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1 **Title:**

2 Toxicity testing of silver nanoparticles in artificial and natural sediments using the benthic  
3 organism *Lumbriculus variegatus*

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26 **ABSTRACT**

27 The increased use of silver nanoparticles (AgNP) in industrial and consumer products worldwide  
28 has resulted in their release to aquatic environments. Previous studies have mainly focused on  
29 the effects of AgNP on pelagic species, while few studies have assessed the risks to benthic  
30 invertebrates despite the sediments acts a large potential sink for nanoparticles. In this study, the  
31 toxicity of sediment-associated AgNP was evaluated with the standard sediment toxicity test for  
32 chemicals provided by the Organization of Economic Cooperation and Development. The  
33 freshwater benthic oligochaete worm *Lumbriculus variegatus* was exposed to sediment-  
34 associated AgNP in artificial and natural sediments at concentrations ranging from 91 to 1098  
35 mg Ag/kg sediment dry weight. Silver nitrate (AgNO<sub>3</sub>) was used as a reference compound for  
36 Ag toxicity. The measured endpoints of toxicity were mortality, reproduction and total biomass.  
37 In addition, the impact of sediment-associated AgNP on the feeding rate of *L. variegatus* was  
38 studied in a similar test set-up as mentioned above. The addition of AgNP into the sediment  
39 significantly affected the feeding rate and reproduction of the test species only in the highest  
40 concentration (1098 mg/kg) of Ag in the natural sediment with the lowest pH. In comparison, the  
41 addition of AgNO<sub>3</sub> resulted in reproductive toxicity in every tested sediment, and Ag was more  
42 toxic when spiked as AgNO<sub>3</sub> than AgNP. In general, sediments were observed to have a high  
43 capacity to eliminate the AgNP derived toxicity. However, the capacity of sediments to eliminate  
44 the toxicity of Ag follows a different pattern when spiked as AgNP than AgNO<sub>3</sub>. This study  
45 emphasizes the importance of sediment toxicity testing and the role of sediment properties when  
46 evaluating the environmental effects and behavior of AgNP in sediments.

47 **Keywords;** Silver nanoparticles, environmental toxicology, nanoecotoxicology, sediment  
48 toxicity

## 49 1. INTRODUCTION

50 Silver nanoparticles (AgNP) used, e.g., in healthcare, textiles, paints, cosmetics and cleaning  
51 agents, have the highest degree of commercialization (as a number of products) of all nanoscale  
52 material due to the unique optical and antibacterial properties (Vance et al. 2015). In surface  
53 waters, AgNP are mainly released through wastewater treatment plants and untreated wastewater  
54 (Gottschalk et al. 2009). Sediment is the final sink for the AgNP, and modeled annual increase of  
55 sediment concentrations varies between 0.15 and 10.18  $\mu\text{g}/\text{kg}/\text{y}$ , resulting in a possible hazard  
56 for aquatic organisms (Gottschalk et al. 2009).

57 In environmental media AgNP may be oxidized, which leads to dissolution and release of Ag  
58 ions ( $\text{Ag}^+$ ) (Loza et al. 2014). Ionic Ag is highly toxic to aquatic organisms, and thus the toxicity  
59 of AgNP may be related to the concentration of dissolved  $\text{Ag}^+$  (Navarro et al. 2008; van Aerle et  
60 al. 2013). However, the concentration of freely dissolved  $\text{Ag}^+$  in environmental media is  
61 typically low due to strong complexation with chloride, sulfide and natural organic matter  
62 (Levard et al. 2013; Loza et al. 2014). Silver nanoparticles also pose nanoparticle-specific  
63 toxicity (Chan and Chiu 2015; Cozzari et al. 2015; García-Alonso et al. 2014). One of the  
64 primary identified toxic mechanisms at the molecular level is the generation of reactive oxygen  
65 species resulting in oxidative stress (Cozzari et al. 2015; Roh et al. 2009).

66 The behavior and toxicity of AgNP in sediment is still poorly understood and there is an urgent  
67 need for studies and standardized test methods. The biggest challenge in studies with  
68 nanomaterials in sediment and other complex environmental media is the lack of proper  
69 characterization methods. As most of the nanomaterials are not stable in water, sediment studies  
70 are still considered to be relevant and sometimes even more representative of environmental

71 exposure than aqueous tests (Petersen et al. 2015). In water-only exposure AgNP with varying  
72 coatings results in LC-50 (lethal concentration to kill 50 % of the test organisms in 96 h) of 0.07-  
73 0.33 mg/L to the benthic organism *Lumbriculus variegatus* (Khan et al. 2015). When the same  
74 species is exposed via sediment, AgNP shows no mortality upon exposure at 367 mg/kg  
75 (Coleman et al. 2013). However in sediment exposure AgNP induces oxidative stress in *Nereis*  
76 *diversicolor* already at concentrations lower than 10 mg/kg (Cozzari et al. 2015). Results indicate  
77 that the AgNP induced toxicity is reduced when particles are introduced into sediment, but the  
78 role of sediment properties has not yet been studied.

79 The aims of this study were: 1) to examine how the sediment properties influence the toxicity of  
80 AgNP, 2) to compare the toxicity of Ag spiked as AgNP to dissolved Ag spiked as silver nitrate  
81 (AgNO<sub>3</sub>), and 3) to evaluate the suitability of the OECD standard test method guideline 225 for  
82 use with nanomaterials. Artificial and two natural sediments that differed in their characteristics  
83 were selected and spiked with polyvinylpyrrolidone-coated AgNP and AgNO<sub>3</sub>. The OECD  
84 standard test guideline 225 “Sediment-Water Lumbriculus Toxicity Test Using Spiked  
85 Sediment” (OECD 2007) was followed, and mortality, reproduction and changes in biomass  
86 were used as indicators of toxicity to the endobenthic aquatic Oligochaeta *Lumbriculus*  
87 *variegatus*. In addition, the feeding rate of *L. variegatus* was used as an endpoint of toxicity for  
88 AgNP.

## 89 2. MATERIAL AND METHODS

### 90 2.1 Silver nanoparticles

91 Silver nanoparticles (polyvinylpyrrolidone coating 0.2 %, NanoAmor) had a nominal reported  
92 surface area of 5–10 m<sup>2</sup>/g and a diameter of 30–50 nm with a purity of 99.9 %. The  
93 characterization of the particles was published in the same year as the experimental part of this  
94 study was done (Heckmann et al. 2011). Particles were stored as dry powder and kept away from  
95 the direct sunlight, as recommended by the manufacturer, to minimize the possible changes in  
96 particle properties during the storage. The characterization of AgNP included: transmission  
97 electron microscopy (Phillips CM20, Phillips/FEI), dynamic light scattering and zeta potential  
98 measurements (Malvern Zetasizer Nano, Malvern Instruments Ltd). The characterization of  
99 AgNP was done in deionized water suspension due to the lack of methods to characterize the  
100 particles in complex environmental media. Characterization in the test water was not considered  
101 to be relevant, as particles were never introduced into the test water. The mean diameter of  
102 AgNP has been reported to be  $82 \pm 2$  nm (n=294) measured from the transmission electron  
103 microscope images and  $235 \pm 4$  nm (n=4) with a zeta potential of  $-28.6 \pm 0.6$  mV (n=8) by the  
104 dynamic light scattering (Heckmann et al. 2011). Agglomeration of the AgNP in water  
105 suspension explains the larger diameter of the particles measured by the dynamic light scattering.  
106 For further details of the characterization, see Heckmann et al. (2011).

### 107 2.2 Test Organisms

108 Endobenthic oligochaeta *Lumbriculus variegatus* originated from the laboratory culture  
109 maintained at the Department of Biology, University of Eastern Finland, Joensuu, Finland.  
110 Worms were cultured in 5 L tanks, containing artificial fresh water (AFW, pH 7, hardness 1.0

111 mM/L as [Ca] + [Mg]) with a constant aeration. The light regime was adjusted to 16 hours light  
112 and 8 hours dark, and temperature was held constant at  $20 \pm 2$  °C. A layer of paper towels was  
113 used as a substrate. Worms were fed twice a week with a Tetramin fish food (Tetrawerke) and  
114 water was renewed once a week. Acclimation phase of 24 h was used to adapt the worms to test  
115 water.

### 116 **2.3 Sediments**

117 One artificial sediment (AS) and two natural sediments collected from Lake Höytiäinen (HS)  
118 and Lake Kuorinka (KS) were used in this study. Both natural sediments have been used as clean  
119 reference sediments in similar experiments, and possible trace amounts of organic chemicals are  
120 low and not believed to have an influence on the outcome of current experiments (Mäenpää et al.  
121 2008; Ristola et al. 1996). The sediment AS was prepared using the OECD guideline 225  
122 (OECD 2007). The exact constituent composition was 5 % peat, 74 % quartz sand (60 % < 0.2  
123 mm, 40 % 0.2–1.0 mm), 20 % kaolin and 51 % water (of total dw). *Urtica dioica* powder (0.5 %)  
124 was added as a food source to AS and pH was adjusted to 6.7 with CaCO<sub>3</sub>.

125 For analyses, natural sediments were sieved through a 1 mm sieve to remove large particles and  
126 debris. Subsamples of the sediments were dried at 105 °C over-night to measure dry weight. The  
127 determination of organic carbon, inorganic carbon and black carbon were done with Analytik  
128 Jena TOC analyzer with a solid sample module (Analytik Jena N/C 2100). Furthermore,  
129 subsamples of the sediments were heated for 2 h at 550 °C in a muffle furnace oven (Naber 2804  
130 L47) to obtain the loss of ignition percent. All analyses were done in three replicates.

131 The heavy metal concentrations of sediments were measured from two different test vessels for  
132 each treatment, and the total Ag concentrations were determined in triplicate for each treatment.



133 The sediment samples were stored frozen at -20 °C prior the extraction. The extraction was as  
134 follows: A subsample of approximately 200 mg (500 mg for total Ag) was taken from dry  
135 sediment, and digested in 1:3 nitric acid:hydrochloric acid (v:v) solution for 9 minutes in three  
136 minute intervals in ultrasound water bath (650 W, 35 kHz, ELMA Transsonic T820/H) at 60 °C.  
137 The sample tubes were shaken between each 3 minute step. The digested sediment samples were  
138 filtered (Whatman No. 41) and diluted to a volume of 20 ml (50 ml for total Ag) with ultrapure  
139 water prior to the analysis. The samples were analyzed with Perkin-Elmer model Optima 8300  
140 inductively coupled plasma optical emission spectrometry. The cyclonic spray chamber equipped  
141 with the GemCone Low-Flow nebulizer was used throughout. The plasma power of 1500 W and  
142 nebulizer flow of 0.6 l/min was used in order to get robust plasma conditions for the accurate  
143 analysis of the elements. Reagent blank samples were used in between of the samples to ensure  
144 the analytical procedure. The accepted relative standard deviation of three replicate  
145 measurements was less than 10 %, and the detection limit was 1.9 µg/L. All the used reagents  
146 were of analytical grade and supplied by Merck.

#### 147 **2.4 Spiking of the sediments**

148 Direct addition of dry AgNP powder to the sediment was chosen as the spiking method due to the  
149 unstable behavior of the particles in the water suspension. The final Ag concentrations were  
150 selected based on the preliminary test (Table 1). The sediments were spiked with AgNP by first  
151 mixing the nanoparticle powder to a small subsample of the sediment with a metal spoon. The  
152 subsample was then mixed to the rest of the sediment. To ensure the homogenous distribution of  
153 the compounds, the sediment was mixed with a rotating metal blade for one hour. Silver nitrate  
154 (high grade: 99.5% purity, supplied by J.T. Baker) was used as a source of dissolved Ag, and

155 added to the sediment in a stock solution dissolved in water (400 g/L). The sediment was treated  
156 in a similar way as the AgNP-spiked sediment.

## 157 **2.5 Toxicity test**

158 The toxicity of AgNP was tested according to the OECD guideline 225, using AgNO<sub>3</sub> as a  
159 reference for Ag<sup>+</sup> toxicity (OECD 2007). The test was conducted in 250 ml beakers (diameter 6  
160 cm) with 4 replicates for each treatment, and 6 replicates for the control treatment. The amount  
161 of the sediment was adjusted to the ratio of 1:50 (dry biomass of worms:total organic carbon of  
162 the sediment, w:w). The sediment-overlying water ratio was adjusted to approximately 1:3 (v:v).  
163 The water hardness of AFW was 2.5 mM/L ([Ca] + [Mg] concentration) and pH was 7.5 (OECD  
164 2007). The sediments were allowed to settle for 7 days with gentle aeration before adding 10  
165 similar-sized *L. variegatus* into the test vessels. The worms were not synchronized for the  
166 toxicity test based on the consistent results with only low variation in reproduction and biomass  
167 of the worms in the preliminary test (data not shown). During the incubation, the temperature  
168 was kept constant at 20 ± 2 °C, and the light regime was 16 h light to 8 h dark. Oxygen and pH  
169 were measured once a week during the test. After the 28-day exposure time, the worms were  
170 removed from the sediment, counted and placed on a petri dish with a small amount of AFW. A  
171 depuration time of 4 h was used to let the worms empty their gut before placing the worms in an  
172 oven at 105 °C for overnight. The dry weight was measured with a microbalance (Sartorius  
173 4503). Missing worms were interpreted as mortality and extra worms as reproduction in the test  
174 vessels after the exposure period.

## 175 **2.6 Feeding rate test**

176 The feeding rate test was done according to Leppänen and Kukkonen (1998) in similar  
177 conditions as described above for the toxicity test. Three replicates were used for each treatment,  
178 and two control treatments were done for each sediment (total n=6). A portion of 23 g of wet test  
179 sediment was added on the bottom of the 50 ml beakers, which were then filled with 2.5 mM/L  
180 ([Ca] + [Mg] concentration) AFW (OECD 2007). The oxygen level and pH in the overlying  
181 water were measured during the test, and the water was renewed using aerated AFW every two  
182 days. Before adding the organisms into the beakers, the sediments were allowed to settle for 2  
183 days. Each beaker received five worms of a similar size. Immediately after the worms buried  
184 themselves into the sediment, a layer of a few millimeters of combusted quartz sand (grain size  
185 1–2 mm) was added on the top of the sediment. The egestion rate of the worms was followed by  
186 collecting fecal pellets every second day for 14 days. The fecal pellets were dried overnight at  
187 105 °C, and the dry weight was measured with a microbalance. On the last day of the  
188 experiment, the worms were removed from the sediment. After a 4 h depuration time in clean  
189 AFW, the worms were counted and dried at 105 °C overnight to measure their dry weight.

## 190 **2.7 Statistical testing**

191 The effective (EC) and inhibition (IC) concentrations were estimated using a three parameter  
192 log-logistic model. The normality of data was tested with Shapiro-Wilk normality test and the  
193 homogeneity of variances with Bartlett's test. Normally distributed data with equal variances  
194 between groups was studied with one-way analysis of variance (ANOVA) followed by pairwise  
195 t-test. When the data was not normally distributed, Kruskal-Wallis rank sum test was used and  
196 multiple comparisons between groups were done according to Siegel & Castellan (1988).

197 One-way ANOVA with Tukey's HSD (honestly significant difference) post-hoc test ( $p < 0.05$ )  
198 was used to compare the amount of fecal pellets in the feeding rate test. The normality of the  
199 data was tested with Shapiro-Wilk normality test, and the homogeneity of variances with  
200 Levene's test. Due to the small sample size ( $n=3$ ) the normality of the treatment groups was  
201 assumed from the normally distributed control groups ( $n=6$ ) in all sediments. Statistical analyses  
202 and graphical illustrations were done with R version 3.0.1.

203

## 204 **3. RESULTS**

### 205 **3.1 Sediment characteristics**

206 The HS sediment had the highest pH (7.10) and organic carbon percent (OC% = 3.12 %) of the  
207 tested sediments (Table 2). In the KS sediment the pH was low (5.10) and OC% (2.22) lower  
208 than in HS but higher than in the AS sediment (0.59 %), which had also higher pH (6.70) than  
209 the KS sediment (Table 2). The visual detection and smaller dw% indicated that the natural  
210 sediments KS and HS consisted of finer material compared to AS (Table 2). The artificial  
211 sediment contained only low levels of heavy metals. The natural sediments had higher  
212 concentrations, the HS sediment containing approximately 2 to 3 fold higher concentrations in  
213 comparison to the KS sediment (Table 2). The determined Ag concentrations were in good  
214 agreement with the nominal concentrations, and standard deviation among the replicates was  
215 relatively small, which indicates homogenous distribution of Ag in the sediments (Table 1.).

### 216 **3.2 Toxicity test**

217 The pH of the overlying water was at acceptable levels (6–9) for *L. variegatus* in AS and HS, but  
218 in the KS sediment the pH was lower than recommended in the guideline (OECD 2007). The  
219 oxygen saturation was over 90 % throughout the experiment in all sediments, but the validity  
220 criteria of an 1.8-fold increase in the number of individuals was only fulfilled in the AS sediment  
221 (OECD 2007).

222 Exposure to AgNP-spiked sediments caused no mortality in any sediment type or exposure  
223 concentration, but reproduction was significantly decreased compared to the control in the  
224 highest concentration in the KS sediment (pairwise *t*-test,  $p < 0.001$ ) (Fig 1a). In this treatment  
225 the worms were also avoiding the sediment. The AgNO<sub>3</sub>-spiked AS sediment was the most toxic

226 to *L. variegatus* and the only sediment where mortality was observed (Fig 1b). Reproduction was  
227 decreasing with increasing AgNO<sub>3</sub> concentration in all of the tested sediments (Fig 1b). The  
228 calculated IC<sub>50</sub> values for the reproduction and EC<sub>50</sub> values for the number of worms  
229 (compared to control) indicate that the Ag spiked as AgNP was only toxic in KS, and Ag spiked  
230 as AgNO<sub>3</sub> showed highest toxicity in AS, followed by HS, and the lowest toxicity was observed  
231 in KS (Table 3).

232 No statistically significant differences in the total dry biomass of the worms were observed  
233 among the exposure concentrations in any of the test sediments spiked with AgNP (Fig 1c). The  
234 total dry biomass decreased with increasing AgNO<sub>3</sub> concentration in AS, but stayed constant in  
235 KS and HS (Fig 1d). Furthermore, the sediment properties affected the total biomass of the test  
236 species among the test sediments. In the beginning of the experiment the total dry biomass of the  
237 worms was 15.69 mg (SD 0.17) in AS and HS, and 19.04 mg (SD 1.78) in KS. After the 28-day  
238 test period the total dry biomass of worms increased in the control groups by on average 49 %  
239 (23.31 mg SD 1.89) in AS but decreased by on average 7 % (14.56 mg SD 1.03) in HS and 16 %  
240 (16.01 mg SD 1.60) in KS.

### 241 **3.3 Feeding rate**

242 Silver nanoparticle exposure had no effect on the fecal pellet production of the worms in HS and  
243 AS (Fig 2a). In the KS sediment, however, the fecal pellet production increased with increasing  
244 exposure concentration of AgNP with an exception that at the highest exposure concentration  
245 (1098 mg/kg) the worms were avoiding the sediment and the pellet production thus decreased  
246 (Fig 2a).

247 In the natural sediments HS and KS the worms reproduced during the 14-day exposure period  
248 (Fig 2b). In the HS sediment the worms reproduced in each concentration somewhat evenly, but  
249 in the KS sediment reproduction was observed only in the controls and in the two lowest  
250 exposure concentrations (Fig 2b). In the AS sediment only few extra worms were found in  
251 occasional test vessels. No significant differences were found in the total dry biomass of the  
252 worms between the different Ag concentrations. The biomass gain was different among the  
253 sediments, as also observed in the toxicity test. The biomass of the worms increased during the  
254 14-day exposure period in AS (71 %), stayed constant in KS, and decreased in HS (17 %) (Fig  
255 2c).

256 **4. DISCUSSION**

257 Silver nitrate and AgNP are known to be extremely toxic to the benthic organisms (Khan et al.  
258 2015; Nair et al. 2013). However, the majority of the toxicity studies have been done using  
259 waterborne exposures, not considering the natural environment of the benthic organisms. In  
260 waterborne exposures the uptake of Ag occurs primarily over the respiratory body surface.  
261 Sediment exposures are more environmentally realistic, as organisms feed on the sediment, and  
262 Ag is also internalized into the organisms through the gut epithelium. Dietary uptake is  
263 especially important when Ag is spiked as AgNP, as particles can be internalized directly via  
264 endocytosis (García-Alonso et al. 2011). Endocytic uptake can lead to nanoparticle-specific  
265 modes of toxicity, which cannot be considered in water-only exposures.

266 In this study, the toxicity of Ag spiked as AgNP and AgNO<sub>3</sub> to *L. variegatus* in sediment  
267 exposures was remarkably lower compared to waterborne exposures in the literature. Khan et al.  
268 (2015) reported the LC50 concentrations in the 96 h acute toxicity test to be 64.6 µg/L for PVP-  
269 coated AgNP and 4.4 µg/L for AgNO<sub>3</sub> in the OECD 225 standard AFW. In the present study no  
270 mortality was observed in any of the tested sediments even in the highest 1098 mg/kg (dw)  
271 concentration of sediment-associated AgNP. The EC50-value for AgNO<sub>3</sub> was 38 mg/kg (dw) in  
272 AS sediment, but no mortality was observed in other test sediments. The decrease of toxicity of  
273 Ag in sediments compared to the waterborne exposures is dramatic, especially when spiked as  
274 AgNP, despite the possible direct uptake of AgNP by endocytosis. The capacity of sediment to  
275 decrease the toxicity of Ag emphasizes the need of sediment toxicity tests when evaluating the  
276 environmental effects of AgNP. Our results indicate that the toxicity to benthic fauna may be  
277 highly overestimated if only waterborne exposures are used.



278 Dissolved Ag spiked as AgNO<sub>3</sub> was more toxic to *L. variegatus* in the artificial sediment than in  
279 the natural sediments. This suggests that the Ag<sup>+</sup> binding capacity is greater in the natural  
280 sediments compared to the AS sediment. The higher OC content of the HS and KS sediments  
281 compared to the AS sediment partly explains the lower toxicity of AgNO<sub>3</sub>, as Ag is known to  
282 form complexes with OC (Erickson et al. 1998). Also the grain size of the natural sediments is  
283 small; 79.0 % (HS) and 77.9 % (KS) of the particles are under 63 µm in diameter (Mäenpää et al.  
284 2003). The high dw% in the AS sediment indicates that the sediment was mainly reconstructed  
285 from coarse quartz sand resulting in a smaller surface area in the AS sediment components to  
286 bind Ag. The concentration of acid volatile sulfides (AVS) in the sediment is often considered to  
287 be the most important individual factor in anoxic sediments, since Ag has a strong affinity  
288 towards organic and inorganic sulfur groups (Bell and Kramer 1999; Berry et al. 1999). In this  
289 study the tested sediments were treated under oxidized conditions, where the concentration of  
290 AVS can be considered negligible (Di Toro et al. 1990). Silver has also high affinity towards Cl<sup>-</sup>  
291 anions (Wingert-Runge and Andren 1993). In our test set-up the amount of Cl anions in the  
292 overlying AFW was theoretically high enough to complex all Ag<sup>+</sup>, but as the Ag compounds  
293 were spiked directly to the sediment, the effect of Cl and other anions is considered small. This is  
294 proved by the toxicity of AgNO<sub>3</sub> in the test sediments despite the complexing anions in overlying  
295 water.

296 The toxicity of Ag increases when pH decreases, due to the increased free Ag<sup>+</sup> concentration in  
297 the media (Erickson et al. 1998). Low pH also increases the dissolution of AgNP, which leads to  
298 a higher free Ag<sup>+</sup> concentration and increased toxicity (Navarro et al. 2008; Peretyazhko et al.  
299 2014; van Aerle et al. 2013). The natural sediment KS had the lowest pH of the tested sediments.  
300 Whereas the toxicity of Ag spiked as AgNO<sub>3</sub> was lowest in KS, it was the only sediment in

301 which the addition of AgNP resulted in reproductive toxicity. This indicates that low pH may  
302 increase the toxicity of AgNP more than that of AgNO<sub>3</sub>. The IC50 values for reproduction were  
303 approximately 2 times higher for AgNP than for AgNO<sub>3</sub> in KS sediment. If the toxicity is  
304 proposed to be solely a function of Ag<sup>+</sup>, around 50 % of the particles would be dissolved. The  
305 partitioning studies done in sediment however show that the bioavailable concentration of Ag<sup>+</sup> in  
306 sediment is higher when added as AgNP than when added as AgNO<sub>3</sub> (Coutris et al. 2012). Direct  
307 comparisons between the toxicity data and the dissolution of AgNP cannot thus be made. AgNP  
308 can also pose nanoparticle-specific toxicity over Ag<sup>+</sup> (Chan and Chiu 2015; Cozzari et al. 2015;  
309 García-Alonso et al. 2014) or “Trojan horse” -type of behavior, leading to the intracellular  
310 release of Ag<sup>+</sup> (Moore 2006; Park et al. 2010; Wang et al. 2013). If these nanoparticle-specific  
311 modes of toxicity would explain the toxicity of AgNP in the KS sediment, the bioavailability of  
312 AgNP should be higher in the KS sediment compared to the other tested sediments, as no toxicity  
313 was observed in the HS or AS sediments. This is unlikely as the relatively low pH in KS actually  
314 suggests lower bioavailability of AgNP compared to the other more alkaline sediments due to a  
315 stronger electrostatic attraction between the negatively charged particles and positively charged  
316 matrix (Cornelis et al. 2014). Considering these facts, we suggest that the AgNP toxicity in KS  
317 was mainly caused by dissolved Ag<sup>+</sup> released from the particles and that the dissolution is  
318 promoted by the low pH of the sediment.

319 The nutritional value of sediment to *L. variegatus* varied between the tested sediments. The AS  
320 sediment was the only sediment where the worms were gaining weight. The total biomass of the  
321 worms was decreasing in the HS and KS sediments despite that the worms ingested both natural  
322 sediments. This indicates the poor nutritional value of the natural sediments compared to the AS  
323 sediment. Especially the KS sediment seems to have a poor nutrient content, since the biomass-

324 normalized ingestion rate was highest among the test sediments but the biomass loss was the  
325 largest. No significant difference in the total biomass was found between the treatments in  
326 toxicity or feeding rate test, despite the significant decrease in the ingested amount of sediment  
327 in the highest concentration of AgNP in KS. The biomass change seems not to be an applicable  
328 endpoint for the acute toxicity tests in the natural sediments with poor nutritional value, as the  
329 worms were losing weight also in the control groups of the HS and KS sediments.

330 The feeding behavior of *L. variegatus* has been shown to give an immediate response to the  
331 exposure, and it is considered to be a more sensitive endpoint than mortality, biomass gain or  
332 reproduction (Leppänen and Kukkonen 1998). Generally the ingestion rate tends to decrease  
333 with increasing concentration of contaminant, but in the KS sediment *L. variegatus* ingested  
334 more sediment with increasing AgNP concentration. We suggest that the antibacterial properties  
335 of AgNP disturbed the microbial growth in the sediment, which impeded adequate nutrition of *L.*  
336 *variegatus*, and thus worms had to compensate for the nutrient-poor food by ingesting more  
337 sediment. In the highest exposure concentration, however, the AgNP-induced stress seemed to  
338 become too high for the *L. variegatus*, as the worms avoided the sediment throughout the test  
339 period and thus the feeding rate was minimal. The increase in the feeding rate was only observed  
340 in the KS sediment. The microbes can be a more important food source in KS compared to the  
341 other test sediments due to the poor nutritional value. Also the low pH of KS is believed to be an  
342 intensifying factor for AgNP toxicity as discussed before.

343 The natural sediments HS and KS used in this study were selected to represent typical unpolluted  
344 Finnish lake sediments from a watershed without industrial influence, and have been used in  
345 studies as clean reference sediments (Mäenpää et al. 2003; Ristola et al. 1996). The geochemical  
346 background level of metals is slightly elevated if compared to the consensus-based threshold

347 effect concentrations (TEC), meaning that these metals possibly cause toxic effects in a  
348 freshwater ecosystem (MacDonald et al. 2000). In HS the Cd, Cr, Cu, Ni, Pb, and Zn  
349 concentrations are above the TEC. In KS the metal concentrations are also elevated but  
350 somewhat lower compared to HS, and only Cd, Cu and Ni are above the TEC values. The  
351 background metal concentrations are however typical for the sediments in this area (Ristola et al.  
352 1996). When a test sediment is amended with Ag, it is possible that Ag<sup>+</sup> and AgNP displace  
353 sediment-bound metals and release them into the sediment pore water. Especially Zn and Ni are  
354 known to be displaced by Ag (Call et al. 1999). The measured toxicity in the natural sediments  
355 may therefore be a mixture effect of metals, Ag being the predominant active substance. Higher  
356 concentration of background metals may therefore explain the higher toxicity of AgNO<sub>3</sub> in HS  
357 sediment compared to the KS sediment. In the AgNP treatments this effect is not pronounced, as  
358 the dissolution of nanoparticles is believed to be more promoted by the lower pH of KS  
359 compared to HS, leading to the higher toxicity in KS. The environmental relevance is often a key  
360 factor when considering the behavior of nanoparticles in the aquatic environment. As the  
361 properties of the natural sediments differ greatly from the artificially prepared standard sediment,  
362 we consider testing in the natural sediments highly important, despite the fact that the  
363 environmental factors apart from the nanoparticle exposure may complicate interpreting the  
364 results.

365 The OECD standard test guideline 225 was successfully applied for use with nanomaterials. The  
366 AS sediment prepared following the OECD standard guideline was the only sediment that  
367 fulfilled the validity criteria of an 1.8-fold increase in the number of individuals, and thus only  
368 this part of the study can be considered as a standardized toxicity test. The low reproduction rate  
369 and pH-related problems in natural sediments advocate the use of artificial sediment in standard

370 testing. The results of the OECD toxicity tests are in line with the feeding rate test, which  
371 increases the reliability of the test. However, the following concerns may have significant effect  
372 on the results of the test and should be properly addressed in the future: 1) The spiking method  
373 of the nanomaterial may have an influence on the outcome of the test. We chose to add the dry  
374 powder of AgNP directly into the sediment, because the amount of nanoparticles was high, and  
375 the particles were unstable in water suspension in such a high concentration. More stable  
376 nanoparticle suspensions could also be spiked as suspension to avoid clumping of the material.  
377 The reduced clumping leads to a higher total surface area of the spiked component and may  
378 possibly lead to elevated toxicity. Indirect addition of the nanoparticles to the overlying water  
379 would be an environmentally more relevant way to spike the nanomaterial, but could decrease  
380 the oral uptake of the substance, since *L. variegatus* burrow into the sediment and feed below the  
381 sediment surface. 2) The characterization of nanomaterial should be carefully considered. Since  
382 we do not currently have proper methods to characterize the sediment-associated nanomaterial,  
383 characterization in this study was done in deionized water before spiking the nanoparticles into  
384 the test media. Despite the fact that the characterization in water does not correspond to the  
385 experimental conditions in the sediment, it is essential to assess the primary structure and  
386 properties of the particles in standard conditions to add comparability between the studies. The  
387 characterization of nanoparticles in the overlying water was not considered relevant, since AgNP  
388 were spiked to the sediment by direct addition and were never present in the water phase. If the  
389 indirect addition is used, the characterization in the overlying water should also be considered, as  
390 the aggregation and dissolution of coated AgNP in the water phase is differently affected by the  
391 presence of sediment (Bone et al. 2012; Unrine et al. 2012). In conclusion, there is an urgent  
392 need to develop reliable and easily achievable methods for the characterization of the

393 nanomaterials in the sediment media. However, the former concerns should not hinder the  
394 toxicity testing of nanomaterials in sediment or other complex environmental matrix. Despite the  
395 methodological challenges, tests give us important information on the possible toxicity of  
396 nanomaterials.

397 **5. CONCLUSIONS**

398 The acute toxicity of Ag spiked as AgNP to *L. variegatus* was greatly decreased in sediments  
399 compared to literature-reported waterborne toxicity. Silver nitrate was significantly more toxic  
400 than AgNP in all of the test sediments, but sediment properties had a different effect on the  
401 toxicity of the two compounds. The toxicity of AgNO<sub>3</sub> was lower in the sediments with fine  
402 grain size and relatively high amount of OC. The low pH of the sediment seemed to overcome  
403 these factors when Ag was spiked as AgNP, and toxicity as reproductive failure, changes in the  
404 feeding behavior and sediment avoidance was only observed in the natural KS sediment with the  
405 lowest pH value of the tested sediments. We suggest that low pH of the KS sediment enhances  
406 the release of Ag<sup>+</sup> from AgNP and thus promotes the toxicity. However, nanoparticle-specific  
407 toxicity or synergistic effect of both Ag<sup>+</sup> and AgNP, and natural heavy metals cannot be  
408 excluded. Finally, we conclude that OECD guideline 225 “Sediment-Water Lumbriculus Toxicity  
409 Test Using Spiked Sediment” can be used for evaluating the toxicity of nanomaterials in  
410 sediments. Further studies considering AgNP in sediments should concentrate on the dissolution  
411 kinetics and the effects of sediment pH on the toxicity of AgNP. In general with nanoparticles,  
412 the development of characterization methods in complex environmental media is the most  
413 essential issue. However, the lack of characterization methods should not hinder the toxicity  
414 testing of nanoparticles in complex environmental media, since nanoparticles are constantly  
415 released into the environment where they are likely to pose a risk to the benthic ecosystems.

416

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

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535 **TABLES**

536

537 **Table 1** Nominal and determined silver concentrations (mg/kg dry weight) spiked as silver538 nitrate (AgNO<sub>3</sub>) and silver nanoparticles (AgNP) in test sediments

Compound	Nominal concentration	Determined concentration in sediments (mg/kg) <sup>a</sup>		
		Artificial	Höytiäinen	Kuorinka
AgNO <sub>3</sub>	10	11.7 (0.6)	10.8 (0.0)	10.1 (0.0)
	40	36.9 (1.0)	42.8 (0.4)	41.9 (0.3)
	100	92.7 (1.7)	98.9 (0.9)	103.1 (3.6)
	200	180.6 (1.6)	199.3 (1.2)	208.9 (2.9)
	400	360.4 (7.1)	394.8 (5.2)	415.9 (4.0)
AgNP	100	90.6 (1.9)	95.7 (10.8)	77.1 (5.0)
	300	262.5 (2.3)	273.8 (18.1)	256.7 (23.5)
	600	537.4 (12.3)	471.1 (9.8)	525.8 (27.4)
	900	725.4 (23.5)	741.2 (20.1)	801.3 (10.5)
	1200	985.1 (17.8)	943.4 (164.0)	1097.9 (7.5)

539 <sup>a</sup> Mean and standard deviation of 3-5 replicates

540 **Table 2** The characteristics of the test sediments. LOI% = Loss of ignition, OC% = organic  
 541 carbon, BC% = black carbon, IC% = inorganic carbon, dw% = dry weight

	Sediment		
	Artificial	Höytiäinen	Kuorinka
pH <sup>a</sup>	6.70 (0.26)	7.10 (0.21)	5.10 (0.11)
LOI% <sup>b</sup>	6.30 (0.06)	10.6 (0.1)	6.03 (0.03)
OC% <sup>b</sup>	0.59 (0.15)	3.12 (0.31)	2.22 (0.14)
BC% <sup>b</sup>	0.04 (0.01)	0.05 (0.00)	0.14 (0.00)
IC% <sup>b</sup>	0.75 (0.37)	-	-
dw% <sup>c</sup>	60.0 (0.2)	18.2 (0.1)	33.2 (0.3)
Cd <sup>d</sup>	0.15 (0.08)	3.34 (0.18)	1.07 (0.07)
Cr <sup>d</sup>	3.79 (0.81)	51.9 (4.8)	20.1 (1.7)
Cu <sup>d</sup>	3.56 (3.15)	53.0 (4.8)	35.8 (3.5)
Ni <sup>d</sup>	2.49 (0.41)	44.2 (3.8)	25.0 (1.7)
Pb <sup>d</sup>	0.77 (0.14)	44.4 (1.9)	18.8 (1.8)
Zn <sup>d</sup>	3.19 (1.60)	130 (7)	56.6 (5.4)

542 <sup>a</sup> Mean (standard deviation) of weekly measures during the 28 d toxicity test (n>55).

543 <sup>b</sup> Percent of the sediment dry weight, mean (standard deviation) of 3 replicates

544 <sup>c</sup> Percent of the sediment total weight, mean (standard deviation) of 3 replicates

545 <sup>d</sup> Metal concentration as mg/kg of dry sediment, mean (standard deviation) of 24 to 26 replicates

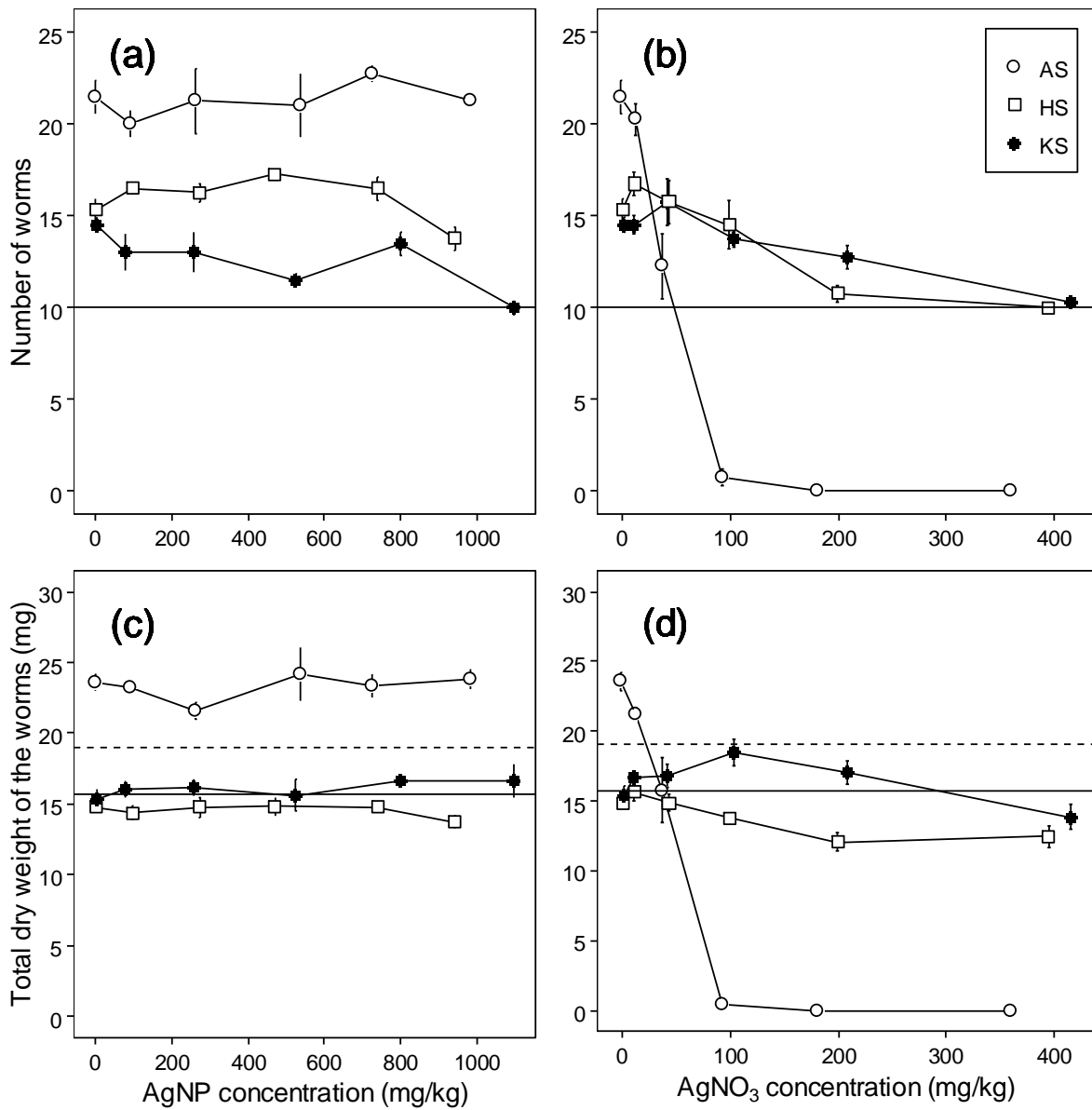
546 **Table 3** Calculated 50 % reproduction-inhibition concentrations (IC50) and 50 % effect  
 547 concentrations (EC50) for decrease in the number of worms compared to control for silver nitrate  
 548 (AgNO<sub>3</sub>) and silver nanoparticle (AgNP) exposed *Lumbriculus variegatus*

	End Point	Compound	Sediment		
			Artificial	Höytiäinen	Kuorinka
Reproduction <sup>a</sup>	IC50	AgNO <sub>3</sub>	23.9 (3.60)	129 (19.9)	213.50 (47.77)
	IC50	AgNP	-	-	442.84 (316.74)
Number of worms <sup>a</sup>	EC50	AgNO <sub>3</sub>	38.0 (1.97)	525 (107.07)	687.59 (187.75)

549 <sup>a</sup>Based on three parameter log-logistic model, Ag (determined concentration) in kg of dry  
 550 sediment (standard deviation)

551

552 **FIGURES**

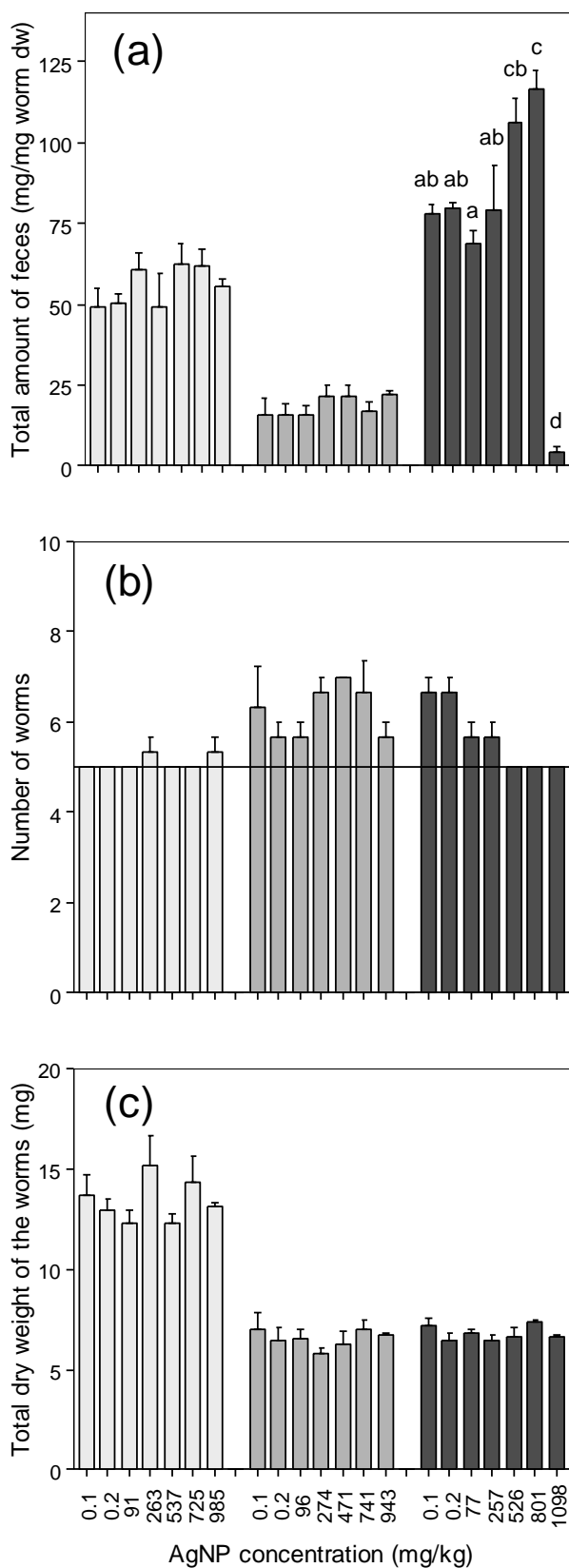


553  
 554 **Fig 1.** Toxicity of Ag spiked as silver nanoparticles (AgNP) and silver nitrate (AgNO<sub>3</sub>) to  
 555 *Lumbriculus variegatus* after a 28-day exposure in spiked artificial (AS), Höytiäinen (HS) and  
 556 Kuorinka (KS) sediments at various Ag concentrations (mg/kg dry weight). Number of worms  
 557 after a 28-day exposure to AgNP (a) and AgNO<sub>3</sub> (b). Dry biomass of worms after exposure to  
 558 AgNP (c) and AgNO<sub>3</sub> (d). Each symbol indicates mean and standard deviation of four replicate  
 559 samples, except for control exposure that had six replicates. Solid line in (a) and (b) indicates the

560 number of worms (10) at the beginning of the experiment. Solid line in (c) and (d) indicates the  
561 starting dry biomass of the worms in the AS and HS sediments (15.69 mg), dashed line in the KS  
562 sediment (19.04 mg).

563





**Fig 2.** Effects of Ag spiked as silver nanoparticles (AgNP) on *Lumbriculus variegatus* in 14-days feeding rate test in artificial (AS), Höytiäinen (HS) and Kuorinka (KS) sediments at various Ag concentrations (mg/kg dry weight). Two control groups without AgNP are marked as 0.1 and 0.2. Mass of dry feces produced during the experiment normalized to the total dry biomass of the worms (a). Identical letters (a-d) indicate groups that do not significantly differ from each other ( $p < 0.05$ ). Number of worms in the end of the experiment; the solid line indicates the number of worms in the beginning of the experiment (b). Total dry biomass of the worms after the experiment (c). Each bar indicates mean and standard deviation of three replicate samples.