



Titre: Title:	Impacts of Continuous Inflow of Low Concentrations of Silver Nanoparticles on Biological Performance and Microbial Communities of Aerobic Heterotrophic Wastewater Biofilm
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Date:	2019
Туре:	Article de revue / Journal article
Référence: Citation:	Alizadeh, S., Abdul Rahim, A., Guo, B., Hawari, J., Ghoshal, S. & Comeau, Y. (2019). Impacts of Continuous Inflow of Low Concentrations of Silver Nanoparticles on Biological Performance and Microbial Communities of Aerobic Heterotrophic Wastewater Biofilm. <i>Environmental Science & Technology</i> , <i>53</i> (15), p. 9148-9159. doi: <u>10.1021/acs.est.9b01214</u>



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Document publié chez l'éditeur officiel

Document issued by the official publisher

Titre de la revue: Journal Title:	Environmental Science & Technology (vol. 53, no 15)
Maison d'édition: Publisher:	ACS Publications
URL officiel: Official URL:	https://doi.org/10.1021/acs.est.9b01214
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Impacts of continuous inflow of low concentrations of silver nanoparticles on biological performance and microbial communities of aerobic heterotrophic wastewater biofilm

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6 Abstract

In this study, two bench-scale moving bed biofilm bioreactors (MBBRs), achieving soluble 7 organic matter removal, were exposed to 10.9 and 109 µg/L polyvinylpyrrolidone (PVP)-coated 8 AgNPs for 9 weeks (64 d). Distribution of continuously added AgNPs were characterized in 9 influent, bioreactor and effluent of MBBRs using single-particle inductively coupled plasma 10 mass spectroscopy (spICP-MS). Continuous exposure to both AgNPinf concentrations 11 significantly decreased soluble chemical oxygen demand (S_{COD}) removal efficiency (11% to 12 31%) and reduced biofilm viability (8% to 30%). Specific activities of both intracellular 13 dehydrogenase (DHA) and extracellular α -glucosidase (α -Glu) and protease (PRO) enzymes 14 were significantly inhibited (8% to 39%) with an observed NP dose-dependent intracellular 15 reactive oxygen species (ROS) production and shift in biofilm microbial community composition 16 by day 64. The release of significant mass of Ag via effluent (<78%), dominantly in NP form 17 due to the limited retention capacity of aerobic heterotrophic biofilm, provide new and useful 18 insight into fate of AgNPs in biofilm-laden engineered biological systems and their 19 20 corresponding inhibitory effects at environmentally representative NP concentrations maintained 21 over an extended period. To our knowledge, this is the first study evaluating chronic inhibitory effect of AgNPs on attached-growth wastewater process efficiency and its microbial 22 23 communities at representative environmental AgNP concentrations by combining biological response and NP characterization. 24

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28 **1. Introduction**

Engineered nanoparticles (ENPs) are manufactured at an estimated rate of 11.5 million tons per 29 year for various industrial and commercial applications¹. Silver nanoparticles (AgNPs) are 30 predominantly used as antimicrobial agents in commercial products, cosmetics, food processing 31 and water industries¹. This rapidly developing nanotechnology market, however, is leading to 32 their environmental exposure, with a significant fraction of the AgNP-laden domestic and 33 industrial waste streams being released in municipal water resource recovery facilities (WRRFs) 34 at an estimated influent concentration ranging from 10 ng/L to 1.5 μ g/L²⁻⁴. Thus, WRRFs serve as 35 a key interface between ENPs releases and their environmental distribution into downstream 36 ecosystems. Previous studies on the inhibitory effects of AgNPs (0.1 to 50 mg/L) in suspended-37 growth systems showed adverse effects on the biological performance and biomass activity 38 caused by oxidative stress, cell membrane damage and inactivation of key enzymes at sufficient 39 AgNP doses (< 1 mg/L)⁵⁻⁹. Yet, the interaction between biofilm processes and ENPs including 40 AgNPs are poorly understood. 41

Attached growth processes, such as moving bed biofilm reactors (MBBRs), are commonly used 42 as an upgrade or replacement for existing biological processes to meet current and new effluent 43 discharge requirements, while minimizing plant footprint and operating costs^{10,11}. In 2014, more than 44 1200 WRRFs in at least 50 countries utilize the MBBR technology in both the municipal and 45 industrial sectors with over 36 in North America^{12,13}. A limited number of studies investigated 46 the impact of a single dose of AgNPs (1 to 200 mg/L) over 24 to 96 h, using mono-species 47 biofilms, particularly P. putida based biofilms or wastewater biofilms in simplified biological 48 media^{11,14-16}. Their findings indicated higher potential of biofilm bacteria than planktonic bacteria 49 50 to withstand the toxic effects of AgNPs, primarily due to the presence of extracellular polymeric substances (EPS), the primary components of biofilm¹⁵, which act to reduce AgNP diffusion in 51

biofilms¹⁷ over short term exposure conditions. Yet, the results of short-term exposure studies 52 may fail to capture the effects of the expected accumulation of AgNPs and higher mass transport 53 of AgNPs by diffusion into deeper layers of the biofilm over extended time intervals, thus 54 underestimating the potential toxicity of AgNPs over long-term exposure scenarios^{18,19}. Further 55 research is thus required first, to understand the interaction mechanisms between ENPs and 56 mature, mixed culture wastewater biofilms and second, to investigate the corresponding AgNP-57 induced inhibitory effects at environmentally representative NP concentrations under conditions 58 that are representative of typical WRRF processes. 59

Rigorous physical and chemical characterization of AgNPs combined with extensive biological and 60 toxicological evaluations in WRRFs are critical in laying the grounds for a better understanding 61 of their environmental fate and for the design of better alternative treatment strategies and future 62 regulations²⁰. The current understanding of the environmental fate and transformation of ENPs is 63 limited due to the limitations of ENP characterization techniques in complex environmental 64 matrices containing ENPs at very low, environmentally relevant concentrations^{21,22}. Single-65 particle inductively coupled plasma-mass spectrometry (spICP-MS) is an emerging powerful 66 technique with the potential to address such limitations, providing quantitative characterization 67 of metal NP size distributions, particle number concentrations and dissolved metal 68 69 concentrations at low NPs concentrations in complex, organic matter-rich, environmental matrices such as wastewaters²³⁻²⁵. 70

In this study, we investigated the impact of continuous injection of low concentrations of AgNPs in an attached growth wastewater treatment process using aerobic heterotrophic wastewater biofilms at nominal influent concentrations of 10.9 and 109 μ g/L AgNPs to approximate environmentally relevant concentrations of AgNPs. Although, these concentrations would still be

at the higher end of the estimated concentration for WRRFs, they would mimic a worst case 75 scenario such as release from biosolid-treated soil or landfills by flooding events or production 76 plant discharge²⁶. The specific objectives of this study were (1) to characterize the interactions 77 and distribution of AgNPs in a biofilm-laden wastewater biological process and (2) to determine 78 the impact of AgNPs on primary biological functions and microbial community of wastewater 79 80 biofilms. Two bench-scale MBBRs were operated for organic matter removal and were fed with a synthetic soluble influent representative of a municipal wastewater. The impacts of AgNPs on 81 the performance of the MBBRs were characterized by monitoring several performance indicators 82 including S_{COD} removal efficiency, effluent quality and enzymatic activity over a 9-week (64 d) 83 exposure period. 84

The biological responses of aerobic heterotrophic biofilm were characterized in terms of 85 (i) biofilm cell membrane integrity using DNA-binding stains, (ii) AgNP-mediated oxidative 86 stress via intracellular ROS measurement and (iii) microbial metabolic functions by intracellular 87 DHA and extracellular α -Glu and PRO specific enzymatic activities using colorimetric assays. 88 The biofilm microbial community compositions, at both influent AgNPs concentrations, were 89 characterized through high-throughput sequencing. The aggregation state, dissolution and 90 distribution of AgNPs were determined between different reactor components (i.e. influent, 91 bioreactor and effluent) using spICP-MS techniques and transmission electron microscopy with 92 energy dispersive X-ray spectroscopy (TEM EDS). To the best of our knowledge, this is the first 93 94 study evaluating the long-term inhibitory effect of AgNPs on attached-growth wastewater process efficiency and its microbial communities at environmentally relevant AgNP 95 concentrations (< 100 μ g/L AgNPs) by combining biological responses and the NP distribution, 96 97 characterization and Ag speciation data.

98 2. Methods

99 2.1 Reactor configuration and AgNPs exposure

Two 1 L bench-scale MBBRs, achieving organic matter removal at a hydraulic retention time 100 (HRT) of 3 hours, operated in parallel under identical conditions, were fed a synthetic soluble 101 influent (Table S1) to ensure constant influent characteristics and well-controlled conditions to 102 characterize the inhibitory effects of the PVP-AgNPs. The concentrated synthetic wastewater 103 $(1.3 \pm 0.2 \text{ g S}_{\text{COD}}/\text{L})$ was pumped and diluted with tap water before entering the reactors to 104 obtain an influent COD concentration of $655 \pm 6 \text{ mg S}_{COD}/L$ at an organic loading rate of $11 \pm$ 105 0.2 g COD m⁻² d⁻¹ of active surface area to be representative of the soluble fraction of a medium 106 to high strength domestic wastewater with typical COD/TKN/TP ratio of 100/12.0/2.0²⁷. The 107 characteristics of the synthetic influent (Table S2) and detailed reactor operation conditions are 108 109 presented as supplementary information (SI). After reaching quasi steady-state conditions with a stable S_{COD} removal efficiency, the reactors were monitored for 85 days as a control period. 110 Influent AgNP suspensions were prepared by dilution of 50 nm PVP-coated AgNPs stock 111 suspension (4.73 mg/mL, Nanocomposix Inc., San Diego, US) in Milli-Q water. The zeta 112 potential and surface area of AgNPs were -37.8 mV (at pH 4) and 9.8 m²/g, respectively, based 113 on the AgNP product description, with a mean diameter of 48 ± 2 nm (SpICP-MS, PerkinElmer 114 NexION 300X). The AgNP influent suspensions were pumped to each reactor from day 125 at a 115 constant flow rate $(2.7 \pm 0.1 \text{ mL/min})$, resulting in an average influent total Ag concentration of 116 $14 \pm 0.5 \ \mu\text{g/L}$ Ag for MBBR₁ and $130 \pm 14 \ \mu\text{g/L}$ Ag for MBBR₂ after dilution. The influent 117 nanoparticle suspensions were replenished every 24 h. The effluent water quality, biofilm 118 biological responses (e.g. viability or enzyme activity) and Ag distribution were monitored over 119 64 days. Chemical oxygen demand (COD), total suspended solids (TSS) and volatile suspended 120 solids (VSS) were measured according to Standard Methods²⁸. 121

122 **2.2 Silver analyses**

The influent, bioreactor and effluent were sampled every 24 h over the first week (day 125-130) 123 and every 3 days afterwards (day 133-189) for a total period of 9 weeks (Figure S1). Bioreactor 124 and effluent samples contained suspended flocs (145 to 480 mg TSS/L) but no K5 carriers. Total 125 Ag concentration was measured in acid-digested homogenized samples using a PerkinElmer 126 NexION 300x ICP-MS in standard mode as described in our previous study¹⁹. The homogenized 127 samples were allowed to settle for about 30 to 45 s and the aqueous supernatant was collected. 128 AgNP concentration and size as well as dissolved Ag were determined simultaneously using 129 spICP-MS, supported by Syngistix nano application module (version 1.1) as described by Azodi 130 et al.²³. Instrumental and data acquisition parameters of the analysis are indicated in SI (Table 131 S3). A cumulative Ag mass distribution in influent ($M_{Ag, inf}$), bioreactor ($M_{Ag,bio}$) and effluent (132 MAgeff) of each MBBR was calculated based on the corresponding total Ag concentrations, 133 obtained from ICP-MS analysis, influent and effluent flow rates, and volume of the bioreactor 134 for each time interval (Δt) as described in our previous study¹⁹. Ag fractionation and the detailed 135 equations and Ag mass balance are presented in SI. 136

137 **2.3 Viability and key enzymatic activities of attached biofilm**

Bacterial viability of attached biofilms was evaluated using the Live/Dead *Baclight* bacterial viability kit (Molecular Probes, Invitrogen, Kit L13152) and a micro plate reader (Synergy-HT, BioTek, USA) as described by Chen et al.²⁹. The specific activities of DHA, α -Glu and PRO enzymes were measured by a colorimetric method³⁰,³¹ using 0.5% 2-(4-iodophenyl)-3-(4nitrophenyl)-5-phenyl-2H-tetrazolium chloride (INT), 1% *p*-nitrophenyl α -D-glucopyranoside and 0.5% azocasein, respectively, as a substrate for the reactions. The intracellular ROS production, as an indicator of oxidative stress, was determined using dichlorodihydrofluorescein diacetate (H₂DCF-DA, Molecular Probes, Invitrogen)³². H₂-DCFDA was used as a cell-permeant
 reagent that measures hydroxyl, peroxyl and other reactive oxygen species activity in cells.
 Details regarding all enzyme activity and ROS assays are provided in SI.

148 **2.4 DNA Extraction, sequencing and microbial community analysis**

Biofilm samples were collected from MBBR₁ and MBBR₂ at the end of the control period 149 (MBBR₁^C, MBBR₂^C) and after exposure to AgNPs for 64 days (MBBR₁⁶⁴, MBBR₂⁶⁴). For each 150 set of the microbial community data, 10 carriers were selected randomly from each reactor; the 151 retained biofilm was homogenized and used for DNA extraction. Genomic DNA was extracted 152 from the biofilm samples using FastDNA[®]spin kit (MP Biomedicals, Santa Ana, CA) following 153 the manufacturer's instructions, and sent for library preparation and sequencing on the Illumina 154 Miseg PE250 platform at McGill University and Génome Québec Innovation Centre (Montréal, 155 Québec). Bacterial universal primers 515F (5'-GTGCCAGCMGCCGCGGTAA-3') and 806R 156 (5'-GGACTACHVGGGTWTCTAAT-3') were used to amplify the V4 variable region of the 16S 157 rDNA. Bioinformatics analysis was performed using QIIME2 pipelines. The de-multiplexed 158 forward and reverse sequences were quality-filtered using DADA2³³ at 100% sequence 159 similarity. Taxonomy was assigned using the 99% operational taxonomic unit (OTU) similarity 160 in the GreenGenes reference database. Alpha-diversity, beta-diversity and their statistical tests 161 were analyzed in QIIME2. Principal coordinates analysis (PCoA) was constructed using 162 weighted UniFrac distance matrix. Heatmap was generated in R using the "gplots" package. 163

164 **2.5 Statistical analysis**

The statistical significance of differences between treatments (p < 0.05), before and after exposure to AgNPs, was evaluated with one-way repeated measures ANOVA using Statistica version 12 (StatSoft Inc., USA).

168 **3. Results and discussion**

169 **3.1 Fate of AgNPs in MBBRs over 64 days**

The influent of MBBR₁ and MBBR₂ contained an average concentration of $10.9 \pm 1.6 \ \mu g/L$ 170 AgNP ([AgNP_{inf}]) and 109.3 \pm 10 μ g/L AgNP_{inf}, with mean diameter (d_{mean}) of 49 \pm 7 nm and 171 48 ± 2 nm, respectively, (Figure S2D), corresponding to a total Ag concentration ([Ag_{inf}]) of 14 172 $\pm 2 \mu g/L Ag 130 \pm 14 \mu g/L Ag$ in corresponding reactors (Figure 1A, B). Each of these quantities 173 was independently measured by spICP-MS and ICP-MS. SpICP-MS analyses showed less than 174 10% variation in dissolved Ag concentrations([dissolved Aginf]), in influent NP stock solutions 175 of both reactors over time (Figure S2A, B). Three distinct trends (Phases) were observed for Ag 176 concentration profiles in both reactors over the 64-day Ag loading (Figure 1A, B). Phase I 177 178 corresponded to the first 15 days of AgNP loading (day 125-140) during which both reactors accumulated Ag in response to the AgNP addition and reached a relatively stable Ag retention 179 efficiency (61% to 72% of $[Ag_{inf}]$). A major fraction of the released total silver ($[Ag_{eff}]$) (40% to 180 63%) was associated with total suspended solids in effluent (TSS_{eff}) in MBBRs. Effluent Ag 181 concentration as NP (AgNP_{eff}) in MBBR₁ (0.2 to 1.4 μ g/L) and MBBR₂ (2.9 to 5.5 μ g/L) 182 corresponded to approximately $10\% \pm 3\%$ of [Ag_{eff}] and $16\% \pm 4\%$ of [Ag_{eff}], respectively, over 183 Phase I. SpICP-MS analyses showed variations in [dissolved Ageff] in the effluent supernatant 184 of MBBR₁ (0.2 to 2.6 μ g/L) and MBBR₂ (4.7 to 11.9 μ g/L) representing about 16% to 49% of 185 [Ageff] as Ag⁺ partially or completely complexed to dissolved organic carbon (DOC) (Figure 1A, 186 B). High concentration of dissolved oxygen $(6 \pm 0.2 \text{ mg/L})$ and pH of 7.2-7.5 in the MBBRs 187 provided thermodynamically favorable conditions for oxidation driven dissolution of AgNPs. 188 Generally, the mean diameter of AgNPeff was up to 7 nm smaller than AgNPinf, despite the 189 minor aggregation over 3 initial days of exposure (Figure 1C). 190

The magnitude of change in the diameters was small because of the short average residence time 191 of the AgNPs in the reactor. The particle size distribution of AgNPeff had slightly higher 192 concentrations of smaller particles compared to AgNPinf (Figure S3). The relatively high amount 193 of [dissolved Ageff] on certain days (e.g., day 133 in MBBR1 and day 154 in MBBR2) cannot be 194 accounted for the changes in particle diameters in the effluent. It is likely that detachment and 195 release of soluble complexes Ag from the biofilm/EPS matrix resulted in relatively high fraction 196 of [dissolved Ageff] relative to [Ageff]. The attached biofilm (Agcarrier) retained about 60% to 71% 197 of cumulative mass of total Ag loading in the influent $(M_{Ag_{inf}})$ in MBBR₁ (0.88 mg Ag/m² of 198 carrier active surface) and MBBR₂ (10.3 mg Ag/m² of carrier active surface) by the end of 199 Phase 1 (day 140), indicating an initial high Ag biofilm retention capacity (Figure 1D, E). The 200 carrier active surface area represents the biofilm covered surface area. 201

Phase II started with a gradual increase of $[Ag_{eff}]$ which reached a maximum concentration of 8.9 202 \pm 1.7 µg/L Ag in MBBR₁ and 118.6 \pm 1.5 µg/L Ag in MBBR₂ over 10 days (day 140 - 150) 203 decreasing the Ag retention efficiency significantly (p < 0.05) compared to Phase I due to 204 biofilm detachment (Figure S5). A larger fraction of $[Ag_{eff}]$ (20% to 37%) was detected as NPs 205 in the effluent supernatant of MBBR₁ (8.9 to 11.2 μ g/L) and MBBR₂ (9.02 to 26.5 μ g/L). The 206 [dissolved Ag_{eff}] in MBBR₁ and MBBR₂ represented about $22\% \pm 3\%$ of [Ag_{eff}] (1.3 to 3.04 207 μ g/L) and 25% ± 8% of [Ag_{eff}] (11.9 to 25.30 μ g/L) respectively, in this phase (Figure 1A, B). 208 209 By the end of Phase II (day 150), the attached biofilm retained only about 44% to 54% of cumulative M_{Agint} in MBBRs (Figure 1D, E), followed by a continuous decrease in retention 210 efficiency (Figure 1A, B). The virgin surface of the biofilm retained much AgNPs during Phase 211 I, likely due to interactions of hydrophobic PVP coatings of AgNPs with heterogeneous 212

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amphiphilic moieties of the biofilm surface³⁴. As the concentration of AgNPs increased inside the reactors during Phase II, the local accumulation of PVP-AgNPs on the outer layer of the biofilm must have blocked further deposition of AgNPs and caused decrease in NPs attachment efficiency onto the biofilm surface³⁴.

Phase III corresponded to the period with Ag release and retention recovery events in both 217 reactors (day 154 - 189). The Ag distribution profile in MBBR₁ consisted of two Ag release 218 events on day 154 (week 4) and day 164 (week 6) with the [AgNP_{eff}] (9.9 ± 1.7 μ g/L to 14.03 ± 219 0.7 μ g/L) constituting 61% to 87% of detected [Ag_{eff}] (Figure 1A, C). Despite a slight Ag 220 recovery in retention by biofilm by day 175 (week 7), significantly higher [Ageff] were released 221 over the last two weeks of exposure, predominantly in the form of AgNPs (81% to 96% of [Ageff 222) likely due to saturation of biofilm outer layers by AgNPs and/or biofilm sloughing off from 223 the surface of the carriers. MBBR₂ demonstrated a lower recovery for Ag retention over a longer 224 time interval as compared to MBBR₁ during the Phase III. A maximum Ag release of 229 μ g/L 225 Ageff was observed at the beginning of Phase III (day 154) in MBBR₂, with a dominant fraction 226 of [Ageff] detected as AgNPeff (52% of [Ageff]) and dissolved Ageff (45% of [Ageff]) in the 227 aqueous phase of the effluent (Figure 1B), suggesting a saturation of the biofilm outer layers. 228 Thereafter, [AgNP_{eff}] (80 to 104 μ g/L) represented an average 40% to 65% of [Ag_{eff}] between 229 day 161 and day 189 (Figure 1B). A relatively smaller mass fraction of $[\mathrm{Ag}_{\mathrm{eff}}]$ was accounted for 230 dissolved Ag_{eff} (11% to 15%) in both reactors in Phase III compared to Phases I and II 231 (Figure 1C). Attached biofilm retained less than 20% of cumulative MAginf in MBBR1 (1.52 mg 232 Ag/m² of carrier active surface) and MBBR₂ (15.2 mg Ag/m² of carrier active surface), 233 respectively, and significant fraction of cumulative $M_{Ag_{inf}} (> 78\%)$ was released via the effluent 234 of the both reactors by the end of Phase III (day 189; Figure 1D, E), indicating poor retention 235

capacity of the biofilm over long term exposure. Similar high bioaccumulation of AgNPs¹¹ and silica-coated iron oxide³⁵ in wastewater biofilms were reported at low NP concentrations over shortterm exposure whereas detachment of NP-rich biofilms became a source of NP-release as NP concentrations increased over time.

A general conclusion from previous studies conducted in batch experiments³⁶, sequencing batch 240 reactors⁷, membrane bioreactors³⁷ and municipal WRRFs⁴ indicated an efficient AgNP removal 241 (72% to 95%) via accumulation in suspended growth activated sludge processes with no 242 extensive AgNP washout as shown here for the MBBRs. The quantitative characterization of 243 nanoparticles in MBBRs, using spICP-MS, indicated an initial adaptation of MBBR to silver 244 addition with an increase in total Ag release over time, predominantly in NP form, and a periodic 245 silver accumulation in biofilm coupled with a biomass concentration increase (Figure S4). Our 246 results imply that there is a limited retention capacity of aerobic heterotrophic biofilm for AgNPs 247 over a long time exposure, compared to the commonly studied activated sludge systems. The 248 observed decrease in [dissolved Ageff] over time in both MBBRs can be attributed to the 249 removal of Ag⁺ via their interaction with functional groups of macromolecules such as cysteine 250 and methionine in the EPS and biofilm matrix and their organic ligands, such as thiols^{38,39,40}. 251 Complete inhibition or significant decrease of AgNPs dissolution was reported in the presence of 252 Cl⁻ ions at low Cl/Ag ratios³⁹. We detected similar changes in the chemical composition of the 253 particles in the effluent of the MBBRs using TEM-EDS analysis (Figure S2). Similar 254 complexation of dissolved silver in wastewater effluents and their significantly reduced 255 bioavailability were reported for a 7 day-experiment⁴⁰. Azodi et al.²³ attributed the decrease in 256 dissolved Ag concentrations in the wastewater effluent to the reformation of the secondary NPs 257 from dissolved Ag. 258

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Figure 1. Fate and retention of Ag in MBBR receiving influent concentration of (A) 10 μ g/L AgNPs and (B) 100 μ g/L AgNPs, (C) dissolution of AgNP_{eff} (%) and difference in mead diameter (d_{mean}) of AgNP_{inf} and AgNP_{eff}, (D) cumulative Ag mass distribution in influent (Inf), attached biofilm (Biofilm) and effluent (Eff) and (E) enlarged Y-scale of panel D.

3.2 Effects of AgNPs on the biological performance of a heterotrophic aerobic biofilm

The biofilm-mediated S_{COD} removal efficiency was determined in the two MBBRs in response to 267 the 64-day continuous exposure to [AgNP_{inf}] of 10.9 and 109 μ g/L (Figure 2A, B). Prior to 268 exposure to the AgNPs, both reactors consisted of $98\% \pm 0.2\%$ viable biofilm over the control 269 period (day 90-124) (Figure 3A) which stabilized the S_{COD} removal efficiency at 93% under 270 quasi steady state conditions (Figure 2A-C). The biodegradation of S_{COD} remained stable, after 271 injection of AgNPs, with an average S_{COD} removal efficiency of 92% \pm 0.7% over the first 35 272 days (day 125-161) in MBBR₁ and $91\% \pm 2\%$ over 23 days (day 125-147) in MBBR₂ indicating 273 an unperturbed primary phase (Figure 2C) with relatively stable biofilm viability (>96%) 274 (Figure 3A). Measured AgNP concentrations in MBBR₁ (0.16 to 0.60 µg/L AgNP_{bio}) and 275 MBBR₂ (0.7 to 5.02 μ g/L AgNP_{bio}) and their corresponding dissolved Ag_{bio} of 0.3 to 1.2 μ g/L 276 and 1.5 to 8.4 μ g/L, respectively, over Phase I (Figure S2) were much lower than previously 277 reported threshold concentrations for toxicity of AgNPs and dissolved Ag for biofilms (IC_{50 PVP-} 278 $_{AgNP@48h} = 114 \ \mu g/L$ and $IC_{50,Ag+@48h} = 44 \ \mu g/L^{41}$). As AgNP concentrations increased in reactors 279 (Figure S2C, D), a secondary phase was observed with higher numbers of inactivated cells, 280 resulting in significant inhibition of viable attached biofilm in MBBR₁ (8%) and MBBR₂ (31%) 281 by day 189 (Figure 3A). The S_{COD} removal efficiency significantly decreased (p < 0.05) by about 282 11% over 29 days (day 160-189) in MBBR₁, and by 31% after 41 days (day 133-189) in MBBR₂ 283 (Figure 2C), corresponding to the observed patterns in Ag_{bio} and Ag_{eff} time profiles. Exposure to 284 both AgNP_{inf} concentrations also induced biofilm detachment from the surface of the carriers, 285 corresponding to the significant increase of TSSeff (Figure S5). Significant detachment of 286 wastewater biofilm and concurrent release of accumulated AgNPs were similarly reported at 287 environmentally relevant AgNPs concentrations (22 and 105 μ g/L AgNPs)¹¹. 288

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Figure 2. Effect of PVP-AgNPs addition on MBBR performance at (A) 10 μ g/L AgNPs (B) 100 μ g/L AgNPs, (C) S_{COD} removal efficiency and (D) correlation between Ag_{carrier} and S_{COD} removal efficiency (error bars are only shown when larger than symbol size). Note: P.I-III refers to three observed phases in Ag distribution profile.

The inhibitory effect of AgNPs on both the S_{COD} removal efficiency (Figure 2D) and the biofilm viability inhibition (Figure 3B) were highly correlated (0.91 < R^2 < 0.97) to the retained mass of Ag in the carriers (Ag_{carrier}). High biomass surface area/volume ratio in attached growth processes (e.g. MBBR) enhances the deposition rate of AgNPs to attached biomass over time, leading to enhanced Ag retention per unit weight of biomass in MBBR. Thus, significant accumulation and associated mass transport of AgNPs into deeper layers of the biofilm can lead to extensive spatial distribution of AgNPs in the biofilm cells, delivering toxic Ag^+ directly to adherent cells via interfacial dissolution of the surface-bound AgNPs and/or partly via direct uptake, leading to an enhanced time exposure and greater toxicity⁴².



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Figure 3. Effect of continuous PVP-AgNP injection on (A) attached cell viability, (B) intracellular ROS generation, (C) correlation between Ag_{carrier} and attached biofilm viability inhibition and (D) correlation between viability inhibition and intracellular ROS generation (error bars are only shown when larger than symbol size). Note: P.I-III refers to three observed phases in Ag distribution profile.

Intracellular ROS did not significantly change in MBBR₁ (10.9 μ g/L AgNP_{inf}), whereas its 322 concentration increased significantly (1.78-fold, p < 0.05) in MBBR₂ (109 μ g/L AgNP_{inf}) over 64 323 days (Figure 2C), consistent with reported concentration-dependent ROS production in activated 324 sludge³². No correlation between biofilm viability inhibition and ROS production was observed 325 in MBBR₁ whereas the cell membrane integrity damage was highly correlated to increased ROS 326 generation in MBBR₂ ($R^2 = 0.97$) (Figure 3D), indicating both ROS-mediated and ROS-327 independent effects of AgNPs on cell membrane integrity³². The interaction between AgNPs/Ag⁺ 328 and the functional groups of proteins, involved in the cell respiratory chain, can lead to 329 intracellular ROS production. Biofilm was able to regulate the ROS production at lower AgNP 330 concentrations (1.4 to 9.04 μ g/L AgNP_{bio}), likely via ROS scavenging enzymes (e.g. superoxide 331 dismutase)⁴³, higher concentrations of AgNPs in MBBR₂ (10.3 to 46.2 μ g/L AgNP_{bio}), however, 332 caused significant overproduction of ROS which can overwhelm the antioxidant systems and 333 induce oxidative damage to cell membranes by for example modification of the unsaturated fatty 334 acids of the membrane phospholipids⁴⁴. 335

336 **3.3 Inhibitory effect of AgNPs on key enzymatic activities of aerobic heterotrophic biofilm**

Average DHA specific activity was inhibited by about 11% and 27% in MBBR₁ and MBBR₂, 337 respectively, after 64 days (Figure 4A₁). The specific activity of α -Glu and PRO were reduced by 338 $16\% \pm 2\%$ and $8\% \pm 1\%$, respectively, in MBBR₁. Higher enzyme activity inhibitions, up to 39% 339 $\pm 2\%$ (α -Glu) and $18\% \pm 2\%$ (PRO), were observed at higher [AgNP_{inf}] in MBBR₂ (Figure 4B₁, 340 C₁), indicating a dose-dependent effect of AgNPs on specific enzymatic activities of 341 biofilms^{45,486}. Significantly different inhibition rates and half-lives of all three enzymes upon 342 exposure to AgNPs (Table S4) indicated the distinct sensitivity of these enzymes to AgNPs due 343 to their different properties and location patterns in the biofilm matrix⁴⁷. 344

The inhibitory effects of AgNPs on specific enzymatic activities of biofilm was highly correlated ($0.80 < R^2 < 0.96$) to the retained mass of Ag in the carriers (Ag_{carrier}) (Figure 4A₂-C₂). The observed pattern highlights the major role of diffusion of retained AgNPs in the biofilm-laden system upon the targeted delivery of both AgNPs and dissolved Ag in close proximity of enzymes and possibly inside the cell in order to reach the Ag concentration needed to exceed inhibitory limits.

Upon the initial contact with enzymes in the biological media, nanoparticles acquire a protein 351 corona leading to substantial structural changes of the enzyme⁴⁸. Extracellular enzymes (e.g. α -352 Glu) interact initially via their functional groups (e.g. carboxyl, hydroxyl, amine, amido, keto) 353 with both the ring and polyvinyl domain of PVP coating and the oxygen atom involved in PVP-354 nanoparticle complex form. The strong bonding of AgNPs and dissolved Ag with electron 355 donors containing sulfur, oxygen, or nitrogen (e.g. thiols, carboxylates, phosphates, hydroxyl, 356 amines, imidazoles, indoles) across the enzymes can form silver complexes which shield the 357 active sites and alter the enzyme's conformation or distort its 3D structure so it no longer retains 358 its full enzymatic activity⁴⁹. Breaking through the barrier of outer membrane permeability, 359 AgNPs and dissolved Ag can strongly associate with specific sequences of amino acids on the 360 DHA active site and thiol (-SH) group of cysteine, by replacing the hydrogen atom to form -S-361 Ag, and irreversibly inactivate dehydrogenase enzymatic functions leading to cellular respiration 362 inhibition^{49,50}. Moreover, the ROS-mediated protein oxidation and microbial community 363 composition alteration can result in loss of function for enzymes associated with biofilms and in 364 a cutback in their production and secretion⁴⁷. 365

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Figure 4. Effect of continuous PVP-AgNP injection on specific activity of (A_1) DHA $(B_1) \alpha$ -Glu and (C_1) PRO, (A_2-C_2) correlation between Ag_{carrier} and enzyme activity inhibition (Error bars are only shown when larger than symbol size).

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374 Seven major phyla (Proteobacteria, Bacteroidetes, Verrucomicrobia, Gemmatimonadetes

375 *Planctomycetes, Actinobacteria*) were identified (abundance > 1%) at the end of the control

period in both reactors (MBBR₁^C, MBBR₂^C) (Figure 5A), as previously reported in microbial 376 composition wastewater biofilm studies^{51,52}. Proteobacteria was the most abundant phylum 377 majorly comprised of Alphaproteobacteria, Betaproteobacteria and Gammaproteobacteria. 378 Certain phyla demonstrated a distinct behavior at two [AgNP_{inf}] after 64 days of exposure (day 379 189) The relative abundance of *Proteobacteria* decreased at lower doses of AgNPs (MBBR₁⁶⁴) 380 but increased at the higher dose of AgNPs (MBBR₂⁶⁴) in contrast to Verrucomicrobia (Figure 381 5A). AgNPs influenced biofilm microbial phylum abundance in a dose-dependent manner, which 382 was confirmed by the principal coordinate analysis (PCoA) based on the weighted UniFrac 383 distance matrix with 75% of total variance on PCoA1 axis (Figure S6). The observed pattern 384 indicated various responses among taxa, including a range from susceptibility towards silver 385 (e.g. Bacteroidetes and Gemmatimonadetes) to tolerance against silver (e.g. Planctomycetes), as 386 reported in previous studies^{53,54}. 387

The heatmap of genera, with total sequence reads higher than 150 in selected phyla, showed the 388 distinct sensitivity of certain genera at both [AgNP_{inf}] (Figure 5B). The relative abundance of 389 Rhodobacter, identified as the most abundant OTU at the genus level (Rhodobacteraceae, a-390 proteobacteria) decreased in both reactors. Higher abundance of Paracoccus other dominant 391 genera from the *Rhodobacteraceae* family and *Zooglea* (β -proteobacteria) at higher [AgNP_{inf}], is 392 likely associated with their heavy metal resistance to enhance their survival in metal-393 contaminated environments⁵⁵. The abundance of *Sphingomonas* genus (*Sphingomonadaceae*) 394 decreased in MBBR164 whereas its abundance increased in MBBR264, possibly due to the 395 presence of signaling molecules (e.g. sphingolipids) in their outer membrane maintaining 396 community diversity at higher silver concentrations⁵⁶. The genera affiliated with 397 398 *Xanthomonadaceae* family such as *Stenotrophomona* (*y-proteobacteria*) are reported as N-acylhomoserine-lactone (AHL) producers in aerobic granular sludge and biofilm, contributing to
 AHL-mediated quorum sensing signaling, integrity and biofilm stability⁵⁷. Thus, the reduction in
 their relative abundance at both [AgNP_{inf}] affected both COD removal efficiency and integrity of
 biofilm structures.

Chronic exposure to both [AgNP_{inf}] greatly decreased the relative abundance of genera affiliated 403 to three dominant orders in the Bacteroidetes phylum (Sphingobacteriales, Flavobacteriales, 404 Cytophagales) known as the core members of microbial communities in WRRFs degrading 405 complex organic materials⁵⁸, resulting in reduced abundance of *Bacteroidetes* and a correlated 406 lower COD removal efficiency in both reactors. Similar differential susceptibilities to AgNPs 407 were observed in other identified phyla with a shift towards silver-tolerant genera (e.g. 408 Gemmata) or more sensitive genera (e.g. Gemmatimonas). Although the alpha diversity of the 409 bacterial community was not significantly impacted (Table S4), the bacterial community 410 composition was clearly changed, as shown in Figure 5B, due to a decrease or loss of silver-411 sensitive species in certain orders (e.g. Burkholderiale or Gemmatimonadales). The observed 412 shift in the microbial community composition can trigger a potential impairment of the biofilm 413 biological functions pertaining to organic matter biodegradation, enzymatic activities and biofilm 414 structural properties. 415

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Figure 5. (A) Taxonomic classification of 16S rDNA paired-end sequencing from the biofilm 423 samples at different AgNP concentrations at phylum level and (B) heatmap of genera with total 424 sequence reads higher than 150 in selected phyla. Note: Superscripts C and 64 represent the 425 biofilms collected at the end of control period and after 64 days of AgNPs exposure in MBBRs, 426 respectively. 427

428

429 4. Environmental implications

The MBBR technology has been successfully used for the treatment of many types of 430 wastewaters from municipalities, paper mills, pharmaceutical industries and fish farms⁵⁹. 431 MBBRs can be easily combined with other pre- or post- treatment technologies such as settling 432 and membrane separation, or used in a series of aerobic and anaerobic MBBRs, thus increasing 433 the likelihood of achieving a 'zero discharge' goal⁶⁰. Our findings, however, suggests that the 434

extended release of AgNPs and Ag-rich biofilm from an MBBR can exert a strong effect on 435 downstream treatments that may lead to membrane fouling, for example, due to significant 436 biofilm detachment, or to potential risks in effluent receiving streams that could impact ecosystems. 437 Our results, corroborated with previous studies^{18,52}, signify that short-term exposure tests may 438 underestimate the inhibitory effects of AgNPs in biofilm-laden environments, especially for 439 treatment processes with long sludge retention times where long-term continuous exposure to 440 AgNPs can result in a cumulative effect of NP-biofilm interaction dynamics leading to a 441 different level of NP-mediated susceptibility in the biofilm. For example, there was no 442 sulfidation of AgNPs in these MBBRs deployed here, due to lack of sulfur in the synthetic 443 wastewater. Sulfidation of AgNPs has been shown to retard dissolution of AgNP due to 444 formation of Ag₂S and organosulfur complexes and could have displayed less^{23, 61}. 445

446 ASSOCIATED CONTENT

447 Supporting Information

Additional information is provided for synthetic wastewater composition (Table S1-2) and 448 reactor operational conditions, characterization of biofilm biological responses, intracellular 449 ROS measurements, sp-ICP-MS instrumentation (Table S3) and TEM-EDS, cumulative total Ag 450 mass balance calculations, biofilm enzyme's half-life and inhibition rates (Table S4), richness 451 and diversity of microbial communities of biofilm (Table S5). In addition, fractionation of Ag in 452 reactor components are provided (Figure S1), Aginf/bio time distribution (Figure S2), AgNPinf/eff 453 particle size distribution (Figure S3), TEM images and EDS analysis for AgNP_{inf/eff} (Figure S4), 454 effluent TSS profile (Figure S5) and principal coordinate analysis (Figure S6) and additional 455 discussion on microbial community analysis. 456

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458 Acknowledgements

The authors thank the Natural Sciences and Engineering Research Council of Canada (Grant no. 459 STPGP 430659–12), Environment and Climate Change Canada, PerkinElmer, Health Sciences 460 Canada, the Fonds de Recherche du Québec Nature et Technologies (FRQNT), the Canadian 461 Water Network (CWN), SNC Lavalin Environment, the City of Calgary and the City of Saint-462 Hyacinthe for their financial support. The authors thank Jean-Philippe Massé, Le Centre de 463 Caractérisation Microscopique des Matériaux of Polytechnique Montreal for TEM/EDS analysis, 464 and Jean-Baptiste Burnet for DNA extraction and the Terrebonne/Mascouche WRRF for 465 assistance with wastewater sampling. 466

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