}\left(d^{-1} ight)$	
Agin ws –	Total Ag	Particulate Ag	Dissolved ions
LAR-AgNW	0.98	0.98	0.97
MAR-AgNW	0.95	0.94	0.87
HAR-AgNW	0.80	0.78	0.87
d-1	= per day		

Table S3.4. Goodness of fit of uptake rate constants for Lactuca sativa exposed todifferent AgNWs at EC25 level over 18 d.



Figure S3.1. Dose–response curves of biomass decrease of *Lactuca sativa* exposed to different concentrations of (A)AgNWs_(total) and AgNO3, and (B) AgNWs_(particulate) expressed as initial exposure concentrations. Data are mean \pm SE (N=3).(LAR-AgNW: 43 nm diameter×1.8 µm, PVP-coated, MAR-AgNW: 65 nm diameter×4.4 µm, PVP coated, HARAgNW: 39 nm diameter×8.4 µm, uncoated