Emerging Contaminant or an Old Toxin in Disguise? Silver Nanoparticle Impacts on Ecosystems

Benjamin P. Colman,*,† Benjamin Espinasse,‡,▽ Curtis J. Richardson,§,‖ Cole W. Matson,‖,⊥,▽ Gregory V. Lowry,*,▽ Dana E. Hunt,‖ Mark R. Wiesner,‡,▽ and Emily S. Bernhardt†,▽

†Department of Biology, Duke University, Durham, North Carolina 27708, United States
‡Civil and Environmental Engineering Department, Duke University, Durham, North Carolina 27708, United States
§Duke University Wetland Center, Nicholas School of the Environment, Duke University, Durham, North Carolina 27708, United States
‖Department of Environmental Science and the Center for Reservoir and Aquatic Systems Research (CRASR), Baylor University, Waco, Texas 76798, United States
*Civil & Environmental Engineering, Carnegie Mellon University, Pittsburgh, Pennsylvania 15213, United States
▽Center for the Environmental Implications of Nanotechnology, Duke University, Durham, North Carolina 27708, United States

Supporting Information

ABSTRACT: The use of antimicrobial silver nanoparticles (AgNPs) in consumer-products is rising. Much of these AgNPs are expected to enter the wastewater stream, with up to 10% of that eventually released as effluent into aquatic ecosystems with unknown ecological consequences. We examined AgNP impacts on aquatic ecosystems by comparing the effects of two AgNP sizes (12 and 49 nm) to ionic silver (Ag⁺; added as AgNO₃), a historically problematic contaminant with known impacts. Using 19 wetland mesocosms, we added Ag to the 360 L aquatic compartment to reach 2.5 mg Ag L⁻¹. Silver treatments and two coating controls were done in triplicate, and compared to four replicate controls. All three silver treatments were toxic to aquatic plants, leading to a significant release of dissolved organic carbon and chloride following exposure. Simultaneously, dissolved methane concentrations increased forty-fold relative to controls in all three Ag treatments. Despite dramatic toxicity differences observed in lab studies for these three forms of Ag, our results show surprising convergence in the direction, magnitude, and duration of ecosystem-scale impacts for all Ag treatments. Our results suggest that all forms of Ag changed solute chemistry driving transformations of Ag which then altered Ag impacts.

INTRODUCTION

The small size of manufactured nanomaterials (1–100 nm) gives them unique physical and chemical properties, which have led to their use in an ever increasing number of consumer products and industrial processes.¹ With this rapid increase in the production and use of nanomaterials, it is inevitable that engineered nanomaterials are entering the environment.¹ However, our ability to create and deploy new nanomaterials far outstrips our understanding of what they may do outside of their intended use, leading to concerns that these novel contaminants will have unpredictable and potentially dire consequences when they enter natural environments.

Silver nanoparticles (AgNPs) are of particular concern because they are produced specifically for their antimicrobial and biocidal properties. The increasing use of AgNPs leads to two obvious questions: At what concentrations are AgNPs entering the environment? What are their impacts on the exposed organisms? A number of risk assessments³,⁴ now suggest that AgNPs are entering wastewaters and are ultimately entering the environment as either wastewater treatment biosolids² or effluent.⁵ At present, there are no empirical data on silver nanoparticle concentrations and no assessments of their impacts in natural environments.

The rapidly expanding toxicological literature on nanomaterial impacts suggests that AgNP pollution has the potential to harm many organisms and that AgNP loading could lead to significant changes in important ecosystem processes, such as primary productivity,⁷ decomposition,⁸ and nitrogen cycling.¹⁰ Toxicological research on the environmental impacts

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of AgNPs, however, has primarily focused on high dose AgNP exposures to single-species cultures grown in well-characterized laboratory media. Although lab studies are essential for elucidating the mechanisms of toxicity, it is quite difficult to extrapolate their findings to natural ecosystems. In natural ecosystems, exposure concentrations are likely to be low, environmental media are both chemically complex and heterogeneous, and the myriad direct and indirect interactions that exist within communities of organisms in natural environments are likely to alter the magnitude and potentially the direction of nanomaterial impacts.12

Up to 10% of the AgNPs entering wastewater treatment plants are predicted to be released in wastewater effluent into aquatic ecosystems,20−6,13 thus we chose to explore the impacts of nanoparticle addition in wetlands. To the water column of triplicate 4.5 m² wetland mesocosms, we added either 12 nm AgNPs, 49 nm polyvinylpyrrolidone coated AgNPs (hereafter, GA-AgNPs and PVP-AgNPs, respectively). We compared the fate and impact of these AgNPs to the addition of ionic silver (Ag⁺) added as AgNO₃. Each Ag treatment consisted of a single dose sufficient to raise initial water column Ag concentrations to 2.5 mg Ag L⁻¹. Silver-treated mesocosms were compared to four replicate control mesocosms, and triplicate GA-coating and PVP-coating controls, which received the same amount of the coatings added in the AgNP treatments, only without AgNPs. With one minor exception described in the results, coatings controls were identical to control mesocosms. Thus, the focus of the work presented here is on comparisons between controls, GA-AgNPs, PVP-AgNPs, and AgNO₃ treatments.

Dissolved silver is a well-studied contaminant with a long history of environmental impact studies,14 and thus a primary goal of our study was to determine the extent to which the ecological impacts of AgNP exposure differ from the better studied effects of Ag⁺ exposure. Direct comparisons within individual studies have led to mixed conclusions: some studies found higher toxicity and/or biouptake for AgNPs than AgNO₃.8,15,16 Other studies report the opposite conclusion.17−19 Much of this discrepancy may be explained by differences in AgNP size or coatings; for that reason, we chose to compare the effects of two very different AgNPs with different published toxicity thresholds: the larger PVP-AgNPs are in general less toxic than the smaller GA-AgNPs, which in turn tend to be less toxic than AgNO₃.18,19 Thus, we expected to find that dissolved silver would reduce microbial and plant biomass and growth to a greater degree than GA-AgNPs, while both GA-AgNPs and AgNO₃ treatments would have stronger impacts than the addition of larger PVP-AgNPs. Furthermore, we expected patterns of bioaccumulation into plant tissues to follow similar trends.

**Materials and Methods**

Setup. The overall design and construction of our slatboard wetland mesocosms were as previously detailed.20 Briefly, treated lumber was used to create 3.66 × 1.22 × 0.81 m boxes. The boxes were partially filled with screening material, and graded to give a flat bottom that was 0.8 m long, connected to a slope upward at an angle of 13° and a run of 2.8 m. Over the screening, an EPDM liner was placed, and over the liner soil was filled to a constant thickness of 22 cm. Mesocosms were built between August of 2009 and March of 2010.

The soil used in this experiment was a blend of three topsoils chosen to give a similar texture and organic matter content to soils used in previous terrestrial and wetland mesocosm experiments.15,20 Soils were blended and provided by Soils and Sand (Durham, U.S.A.), and were a blend of three soils (“Sandhills”, from Lemon Springs, NC, U.S.A.; “Clay” and “Topsoil”, from Durham, NC, U.S.A.). The final soil had a texture of 63.9% sand, 28.3% silt, and 13.0% clay, with 5.1% loss on ignition.

The mesocosms consist of the following: a permanently flooded portion (aquatic zone), a periodically flooded portion (transition zone), and a rarely flooded portion (upland zone). Mesocosms were planted in March of 2010. In the aquatic compartment, six stems of Egeria densa were planted in a two by three grid, Potamogeton diversifolius was unintentionally introduced with the E. densa, and 2 g of the duckweed Landoltia punctata was floated on the water’s surface (Carolina Biological, Burlington, U.S.A.). The transition and upland zones were planted as described in the Supporting Information, SI. Algae and zooplankton were added in 250 mL of unfiltered water per mesocosm from a local wetland. Other organisms were added or were allowed to colonize the mesocosms, as described in the SI.

To normalize the water chemistry and algal composition in the mesocosms, a submersible pump was used to circulate water between the 19 mesocosms. Water was circulated using a combination of pumping and siphoning twice prior to dosing in order to ensure reasonable homogenization of the water, as verified by convergence of water chemistry.

**Experimental Treatments.** There were six different treatments overall: Control, received DI water; PVP-coating control, received PVP equivalent to that added with the PVP-AgNPs; GA-coating control received gum arabic equivalent to that added with the GA-AgNPs; AgNO₃ received sufficient NO₃⁻ to bring watercolumn concentration to 2.5 mg Ag L⁻¹; GA-AgNPs received sufficient 12.9 ± 9.2 nm (mean ± standard deviation) gum arabic coated AgNPs to bring the watercolumn Ag concentrations to 2.5 mg L⁻¹; PVP-AgNPs received sufficient 49.3 ± 22.5 nm PVP-coated AgNPs to reach 2.5 mg Ag L⁻¹. Silver-treated mesocosms all had 900 mg Ag added to bring the watercolumn to 2.5 mg Ag L⁻¹. Since the AgNO₃ treatment received a sizable NO₃⁻ input, we added an equivalent amount of NO₃⁻ to all other treatments in the form of KNO₃. The characterization of the AgNP stocks used in this experiment have been previously described.21,22 Treatments were applied mid-day on August 17, 2010, (Day 0) and were applied by pouring treatment water into a funnel with a 10 mL pipet tip on the end of the funnel to slow the flow into the watercolumn. The tip was submerged, but kept within 1 cm of the surface and moved in a grid to add material as uniformly as possible to the watercolumn.

The concentration and rate of addition of AgNPs in this experiment are admittedly high, more consistent with existing toxicological studies than they were with predicted environmental exposure scenarios. Additionally, rather than Ag⁰NPs, what has been observed in waste streams has been Ag₅S-NPs, which have been shown to have diminished toxicity in lab studies5−7,27 when compared to Ag⁰NPs. While the 2.5 mg Ag L⁻¹ concentration was easily at the low end of the range used in many toxicological studies and is quite low in comparison to concentrations used in many other nanoparticle impact studies conducted to date, it is more similar to untreated photoprocessing effluents than it is to expected concentrations in wastewater effluent, let alone wetlands and streams.25,26,28 In this experiment, these high concentrations were a compromise...
between expected analytical tractability—measuring the fate, transformation, and impacts of AgNPs in a complex environment over the full course of the 15 month experiment—and environmental realism. While actual loading rates and water-column concentrations are likely to be much lower, it should also be mentioned that those exposures are likely to be in the form of chronic addition of silver, and so the cumulative load over time may be similar. Additionally, by studying AgNPs in comparison to Ag⁺, we are able to better bound the range of what is possible in this worst case scenario, and describe the similarities and differences of AgNP impacts and Ag⁺ in this complex wetland ecosystem environment.

**Sampling.** For the first 30 days of this experiment, we focused our analyses on the biogeochemistry of the aquatic zone and the treatment effects on plants, phytoplankton, and microbes; these were the organisms that were most abundant, most directly influencing biogeochemistry, and directly affected by the addition of silver. Full details on sampling protocols and sample handling and treatment are available as SI. Briefly, water was collected from 10 cm depth for dissolved solutes, dissolved organic carbon (DOC), and silver concentration. Water for dissolved gases was sampled at the surface using established protocols with modifications. Vegetation in the aquatic compartment was photographed on Days 2, 7, and 16 from approximately 1 m above the aquatic end of the mesocosms.

**Analyses.** While a subset of water chemistry parameters are highlighted in this paper, water chemistry parameters measured included: the anions $\text{Cl}^-$, $\text{SO}_4^{2-}$, $\text{Br}^-$, and $\text{NO}_3^-$, measured on a Dionex ICS 2000 Ion Chromatograph; $\text{NH}_4^+$ and $\text{PO}_4^{3-}$ measured on a Lachat QuickChem 8500; nonpurgeable dissolved organic carbon (our metric of dissolved organic carbon, DOC) and total dissolved nitrogen (TDN) measured on a Shimadzu TOC-VCPH Analyzer with a TNM-1 module. Dissolved gas samples from headspace equilibrations (see SI) were injected into a Shimadzu 17A Gas Chromatograph using a Tekmar 7050 Headspace Autosampler, as has been previously described. Water column silver was measured on a PerkinElmer 3100 flame-AAS up to Day 8, after which point samples were measured on a PerkinElmer 5100 GF-AAS. Ultrafiltered samples were all measured by GF-AAS. Phytoplankton were enumerated using a BD FACSCalibur Flow Cytometer as previously described. Non-photosynthetic prokaryotes were quantified by flow cytometry by first staining the samples with Sybr Green-I as has been previously described. Silver speciation was estimated using Visual MINTEQ V3.0 (KTH Royal Institute of Technology, Stockholm, Sweden), which was parameterized using measured solute concentrations and pH.

**Statistics.** In examining solute and gas concentrations over time, we had three questions. First, were there any differences between the true control treatment and the coating controls over time? Second, were there differences between the control(s) and silver treatments over time? Third, were there differences among the silver treatments over time? To answer all of these questions, we conducted two-way ANOVA with time and treatment as factors, and using Holm-Sidak posthoc tests for multiple comparisons within an individual factor.

## RESULTS

**Silver Concentrations in the Water Column.** Within 24 h of dosing the water column of each mesocosm, total silver concentrations dropped from our target of 2.5 mg/L down to 2.02 ± 0.12, 1.99 ± 0.07, and 0.85 ± 0.24 mg Ag L⁻¹ for the GA-AgNPs, PVP-AgNPs, and AgNO₃, respectively (Figure 1A, SI Table S1; mean ± standard error of the mean). Total silver concentrations declined most rapidly in the AgNO₃ treatment, where Ag concentrations were significantly lower in the first 4 days following dosing ($p < 0.05$, days 1–4). Despite their size and coating differences, there were no significant differences in water column silver concentration between the two AgNP treatments on any individual date. By day 6, silver concentrations in all Ag treatments had converged to 0.44 ± 0.06 mg Ag L⁻¹. Total silver concentrations were nearly identical to samples filtered through 0.7 μm filters for all three silver forms across all days, with a regression between filtered and ultrafiltered silver yielding an $r^2 = 0.993$. Between days 8 and 28, Ag concentrations in all Ag treatments remained fairly stable and were similar across treatments (averaging 0.13 ± 0.02 mg Ag L⁻¹). Silver concentrations in the <3 kDa ultrafiltrate were 3 orders of magnitude lower than total or filtered silver concentrations. These concentrations did not differ significantly between Ag treatments on any date. Concentrations declined to below our detection limit (<0.0005 mg Ag L⁻¹) by Day 6, at which point we stopped measuring the ultrafiltrate fraction.

**Organismal and Ecosystem Responses.** Within 2 days of adding silver, we observed widespread leaf loss and browning of the submerged and floating aquatic plants within all Ag dosed mesocosms. All three of the dominant macrophytes (P. diversifolius, E. densa, and L. punctata) showed visual evidence of toxicity, with a loss of photosynthetic pigments from the foliage. These qualitative changes were most apparent in the GA-AgNP and AgNO₃ treatments, with a smaller fraction of foliage showing evidence of toxicity in the PVP-AgNP treatment (Figure 2), and stood in marked contrast to the controls. We observed a similar trend for phytoplankton, with cell densities declining in the three Ag treatments over the first 4 days (Figure 3).
This decline in macrophyte biomass was accompanied by a short-lived bloom of water column prokaryotes (bacteria and archaea), with water column cell densities increasing by three to 4-fold within a day of dosing in all silver treatments (Figure 3, SI Table S2; \( p < 0.05 \) as compared to controls on: day 1, GA-AgNPs and PVP-AgNPs; day 2, all treatments; day 3, AgNO\(_3\) and GA-AgNPs). The largest increases in prokaryote abundance were observed in the GA-AgNP treatment, which maintained significantly higher prokaryote cell densities through day 4. A large increase in prokaryote cell numbers in all mesocosms on day 6 is attributed to an intense rain event (24 mm in 18 min; SI Figure S1), which caused considerable erosion of soil particles and associated microbiota from the adjacent upland into the water column. No consistent treatment effects were observed subsequent to this event.

We observed new growth of macrophytes in all silver treatments by Day 16. While we lack quantitative measures of total plant biomass, we documented that \( P. \) diversifolius began producing new surface foliage as early as Day 9 in all Ag treatments, while \( E. \) densa was observed to have new shoots from old growth by Day 16 in all silver treatments. The phytoplankton also began increasing in abundance in all silver treatments (Figure 3B) giving significantly higher biomass for some portion of the time from days 8 to 28 compared to controls. The phytoplankton in the GA-AgNP treatment increased, but that increase was only marginally different from controls and converged with controls by day 28. Phytoplankton abundance in the PVP-AgNP treatments showed a still different pattern, slowly increasing in abundance through day 28, becoming significantly different from controls starting on day 20.

In all three Ag treatments, tissue concentrations ranged from 93 ± 7 to 4180 ± 250 mg Ag kg\(^{-1}\) for macrophytes growing in the water column (\( E. \) densa, \( P. \) diversifolius, and \( L. \) punctata; Figure 4). Each species accumulated the highest concentrations of tissue Ag in the AgNO\(_3\) treatment, followed by GA-AgNPs, and then PVP-AgNPs, though the concentrations were not always significantly different between treatments (Figure 4). Treatment differences were most pronounced for \( E. \) densa, which consistently accumulated higher biomass Ag in all treatments over the first week (Figure 4). By day 28, tissue Ag concentrations had increased in \( P. \) diversifolius and \( L. \) punctata and declined in \( E. \) densa so that treatment and species differences were less apparent.

**Dissolved Solutes and Gases.** Within the first 24 h following dosing, dissolved organic carbon (DOC) increased 3-fold in the AgNO\(_3\) and GA-AgNP treatments, rising by 20 mg
C/L over the background concentration of 12.5 ± 0.4 mg C/L in both treatments. Concentrations of DOC were also elevated in the PVP-AgNP treatment, increasing by ∼50% above pretreatment concentrations. DOC concentrations were unchanged in the control mesocosms over this initial time period. These elevated DOC concentrations began to decline in all silver treatments on day 2, finally returning to a level equivalent to control mesocosms by day 13 (Figure 5, SI Table S3). From day 13 to day 28, DOC concentrations increased in all mesocosms due to evaporative losses of water, but there were no differences between treatments.

Water column chloride (Cl⁻) also increased in all Ag treatments (Figure 5). The increase was similar for the AgNO₃ and GA-AgNP treatments (2.2 ± 0.24 mg Cl⁻ L⁻¹), and a less dramatic, but still significant increase in the PVP-AgNP treatment (0.77 ± 0.48 mg Cl⁻ L⁻¹). Chloride concentrations varied very little 0.30 ± 0.12 in control mesocosms over the same period. Chloride concentrations subsequently declined in the Ag treatments from day 3 until day 13. At that point, Cl⁻ concentrations began to increase in all mesocosms through evaporative concentration. Treatment effects persisted to day 22 for GA-AgNPs, and through to day 28 for the AgNO₃ treatment.

Water column dissolved oxygen concentration (DO) declined precipitously in both the AgNO₃ and GA-AgNP treatments, dropping to a minimum of 0.31 ± 0.02 mg O₂ L⁻¹ in the AgNO₃ on day 3 and 0.41 ± 0.06 mg O₂ L⁻¹ in the GA-AgNPs on day 2 (Figure 6A, SI Table S4). In contrast, DO concentrations in control mesocosms never fell below 4.4 mg L⁻¹. Although DO declined slightly in the PVP-AgNP treatments, DO concentrations were never significantly different from controls. Dissolved oxygen concentrations remained significantly depressed through day 22 in GA-AgNPs and day 28 in AgNO₃.

The decrease in DO was accompanied by dramatic increases in carbon dioxide (CO₂) concentrations in all three silver treatments. CO₂ concentrations increased by ∼4000 ppmv in the AgNO₃ and GA-AgNP treatments, and by 2300 ppmv in the PVP-AgNP treatments by day 2, while concentrations in our controls remained <2000 ppmv of their pre-experiment concentrations (Figure 6, SI Table S4). The increase in dissolved CO₂ was short-lived in the PVP-AgNP treatments, converging on controls by day 3, but increases relative to controls were sustained in the GA-AgNP treatment through day 8 and in the AgNO₃ treatment through day 20. There was also a small and short-lived but significant increase in CO₂ associated with our GA-Control treatment on day 2. Among all measured parameters, this is the only one in which there was a divergence of a coating control from the control treatment, likely due to the respiration of gum arabic by microbes.

Produced by methanogenic archaea under anaerobic conditions, methane (CH₄) was present at low but measurable concentrations in all mesocosms prior to treatment (averaging 37 ppmv) but increased to extremely high concentrations of ∼2000 ppmv in all three silver treatments by the third day following dosing (Figure 6). In contrast to all other measured solutes and gases—which had similar trends between the three silver treatments but divergent magnitudes—both the magnitude and duration of increases in CH₄ concentrations were similar for all three Ag treatments. All Ag treatments converged on the control concentration (averaging 52 ± 16 ppmv) by Day 6.

■ DISCUSSION

There were dramatic impacts at the organismal and ecosystem level in what is one of the most complete and realistic fully replicated environmental exposure experiments with nanomaterials to date. Both GA-AgNPs and their dissolved counterpart, AgNO₃, caused several effects including: a rapid fold increase in DOC, CO₂, and Cl⁻; and rapid depletion of O₂ in the water column. The duration and direction of impacts were similar for the PVP-AgNP treatment; however, the magnitude of these changes were lower. However, in all three Ag treatments, there was a 40-fold increase in CH₄.
Silver. On the basis of previous work, we expected that ionic Ag (added as AgNO₃) would persist in the water column longer than AgNP-Ag, and that the GA-AgNPs would be more stable than the larger PVP-AgNPs, given their divergent distribution coefficients. Instead, water column Ag concentrations in the AgNO₃ treatment decreased rapidly, falling by 67% of initial concentrations within 24 h. In contrast, in the GA-AgNP and PVP-AgNP treatments, water column silver concentrations dropped by only 20% over 24 h. In fact, Ag concentration over time of these two very different nanoparticles was nearly identical throughout the experiment.

We suggest that the discrepancy between the predicted and observed behavior of AgNO₃ and AgNPs alike can be explained by the effects of all Ag treatments on submersed and floating macrophytes, which resulted in the release of large quantities of tissue derived Cl⁻ and DOC to the water column. The extremely rapid decrease in Ag concentration in the first 24 h following AgNO₃ addition could easily be attributed to the 13-fold increase in water column Cl⁻ concentrations in the AgNO₃ treatments. While initial speciation modeling with background Cl⁻ concentrations suggested that very little AgCl(s) would be formed, the same model suggests that at post-treatment Cl⁻ concentrations, nearly 82% of Ag would be expected to form AgCl(s) (SI Figure S2) either as colloids or as a precipitate. The release of the micronutrient Cl⁻ from plant tissues was an unexpected treatment effect, but this explanation is realistic given that Cl⁻ can be found at concentrations from 0.2 to 2% of dry plant mass.38 The increase in dissolved organic matter (DOM; quantified as DOC) in both AgNP treatments likely lead to their nearly identical concentrations during the first 30 days of this experiment ($r^2 = 0.997$). The released DOM likely stabilized the relatively unstable PVP-AgNPs as primary particles.21

For all these silver treatments, additional insight can be gained by calculating the rate constant for removal of silver for all three treatments, assuming first-order kinetics from day 1 through day 8 using the following equation:

$$[\text{Ag}]_t = [\text{Ag}]_0 e^{-kt}$$

where $[\text{Ag}]_t$ is the Ag concentration at time $t$, $[\text{Ag}]_0$ is the calculated initial Ag concentration, and $k$ is the removal rate constant which is itself a composite of: aggregation; sedimentation; sorption/uptake by plants; and sorption/uptake by biofilms. Surprisingly, there were no significant differences in the rate constant between AgNO₃, GA-AgNPs, and PVP-AgNPs during this time period (1-way ANOVA, $p = 0.629$), and $k$ averaged 0.21 ± 0.066 day⁻¹ across all three silver treatments. This suggests that the processes driving Ag concentration may have been similar at later time points, despite having been dramatically different in the first 24 h between the AgNO₃ and AgNP treatments. There are several hypotheses that could explain this: Ag⁺ was likely rapidly transformed by Cl⁻ released from macrophytes into small AgCl(s) colloids, which may have been stabilized by DOM and behaved similarly to the AgNPs; Ag⁺ in the presence of Cl⁻ and light or high concentrations of freshly derived plant DOM may have been reduced to Ag(0) nanoparticles in the watercolumn; or the factors driving Ag⁺ removal may have been mechanistically distinct from those driving AgNP removal, but may have been kinetically similar. However, with the data we have, it is not possible to readily distinguish between these hypotheses.

Plants accumulated high concentrations of silver after 7 days, with biomass silver concentrations ranging from a low of 93 ± 7 mg Ag kg⁻¹ in the floating L. punctata in the PVP-AgNP treatment, to a high of 4180 ± 250 mg Ag kg⁻¹ for E. densa in the AgNO₃ treatment. The highest silver concentration for AgNPs was for the GA-AgNP treatments with E. densa at 3060 ± 250 mg Ag L⁻¹. Concentrations of Ag in plants in PVP-AgNP were generally half those found for the GA-AgNP, which may be due to higher bioavailability of the smaller GA-AgNPs. Regardless of whether it was due to uptake by plants, epiphytic biofilms, or sorption,39 plants were an important early sink of Ag.

Ecosystem Responses. On the basis of previous experiments, we had expected that the addition of AgNPs and AgNO₃ would have divergent impacts with the effects of AgNO₃ > GA-AgNPs > PVP-AgNPs. Furthermore, these effects were expected to be modest given our 2.5 mg L⁻¹ concentration pulse exposure.7,16,17 As predicted, the effects of PVP-AgNPs were more modest than either GA-AgNPs or AgNO₃. However, rather than the expected modest but divergent effects of the AgNO₃ and GA-AgNP treatments, both led to a series of dramatic ecosystem level effects which were virtually identical for these two different forms of silver.

In both the AgNO₃ and GA-AgNP treatments, there was a cascade of ecosystem level impacts, all with nearly identical magnitude. Initially, there was leaf senescence for the dominant macrophytes, together with declines in phytoplankton biomass. As autotrophs released an estimated 10% of their C into the water column as DOC in each of these two treatments, heterotrophic prokaryotes rapidly increased in abundance and consumed dissolved oxygen. Coupled with the loss of photosynthetic capacity, this drove the water column to hypoxia and significantly increased the water column concentrations of carbon dioxide. These ecological impacts were remarkably similar in both magnitude and timing between the GA-AgNP and AgNO₃ treatments despite the fact that AgNO₃ had a much more rapid initial decline in water column silver concentrations.

While PVP-AgNPs had modest or nonsignificant impacts on DOC, CO₂, and O₂, those impacts were always in the same direction as GA-AgNPs and AgNO₃. Strangely, the most striking ecosystem level impact was nearly identical in all three silver treatments: dissolved methane increased 40-fold in all three silver treatments at 2–3 days after Ag addition. While conditions were primed for increased methane in the AgNO₃ and GA-AgNP treatments (elevated DOC and CO₂, depleted O₂), in the PVP-AgNP treatment they were not, yet all three treatments had the same high elevated concentration of methane. These results suggest that the water column CO₂, DOC, and DO concentrations were not the primary drivers of methane concentrations. We therefore speculate that the patterns in methane could have been the result of several different mechanisms: a direct effect of Ag on sensitive membrane bound methane monoxygenase enzymes used by the microbes that metabolize methane;18 an indirect effect of Ag on methanotrophs mediated by sediment dissolved oxygen and substrate concentrations; or the release of dissolved CH₄ from plant biomass.40

Through our use of a mixed community of organisms in realistic environmental media and field conditions, our attempt to understand the impacts of nanoparticles on ecosystems led to surprising conclusions. Despite great dissimilarities in the form of silver added in each of the three silver treatments,
overall they had similar effects, and all were removed rapidly from the water column, ending up in plants and sediment. The convergence we observed in terms of the impacts and fate of Ag was driven in large part by the cascade of biotic impacts and interactions that resulted from the Ag additions. Our most interesting findings, therefore, are the similarities between AgNO₃ and AgNPs, and the surprising and dramatic impacts of the environment on Ag fate. While we expected the environment to impact the AgNPs, we did not anticipate the nature of that interaction to be a result of the original impacts of Ag on the environment.

ASSOCIATED CONTENT

Supporting Information

Figures showing cumulative rainfall, and modeled silver speciation; additional details about mesocosm setup, sampling, sample handling, and sample analyses; tables detailing silver concentrations, phytoplankton and prokaryote densities, dissolved solutes, and watercolumn dissolved gases. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

*Phone: +1 (919) 660-7262; fax: +1 (919) 660-7293; e-mail: bpc8@duke.edu.

Notes

The authors declare no competing financial interest.

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