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1 Effects of nano sized- zero-valent iron (nZVI) on DDT degradation in soil and its toxicity to
2 collembola and ostracods

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41

42 **Abstract**

43 Nano sized zero valent iron (nZVI) has been studied for *in-situ* remediation of contaminated soil
44 and ground water. However, little is known about its effects on organisms in soil and aquatic
45 ecosystems. In this study, the effect of nZVI on degradation of DDT and its ecotoxicological effects
46 on collembola (*Folsomia candida*) and ostracods (*Heterocypris incongruens*) were investigated.
47 Two soils were used in suspension incubation experiments lasting for 7 and 30 days; a spiked (20
48 mg DDT kg⁻¹) sandy soil and an aged (>50 yrs) DDT-polluted soil (24 mg DDT kg⁻¹). These were
49 incubated with 1 or 10 g nZVI kg⁻¹, and residual toxicity in soil and the aqueous phase tested using
50 ecotoxicological tests with collembola or ostracods. Generally, addition of either concentration of
51 nZVI to soil led to about 50 % degradation of DDT in spiked soil at the end of 7 and 30 d incubation,
52 while the degradation of DDT was less in aged DDT-polluted soil (24 %). Severe negative effects
53 of nZVI were observed on both test organisms after 7 d incubation, but prolonged incubation led
54 to oxidation of nZVI which reduced its toxic effects on the tested organisms. On the other hand,
55 DDT had significant negative effects on collembolan reproduction and ostracod development. We
56 conclude that 1g nZVI kg⁻¹ was efficient for significant DDT degradation in spiked soil, while a
57 higher concentration was necessary for treating aged pollutants in soil. The adverse effects of nZVI
58 on tested organisms seem temporary and reduced after oxidation.

59 **Key words:** Nano-remediation, DDT, Nano-ecotoxicity, aged-polluted soil, chlorinated organic
60 pollutants, nanoparticles, nZVI, ostracods, collembola.

61

62 **1. Introduction**

63 Organo-chlorine insecticides such as DDT [1,1,1-trichloro-2,2-bis(*p*-chlorophenyl) ethane]
64 have been extensively used throughout the world since the 1940-ies to control pests in agriculture
65 and vectors of diseases such as malaria mosquitos (WHO, 1979). DDT has been recognized as
66 potentially toxic to humans and animals because of its persistence, bioaccumulation, and
67 biomagnification in food chains (Behrooz et al., 2009). Therefore, during the 1970-ies the
68 production of DDT was banned in most Western countries and the usage of DDT as insecticides
69 was restricted in many developing countries (Yang et al., 2008). With regard to its persistence, the
70 half-life of DDT in nature has been estimated to be between 4 to 30 years (Tomlin, 2005). Even
71 though it was banned three decades ago, its residues and metabolites can still be detected in the
72 environment (Guo et al., 2009).

73 Several techniques or approaches have been developed for remediation of DDT, including
74 biodegradation treatments (Li et al., 2010), soil excavation and incineration or thermal degradation
75 at high temperatures (Rodante et al., 1992), washing soil with surfactants (Smith et al., 2004), and
76 advanced oxidation technologies, such as photochemical reactions using nano-sized TiO₂/UV (Lin
77 and Lin, 2007) and metal-catalyzed reactions (Pd/C catalysts) (Zinovyev et al., 2005). Both the
78 latter have been shown to be effective for DDT degradation, but they are also expensive treatment
79 methods. As a powerful, inexpensive and environmentally friendly reducing agent, zero-valent iron
80 has been used for DDT degradation in water and soil (Sayles et al., 1997; Eggen and Majcherczyk,
81 2006; Yang et al., 2010).

82 Recently, nanotechnology has offered a new generation of environmental remediation
83 technologies that can provide cost effective solutions to some of the most challenging
84 environmental cleanup problems. Nanoscale zero-valent iron (nZVI) has smaller particle size than
85 traditional ZVI and a very high reactivity, well suited for injection and transport in porous media.

86 nZVI has been tested for remediation of several contaminant groups, including chlorinated organic
87 contaminants (Wang and Zhang, 1997; Karn et al., 2009). Most published studies targeting
88 dechlorination of chlorinated organics have however used bimetallic nZVI, containing small
89 amounts of Palladium (Pd) or Nickel (Ni). These include organochlorine pesticides (Zhang, 2003),
90 polychlorinated biphenyls (PCBs), trichloroethylene (TCE) (Schrick et al., 2002),
91 pentachlorophenol (PCP) (Zhang and Elliott, 2006), atrazine (Zhang et al., 2011) and DDT
92 degradation in water (Tian et al., 2009). Even though bimetallic nZVI is effective for pollutant
93 degradation, it has drawbacks regarding cost efficiency and environmental compatibility due to
94 spreading of other metals than Fe (Mueller et al., 2012).

95 So far, *in situ* nZVI applications have mainly targeted contaminants in aqueous systems and
96 groundwater. For instance, most of field applications carried out in Europe, and about 80% of the
97 sites treated in USA until now, have targeted contaminated groundwater only (Karn et al., 2009;
98 Mueller et al., 2012). Few studies have reported on the use of nZVI in soil. Nevertheless, the
99 application of nZVI in soil is important because the residues of the contaminants mostly remain
100 within the soils above the groundwater (Reddy, 2010). Usually, nZVI reactivity and degradation
101 efficiency is less in soils than in aqueous solutions due to limited desorption or solubilization of
102 the contaminants in soil (Wang and Zhang, 1997; Varanasi et al., 2007). Further, degradation of
103 chlorinated compounds which have aged in soil for many years is far slower than for recently
104 polluted and spiked soil due to lower bioavailability of the former.

105 Nanoecotoxicology is a recent branch within toxicology which has focused on measuring
106 toxicity of nanoparticles entering in contact with organisms like plants, bacteria, fish and
107 invertebrates (Handy et al., 2008). Nanoscale ZVI is considered the single largest source of
108 engineered nanoparticles entering the environment (Nowack and Bucheli, 2007). Further, the same
109 properties which make nZVI potentially useful for environmental remediation, such as its small

110 size and high reactivity, may also make it potentially harmful to living organisms (Sevcu et al.,
111 2011; Crane and Scott, 2012). Yet, its ecotoxicity has evoked little research interest until recently.
112 A few studies have been conducted using terrestrial species (earthworms, microorganisms and
113 plants) (Sevcu et al., 2011; El-Temsah and Joner, 2012a), but the lack of ecotoxicological data and
114 unknown potential effects of nZVI on organisms and the environment is currently hampering the
115 use of the nZVI technology in Europe.

116 The aims of this study were a) to test the efficiency of nZVI on DDT degradation in a spiked
117 sandy loam soil compared to that in a historically contaminated soil and b) to test the toxicity of
118 nZVI in DDT-containing soil on ostracods and collembola. The selected test organisms were
119 chosen because they represent key organism groups in their respective environment. [collembola
120 are among the most abundant soil arthropods, they feed on soil microorganisms (Crouau et al.,
121 1999), they play an important role in soil organic matter degradation, and they even contribute to
122 remediation processes in soil. Ostracods are crustaceans that feed on settled organic materials either
123 as particles or as larger fragments (Baun et al., 2008), and which are considered one of the
124 important food sources for fish larvae].

125

126 **2. Materials and methods**

127 **2.1. Nanosized zero-valent iron preparation**

128 Nanosized zero-valent iron was prepared using a modified borohydride method according
129 to He et al. (2010). Briefly, nZVI was prepared by dissolving 50 g of $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ in 450 mL water
130 immediately before use and mixing with 450 mL of an aqueous solution of 1 % carboxymethyl
131 cellulose (CMC; non-toxic according to Chen et al. 2011). ZVI nanoparticles were formed by
132 reducing the ferrous sulphate using a 1.9 M borohydride solution (30 mL, introduced at 5 ml min⁻¹
133 ¹) and adjust the volume to 1L. The size of the resulting nZVI particles, measured using high

134 resolution transmission electron microscopy (JEM-2011; Jeol, Japan, operating at 200 keV), was
135 in the range 20–100 nm. The hydrodynamic diameter and zeta potential, measured by dynamic
136 light scattering (DLS) and phase analysis light scattering (PALS), respectively, using a Malvern
137 Zetasizer Nano ZS (Malvern Instruments Ltd., England) showed particle size between 178 and 424
138 nm and a zeta potential of -42.8 mV (previously described in El-Temsah and Joner, 2012a).

139 **2.2. DDT degradation experiment**

140 Fifty grams (dry weight) of a sandy loam soil (sieved <2 mm, sand 85 %, silt 11%, clay 4
141 %, organic matter 1.1% and pH_{water} 5.8) amended with 20 mg DDT kg^{-1} (PS-74, Chem Service
142 Inc., West Chester, PA, USA; containing 18 % o,p' DDT and 77 % p,p' DDT) was incubated with
143 100 ml of an nZVI suspension at 1 or 10 g kg^{-1} soil in 250 ml glass bottles at room temperature.
144 Controls without nZVI were included, and all treatments prepared in triplicate. Bottles were shaken
145 at 175 rpm on a horizontal shaker for 7 or 30 days, and during the incubation bottles were opened
146 for 1 minute once per day for aeration. At the end of each shaking period, the slurry samples were
147 separated into a solid and a liquid phase by centrifugation ($3622 \times g$) and the water phase filtered
148 using Whatman No. 5 paper filters. Samples of the solid and water phases were taken for DDT
149 analysis. The same procedure was used with a historically polluted soil rich in organic matter
150 (organic silty clay soil, clay 11 %, silt 49 %, sand 40 %, organic matter 8.8 %, pH_{water} 5.2)
151 containing 23.1 mg DDT kg^{-1} (sampled at a fruit farm at the west coast of Norway, approx. 50
152 years since contamination).

153 **2.3. DDT extraction and analysis**

154 Soil samples (3 g dry weight) were extracted in 50 ml glass bottles by using 10 ml hexane
155 and 10 ml acetone. The suspension was shaken at 175 rpm on a horizontal shaker for 1h (adapted
156 from Tian et al 2009). After shaking, 15 ml of deionized water were added and the emulsion shaken
157 for another 5 min. The emulsion was centrifuged ($671 \times g$, 5 min) to obtain phase separation, and

158 1.5 ml of the hexane phase was transferred to GC glass vials and analyzed by GC-MS (GC 6890N
159 and MS 5973N, Agilent, USA) using a 0.2 mm x 50 m (0.25 μ m film thickness) Varian CP7482
160 capillary column and 1 ml/min He as carrier gas. A 2 μ l sample was injected into a split/split less
161 injector (Agilent) at an initial temperature of oven 80 °C, injector temperature of 250°C and column
162 temperature of 325°C. Partially due to difficulties in separating isomers, and partly because
163 differences in concentrations of the separated isomers were never significantly different between
164 treatments, we present the combined isomers only; the o,p'-DDT+ p',p' DDT as DDT, the o,p'-
165 DDD+ p',p' DDD as DDD (1,1-dichloro-2,2-bis(p-chlorophenyl)ethane), and the o,p'-DDE+ p',p'
166 DDE as DDE ([1,1-dichloro-2,2-bis(p-chlorophenyl)ethylene). The recovery of total DDT from
167 soil was 93.6 \pm 4.8 %. DDD and DDE were quantified as DDT metabolites, but not subjected to
168 individual toxicity measurements, as their toxicity are inherently lower than that of DDT
169 (Richardson and Gangolli, 1994)

170 **2.4. Toxicity to ostracods**

171 A 6-day direct contact ostracod (*Hetrocypris. incongruens*) toxicity test was performed
172 according to the standard operational procedure of the Ostracodtoxkit F (Micro-BioTests,
173 Nazareth, Belgium). The test was carried out in twenty-four well trays. Three replicates were
174 prepared from each treatment for both soil and filtrated aqueous samples taken immediately after
175 separation. For soil samples, 0.4 g dry weight soil was added to each well and mixed gently with 1
176 ml of medium-hard EPA water (Ostracodtoxkit F) and left for the soil to settle. One ml of algae
177 suspension and 5 neonate ostracods were added to each well. The same procedure was made with
178 water samples where 0.2 ml sample per well was added instead of soil. The test plate was sealed
179 with parafilm, covered by a lid and incubated in darkness at 25 °C for 6 d. The contents of each
180 well were then microsieved to retain the ostracods, which were transferred to small Petri dishes.
181 Mortality and growth of the surviving ostracods were determined. The measurement of length was

182 carried out by using of a micrometric strip placed under the well. The growth inhibition test was
183 considered valid when the mean death of ostracods concurrently exposed to a reference sediment
184 was less than 30 %. Growth inhibition (GI) of *H. incongruens* was calculated as: $GI = 100 -$
185 $(A/B * 100)$, where A is growth increment of ostracods in the reference sediment, B is increment of
186 the ostracods in the treatment.

187 **2.5. Collembolan tests**

188 Collembola (*Folsomia candida*) were synchronized to 11 to 13 days of age according a
189 standard protocol (OECD, 2008), and ten collembola were exposed to treated and untreated soil
190 immediately after the 7 or 30 d incubations. Approx. 29 g soil was transferred into plastic cylinders
191 measuring 5.5 cm height and 4 cm inner diameter with small a space in the plastic lids for
192 collembolan respiration. Dried baker's yeast (15 mg) was spread onto the soil surface to serve as
193 food source. The tests were carried out at about 50 % of the soil's water holding capacity. The test
194 containers were kept at 20 °C with a light–dark cycle of 16:8 h at 400–800 lux. The reproduction
195 of the test species took 4 weeks to complete, and at the end of the incubation period, adults and
196 juveniles were counted after flotation (Skovlund et al., 2006).

197 **2.6. Fe (II) extract from soil**

198 Immediately after the 7 and 30 d incubation periods, approx. 0.5 g of soil was transferred
199 to 5 ml of 0.5 M HCl in a glass vial and mixed gently for 30 s. After 1 h at room temperature, a 0.1
200 ml sample of the extract was added to 5 ml of ferrozine (1 g l⁻¹) in 50 mM HEPES (N-2-
201 hydroxyethylpiperazine-N'-2-ethanesulfonic acid) buffered to pH 7 using NaOH. The amount of
202 Fe(II) was determined spectrophotometrically by measuring the absorbance of the supernatant at
203 562 nm. Fe(II) was not oxidized and FeIII was not reduced during the extraction. Another sample
204 of the soil was extracted by the same procedure as that described above with the exception that the
205 extractant was 5 ml of 0.25 M hydroxylamine hydrochloride in 0.25 M HCl. Under acidic

206 conditions, hydroxylamine reduces Fe(III) to Fe(II). The amount of hydroxylamine-reducible
207 Fe(III) was calculated as the difference between the Fe(II) measured in the hydroxylamine and
208 HCl extractions (Lovley and Phillips, 1986).

209 Standard toxicity curves were established for ostracods to determine LC₅₀ and EC₅₀ by
210 using a wide range of concentrations of Fe(II) (FeSO₄), Fe(III) (FeCl₃), DDT and nZVI. Ostracods
211 were exposed to DDT added to soil and Fe(II), Fe(III) and nZVI were tested in water. pH and Eh
212 for all water samples were determined and Fe(II) and Fe(III) determined in treatment samples and
213 in nZVI suspensions.

214 For statistical analysis, one way analysis of variance (ANOVA) followed by Student t-tests
215 were used for comparisons of toxic effects between nZVI treatments and controls. Probit regression
216 analysis (EPA Probit analysis, v. 1.5) was used to determine EC₅₀ and LC₅₀ values (50 % effect or
217 lethal concentration) using % mortality or growth inhibition at the different exposure
218 concentrations.

219

220 **3. Results**

221 **3.1. DDT degradation**

222 DDT degradation in soil with 1 or 10 g kg⁻¹ of nZVI after incubation for 7 or 30 days
223 is shown in Table 1. Significant amounts of DDT were degraded in the nZVI treatments compared
224 with controls without nZVI. Generally, the lowest concentration of DDT was found in treatments
225 with the lowest concentration of added nZVI for the first incubation period of 7 days. Adding 1g
226 nZVI kg⁻¹ soil thus resulted in 56 % degradation of DDT as the sum of DDT and its degradation
227 products after 7 d, while it was only 5 % with 10 g nZVI kg⁻¹ soil (not significantly different from
228 the control). Longer incubation time (30 days) led to continued DDT degradation at the highest

229 dose of nZVI, whereas the lowest dose of nZVI gave no additional degradation during the period
230 from 7 to 30 d. After 30 d, the concentrations of remaining DDT also differed significantly between
231 the 1g and the 10 g nZVI kg⁻¹ soil treatments, but the difference was smaller than after 7 days. The
232 nZVI treatment did not reduce DDE levels significantly after 7 d, whereas DDD levels were
233 enhanced in the 10 g nZVI kg⁻¹ at both incubation times compared to controls.

234 The effect of nZVI on DDT degradation in historically contaminated soil is shown in Table
235 1. The untreated controls contained 23.1 mg kg⁻¹ of total DDT after 7 d incubation, whereas the
236 soils treated with 1 and 10 g nZVI kg⁻¹ soil contained 17.6 and 16.7 mg kg⁻¹ of total DDT,
237 respectively, corresponding to 24 % and 28 % reduction of total DDT. No significant changes in
238 DDD or DDE were found for any of the treatments for the historically contaminated soil. The pH
239 did not change significantly in either of the soils due to nZVI additions.

240 **3.2.Toxicity effects on collembola**

241 Toxicity of nZVI at 1 and 10 g kg⁻¹ to collembola measured as mortality and production of
242 juveniles after 21 days test exposure are shown in the Table 2. There were significant ($p<0.05$)
243 negative effects on both adult and juvenile collembola compared with control soil without nZVI
244 and DDT. Toxicity effects of nZVI at 1 and 10 g kg⁻¹ concentration on collembola after 30 days
245 incubation was significantly lower than after 7 days incubation. Mortality reached 100 % when
246 adults were exposed to either nZVI concentration in soil after 7 days incubation, while after 30
247 days incubation toxicity was reduced and resulted in about 60 % and 80 % mortality for 1 and 10
248 g nZVI kg⁻¹, respectively. Additionally, pristine nZVI suspensions had highly negative effect on
249 adult and juvenile collembola. On the other hand, DDT alone had no significant negative effect on
250 adults, whereas no juveniles were observed in soil spiked with DDT alone. Generally, there were
251 no observed juveniles in either of the soils treated with the two nZVI concentrations, irrespective
252 of the length of the nZVI incubation period.

253 3.3. Toxicity effects on ostracods

254 The toxicity effects of nZVI residues in solid and aqueous soil fractions from treated soil
255 after 7 and 30 days incubation on ostracod mortality and growth inhibition are shown in Table 3
256 and 4. Toxicity effects of nZVI in soil and leachates decreased with increased incubation time.
257 There was a highly negative effect of nZVI in soil and its leachate on mortality and growth of
258 ostracods compared to controls consisting of untreated soil and a standard sediment included in the
259 Ostracod tox kit. The liquid phase of slurries from soil contaminated with DDT alone had negative
260 effects on ostracod mortality and growth inhibition which resulted in 27 % mortality and 56 %
261 growth inhibition after 7 days incubation. Similarly, 33 % mortality and 72 % growth inhibition
262 were observed in the solid phase of the DDT-treated soil without nZVI. Both nZVI treatments had
263 strong negative effects on ostracod mortality after 7 d incubation, and no ostracod survival was
264 observed with any concentrations in soil or its leachate (Table 3). After 30 days incubation with
265 nZVI there was less negative effects of nZVI on ostracod development. There was a similar weak
266 toxicity of 1 g nZVI kg⁻¹ soil and soil treated with DDT alone. Soil treated with 10 g nZVI kg⁻¹ soil
267 still caused 100 % mortality after 30 d. Leachates from DDT-spiked soil enhanced the ostracod
268 growth about 12 %. On the other hand, 100% mortality was observed in the soil and leachates of
269 10 g kg⁻¹ of nZVI treatment. EC₅₀ and LC₅₀ values for mortality and growth inhibition of ostracods
270 after exposure to serial dilutions of DDT, Fe(II) and nZVI in soil and water are presented in Table
271 5. EC₅₀ calculated from the data on growth inhibition was 11.5 mg kg⁻¹ soil for DDT, while in
272 water it was 36 and 19 mg l⁻¹ for nZVI and Fe(II), respectively. LC₅₀ was 77 and 13 mg l⁻¹ for nZVI
273 and Fe(II), respectively.

274 The effects of aged DDT-contaminated soil on ostracods are shown in Table 3. The liquid
275 phase from slurries of soil without nZVI had a low effect on mortality (7 %) and soil treated with
276 1 g kg⁻¹ of nZVI showed three times higher effects on ostracod mortality. Hundred percent

277 mortality was observed in the liquid phase of slurries from soil treated with 10 g kg⁻¹ of nZVI.
278 Liquid phase from soil without nZVI and soil treated with 1 g kg⁻¹ showed significant negative
279 effects on growth inhibition, which was 42 and 75 %, respectively. Untreated DDT contaminated
280 soil had negative effects on both growth and mortality of ostracods, with 33 % mortality and 51 %
281 growth inhibition. When treated with 1 or 10 g kg⁻¹ nZVI, 100 % mortality was observed.

282

283 **3.4. Effects of individual components on ostracods**

284 From dilution series experiments with iron, Fe(II) (as FeSO₄) showed negative effects on
285 ostracods at higher concentrations. Eight concentrations of Fe(II) from 0.1 to 100 µg Fe mL⁻¹ water
286 were used. No effects on growth inhibition or mortality were observed at lower concentrations (0.1,
287 0.5 and 1 µg mL⁻¹). Significant negative effects on ostracod development were observed at 5 and
288 10 µg Fe mL⁻¹. About 45 % growth inhibition and 40 % mortality was observed at 20 µg Fe mL⁻¹.
289 Increasing the concentration to 50 µg Fe(II) mL⁻¹ resulted in 100 % mortality. pH of all samples
290 was between 6.5 and 7.

291 Freshly prepared nZVI suspensions were used for ostracod test at 4 concentrations (10, 100,
292 500, and 1000 µg Fe mL⁻¹ water). There was a weak negative effect at 10 µg Fe mL⁻¹, whereas 100
293 µg Fe mL⁻¹ had a strong negative effect on ostracod development and mortality and caused 90 %
294 growth inhibition and 45 % mortality. For higher concentrations, 100% mortality was observed.

295 Effects of DDT on ostracods were studied using 10, 20 and 50 mg kg⁻¹ DDT added to soil.
296 DDT at 10 mg kg⁻¹ had a negative effect on growth of ostracods (50 % growth inhibition), but no
297 effects on mortality. At 20 mg DDT kg⁻¹, 64 % growth inhibition and 7 % mortality were observed.
298 When 50 mg DDT kg⁻¹ was tested, 100 % mortality of ostracods was observed. Growth inhibition
299 of ostracods was a more sensitive end point than mortality.

300

301 The concentrations of iron in aqueous and solid phase of spiked and aged DDT soil
302 suspension after 7 and 30 days incubation are shown in Table 5. Both concentrations of nZVI (1
303 and 10 g kg⁻¹) slightly increased Fe(II) concentrations (7.2 and 10 mg L⁻¹, respectively) in the
304 aqueous phase of treated DDT spiked soil compared to control (6.6 mg L⁻¹), while the addition of
305 10 g nZVI kg⁻¹ to spiked soil significantly increased Fe(II) and Fe(III) concentrations in the solid
306 phase (to 102 and 1042 mg kg⁻¹, respectively). In aged DDT-contaminated soil a similar minor
307 increase in Fe(II) and Fe(III) concentrations were seen in the aqueous phase after addition of 1 or
308 10 g nZVI kg⁻¹ compared to the control. A strong increase in Fe(II) and Fe(III) concentrations
309 (reaching 205 and 1471 mg kg⁻¹, respectively) was observed in the solid phase of soil treated with
310 10 g nZVI kg⁻¹ after 7 d incubation. While soil receiving 1 g nZVI kg⁻¹ had increased from 59 to
311 99 mg Fe(II) kg⁻¹ and from 242 to 348 mg Fe(III) kg⁻¹.

312

313 **4. Discussion**

314 *4.1. Effects of nZVI on DDT*

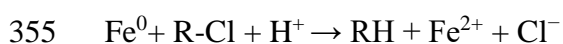
315 In this study, the potential of nano-sized zero-valent iron (nZVI) on degradation of DDT in
316 sandy loam and historically contaminated soil was investigated. Also, the toxicity effects of the
317 complex system (nZVI, DDT and DDT byproducts) on ostracods and collembola was examined.
318 The results showed that nZVI has a potential for degradation of DDT, even in historically
319 contaminated soil where DDT bioavailability for degradation is low. It has been documented that
320 nZVI can be used for remediation of both organic and inorganic pollutants in the aqueous phase,
321 and that it is particularly suited for *in-situ* remediation of contaminated soil and ground water
322 (Wang and Zhang, 1997; Joo and Zhao, 2008). However, very little is known about its efficiency
323 for treating chlorinated pollutants in topsoil or about effects on ecosystems and organisms in soil

324 and freshwater. DDT is recalcitrant and persistent in soils, and our control treatments showed no
325 significant changes in DDT concentrations even in spiked soil without nZVI additions during
326 incubated for up to 30 days. The DDT degradation in treated soils was thus clearly caused by nZVI.
327 The data also showed that after 7 d incubation, nZVI at 1 g kg⁻¹ reduced DDT levels faster than at
328 10 g kg⁻¹. This could be due to extensive and more complete oxidation and enhanced electron
329 release from nZVI which is required for reduction reactions of chlorinated compounds. On the
330 other hand, the degradation capacity with 1 g kg⁻¹ of nZVI did not change much from 7 to 30 d
331 incubation. This may be caused by the oxidation process and production of hydroxide substance
332 on the nZVI particles surface and a general depletion of reactive nZVI due to such oxidation. In
333 contrast, for the soil with aged DDT having a lower bioavailability in soil, adding either 1 or 10 g
334 nZVI kg⁻¹ resulted in similar rates of DDT degradation after 7 d incubation. Eggen and
335 Majcherczyk (2006) found that DDT degradation in aged sediment was difficult using macro-sized
336 zero-valent iron, even when they added high concentrations of ZVI (1.7 g ZVI g⁻¹ sediment for 10
337 and 40 weeks resulting in 64 and 93 % degradation of DDT, respectively). Adsorption of organic
338 matter such as humic acid onto the nZVI surface is known to decrease its activity due to
339 accumulation of humic acid on the active surface sites of the nZVI interface with water and soil
340 (Giasuddin et al., 2007).

341 DDE is known as the DDT metabolite that degrades slowest among DDE and DDD, and it
342 is therefore often recovered in higher concentrations during degradation studies (Sayles et al.,
343 1997). Thus, DDE was recovered in higher concentrations than DDD in both soils after incubation
344 for 7 days. The control treatment also contained significant amounts of DDE, indicating that it was
345 already present in the DDT added to soil at the start of the experiment. DDE is an aerobic
346 dehalogenation product of DDT, and DDE has been reported as difficult to decompose further
347 (Wang et al., 2006). The fact that no DDE accumulated still shows that nZVI lead to proper

348 dechlorination and that its efficiency towards DDE dechlorination is comparable to that towards
349 DDT.

350 Generally, the degradation efficiency of nZVI in spiked soil was higher than in historically
351 contaminated soil, and degradation of DDT depended on both nZVI concentration and time. This
352 may be due to complex relationships between DDT and soil, such as desorption, solubilization and
353 dissolution of DDT. Dombek et al. (2001) showed in their study that the dechlorination reaction
354 between iron and organochlorine compounds in water solution was as follows:



356 This is consistent with reduction reactions occurring by electron transfer at the iron surface to
357 degrade these chlorinated compounds. Usually the reaction between pollutants and nZVI is carried
358 out in the water phase; therefore the DDT solubility in water has significant impact on the
359 efficiency of the degradation. Water solubility of DDT, DDD and DDE are $25 \mu\text{g L}^{-1}$, $120 \mu\text{g L}^{-1}$
360 and $90 \mu\text{g L}^{-1}$ (25°C), respectively (ATSDR, 2012). The low solubility of DDT makes transfer of
361 DDT into an aqueous solution, where it can react with iron, a rate limiting factor. This may explain
362 why spiked DDT was easier to reduce than aged DDT when either of these reacted with limited
363 amounts of iron (low dose of nZVI) during a short incubation period.

364 4.2. Toxicity effects on collembola

365 Nanosized ZVI is considered the largest stream of engineered nanoparticles entering the
366 environment, and existing ecotoxicological data are not conclusive. NZVI application mainly
367 targets treatment of pollutants in the subsoil under saturated conditions (Mueller et al., 2012).
368 Supposedly, the beneficial effects of nZVI degrading pollutants should largely outweigh any
369 potential harmful effects (Karn et al., 2009; Grieger et al., 2010). However, mobility aspects are
370 not resolved, and there is still a lack of knowledge in most soil remediation approaches regarding
371 the impact of soil treatment processes on soil biota.

372 Collembola, or springtails, are one of the most abundant groups of soil arthropods.
373 Ecotoxicity tests using the collembola *Folsomia candida*, which assess its population development,
374 has been standardized for use in Europe (ISO 11267), and was originally designed to test the effects
375 of individual chemicals on a soil arthropod (Crouau et al., 2002). There is a risk that when nZVI is
376 applied to subsurface soil, it will come in contact with organisms of terrestrial ecosystems. It is
377 therefore useful to know if the ecotoxicity of nZVI can be measured by standardized tests based on
378 for example soil arthropods. The results of the present study indicate that the collembola test can
379 be applied also for the evaluation of the toxicity of nZVI in soil, even in combination with a
380 complex insecticide like DDT.

381 Our results showed severe toxic effects on collembola, particularly short time (7 days) after
382 addition to soil with a high mortality of adults at both nZVI application rates. After 30 days, adults
383 could survive in treated soils, while no juveniles were produced. Thus, collembolan reproduction
384 was a more sensitive parameter than survival, and supplies more detailed information on toxicity,
385 as recognized for other environmental toxicants (Krogh and Petersen, 1995; Crouau et al., 2002).
386 DDT alone showed negative effects only on reproduction, but contrary to nZVI, the toxicity of
387 DDT in untreated soil is not likely to decline nearly as fast.

388 Iron toxicity studies have primarily focused on Fe(II) and its oxides, and little is known
389 about the toxicity specific to nZVI or macroscale ZVI. However, ZVI produces Fe(II) and iron
390 oxides through oxidation, and nZVI can produce free radicals which are highly reactive and cause
391 oxidative stress (Li et al., 2009). This could be one of the mechanisms behind the toxic effects of
392 nZVI on soil organisms. Cullen et al. (2011) assessed the effect of micro- and nano-sized ZVI at
393 10 g kg⁻¹ on soil enzymes under aerobic conditions, and did not observe any significant effects on
394 soil enzyme activity. Fajardo et al. (2012) assessed the impact of nZVI at 34 g kg⁻¹ on soil microbial
395 community structure and functionality using a molecular approach. They did however observe little

396 negative effects on microbial cellular viability and biological activity in soil. Recently, we reported
397 that nZVI has a negative impact on plants and earthworms in soil and water (El-Temsah and Joner,
398 2012b), which support the present results.

399 4.3. Toxicity effects on ostracods

400 The majority of the ecotoxicological studies of engineered nanoparticles in water have used
401 the crustacean *Daphnia magna* as test organism (Baun et al., 2008). A few studies have also
402 examined the impact of nZVI on aquatic organisms such as zebra fish (Li et al., 2009; Chen et al.,
403 2011) and river water bacteria (Barnes et al., 2010). To the best of our knowledge, there is no data
404 in the literature on the effects of nZVI or DDT on the ostracod *H. incongruens* as fresh water
405 organism and sediment dweller. The ostracod assay (mortality and growth inhibition) is both rapid,
406 sensitive, relatively inexpensive, and demand small sample volumes compared to e.g. collembolan
407 tests. By using soil as if it was a sediment, the ostracod test makes it possible to measure the
408 combined toxicity of DDT and nZVI and simultaneously evaluate both acute and chronic endpoints
409 (mortality and growth inhibition). It thereby gives a reflection of the toxicity of the whole sample
410 taking into account bioavailability of the contaminants present in the matrix. Ostracods (*H.*
411 *incongruens*) have previously been used successfully as test organism for soil toxicity assessment
412 in this way (Joner et al., 2004; Manzo et al., 2011). Manzo et al. (2011) has used *H. incongruens*
413 to evaluate the toxicity effects of ZnO nanoparticles in soil, and the results indicated that ostracods
414 were the most sensitive organisms to ZnO nanoparticles in soil. In our study, ostracods were also
415 very sensitive to nZVI in water and soil, which was probably due to low oxygen levels resulting
416 from initial oxidation of nZVI. This assumption is supported by the fact that mortality declined
417 strongly with prolonged slurry incubation prior to the test which allowed oxygenation, and that
418 mortality was higher in treatments receiving higher doses of nZVI where more oxygen would be
419 needed to reach aerobic conditions. EC50 thresholds established for the individual components

420 showed that nZVI is less toxic than either DDT or Fe(II). The negative effects of nZVI on ostracod
421 mortality could thus also be indirect, due to release of Fe(II). This is in agreement with the findings
422 of Chen et al. (2011) who studied the toxic effects of nZVI and its oxidation products in medaka
423 fish larvae. They concluded that nZVI causes hypoxia due to O₂ consumption, and that nZVI
424 released excess Fe(II) which caused toxicity due to production of reactive oxygen species (ROS).
425 Indirect effects on food depletion has also been used to explain nanoparticle toxicity (Manzo et al.,
426 2011), but this seem less relevant in the case of ostracods under the test conditions used here.

427 **5. Conclusion**

428 In this study, nanosized zero-valent iron was used to degrade DDT in spiked and aged
429 contaminated soil. Toxicity of aqueous and solid phases of soil slurries after incubation with nZVI
430 were measured on collembola and ostracods. The results showed that the degradation rates of DDT
431 in spiked soil were higher than in historically contaminated soil. nZVI had severe effects on
432 collembola and ostracods, while DDT had weaker negative effects on the reproduction of
433 collembola and development of ostracods. We also observed that increasing the incubation time or
434 reaction period alleviate the toxicity effects of nZVI on collembola and ostracods. The addition of
435 nZVI increased the concentration of Fe(II) and Fe(III) after incubation in soil, and particularly
436 Fe(II) was more toxic to ostracods than nZVI. Further studies are needed to optimize the use of
437 nZVI in different types of soils to ensure high degradation of DDT, and at the same time take into
438 account the extent and duration of negative effects on soil biota.

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442 analyses.

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447 **6. References**

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578 Table 1. Residual concentrations of DDT in sandy loam soil spiked with 20 mg DDT kg⁻¹ and soil
 579 contaminated with DDT >50 years ago incubated with 1 or 10 g nZVI kg⁻¹ for 7 and 30 days in soil slurries.
 580 Means associated with the same letter in each column are not significantly different (one way ANOVA,
 581 p<0.05, n=3)

7 days incubation of spiked soil				
	o+p DDT (mg kg ⁻¹)	DDD (mg kg ⁻¹)	DDE (mg kg ⁻¹)	Total DDT (mg kg ⁻¹)
Control	15.8 a	0.2 c	2.3 a	18.3 a
1 g nZVI	5.1 d	0.2 c	2.8 a	8.1 c
10 g nZVI	15.2 a	0.4 b	1.9 a	17.4 a
30 days incubation of spiked soil				
Control	16.1 a	0.4 b	0.04 c	17 a
1 g nZVI	7.6 c	0.2 c	0.02 c	7.9 c
10 g nZVI	10 b	0.9 a	0.04 c	10.9 b
7 days incubation of aged DDT-contaminated soil				
Control	16.3 x	0.2 x	6.6 x	23.1 x
1 g nZVI	11.3 y	0.2 x	6.0 x	17.6 y
10 g nZVI	10.7 y	0.3 x	5.7 x	16.7 y

582

583

584 Table 2. Effects of DDT and/or nZVI on collembolan mortality (adult survival) and reproduction (juvenile
 585 numbers) in spiked soil after 7 and 30 days of incubations with 1 or 10 g nZVI kg⁻¹.

	7 days		30 days	
	Adults	Juveniles	Adults	Juveniles
Soil without DDT (control)	7±1.4	68±12	8± 0.9	138±13
Soil with DDT	9±0.7	0	8±1.2	3±0.7
Soil with 1g nZVI, no DDT	0	0	4±0	0
Soil with 10 g nZVI, no DDT	0	0	2±0.7	0
Soil with 1g nZVI and DDT	0	0	4±0.7	0
Soil with 10g nZVI and DDT	0	0	2±0.7	0

586

587

588 Table 3. Effects of exposure to the aqueous phase or solid phase of soil suspension made of spiked or
 589 historically DDT-contaminated soil treated with nZVI for 7 or 30 d on ostracod mortality and growth
 590 inhibition (GI), (n=3).

591

	Water phase		Solid phase	
	Mortality %	GI %	Mortality %	GI %
Spiked soil after 7 days slurry incubation				
Soil without DDT	0	0	7 c	39 c
Soil with DDT	27 c	56 b	33 b	72 b
Soil with 1g nZVI kg ⁻¹	67 b	**	100 a	100 a
Soil with 10 g nZVI kg ⁻¹	100 a	100 a	100 a	100 a
Soil with 1g nZVI kg ⁻¹ and DDT	100 a	100 a	100 a	100 a
Soil with 10 g nZVI kg ⁻¹ and DDT	100 a	100 a	100 a	100 a
Spiked soil after 30 days slurry incubation				
Soil without DDT	7 c	17 c	7 c	7 b
Soil with DDT	27 b	-12 c	27 b	28 c
Soil with 1g nZVI	27 b	58 b	27 b	27 c
Soil with 10g nZVI	100 a	100 a	100 a	100 a
Soil with 1g nZVI kg ⁻¹ and DDT	20 b	60 b	20 b	57 b
Soil 10g nZVI kg ⁻¹ and DDT	100 a	100 a	100 a	100 a
Aged DDT-contaminated soil after 7 days slurry incubation				
Soil without nZVI	7 c	42 c	33 b	51 b
Soil with 1 g nZVI kg ⁻¹	23 b	75 b	100 a	100 a
Soil with 10 g nZVI kg ⁻¹	100 a	100 a	100 a	100 a

592 ** High mortality invalid for inhibition measurement

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596

597 Table 4. EC₅₀ and LC₅₀-values of components from nZVI contributing to ostracod toxicity.

598

	EC ₅₀	LC ₅₀
DDT (mg kg ⁻¹)	11.5	-
nZVI (mg L ⁻¹)	36	77
Fe ²⁺ (mg L ⁻¹)	19	13

599

600
 601 Table 5. Iron in the aqueous and solid phases of slurries from spiked or historically DDT-contaminated soils
 602 after 7 and 30 days incubation with nZVI (mean \pm SD, n=3).

	Water phase		Solid phase	
	Fe ^{II} (mg L ⁻¹)	Fe ^{III} (mg L ⁻¹)	Fe ^{II} (mg kg ⁻¹)	Fe ^{III} (mg kg ⁻¹)
7 days				
Control	6.6 \pm 0.5	0.33 \pm 0.6	31 \pm 2.5	82 \pm 30
1 g nZVI kg ⁻¹	7.2 \pm 0.1	1.4 \pm 0.2	34 \pm 4.4	136 \pm 39
10 g nZVI kg ⁻¹	10 \pm 0.2	2.3 \pm 0.1	102 \pm 7	1042 \pm 61
30 days				
Control	4.9 \pm 0.2	0,18 \pm 0.3	37 \pm 2	175 \pm 28
1 g nZVI kg ⁻¹	6.1 \pm 0.1	3.1 \pm 0.7	45 \pm 8	193 \pm 46
10 g nZVI kg ⁻¹	7 \pm 0.8	2.8 \pm 0.03	148 \pm 24	1102 \pm 72
DDT-contaminated soil after 7 days				
Control	6.8 \pm 0.3	0.26 \pm 0.2	59 \pm 1.3	242 \pm 15
1 g nZVI kg ⁻¹	6.9 \pm 0.1	0.62 \pm 0.15	99 \pm 2	348 \pm 5
10 g nZVI kg ⁻¹	8.1 \pm 0.4	2.2 \pm 0.2	205 \pm 11	1471 \pm 244

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 604