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Nanosilver and the Microbiological Activity of the Particulate Solids versus the Leached Soluble Silver

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Abstract

Nanosilver (Ag NPs) is currently one of the most commercialized antimicrobial nanoparticles with as yet, still unresolved cytotoxicity origins. To date, research efforts have mostly described the antimicrobial contribution from the leaching of soluble silver, while the undissolved solid Ag particulates are often considered as being microbiologically inert, serving only as source of the cytotoxic Ag ions. Here, we show the rapid stimulation of lethal cellular oxidative stress in bacteria by the presence of the undissolved Ag particulates. The cytotoxicity characteristics are distinct from those arising from the leached soluble Ag, the latter being locked in organic complexes. The work also highlights the unique oxidative stress-independent bacterial toxicity of silver salt. Taken together, the findings advocate that future enquiries on the antimicrobial potency and also importantly, the environmental and clinical impact of Ag NPs use, should pay attention to the potential bacterial toxicological responses to the undissolved Ag particulates, rather than just to the leaching of soluble silver. The findings also put into question the common use of silver salt as model material for evaluating bacterial toxicity of Ag NPs.

Keywords: silver nanoparticles; Ag solids; silver leaching; toxicity; reactive oxygen species

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Introduction

The rapid development in nanotechnology has seen inorganic nanomaterials such as nanosilver, copper oxide and zinc oxide, subjected to advanced physicochemical manipulation to exhibit powerful antimicrobial activity (Gunawan et al. 2009, 2011, 2013a, Hajipour et al. 2012).

Among these materials, nanosilver (silver nanoparticles, Ag NPs) is currently one of the most commercialized due to its potent and broad-spectrum antimicrobial characteristics (Consumer Products Inventory – Project on Emerging Nanotechnologies). Along with applications as core or co-antimicrobial ingredients in wound dressings and internal catheters (Ge et al. 2014), Ag NPs have also been incorporated in an increasing array of consumer products (Deardorff 2014), ranging from personal care products, textiles and household appliances to food and beverages and even children’s products (Benn et al. 2010, Quadros et al. 2013). The widespread use is despite the ill-defined antimicrobial mechanisms of Ag NPs, in particular the lack of knowledge regarding the origins of cytotoxicity. The controversy has been at least one of the underlying reasons for regulatory bodies to still classify and regulate Ag NPs as regular bulk silver.

Therefore, the nanoparticles are subjected to the same reporting requirements, threshold levels and toxicity tests as bulk silver, despite the mounting evidence indicating differences in their antimicrobial potency and properties (Faunce and Watal 2010). The antimicrobial activity of Ag NPs is influenced by the particles’ physicochemical characteristics (e.g. size, shape, surface functional groups) as well as interactions with the particles’ environment. In real-world settings of Ag NPs antimicrobial applications, the almost inevitable contact of the nanoparticles with aqueous environments, including those in the environment and in the human body, will lead to leaching of soluble silver species through oxidative dissolution of the silver metal (Trop et al. 2006, Benn and Westerhoff 2008, Liu et al. 2012, Sotiriou et al. 2012). Considerable research efforts have described the cytotoxic activity of the leached soluble silver on bacteria, even in their various forms, such as the soluble Ag(I)-chloride anionic complexes (Levard et al. 2013) and organo complexes (Gunawan et al. 2009), as a result of potential interactions of the released
silver with the ubiquitous presence of halides (Cl\textsuperscript-, Br\textsuperscript-, I\textsuperscript-) and biomolecules in the environment and in body fluids (Silver 2003, Liu et al. 2012, Eckhardt et al. 2013). Uncertainty however, still lingers as to the bacterial toxicological responses to the undissolved Ag residue (Gunawan et al. 2009, Sotiriou and Pratsinis 2010, Xiu et al. 2012), that remains after leaching of silver. The solid Ag particulates have been indicated to physically interact with cellular membranes of bacteria (Sondi and Salopek-Sondi 2004, Mirzajani et al. 2011), but otherwise are often regarded as being inert, indirectly contributing to the antimicrobial activity as a source of the cytotoxic Ag ions. This view is inclusive of the hypothesized Trojan-horse type of Ag NPs cytotoxicity, whereby leaching occurs intracellularly following uptake of particles, or, the suggested cell-particle contact to cause additional leaching at the cell-particle interface and in turn, increasing the uptake of Ag ions by bacteria (Lemire et al. 2013, Bondarenko et al. 2013). The elucidation of the source of Ag NPs cytotoxicity will not only clarify the nanoparticles’ ‘true’ antimicrobial potency in real-world applications, but will also contribute to more accurate assessments of their long-term impact on the environment and human health.

Here, we investigated the origins of Ag NPs cytotoxicity through detailed investigations of bacterial toxicological responses to the ‘overall’ presence of nanosilver (i.e. both leached soluble Ag and Ag particulate residue are present in the systems), as compared to those of the corresponding pre-leached filtered Ag leachate samples. Nanosilver in products can be in the forms of nano-sized Ag(I) or metallic Ag\textsuperscript{0} coated on or impregnated in support materials (Gunawan et al. 2017). As model material, the current work used nanosilver in the form of nano-sized Ag\textsubscript{2}O deposits (d\textsubscript{TEM} = 2 nm (Gunawan et al. 2009)) homogenously dispersed on the surface of inert TiO\textsubscript{2} support (d\textsubscript{TEM} = 30 nm (Gunawan et al. 2009)). It is noteworthy to point out that studies have observed discrepancies on the leaching behaviour as well as capability of cellular oxidative stress stimulation of Ag(I) versus Ag\textsuperscript{0} nanoparticles (Gunawan et al. 2009, Gunawan et al. 2013b). Nonetheless, the generated knowledge of cellular responses to the two
fundamental forms of nanosilver-derived microbiologically active components, that is, the
leached soluble silver and the solid Ag particulates in the present study, is relevant to the
countless nanosilver design with variation in the particle’s properties (e.g. size, shape and
oxidation states). This facile approach enables unambiguous elucidation of the source of
nanoparticulate cytotoxicity without the need to employ simulation materials, such as soluble Ag
salt (Gunawan et al. 2009, Sotiriou and Pratsinis 2010, Gunawan et al. 2011, Bondarenko et al.
2013, Ivask et al. 2014), which, as also shown in the current work, exhibits different cytotoxicity
characteristics. We report cytotoxic activity of the solid Ag particulates on bacteria, distinct from
the leached soluble silver.

Methods

Synthesis of Ag NPs and Preparation of Ag leachate from NPs

The 5 at% Ag/TiO$_2$ nanoparticles as finely dispersed Ag$_2$O on inert TiO$_2$ support were
synthesized using the flame spray pyrolysis (FSP) technique as earlier described (Gunawan et al.
2013b, note that at% refers to the percentage of Ag atom relative to the total number of atoms in
the particle). TEM images of the particles and XPS spectra that confirm the presence of silver (I)
oxide are available (Gunawan et al. 2009). The Ag-leachate was prepared by aseptically pre-
dissolving known amounts of Ag NPs (3, 6, 8, 10 mg Ag L$^{-1}$) in sterile Luria Bertani (LB) broth
(5 g L$^{-1}$ yeast extract, 10 g L$^{-1}$ tryptone, 5 g L$^{-1}$ NaCl in deionized water) at 37°C, 280 rpm under
dark conditions for 6 h, unsonicated. The undissolved particulates (mean aggregate size = 1.09 ±
0.03 µm by dynamic light scattering (Gunawan et al. 2009)) were removed by centrifugation
(5,000 rpm) followed by filtration of the leachate with 0.22 µm polyethersulfone membrane
(Millipore Express). Comparable light scattering intensity of the filtered Ag leachate to that of
the filtered LB medium confirmed the removal of the solid Ag residue (data not shown). The
concentration of soluble silver in the filtered Ag leachate was determined by inductively coupled
plasma mass spectrometry (ICP-MS) (Nexion 300D, PerkinElmer). ICP-MS analysis was also performed on the undissolved Ag residue (3-4 h digestion with 70% (v/v) HNO$_3$ to dissolve the Ag solid). This Ag solid concentration (no cells) reflected, at least in approximation, the presence of the undissolved Ag fraction in the nanoparticle-bacteria exposure systems (note the comparable leaching of Ag NPs in the presence and absence of bacteria, Figure 1 and S1, Supplementary Data). The ICP-MS analysis of the (digested) solid Ag residue and the corresponding Ag leachate fractions (undigested) found that their concentrations added up (within 10-15%) to the nominal total Ag concentrations of the nanoparticles (Figure S1). Finally, the ICP-MS analysis of digested leachate samples found comparable Ag concentrations before and after digestion, which further validated the removal of the solid Ag residue. Suspended Ag/TiO$_2$ particulates in the growth medium is expressed as mg L$^{-1}$ to reflect their heterogeneous presence, while the homogeneous nature of soluble Ag is referred to in ppm.

**Bacterial Growth Studies with Ag NPs, Ag leachate and AgNO$_3$ salt**

The growth experiments on *Bacillus subtilis* strain UNSW 448700 were carried out in triplicate in LB culture medium at 37°C, 280 rpm under dark conditions for 6 h. To prepare the bacterial inoculum, a single agar plate colony was cultured overnight at 30°C, 220 rpm in LB broth. A measured volume of 1-2 mL of the overnight culture (typical OD$_{600}$ of 6-8) was transferred into 50 mL fresh LB broth for a further 0.5-1 h conditioning at 37°C, 280 rpm. For the Ag NPs and AgNO$_3$ exposure, pre-weighed Ag NPs (1.1x of the intended dosage) and 0.5 mL (110x concentrated of the intended dosage) solution of AgNO$_3$ were aseptically added into 50 mL and 49.5 mL LB respectively. The experiments were initiated by the addition of 5 mL bacterial inoculum into the 50 mL broth containing suspended Ag NPs or dissolved silver salt (OD$_{600}$ bacteria initial = 0.04, corresponding to $\sim$2 x $10^7$ cfu mL$^{-1}$). For the Ag leachate exposure, 5 mL of the bacterial inoculum was added into 50 mL LB containing 1.1x concentrated pre-leached Ag...
NPs (particle-free). The growth profiles were determined by OD$_{600}$ measurement of the biomass (UV/Vis spectrophotometer, Hitachi U-1100) and the growth inhibiting effects were assessed relative to controls with no added silver. A cell-free silver control (particulates or soluble silver) was employed as a reference to obtain the OD$_{600}$ corresponding to the bacteria. The corresponding leaching profile of Ag NPs during the bacterial exposure was measured by ICP-MS (Nexion 300D, PerkinElmer). For this purpose, a measured volume was sampled from the NPs-exposed culture, centrifuged (5,000 rpm) then filtered with the 0.22 µm membrane to remove the bacteria and Ag solid. The resulting solution was 100x diluted in deionized water and subjected to the ICP-MS analysis.

Detection of Intracellular ROS and Cell Viability

The measurement of cellular ROS generation was performed using the cell permeable oxidative reporter dye H$_2$DCFDA (2′,7′-dichlorodihydrofluorescein diacetate, Sigma-Aldrich). Following its uptake, cellular esterases cleave the diacetate moieties of H$_2$DCFDA to form H$_2$DCF, which readily transforms to the fluorescent DCF when reacts with ROS. The cell viability assay was based on the fluorescent nucleic acid dye propidium iodide (Sigma-Aldrich) staining. PI enters cells with damaged cytoplasmic membrane, while being excluded by healthy cells. Following removal of the culture medium by centrifugation, samples from the Ag NPs, Ag leachate and AgNO$_3$ exposure systems (and the silver-free controls) were washed and re-suspended in sterile saline (8 g L$^{-1}$ NaCl, 0.2 g L$^{-1}$ KCl) at 2.5 x 10$^8$ CFU mL$^{-1}$. Independent cellular ROS and cell viability assays were carried out with 10 µM H$_2$DCFDA and 30 µM PI for 1 h and 15 min respectively, at room temperature under dark conditions. The stained cells were washed with saline and analysed by flow cytometry (FACSCanto™ II, BD Bioscience) at 488 nm excitation with 530 nm and 670 nm emission filter settings for the detection of DCF and PI fluorescence respectively. DCF fluorescence was also measured using a microplate reader (Ensight™ Multimode, Perkin Elmer) at 492 nm and 520 nm excitation and emission filter settings.
respectively. The stained cells were also visualized with a BX51WI fluorescence microscope (Olympus) with 460–490 nm excitation filter settings.

Results and discussion

**Bacterial growth inhibition: Activity of the solid Ag particulates, the leached soluble Ag and silver salt**

To distinguish the cytotoxicity or antimicrobial contribution of the leached soluble Ag and the undissolved Ag particulates, we exposed a model bacteria *B. subtilis* UNSW 448700 to 0 – 10 mg Ag L\(^{-1}\) NPs (Ag/TiO\(_2\)) and compared the bacterial growth to that of the corresponding leachate-only systems, as a function of soluble silver detected in the exposure systems. The leachate samples were prepared by aseptically pre-dissolving Ag NPs in culture medium followed by removal of the solid Ag residue. Firstly, as shown in Figure 1a, the dose-response growth inhibiting effects of the Ag NPs correlates with the increasing concentration of soluble silver that leached from the NPs. The extent of growth of *B. subtilis* was reduced to ∼80% upon exposure to 3 mg Ag L\(^{-1}\) NPs (1.3 ppm silver leached into the culture medium at equilibrium) relative to silver-free control cultures after 6 h. The control cultures were characterized by a relatively short lag phase of 1 h, followed by 3 to 4 h active exponential growth phase before entering the stationary phase at 6 h (Figure 1b). Increasing the NPs dosage to 6 mg Ag L\(^{-1}\) (2.7 ppm leached Ag) saw 50% bacterial growth, while almost complete growth suppression was observed at MIC\(_{95}\) 10 mg Ag L\(^{-1}\) NPs exposure (4 ppm leached Ag, see Figure 1b for growth profile, MIC\(_{95}\) is minimum inhibitory concentration that cause 5% growth relative to the control).

At all of the tested Ag NPs loading, leaching of Ag from NPs was rapid, with detection of ∼70% soluble Ag (relative to the leached Ag concentration detected at equilibrium) within 5 min of the Ag NPs-bacterial exposure (see Figure 1c inset for leaching profile of 10 mg Ag L\(^{-1}\) NPs).

Equilibrium was reached in 1 h with the soluble Ag concentration remained constant afterwards, indicating absence of the Ostwald ripening phenomenon that refers to re-deposition of the

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leached Ag on larger particulates (Sotiriou et al. 2012). Increasing the Ag NPs loading saw detection of elevated soluble Ag concentration at equilibrium, with the extent of leaching essentially comparable at 38 – 40% relative to the total added Ag (Figure 1c). This is consistent to earlier studies under comparable conditions (Gunawan et al. 2009, Sotiriou and Pratsinis 2010) with the relatively high degree of leaching was due to, at least in part, the presence of organics in the culture medium as shown later in this study. Note that at all of the tested Ag NPs loadings, similar extent of leaching were observed in the absence of bacteria, therefore excluding the possibility of microbial-induced leaching of Ag (Figure S1).

Despite the correlation between Ag NPs growth inhibiting effects and Ag leaching, a comparison with bacterial growth in the corresponding leachate-only systems yields an interesting observation. Exposure of *B. subtilis* to the pre-leached soluble Ag in fact resulted in much less growth inhibition when compared to those of the corresponding Ag NPs samples (Figure 1a). The presence of ~1.3 ppm Ag leachate for example, was benign to the cultures as they grew to a similar extent as the silver-free control cultures after 6 h. This was in contrast to the ~20% growth reduction of the bacteria when exposed to the corresponding 3 mg Ag L\(^{-1}\) NPs with comparable leached soluble Ag content. At higher exposure, the bacterial growth in 4 ppm Ag leachate system was ~85% relative to the control cultures (refer to Figure 1b for growth profile), in contrast to the near complete growth suppression observed in the corresponding 10 mg Ag L\(^{-1}\) NPs system. Even doubling the concentration of Ag leachate to 8.3 ppm only slightly reduced the bacterial growth to ~75%. The findings suggest predominant cytotoxicity contribution from the undissolved Ag particulates, rather than that arising from the leached soluble Ag. Further antimicrobial simulation with an equivalent concentration of soluble Ag from AgNO\(_3\) salt as shown in Figure 1a, saw more severe growth inhibiting activity of the salt. In the presence of 4 ppm soluble Ag from AgNO\(_3\) for example, ~25% *B. subtilis* growth was
observed relative to the control cultures after 6 h (growth profile is shown in Figure 1b), in contrast to the ~85% growth in the leachate system with comparable Ag concentration. Such differences in cytotoxicity may arise from unique cellular physiological responses to the different silver species; the leached soluble Ag and the undissolved Ag particulates from Ag NPs, and the soluble silver from silver salt, as investigated in the following.

**Dynamic stimulation of cellular oxidative stress and cell death**

We carried out dynamic tracking of intracellular reactive oxygen species (ROS) generation (measured by H$_2$DCFDA assay) and cell viability (measured by propidium iodide assay, whereby PI enters cells with damaged cytoplasmic membrane, which is indicative of cell death) over the 6 h growth course of *B. subtilis* in the presence of the various forms of silver; the Ag NPs (MIC$_{95}$ 10 mg Ag L$^{-1}$ as reference point, contained 4 ppm leached Ag), its corresponding Ag leachate system (4 ppm Ag) and the equivalent AgNO$_3$ system (4 ppm Ag).

*The solid Ag particulates and the leached soluble Ag*

At 5 min exposure to 10 mg Ag L$^{-1}$NPs, a 3-fold higher cellular ROS level was detected in *B. subtilis* relative to the basal ROS levels of the silver-free control cultures, which are by-products of aerobic metabolism in bacteria (Choi and Hu 2008, Gunawan et al. 2011, Eckhardt et al. 2013) (Figure 2a, 2b, 4a). Within 30 min of Ag NPs exposure, the cellular ROS level doubled to ~6-fold of the control. A secondary oxidative stress response, the cellular ROS stimulation has been increasingly realized as one of the major cellular toxicological responses to Ag NPs in bacteria (Choi and Hu 2008, Hwang et al. 2008, Lemire et al. 2013, Gunawan et al. 2013b). The ROS generation is thought to result from destruction of the iron-sulfur [4Fe-4S] clusters of proteins by Ag metal (Xu and Imlay 2012, Lemire et al. 2013) and in turn, releasing the Fenton-active free Fe into the cytoplasm for subsequent reaction with cellular H$_2$O$_2$ to produce hydroxyl...
radicals (OH$^-$) (Imlay et al. 1988). Alternatively, indirect destruction of the iron-sulfur clusters could result from inhibition of respiratory enzymes by Ag NPs in bacteria (Li et al. 2010, 2011). The resulting premature leakage of electrons to oxygen will generate superoxide radicals (O$_2^-$) (Imlay 2003) that in turn again, induces the release of free Fe from iron-sulfur clusters in proteins (Kohanski et al. 2007). Indeed, there have been reports on the cytoplasmic presence of the solid Ag particulates upon bacterial exposure to Ag NPs, as well as the presence of the solids within the bacterial membrane layers (Morones et al. 2005, Grigor'eva et al. 2013, Pal et al. 2007). Here, 75-90% PI-positive non-viable bacteria had been detected within 5 to 30 min exposure to Ag NPs, then close to 100% bactericidal or cell death toxicity at as early as 1 h exposure (Figure 2a, 2b, 4b), which indicates cytoplasmic membrane as one of the target destruction sites of the Ag NPs-stimulated cellular ROS (1-8% non-viable cells were detected in the control cultures over the 6 h growth course) (D'Autreaux et al. 2007, Lemire et al. 2013). As expected, the levels of cellular ROS drastically dropped following the rapid high level stimulation, with the majority if not all of the bacterial population were already killed (Sintubin et al. 2011, Gunawan et al. 2013b). Up to this stage, the data suggest that the generation of high levels of cellular ROS and associated bacteria killing was likely to be responsible for the near complete suppression of *B. subtilis* growth (Figure 1a, 1b).

Interestingly, such cellular ROS stimulation was absent in the bacteria when studied in the corresponding 4 ppm Ag leachate system. Over the 6 h growth course, only basal ROS levels, comparable to those of the silver-free control cultures were detected (Figure 2a, 3a, 4a) and not surprisingly, the little to no changes in the fraction of non-viable cells relative to the control (Figure 2a, 3a, 4b). The stimulation of lethal levels of cellular oxidative stress by the presence of solid Ag particulates therefore suggests their substantial contribution to the cytotoxicity effects observed in the growth studies. Recalling the observed ~15% growth inhibition of the bacteria in
the presence of 4 ppm Ag leachate (Figure 1a, 1b), it would be reasonable to deduce that the exposure only resulted in sub-lethal cytotoxicity, causing a minor fraction of the viable cells uncultivable or slowly proliferating, as further indicated by our growth prediction based on the fraction of viable cells (Figure S2). Indeed, doubling the Ag leachate concentration to 8.3 ppm still saw typical cellular ROS (Figure 3c, 4a inset) and dead cells (Figure 3c, 4b inset) detection as those of the control cultures, despite the slightly higher growth suppression, at ∼25% (Figure 1a).

The leached soluble Ag and silver salt

The minimal cellular ROS stimulation was also seen upon exposure of *B. subtilis* to the equivalent 4 ppm soluble Ag from AgNO₃. Similar to the 4 ppm Ag leachate system, no elevated level of cellular ROS was observed over the 6 h growth course relative to the control cultures (Figure 2a, 3b, 4a). Unlike the leachate samples however, up to ∼40% non-viable cells were detected in the salt system (Figure 3b, 4b), indicating attacks on cytoplasmic membrane (Eckhardt et al. 2013). Considering the comparable Ag content, such discrepancies in cytotoxicity are most likely to result from differences in the chemical speciation of the soluble silver, as herein described. Our Ag NPs leaching study (at the MIC₉₅ 10 mg Ag L⁻¹ NPs) in the individual culture medium components revealed a characteristic trend of complexation-assisted dissolution of nanoparticles (Gunawan et al. 2011), with higher extent of Ag leaching in the peptide-rich components, in particular tryptone (90% leaching relative to the total added Ag), compared to those in the deionized water (60% leaching) or NaCl (10% leaching) (Figure 4c). A soft Lewis acid, Ag(I) forms silver-peptide complexes upon its release from NPs (Bolea et al. 2014), which is most likely to result from its strong affinity to the NH₄ donor groups of histidine (NH₄⁺), arginine (-NH₂⁻) and lysine (-NH₃⁻) amino acids and also to the thiol (-S⁻) donor groups of cysteine and methionine amino acids (Eckhardt et al. 2013). Silver-peptide complexes also form with AgNO₃ (Bolea et al. 2014), with a fraction of silver is thought to remain as free ions in
the organic-rich medium (Percival et al. 2005). Thermodynamically feasible, the co-existence of free metal ions and organo metal complexes has been reported for the chemical speciation of soluble copper salts, also a soft Lewis acid metal, in similar culture medium as that used here (Gunawan et al. 2011) (note that the current technology for elemental analysis does not differentiate free Ag ions to those locked in organo complexes (Eckhardt et al. 2013)). When compared to free Ag ions, the hindered transport of the bulkier silver-peptide complexes into bacteria (Solioz and Odermatt 1995) is thought to be at least in part, responsible for the passivated, in this case, sub-lethal cytotoxicity of the Ag leachate. Unlike free Ag ions, research indicates that soluble organo Ag complexes are not recognized by the P-type ATPase transporter present in bacteria (Luoma 2008). As also observed in the current study with the AgNO₃ systems, exposure of bacteria to Ag ions has been reported to suppress their proliferation, which was indicated to result from a ROS-independent inhibition of metabolic enzymes (dehydratases) (Xu and Imlay 2012), the lack of cellular ROS stimulation also apparent in this work. Further, complete suppression of *B. subtilis* growth was seen at 8.3 ppm Ag from AgNO₃ (Figure 1a), despite there being no change in the fraction of non-viable cells when compared to the 4 ppm Ag exposure (Figure 3d, 4b inset). Our growth prediction based on the fraction of viable cells indicates major presence of non- or slowly proliferating viable cells with the AgNO₃ exposure (Figure S2). This loss in replication could also result from the known interactions of Ag ions with DNA in bacteria (most likely with the phosphorus moieties) causing DNA condensation (Feng et al. 2000). The seemingly higher cytotoxic effects of Ag ions as compared to the organo Ag complexes are in agreement with other bacterial studies, whereby extracellular presence of thiol-containing reduced glutathione (GSH) as silver complexing agent lowered the antimicrobial activity of Ag ions on the Gram-positive *Staphylococcus aureus* and the Gram-negative *Escherichia coli* and *Pseudomonas aeruginosa* (Mulley et al. 2014). Finally, the detection of only basal cellular ROS levels in the AgNO₃ exposure systems, even at the double 8.3 ppm Ag (Figure 3d, 4a inset), rules out the oxidative stress stimulation as the main...
mechanisms of AgNO₃ cytotoxicity. Indeed, studies have found no differences in the antimicrobial activity of Ag ions under aerobic and anaerobic conditions on bacteria (Sintubin et al. 2011).

Conclusions

Here, we report multiple cytotoxicity origins of Ag NPs towards bacteria. Presence of undissolved Ag particulates in a biological environment is not inert. In their presence, rapid generation of lethal cellular ROS levels were detected in bacteria, while the corresponding leached soluble Ag, being locked in organo complexes, only imparts sub-lethal cytotoxicity. The observed differences in bacterial toxicological responses to the solid versus soluble Ag corroborate earlier reports on the distinct extent of growth inhibiting activity of the Ag NPs’ soluble and solid components (Gunawan et al. 2009, Sotiriou and Pratsinis 2010). With regard to the widespread use of Ag NPs, the resolved unique toxicological responses are expected to result in better recognition of the antimicrobial potency of the nanoparticles in real-world settings and importantly, the long-term impact. Research inquiries have shown elevated and persistent presence of silver in wounds, bladder and even in sewage and estuaries, being associated with the intended or in some cases, accidental release from nanosilver applications; the use of wound dressings, pesticides and washing machines are among the examples (Chen et al. 2004, Trop et al. 2006, Reidy et al. 2013, Donner et al. 2015, Beddow et al. 2017). The current findings imply bacterial toxicological responses to not only the leached soluble Ag, but also the Ag particulates in the microbial habitats. Indeed, studies have observed disruptions in the dynamic and balance of microbial communities from natural aquatic waters upon exposure to nanosilver (Das et al. 2012, Beddow et al. 2017), with the work also detecting presence of soluble Ag and aggregates of Ag from nanosilver in these environmental samples (Beddow et al. 2017). The resolved toxicological responses is key to the elucidation of the recently discovered bacterial potential for adaptation to Ag NPs cytotoxicity (Das et al. 2012, Gunawan et al. 2013b). Finally, the work
highlights the unsuitability of soluble silver salt as model material for Ag NPs cytotoxicity in biological environments, noting a distinct ROS-independent antimicrobial characteristic of soluble Ag when supplied as AgNO₃ salt.

Acknowledgments

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Declaration of interest

The authors declare no conflict of interest.

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Supplementary material is available: Supplementary Figure S1, S2 and S3.

Figure captions

Figure 1. Bacterial growth in the presence of Ag NPs, Ag NPs leachate, silver salt and leaching of Ag NPs. (a) Growth of B. subtilis (6 h) relative to cell-only control upon exposure to Ag NPs (3, 6, 8, 10 mg Ag L\(^{-1}\)), Ag leachate from NPs and AgNO\(_3\) as a function of soluble silver detected in the bacterial exposure systems (the growth studies were performed in LB medium). (b) Growth profiles of the bacteria in the presence of 10 mg Ag L\(^{-1}\) NPs (4 ppm Ag leached into medium at equilibrium), 4 ppm Ag leachate from NPs and 4 ppm Ag from AgNO\(_3\). Also shown is the cell-only control growth profile (dashed line). The growth in the presence of Ag was normalised to the extent of growth of the control (in colony forming units, cfu). (c) The corresponding equilibrium leaching of Ag NPs in the bacterial exposure systems, shown in the inset is the leaching profile for 10 mg Ag L\(^{-1}\) NPs. Each data point in (a), (b), (c) is the average of triplicate experiments with error bars representing the maximum and minimum values detected. The growth studies were performed under dark conditions to render the TiO\(_2\) support
photocatalytically inactive and the benign effect of the TiO$_2$ support on $B. subtilis$ growth had been confirmed (Gunawan et al. 2013b). The growth studies were reproduced on different days with unique bacterial inoculum and particle preparations.

Figure 2. Detection of cellular reactive oxygen species (ROS, H$_2$DCFDA staining, green cells) and cell death (PI staining, red cells) of $B. subtilis$ over its growth course: (a) cell-only control and (b) in the presence of 10 mg Ag L$^{-1}$ NPs. All stained samples were imaged at comparable cell concentrations (scale bars = 50 µm).

Figure 3. Detection of cellular reactive oxygen species (ROS, H$_2$DCFDA staining, green cells) and cell death (PI staining, red cells) of $B. subtilis$ over its growth course, in the presence of: (a) 4 ppm Ag leachate from NPs (equivalent leachate to 10 mg Ag L$^{-1}$ NPs exposure), (b) 4 ppm Ag from AgNO$_3$, (c) 8.3 ppm Ag leachate from NPs and (d) 8.3 ppm Ag from AgNO$_3$. All stained samples were imaged at comparable cell concentrations (scale bars = 50 µm).

Figure 4. (a) Dynamic stimulation of cellular ROS in $B. subtilis$ measured by H$_2$DCFDA assay over its growth course (5, 30 min and 1, 3, 4.5, 6 h) upon exposure to 10 mg Ag L$^{-1}$ NPs (4 ppm Ag leached into medium at equilibrium), 4 ppm Ag leachate from NPs and 4 ppm Ag from AgNO$_3$. The detected cellular ROS was normalised to the basal ROS levels of the cell-only control growth. Shown in the inset is cellular ROS detected in the presence of 8.3 ppm Ag leachate from NPs and 8.3 ppm Ag from AgNO$_3$. (b) The corresponding dynamic cell death detection probed by PI staining of $B. subtilis$ throughout its growth course. Also shown is the fraction of dead cells detected for the cell-only control. Statistical analysis of the data was performed with one-way ANOVA followed by Dunnett’s posthoc analysis in Prism (GraphPad) (Figure S3). The experiments in (a) and (b) were reproduced on different days with unique bacterial inoculum and particle preparations. (c) Leaching profile of 10 mg Ag L$^{-1}$ NPs in cell-

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free individual components of the Luria Bertani (LB) culture medium (5 g L\(^{-1}\) NaCl, 5 g L\(^{-1}\) yeast extract, 10 g L\(^{-1}\) tryptone dissolved in deionized water). Each data point in (a), (b), (c) is the average of triplicate batches with the error bars representing the maximum and minimum.
Figure 1
Figure 2
Figure 3
Figure 4
Nanosilver and the Microbiological Activity of the Particulate Solids \textit{versus} the Leached Soluble Silver

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\textbf{Supplementary Data}

URL: http://mc.manuscriptcentral.com/tnan
Quantification of the soluble and solid Ag fractions

ICP-MS analysis was performed to determine the concentration of the leached soluble Ag and the undissolved Ag particulate residue of the Ag NPs-culture medium systems (no cells).

**Figure S1.** Concentration of the soluble and solid Ag fractions of the Ag NPs-culture medium systems (no bacteria), samples were taken at 6 h following NPs addition.
First order kinetic prediction of growth upon exposure to Ag leachate from NPs and AgNO₃

The presence of non- or slowly proliferating viable cells as a result of exposure of *B. subtilis* to 4 and 8.3 ppm Ag leachate as well as to the equivalent 4 and 8.3 ppm soluble Ag from AgNO₃ salt is validated by overestimation of the predicted biomass growth based on first order kinetic, *i.e.*

\[
\frac{[A_n]}{[A_{n-1}]} = e^{kt},
\]

where \([A_{n-1}]\) is the concentration of PI-negative (viable) cells at previous time interval. The rate coefficient (k) is estimated from bacteria-only control growth.

**Figure S2.** First order kinetic growth prediction of *B. subtilis* upon exposure to 4 & 8.3 ppm Ag NPs leachate and to 4 & 8.3 ppm Ag from AgNO₃ as compared to the actual growth.
Statistical analysis of silver-induced cellular reactive oxygen species generation and cell death detection

The statistical significance of cytotoxicity of Ag NPs, Ag NPs leachate and AgNO₃ relative to the bacteria-only control was analyzed by using a one-way ANOVA followed by Dunnett’s posthoc analysis in Prism (GraphPad).

![Figure S3](http://mc.manuscriptcentral.com/tnan)

**Figure S3.** Statistical analysis of dynamic cellular ROS and cell death detection over the course of *B. subtilis* growth upon exposure to (a, b) 10 mg Ag L⁻¹ NPs (4 ppm Ag leached into medium
at equilibrium) (c, d) 4 ppm Ag leachate from NPs (inset is 8.3 ppm Ag leachate from NPs) (e, f) 4 ppm Ag from AgNO$_3$ (inset is 8.3 ppm Ag from AgNO$_3$). The asterisks (*, **) correspond to p values of $\leq 0.01, 0.001$ respectively, and indicate statistical significance relative to the bacteria-only control (dashed lines).
Nanosilver and the Microbiological Activity of the Particulate Solids versus the Leached Soluble Silver

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Nanosilver and the Microbiological Activity of the Particulate Solids versus the Leached Soluble Silver

Abstract

Nanosilver (Ag NPs) is currently one of the most commercialized antimicrobial nanoparticles with as yet, still unresolved cytotoxicity origins. To date, research efforts have mostly described the antimicrobial contribution from the leaching of soluble silver, while the undissolved solid Ag particulates are often considered as being microbiologically inert, serving only as source of the cytotoxic Ag ions. Here, we show the rapid stimulation of lethal cellular oxidative stress in bacteria by the presence of the undissolved Ag particulates. The cytotoxicity characteristics are distinct from those arising from the leached soluble Ag, the latter being locked in organic complexes. The work also highlights the unique oxidative stress-independent bacterial toxicity of silver salt. Taken together, the findings advocate that future enquiries on the antimicrobial potency and also importantly, the environmental and clinical impact of Ag NPs use, should pay attention to the potential bacterial toxicological responses to the undissolved Ag particulates, rather than just to the leaching of soluble silver. The findings also put into question the common use of silver salt as model material for evaluating bacterial toxicity of Ag NPs.

Keywords: silver nanoparticles; Ag solids; silver leaching; toxicity; reactive oxygen species

Word count: 6472
Introduction

The rapid development in nanotechnology has seen inorganic nanomaterials such as nanosilver, copper oxide and zinc oxide, subjected to advanced physicochemical manipulation to exhibit powerful antimicrobial activity (Gunawan et al. 2009, 2011, 2013a, Hajipour et al. 2012).

Among these materials, nanosilver (silver nanoparticles, Ag NPs) is currently one of the most commercialized due to its potent and broad-spectrum antimicrobial characteristics (Consumer Products Inventory – Project on Emerging Nanotechnologies). Along with applications as core or co-antimicrobial ingredients in wound dressings and internal catheters (Ge et al. 2014), Ag NPs have also been incorporated in an increasing array of consumer products (Deardorff 2014), ranging from personal care products, textiles and household appliances to food and beverages and even children’s products (Benn et al. 2010, Quadros et al. 2013). The widespread use is despite the ill-defined antimicrobial mechanisms of Ag NPs, in particular the lack of knowledge regarding the origins of cytotoxicity. The controversy has been at least one of the underlying reasons for regulatory bodies to still classify and regulate Ag NPs as regular bulk silver.

Therefore, the nanoparticles are subjected to the same reporting requirements, threshold levels and toxicity tests as bulk silver, despite the mounting evidence indicating differences in their antimicrobial potency and properties (Faunce and Watal 2010). The antimicrobial activity of Ag NPs is influenced by the particles’ physicochemical characteristics (e.g. size, shape, surface functional groups) as well as interactions with the particles’ environment. In real-world settings of Ag NPs antimicrobial applications, the almost inevitable contact of the nanoparticles with aqueous environments, including those in the environment and in the human body, will lead to leaching of soluble silver species through oxidative dissolution of the silver metal (Trop et al. 2006, Benn and Westerhoff 2008, Liu et al. 2012, Sotiriou et al. 2012). Considerable research efforts have described the cytotoxic activity of the leached soluble silver on bacteria, even in their various forms, such as the soluble Ag(I)-chloride anionic complexes (Levard et al. 2013) and organo complexes (Gunawan et al. 2009), as a result of potential interactions of the released
silver with the ubiquitous presence of halides (Cl\(^-\), Br\(^-\), I\(^-\)) and biomolecules in the environment and in body fluids (Silver 2003, Liu et al. 2012, Eckhardt et al. 2013). Uncertainty however, still lingers as to the bacterial toxicological responses to the undissolved Ag residue (Gunawan et al. 2009, Sotiriou and Pratsinis 2010, Xiu et al. 2012), that remains after leaching of silver. The solid Ag particulates have been indicated to physically interact with cellular membranes of bacteria (Sondi and Salopek-Sondi 2004, Mirzajani et al. 2011), but otherwise are often regarded as being inert, indirectly contributing to the antimicrobial activity as a source of the cytotoxic Ag ions. This view is inclusive of the hypothesized Trojan-horse type of Ag NPs cytotoxicity, whereby leaching occurs intracellularly following uptake of particles, or, the suggested cell-particle contact to cause additional leaching at the cell-particle interface and in turn, increasing the uptake of Ag ions by bacteria (Lemire et al. 2013, Bondarenko et al. 2013). The elucidation of the source of Ag NPs cytotoxicity will not only clarify the nanoparticles’ ‘true’ antimicrobial potency in real-world applications, but will also contribute to more accurate assessments of their long-term impact on the environment and human health.

Here, we investigated the origins of Ag NPs cytotoxicity through detailed investigations of bacterial toxicological responses to the ‘overall’ presence of nanosilver (i.e. both leached soluble Ag and Ag particulate residue are present in the systems), as compared to those of the corresponding pre-leached filtered Ag leachate samples. Nanosilver in products can be in the forms of nano-sized Ag(I) or metallic Ag\(^0\) coated on or impregnated in support materials (Gunawan et al. 2017). As model material, the current work used nanosilver in the form of nano-sized Ag\(_2\)O deposits (d\(_{\text{TEM}} = 2\) nm (Gunawan et al. 2009)) homogenously dispersed on the surface of inert TiO\(_2\) support (d\(_{\text{TEM}} = 30\) nm (Gunawan et al. 2009)). It is noteworthy to point out that studies have observed discrepancies on the leaching behaviour as well as capability of cellular oxidative stress stimulation of Ag(I) versus Ag\(^0\) nanoparticles (Gunawan et al. 2009, Gunawan et al. 2013b). Nonetheless, the generated knowledge of cellular responses to the two...
fundamental forms of nanosilver-derived microbiologically active components, that is, the leached soluble silver and the solid Ag particulates in the present study, is relevant to the countless nanosilver design with variation in the particle’s properties (e.g. size, shape and oxidation states). This facile approach enables unambiguous elucidation of the source of nanoparticulate cytotoxicity without the need to employ simulation materials, such as soluble Ag salt (Gunawan et al. 2009, Sotiriou and Pratsinis 2010, Gunawan et al. 2011, Bondarenko et al. 2013, Ivask et al. 2014), which, as also shown in the current work, exhibits different cytotoxicity characteristics. We report cytotoxic activity of the solid Ag particulates on bacteria, distinct from the leached soluble silver.

Methods

Synthesis of Ag NPs and Preparation of Ag leachate from NPs

The 5 at% Ag/TiO$_2$ nanoparticles as finely dispersed Ag$_2$O on inert TiO$_2$ support were synthesized using the flame spray pyrolysis (FSP) technique as earlier described (Gunawan et al. 2013b, note that at% refers to the percentage of Ag atom relative to the total number of atoms in the particle). TEM images of the particles and XPS spectra that confirm the presence of silver (I) oxide are available (Gunawan et al. 2009). The Ag-leachate was prepared by aseptically pre-dissolving known amounts of Ag NPs (3, 6, 8, 10 mg Ag L$^{-1}$) in sterile Luria Bertani (LB) broth (5 g L$^{-1}$ yeast extract, 10 g L$^{-1}$ tryptone, 5 g L$^{-1}$ NaCl in deionized water) at 37°C, 280 rpm under dark conditions for 6 h, unsonicated. The undissolved particulates (mean aggregate size = 1.09 ± 0.03 µm by dynamic light scattering (Gunawan et al. 2009)) were removed by centrifugation (5,000 rpm) followed by filtration of the leachate with 0.22 µm polyethersulfone membrane (Millipore Express). Comparable light scattering intensity of the filtered Ag leachate to that of the filtered LB medium confirmed the removal of the solid Ag residue (data not shown). The concentration of soluble silver in the filtered Ag leachate was determined by inductively coupled
plasma mass spectrometry (ICP-MS) (Nexion 300D, PerkinElmer). ICP-MS analysis was also performed on the undissolved Ag residue (3-4 h digestion with 70% (v/v) HNO₃ to dissolve the Ag solid). This Ag solid concentration (no cells) reflected, at least in approximation, the presence of the undissolved Ag fraction in the nanoparticle-bacteria exposure systems (note the comparable leaching of Ag NPs in the presence and absence of bacteria, Figure 1 and S1, Supplementary Data). The ICP-MS analysis of the (digested) solid Ag residue and the corresponding Ag leachate fractions (undigested) found that their concentrations added up (within 10-15%) to the nominal total Ag concentrations of the nanoparticles (Figure S1). Finally, the ICP-MS analysis of digested leachate samples found comparable Ag concentrations before and after digestion, which further validated the removal of the solid Ag residue. Suspended Ag/TiO₂ particulates in the growth medium is expressed as mg L⁻¹ to reflect their heterogeneous presence, while the homogeneous nature of soluble Ag is referred to in ppm.

**Bacterial Growth Studies with Ag NPs, Ag leachate and AgNO₃ salt**

The growth experiments on *Bacillus subtilis* strain UNSW 448700 were carried out in triplicate in LB culture medium at 37°C, 280 rpm under dark conditions for 6 h. To prepare the bacterial inoculum, a single agar plate colony was cultured overnight at 30°C, 220 rpm in LB broth. A measured volume of 1-2 mL of the overnight culture (typical OD₆₀₀ of 6-8) was transferred into 50 mL fresh LB broth for a further 0.5-1 h conditioning at 37°C, 280 rpm. For the Ag NPs and AgNO₃ exposure, pre-weighed Ag NPs (1.1x of the intended dosage) and 0.5 mL (110x concentrated of the intended dosage) solution of AgNO₃ were aseptically added into 50 mL and 49.5 mL LB respectively. The experiments were initiated by the addition of 5 mL bacterial inoculum into the 50 mL broth containing suspended Ag NPs or dissolved silver salt (OD₆₀₀ bacteria initial = 0.04, corresponding to ~2 x 10⁷ cfu mL⁻¹). For the Ag leachate exposure, 5 mL of the bacterial inoculum was added into 50 mL LB containing 1.1x concentrated pre-leached Ag.
NPs (particle-free). The growth profiles were determined by OD$_{600}$ measurement of the biomass (UV/Vis spectrophotometer, Hitachi U-1100) and the growth inhibiting effects were assessed relative to controls with no added silver. A cell-free silver control (particulates or soluble silver) was employed as a reference to obtain the OD$_{600}$ corresponding to the bacteria. The corresponding leaching profile of Ag NPs during the bacterial exposure was measured by ICP-MS (Nexion 300D, PerkinElmer). For this purpose, a measured volume was sampled from the NPs-exposed culture, centrifuged (5,000 rpm) then filtered with the 0.22 µm membrane to remove the bacteria and Ag solid. The resulting solution was 100x diluted in deionized water and subjected to the ICP-MS analysis.

Detection of Intracellular ROS and Cell Viability

The measurement of cellular ROS generation was performed using the cell permeable oxidative reporter dye H$_2$DCFDA (2',7'-dichlorodihydrofluorescein diacetate, Sigma-Aldrich). Following its uptake, cellular esterases cleave the diacetate moieties of H$_2$DCFDA to form H$_2$DCF, which readily transforms to the fluorescent DCF when reacts with ROS. The cell viability assay was based on the fluorescent nucleic acid dye propidium iodide (Sigma-Aldrich) staining. PI enters cells with damaged cytoplasmic membrane, while being excluded by healthy cells. Following removal of the culture medium by centrifugation, samples from the Ag NPs, Ag leachate and AgNO$_3$ exposure systems (and the silver-free controls) were washed and re-suspended in sterile saline (8 g L$^{-1}$ NaCl, 0.2 g L$^{-1}$ KCl) at 2.5 x 10$^8$ CFU mL$^{-1}$. Independent cellular ROS and cell viability assays were carried out with 10 µM H$_2$DCFDA and 30 µM PI for 1 h and 15 min respectively, at room temperature under dark conditions. The stained cells were washed with saline and analysed by flow cytometry (FACSCanto™ II, BD Bioscience) at 488 nm excitation with 530 nm and 670 nm emission filter settings for the detection of DCF and PI fluorescence respectively. DCF fluorescence was also measured using a microplate reader (Ensight™ Multimode, Perkin Elmer) at 492 nm and 520 nm excitation and emission filter settings.
respectively. The stained cells were also visualized with a BX51WI fluorescence microscope (Olympus) with 460–490 nm excitation filter settings.

Results and discussion

Bacterial growth inhibition: Activity of the solid Ag particulates, the leached soluble Ag and silver salt

To distinguish the cytotoxicity or antimicrobial contribution of the leached soluble Ag and the undissolved Ag particulates, we exposed a model bacteria \textit{B. subtilis} UNSW 448700 to 0 – 10 mg Ag L\(^{-1}\) NPs (Ag/TiO\(_2\)) and compared the bacterial growth to that of the corresponding leachate-only systems, as a function of soluble silver detected in the exposure systems. The leachate samples were prepared by aseptically pre-dissolving Ag NPs in culture medium followed by removal of the solid Ag residue. Firstly, as shown in Figure 1a, the dose-response growth inhibiting effects of the Ag NPs correlates with the increasing concentration of soluble silver that leached from the NPs. The extent of growth of \textit{B. subtilis} was reduced to \(\sim 80\%\) upon exposure to 3 mg Ag L\(^{-1}\) NPs (1.3 ppm silver leached into the culture medium at equilibrium) relative to silver-free control cultures after 6 h. The control cultures were characterized by a relatively short lag phase of 1 h, followed by 3 to 4 h active exponential growth phase before entering the stationary phase at 6 h (Figure 1b). Increasing the NPs dosage to 6 mg Ag L\(^{-1}\) (2.7 ppm leached Ag) saw 50\% bacterial growth, while almost complete growth suppression was observed at MIC\(_{95}\) 10 mg Ag L\(^{-1}\) NPs exposure (4 ppm leached Ag, see Figure 1b for growth profile, MIC\(_{95}\) is minimum inhibitory concentration that cause 5\% growth relative to the control).

At all of the tested Ag NPs loading, leaching of Ag from NPs was rapid, with detection of \(\sim 70\%\)
soluble Ag (relative to the leached Ag concentration detected at equilibrium) within 5 min of the Ag NPs-bacterial exposure (see Figure 1c inset for leaching profile of 10 mg Ag L\(^{-1}\) NPs).

Equilibrium was reached in 1 h with the soluble Ag concentration remained constant afterwards, indicating absence of the Ostwald ripening phenomenon that refers to re-deposition of the
leached Ag on larger particulates (Sotiriou et al. 2012). Increasing the Ag NPs loading saw
detection of elevated soluble Ag concentration at equilibrium, with the extent of leaching
essentially comparable at 38 – 40% relative to the total added Ag (Figure 1c). This is consistent
to earlier studies under comparable conditions (Gunawan et al. 2009, Sotiriou and Pratsinis
2010) with the relatively high degree of leaching was due to, at least in part, the presence of
organics in the culture medium as shown later in this study. Note that at all of the tested Ag NPs
loadings, similar extent of leaching were observed in the absence of bacteria, therefore excluding
the possibility of microbial-induced leaching of Ag (Figure S1).

Despite the correlation between Ag NPs growth inhibiting effects and Ag leaching, a
comparison with bacterial growth in the corresponding leachate-only systems yields an
interesting observation. Exposure of *B. subtilis* to the pre-leached soluble Ag in fact resulted in
much less growth inhibition when compared to those of the corresponding Ag NPs samples
(Figure 1a). The presence of ~1.3 ppm Ag leachate for example, was benign to the cultures as
they grew to a similar extent as the silver-free control cultures after 6 h. This was in contrast to
the ~20% growth reduction of the bacteria when exposed to the corresponding 3 mg Ag L⁻¹ NPs
with comparable leached soluble Ag content. At higher exposure, the bacterial growth in 4 ppm
Ag leachate system was ~85% relative to the control cultures (refer to Figure 1b for growth
profile), in contrast to the near complete growth suppression observed in the corresponding 10
mg Ag L⁻¹ NPs system. Even doubling the concentration of Ag leachate to 8.3 ppm only slightly
reduced the bacterial growth to ~75%. The findings suggest predominant cytotoxicity
contribution from the undissolved Ag particulates, rather than that arising from the leached
soluble Ag. Further antimicrobial simulation with an equivalent concentration of soluble Ag
from AgNO₃ salt as shown in Figure 1a, saw more severe growth inhibiting activity of the salt. In
the presence of 4 ppm soluble Ag from AgNO₃ for example, ~25% *B. subtilis* growth was
observed relative to the control cultures after 6 h (growth profile is shown in Figure 1b), in contrast to the ~85% growth in the leachate system with comparable Ag concentration. Such differences in cytotoxicity may arise from unique cellular physiological responses to the different silver species; the leached soluble Ag and the undissolved Ag particulates from Ag NPs, and the soluble silver from silver salt, as investigated in the following.

Dynamic stimulation of cellular oxidative stress and cell death

We carried out dynamic tracking of intracellular reactive oxygen species (ROS) generation (measured by H$_2$DCFDA assay) and cell viability (measured by propidium iodide assay, whereby PI enters cells with damaged cytoplasmic membrane, which is indicative of cell death) over the 6 h growth course of $B$. subtilis in the presence of the various forms of silver; the Ag NPs (MIC$_{95}$ 10 mg Ag L$^{-1}$ as reference point, contained 4 ppm leached Ag), its corresponding Ag leachate system (4 ppm Ag) and the equivalent AgNO$_3$ system (4 ppm Ag).

The solid Ag particulates and the leached soluble Ag

At 5 min exposure to 10 mg Ag L$^{-1}$NPs, a 3-fold higher cellular ROS level was detected in $B$. subtilis relative to the basal ROS levels of the silver-free control cultures, which are by-products of aerobic metabolism in bacteria (Choi and Hu 2008, Gunawan et al. 2011, Eckhardt et al. 2013) (Figure 2a, 2b, 4a). Within 30 min of Ag NPs exposure, the cellular ROS level doubled to ~6-fold of the control. A secondary oxidative stress response, the cellular ROS stimulation has been increasingly realized as one of the major cellular toxicological responses to Ag NPs in bacteria (Choi and Hu 2008, Hwang et al. 2008, Lemire et al. 2013, Gunawan et al. 2013b). The ROS generation is thought to result from destruction of the iron-sulfur [4Fe-4S] clusters of proteins by Ag metal (Xu and Imlay 2012, Lemire et al. 2013) and in turn, releasing the Fenton-active free Fe into the cytoplasm for subsequent reaction with cellular H$_2$O$_2$ to produce hydroxyl...
radicals (OH•) (Imlay et al. 1988). Alternatively, indirect destruction of the iron-sulfur clusters could result from inhibition of respiratory enzymes by Ag NPs in bacteria (Li et al. 2010, 2011). The resulting premature leakage of electrons to oxygen will generate superoxide radicals (O2•−) (Imlay 2003) that in turn again, induces the release of free Fe from iron-sulfur clusters in proteins (Kohanski et al. 2007). Indeed, there have been reports on the cytoplasmic presence of the solid Ag particulates upon bacterial exposure to Ag NPs, as well as the presence of the solids within the bacterial membrane layers (Morones et al. 2005, Grigor'eva et al. 2013, Pal et al. 2007). Here, 75-90% PI-positive non-viable bacteria had been detected within 5 to 30 min exposure to Ag NPs, then close to 100% bactericidal or cell death toxicity at as early as 1 h exposure (Figure 2a, 2b, 4b), which indicates cytoplasmic membrane as one of the target destruction sites of the Ag NPs-stimulated cellular ROS (1-8% non-viable cells were detected in the control cultures over the 6 h growth course) (D'Autreaux et al. 2007, Lemire et al. 2013). As expected, the levels of cellular ROS drastically dropped following the rapid high level stimulation, with the majority if not all of the bacterial population were already killed (Sintubin et al. 2011, Gunawan et al. 2013b). Up to this stage, the data suggest that the generation of high levels of cellular ROS and associated bacteria killing was likely to be responsible for the near complete suppression of B. subtilis growth (Figure 1a, 1b).

Interestingly, such cellular ROS stimulation was absent in the bacteria when studied in the corresponding 4 ppm Ag leachate system. Over the 6 h growth course, only basal ROS levels, comparable to those of the silver-free control cultures were detected (Figure 2a, 3a, 4a) and not surprisingly, the little to no changes in the fraction of non-viable cells relative to the control (Figure 2a, 3a, 4b). The stimulation of lethal levels of cellular oxidative stress by the presence of solid Ag particulates therefore suggests their substantial contribution to the cytotoxicity effects observed in the growth studies. Recalling the observed ~15% growth inhibition of the bacteria in
the presence of 4 ppm Ag leachate (Figure 1a, 1b), it would be reasonable to deduce that the exposure only resulted in sub-lethal cytotoxicity, causing a minor fraction of the viable cells uncultivable or slowly proliferating, as further indicated by our growth prediction based on the fraction of viable cells (Figure S2). Indeed, doubling the Ag leachate concentration to 8.3 ppm still saw typical cellular ROS (Figure 3c, 4a inset) and dead cells (Figure 3c, 4b inset) detection as those of the control cultures, despite the slightly higher growth suppression, at ~25% (Figure 1a).

The leached soluble Ag and silver salt

The minimal cellular ROS stimulation was also seen upon exposure of *B. subtilis* to the equivalent 4 ppm soluble Ag from AgNO$_3$. Similar to the 4 ppm Ag leachate system, no elevated level of cellular ROS was observed over the 6 h growth course relative to the control cultures (Figure 2a, 3b, 4a). Unlike the leachate samples however, up to ~40% non-viable cells were detected in the salt system (Figure 3b, 4b), indicating attacks on cytoplasmic membrane (Eckhardt et al. 2013). Considering the comparable Ag content, such discrepancies in cytotoxicity are most likely to result from differences in the chemical speciation of the soluble silver, as herein described. Our Ag NPs leaching study (at the MIC$_{95}$ 10 mg Ag L$^{-1}$ NPs) in the individual culture medium components revealed a characteristic trend of complexation-assisted dissolution of nanoparticles (Gunawan et al. 2011), with higher extent of Ag leaching in the peptide-rich components, in particular tryptone (90% leaching relative to the total added Ag), compared to those in the deionized water (60% leaching) or NaCl (10% leaching) (Figure 4c). A soft Lewis acid, Ag(I) forms silver-peptide complexes upon its release from NPs (Bolea et al. 2014), which is most likely to result from its strong affinity to the NH$_x$ donor groups of histidine (NH$^+$), arginine (-NH$_2^-$) and lysine (-NH$_3^+$) amino acids and also to the thiol (-S$^-$) donor groups of cysteine and methionine amino acids (Eckhardt et al. 2013). Silver-peptide complexes also form with AgNO$_3$ (Bolea et al. 2014), with a fraction of silver is thought to remain as free ions in
the organic-rich medium (Percival et al. 2005). Thermodynamically feasible, the co-existence of
free metal ions and organo metal complexes has been reported for the chemical speciation of
soluble copper salts, also a soft Lewis acid metal, in similar culture medium as that used here
(Gunawan et al. 2011) (note that the current technology for elemental analysis does not
differentiate free Ag ions to those locked in organo complexes (Eckhardt et al. 2013)). When
compared to free Ag ions, the hindered transport of the bulkier silver-peptide complexes into
bacteria (Solioz and Odermatt 1995) is thought to be at least in part, responsible for the
passivated, in this case, sub-lethal cytotoxicity of the Ag leachate. Unlike free Ag ions, research
indicates that soluble organo Ag complexes are not recognized by the P-type ATPase transporter
present in bacteria (Luoma 2008). As also observed in the current study with the AgNO₃ systems,
exposure of bacteria to Ag ions has been reported to suppress their proliferation, which was
indicated to result from a ROS-independent inhibition of metabolic enzymes (dehydratases) (Xu
and Imlay 2012), the lack of cellular ROS stimulation also apparent in this work. Further,
complete suppression of B. subtilis growth was seen at 8.3 ppm Ag from AgNO₃ (Figure 1a),
despite there being no change in the fraction of non-viable cells when compared to the 4 ppm Ag
exposure (Figure 3d, 4b inset). Our growth prediction based on the fraction of viable cells
indicates major presence of non- or slowly proliferating viable cells with the AgNO₃ exposure
(Figure S2). This loss in replication could also result from the known interactions of Ag ions
with DNA in bacteria (most likely with the phosphorus moieties) causing DNA condensation
(Feng et al. 2000). The seemingly higher cytotoxic effects of Ag ions as compared to the organo
Ag complexes are in agreement with other bacterial studies, whereby extracellular presence of
thiol-containing reduced gluthathione (GSH) as silver complexing agent lowered the
antimicrobial activity of Ag ions on the Gram-positive Staphylococcus aureus and the Gram-
negative Escherichia coli and Pseudomonas aeruginosa (Mulley et al. 2014). Finally, the
detection of only basal cellular ROS levels in the AgNO₃ exposure systems, even at the double
8.3 ppm Ag (Figure 3d, 4a inset), rules out the oxidative stress stimulation as the main
mechanisms of AgNO$_3$ cytotoxicity. Indeed, studies have found no differences in the antimicrobial activity of Ag ions under aerobic and anaerobic conditions on bacteria (Sintubin et al. 2011).

**Conclusions**

Here, we report multiple cytotoxicity origins of Ag NPs towards bacteria. Presence of undissolved Ag particulates in a biological environment is not inert. In their presence, rapid generation of lethal cellular ROS levels were detected in bacteria, while the corresponding leached soluble Ag, being locked in organo complexes, only imparts sub-lethal cytotoxicity. The observed differences in bacterial toxicological responses to the solid versus soluble Ag corroborate earlier reports on the distinct extent of growth inhibiting activity of the Ag NPs’ soluble and solid components (Gunawan et al. 2009, Sotiriou and Pratsinis 2010). With regard to the widespread use of Ag NPs, the resolved unique toxicological responses are expected to result in better recognition of the antimicrobial potency of the nanoparticles in real-world settings and importantly, the long-term impact. Research inquiries have shown elevated and persistent presence of silver in wounds, bladder and even in sewage and estuaries, being associated with the intended or in some cases, accidental release from nanosilver applications; the use of wound dressings, pesticides and washing machines are among the examples (Chen et al. 2004, Trop et al. 2006, Reidy et al. 2013, Donner et al. 2015, Beddow et al. 2017). The current findings imply bacterial toxicological responses to not only the leached soluble Ag, but also the Ag particulates in the microbial habitats. Indeed, studies have observed disruptions in the dynamic and balance of microbial communities from natural aquatic waters upon exposure to nanosilver (Das et al. 2012, Beddow et al. 2017), with the work also detecting presence of soluble Ag and aggregates of Ag from nanosilver in these environmental samples (Beddow et al. 2017). The resolved toxicological responses is key to the elucidation of the recently discovered bacterial potential for adaptation to Ag NPs cytotoxicity (Das et al. 2012, Gunawan et al. 2013b). Finally, the work...
highlights the unsuitability of soluble silver salt as model material for Ag NPs cytotoxicity in biological environments, noting a distinct ROS-independent antimicrobial characteristic of soluble Ag when supplied as AgNO₃ salt.

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Declaration of interest

The authors declare no conflict of interest.

References


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Supplementary material is available: Supplementary Figure S1, S2 and S3.

Figure captions

Figure 1. Bacterial growth in the presence of Ag NPs, Ag NPs leachate, silver salt and leaching of Ag NPs. (a) Growth of B. subtilis (6 h) relative to cell-only control upon exposure to Ag NPs (3, 6, 8, 10 mg Ag L\(^{-1}\)), Ag leachate from NPs and AgNO\(_3\) as a function of soluble silver detected in the bacterial exposure systems (the growth studies were performed in LB medium). (b) Growth profiles of the bacteria in the presence of 10 mg Ag L\(^{-1}\) NPs (4 ppm Ag leached into medium at equilibrium), 4 ppm Ag leachate from NPs and 4 ppm Ag from AgNO\(_3\). Also shown is the cell-only control growth profile (dashed line). The growth in the presence of Ag was normalised to the extent of growth of the control (in colony forming units, cfu). (c) The corresponding equilibrium leaching of Ag NPs in the bacterial exposure systems, shown in the inset is the leaching profile for 10 mg Ag L\(^{-1}\) NPs. Each data point in (a), (b), (c) is the average of triplicate experiments with error bars representing the maximum and minimum values detected. The growth studies were performed under dark conditions to render the TiO\(_2\) support
photocatalytically inactive and the benign effect of the TiO$_2$ support on _B. subtilis_ growth had been confirmed (Gunawan et al. 2013b). The growth studies were reproduced on different days with unique bacterial inoculum and particle preparations.

Figure 2. Detection of cellular reactive oxygen species (ROS, H$_2$DCFDA staining, green cells) and cell death (PI staining, red cells) of _B. subtilis_ over its growth course: (a) cell-only control and (b) in the presence of 10 mg Ag L$^{-1}$ NPs. All stained samples were imaged at comparable cell concentrations (scale bars = 50 µm).

Figure 3. Detection of cellular reactive oxygen species (ROS, H$_2$DCFDA staining, green cells) and cell death (PI staining, red cells) of _B. subtilis_ over its growth course, in the presence of: (a) 4 ppm Ag leachate from NPs (equivalent leachate to 10 mg Ag L$^{-1}$ NPs exposure), (b) 4 ppm Ag from AgNO$_3$, (c) 8.3 ppm Ag leachate from NPs and (d) 8.3 ppm Ag from AgNO$_3$. All stained samples were imaged at comparable cell concentrations (scale bars = 50 µm).

Figure 4. (a) Dynamic stimulation of cellular ROS in _B. subtilis_ measured by H$_2$DCFDA assay over its growth course (5, 30 min and 1, 3, 4.5, 6 h) upon exposure to 10 mg Ag L$^{-1}$ NPs (4 ppm Ag leached into medium at equilibrium), 4 ppm Ag leachate from NPs and 4 ppm Ag from AgNO$_3$. The detected cellular ROS was normalised to the basal ROS levels of the cell-only control growth. Shown in the inset is cellular ROS detected in the presence of 8.3 ppm Ag leachate from NPs and 8.3 ppm Ag from AgNO$_3$. (b) The corresponding dynamic cell death detection probed by PI staining of _B. subtilis_ throughout its growth course. Also shown is the fraction of dead cells detected for the cell-only control. Statistical analysis of the data was performed with one-way ANOVA followed by Dunnett’s posthoc analysis in Prism (GraphPad) (Figure S3). The experiments in (a) and (b) were reproduced on different days with unique bacterial inoculum and particle preparations. (c) Leaching profile of 10 mg Ag L$^{-1}$ NPs in cell-
free individual components of the Luria Bertani (LB) culture medium (5 g L$^{-1}$ NaCl, 5 g L$^{-1}$ yeast extract, 10 g L$^{-1}$ tryptone dissolved in deionized water). Each data point in (a), (b), (c) is the average of triplicate batches with the error bars representing the maximum and minimum.