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Mercury accumulation from food decreases collembolans' growth

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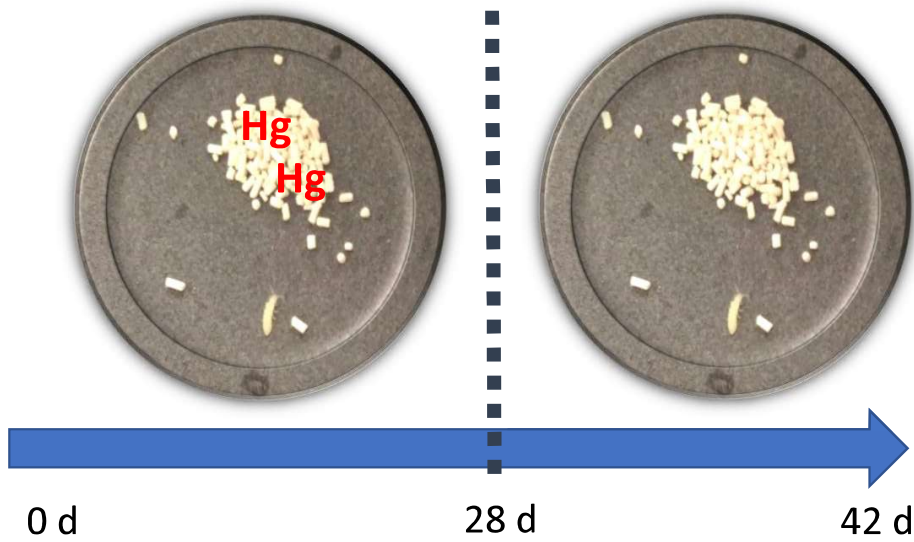
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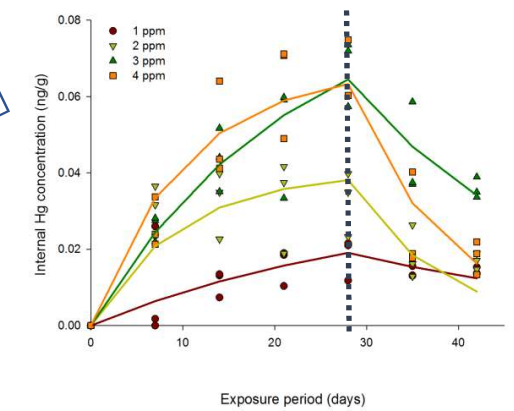
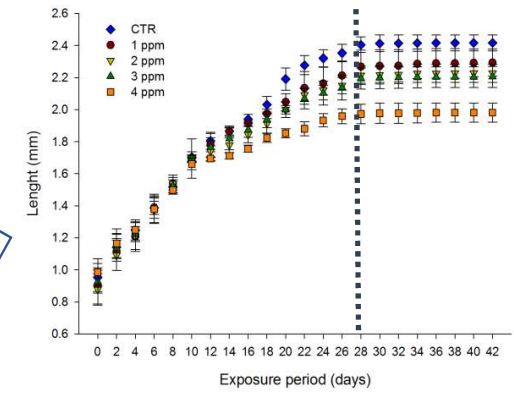
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*Graphical Abstract



growth

bioaccumulation



Highlights

- Exposure to mercury through food reduces the growth of collembolans;
- Mercury uptake in the organisms through a dietary exposure increases Hg inside organisms;
- A possible maximum sub lethal concentration of 0.07 ng Hg/g for collembolans was found;
- Food avoidance may occur in the presence of contaminated food at high Hg levels;
- At higher mercury levels, collembolans presented a higher elimination rate.

1 **Mercury accumulation from food decreases collembolans' growth**

2

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25 **Abstract**

26 In the terrestrial environment mercury (Hg) is redistributed and transformed into different
27 inorganic and metal - organic species that are deposited into soils. In the present study, the
28 effects of contaminated food with environmentally relevant concentrations of Hg in the form
29 of HgCl₂ on the soil-dwelling collembolan *Folsomia candida* were assessed. Changes in growth
30 rate and Hg bioaccumulation levels were observed at different concentrations of Hg in food
31 complementing data on the effects of Hg on reproduction and survival using standardized
32 protocols. Collembolan growth was recorded every two days, and a Von Bertalanffy's growth
33 curve was derived along with the growth rate. Collembolan growth was dependent on the Hg
34 food concentration. Also, the final length of animals was affected by the presence of Hg in
35 food, with differences in all treatments comparing to non-exposed organisms. Toxicokinetic
36 patterns from Hg exposure in food at different concentrations were not significantly different
37 from each other in the uptake, but differences were found in the depuration phase. Combining
38 the two approaches, collembolans seems to invest their effort in the depuration process,
39 neglecting their efforts in other vital processes, such as growth. Also, metal contaminated food
40 avoidance possibly occurred, a behavior already reported in the literature, thus decreasing
41 their feeding and contaminant intake. Therefore, growth tests can act an important asset to fill
42 the gaps of bioaccumulation tests and reproductive assays, towards a mechanistic
43 understanding. Changes in growth rate, even at low and environmentally relevant
44 concentrations, could be a warning signal when occurring in species with key roles in
45 ecosystems. Also, this study highlights the importance of these complementary tests for a
46 better and complete approach to risk assessment studies.

47

48 Key-words: Mercury, contaminated food, bioaccumulation, Von Bertalanffy's growth curve

49

50 **Capsule**

51 This study is a step forward on the understanding of mercury exposure effects to soil
52 organisms, using growth and bioaccumulation in time as endpoints.

53

54 **1 - Introduction**

55 Mercury (Hg) pollution is a worldwide problem, posing a serious threat to ecosystems and
56 consequently to humans, due to its ability to biomagnify along food chains (Boening, 2000).
57 Being a class B metal (Nieboer and Richardson, 1980), Hg has high affinity for reduced sulphur
58 atoms such as those from proteins and peptides containing thiol, which could lead to a
59 disruption of the tertiary structure of proteins, necessary for optimal function and condition of
60 organisms (Valko et al., 2005). Furthermore, Hg can also affect organisms at a cellular level,
61 depleting cellular antioxidation systems, which can lead to the production of reactive oxygen
62 species inducing oxidative stress damage - (e.g., peroxidation of membrane lipids) - (Stohs and
63 Bagchi, 1995). Since Hg is a widely-distributed contaminant, it is transported by air and
64 deposited in areas far from emission sources (Pacyna et al., 2009), on the top-soil layer, and a
65 consequent redistribution and further uptake by vegetation may occur (Miller et al., 2005).
66 Besides that, Hg can be retained in soils for a period of 500 to 1000 years (Pendias et al., 2011),
67 and suffer transformations into organic forms (most commonly methylmercury), through
68 microbial activity, thus becoming more toxic to organisms (Wang et al., 2003). Organic and
69 inorganic forms of Hg will eventually enter food webs and bioaccumulate and biomagnify in
70 soil invertebrates (Pedrini-Martha et al., 2012; Wiener et al., 2006).

71 Despite the growing concern about the potential adverse effects of elevated mercury
72 concentrations in the environment, the toxicity data available for soil invertebrates is scarce.
73 Some examples are studies in earthworms (Abbasi and Soni, 1983; Beyer et al., 1985; Buch et
74 al., 2017; Fischer and Koszorus, 1992; Mahbub, et al., 2017.), millipedes (Buch et al., 2018),

75 collembolans and enchytraeids (Buch et al., 2016; Lock and Janssen, 2001). Since those studies
76 were conducted in different substrates, some toxicity differences were observed, as soil
77 characteristics have a significant influence on the bioavailability and toxicity of metals. High pH
78 values and organic matter content could lead to an increase in chemical sorption and
79 consequently a decrease in bioavailability and toxicity (Lock and Janssen, 2001; Sandifer and
80 Hopkin, 1996). The interactions of Hg with temperature were also reported, reducing the cold
81 tolerance of earthworms and collembolans in the presence of Hg, with a dominant synergistic
82 pattern between these two stressors (Bindesbøl et al., 2009; Holmstrup et al., 2008)

83 Current recommended ecotoxicological tests with the springtail *Folsomia candida* provide
84 useful data regarding their reproduction output and survival, upon a 28-day exposure (ISO,
85 1999; ISO, 2014). Xenobiotics exposure to collembola is mainly carried out and assessed
86 through the interstitial soil pore water (Fountain and Hopkin, 2005), but the evaluation of
87 other routes of exposure are essential in risk assessment studies. Contaminated food exposure
88 can, therefore, complement the information provided by the already existing standardized
89 tests, facilitating, for example, the observation of growth, and providing additional information
90 on xenobiotics' effects on organisms (Roex et al., 2003). It is widely known that the primary
91 dietary component of most springtails is fungi (Van Straalen and Van Meerendonk, 1987), and
92 most fungal species accumulate metals (e.g., mercury) in their hyphae (Bengtsson et al., 1983),
93 being food a pertinent source of contaminants. Also, using growth as an endpoint is of utmost
94 importance for the equilibrium and sustainability of any species, as a feature closely related to
95 the organism reproduction (Fountain and Hopkin, 2001). In addition, the exposure of
96 xenobiotics through food can also provide new insight on toxicity effects to soil invertebrates
97 as their central role in the ecosystem is related to decomposition processes (Crouau and Moia,
98 2006). In an organism, growth is dependent upon food quality availability and the metabolism
99 efficiency, which can be affected by the presence of xenobiotics/contaminants, being,

100 therefore, a relevant endpoint to assess environmental hazard (Crommentuijn et al., 1993;
101 Folker-Hansen et al., 1996).

102 Considering the above mentioned, this study aimed at assessing the effects of Hg on *Folsomia*
103 *candida* growth and to infer if Hg bioaccumulation patterns were related to the observed
104 growth patterns. For that, organisms were exposed to Hg-contaminated food at different
105 relevant concentrations (1 to 4 ppm) for a 28 days' exposure period. During this period
106 collembolan growth was recorded every two days, and a Von Bertalanffy growth curve derived
107 along with the correspondent growth rate. At the same time, a similar experimental setup was
108 conducted to derive bioaccumulation patterns and toxicokinetic parameters (uptake and
109 elimination rates) also at different exposure concentrations. These two experiments
110 complemented each other to understand the mechanisms of toxicity that Hg can induce in *F.*
111 *candida*.

112

113 **2 – Material and methods**

114 **2.1 - Test species and test chemical**

115 Collembolans from the species *F. candida* were obtained from synchronized laboratory
116 cultures maintained at the University of Aveiro, Portugal. Cultures are kept in plastic boxes
117 lined with a mixture of plaster of Paris and activated charcoal in a ratio of 9:1 (ISO, 2014), in
118 the dark, under a constant temperature regime (20 ± 2 °C). Once a week, granulated dry yeast
119 was added as a food source (Fermipan, Setúbal, Portugal).

120 The same yeast provided in cultures was contaminated with mercury (II) chloride (HgCl_2 - CAS
121 no: 7487-94-7) purchased from Merck Millipore (99.5 % purity).

122

123 **2.2 – Growth test with *Folsomia candida***

124 The growth test was run in two phases: 1) an exposure phase, where organisms were exposed
125 to contaminated yeast for 28 days and 2) a post-exposure phase, where organisms were
126 moved to clean vials with clean food for 14 days. The test started with 10 days old
127 collembolans that were exposed to Hg via food, provided as contaminated yeast with 1, 2, 3
128 and 4 ppm of Hg. The chosen concentrations are environmental relevant since concentrations
129 ranging from 0.003 to 4.6 mg/kg of Hg have been reported in soils worldwide (E. Steinnes,
130 1997). Also, the mean content of Hg in soils was estimated to be 1.1 mg/kg, with an average
131 background content in different soil types ranging from 0.58 to 1.8 mg/kg (Xu et al., 2015).
132 Each replicate included one organism in a cylindrical plastic pot (30 cm³), on a moist substrate
133 of 9:1 (w/w) plaster of Paris:charcoal mixture. Granulated dry yeast was previously spiked with
134 the respective Hg solutions and dried by a lyophilization process. Afterward, it was supplied *ad*
135 *libitum* during the exposure phase and being replaced every week or in the presence of fungi.
136 A total of 10 replicates were used per control, Hg treatments and sampling time. Digital
137 photographs were taken every two days for 42 days, and the collembolan's length was
138 recorded from the end of the posterior abdominal segment to the anterior margin of the head,
139 using the Image J analysis software program (Schneider et al. 2012). Calibration run by using a
140 small millimeter paper placed near the measured organism.

141 Considering that collembolan finished the test with 52 days old, the appearance and number
142 of new-born juveniles were also recorded.

143

144 **2.3 – Bioaccumulation test with *Folsomia candida***

145 Running at the same time as the growth test, 18 replicates with 20 organisms each per Hg
146 concentration were also exposed to the same Hg concentrations following similarly the
147 procedures described in section 2.2. Briefly, organisms were exposed to contaminated food (1,
148 2, 3 and 4ppm Hg) for 28 days, followed by a non-contaminated yeast exposure for an extra 14

149 days' period. Organisms were sampled after 7, 14, 21, 28 days during the exposure period and
150 at day 35 and 42 (belonging to the post-exposure test), sacrificing 3 replicates in each sampling
151 time for bioaccumulation analysis (3 replicates used in each one of the 6 sampling times, a
152 total of 18 replicates initially performed). Samples were immediately frozen at -80 °C for
153 following chemical analysis. Before the measurements of total Hg inside the organisms were
154 dried by a lyophilization process.

155

156 **2.4 - Mercury analysis**

157 Total mercury (THg) concentration in yeast (food source) and collembolans was quantified by
158 atomic absorption spectrophotometry with thermal decomposition using an Advanced
159 Mercury Analyser (AMA) LECO 254 (Costley et al., 2000). The analytical procedure was adapted
160 as described by Cabecinhas et al. (2015) and Vieira et al. (2014): drying time of 60 s,
161 decomposition time of 150 s and waiting time of 45 s. The detection limit of the procedure was
162 0.02 ng Hg, and first and second working range of analysis (automatically switched) varied
163 between 0.05 – 40 ng Hg and 40-600 ng Hg, respectively.

164 Mercury determinations were all done in triplicate and blank run in parallel before and after
165 each sample to ensure the equipment was clean from any internal Hg contamination. All blank
166 measurements were run, at least, in triplicate until values obtained were consistently inferior
167 to the equipment detection limit (0.02 ng of Hg); additional blanks were, always performed at
168 the beginning and end of the day.

169 Accuracy and precision of the analytical method were assessed by the analysis of certified
170 reference materials (NRC TORT-2 Lobster hepatopancreas, and NRC DOLT-3 Dogfish liver) in
171 parallel with samples (Pereira et al., 2008). The average recovery of TORT-2 and DOLT-3 were
172 101 and 104%, respectively.

173

174 2.5 - Data analysis

175 Growth was described by the Von Bertalanffy expression:

$$176 L_t = L_\infty (1 - e^{-k(t-t_0)}) \quad (1)$$

177 Where L_t = length at time t (mm), L_∞ = the asymptotic length (mm), k = growth rate and t_0 =
178 the hypothetical negative time estimated from hatching date for an individual with length 0
179 (days) (Bertalanffy, 1960). The theoretical growth curve was fitted to the average length of
180 individuals per replicate and the parameters L_∞ and k were estimated. For comparisons
181 between the effects of Hg treatments and the negative control, slopes of the regression and
182 growth rate values (k) were compared by a generalized likelihood ratio test. Differences in
183 collembolans' length exposed to different Hg treatments, in different days of exposure were
184 analyzed using a one-way ANOVA, followed by post hoc Dunnett's test ($p < 0.05$), comparing to
185 the correspondent negative control (in time). To infer on the possible effects of Hg exposure
186 in collembolan's reproduction output, the day at which newborn juveniles were first observed
187 in each test box were compared between treatments and the control by using a Kruskal-Wallis
188 test followed by Dunn's *post hoc* test. Differences in the weight of collembolans at different
189 sampling times were derived by a one-way ANOVA, followed by post hoc Dunnett's test
190 ($\alpha=0.05$). A two-way ANOVA followed by a Tukey Test ($\alpha=0.05$) was also used to observe
191 statistical differences in the weight of collembolans in the different concentrations of Hg, at
192 different sampling times, also looking at the interaction between these two factors. The R-
193 squared (r^2) was obtained by dividing the sum of squares of each factor and of their interaction
194 by the total sums of squares of the two-way ANOVA assessing the percentage of variance
195 accounted for each factor in the ANOVAs.

196 For the bioaccumulation test, the toxicokinetics of Hg was described by applying a one-
197 compartment model. Uptake and elimination equations were fitted simultaneously. In the

198 model, body concentration of Hg was used, and the initial Hg concentration in the organisms
199 ($Q_{(0)}$) was considered to be zero.

200 For the uptake phase, the following equation was used:

201

$$202 \quad Q_{(t)} = \frac{k_1}{k_2} \cdot C_e \cdot (1 - e^{-k_2 \cdot t}) \quad (2)$$

203

204 Where $Q_{(t)}$ - Hg body concentrations at t days (ng/mg); k_1 – uptake rate constant ($\text{kg}_{\text{food}}/\text{kg}_{\text{org}}$
205 $\cdot \text{day}^{-1}$); k_2 – elimination rate constant (day^{-1}); C_e – Hg exposure concentration (ng/mg); and t -
206 time (days).

207 For the elimination phase, the metal body burden was described as:

208

$$209 \quad Q_{(t)} = \frac{k_1}{k_2} \cdot C_e \cdot (e^{-k_2 \cdot (t-t_c)} - e^{-k_2 \cdot t}) \quad (3)$$

210

211 where t_c - time the animals are transferred to clean medium (days).

212 Differences in k_1 and k_2 values between concentrations were tested by a Generalized
213 Likelihood Ratio test (GLR test). The assimilation rate (a) was calculated as $k_1 \cdot C_e$, and the
214 bioaccumulation factor (BAF) was calculated as k_1/k_2 . Also, $\text{BAF}_{\text{org/food}}$ was also calculated as the
215 Concentration of Hg in organisms/concentration of Hg in food although no equilibrium was
216 reached (plateau) during the uptake phase. Hg half-lives (DT_{50}) in the collembolans were
217 calculated as $\ln 2/k_2$. All calculations were performed using SPSS (version 20).

218

219 **3 – Results and discussion**

220 **3.1 Hg measurements in food**

221 The measured concentrations in yeast were 0.918, 2.03, 2.87 and 3.9 ppm of Hg for the
222 nominal concentrations chosen of 1, 2, 3, and 4 ppm, respectively. Since differences between
223 nominal and measured values were lower than 10%, nominal concentrations were used
224 throughout the manuscript. However, measured concentrations were used in the data
225 analysis.

226

227 **3.2 - Growth test with *Folsomia candida***

228 Collembolan growth was dependent on Hg concentration in food as observed in Figure 1. A
229 Von Bertalanffy growth curve was fitted to the data in all cases, and the values of L_{∞} and K
230 derived (Table 1). For each Hg treatment and control, two curves were settled, and data are
231 presented separately in Table 1-A for data for the exposure phase only (first 28 days of the
232 experiment), and in Table 1-B for the 42-day period, including both exposure and post-
233 exposure phases (28 days of exposure to Hg-contaminated food and 14 days of recovery with
234 non-contaminated food).

235

236 Table 1 - Parameters in von Bertalanffy growth curves. Estimates of mean maximum length (L_{∞}) and growth rate (k)
237 with asymptotic confidence intervals (C.I.) A- using growth measurements for 28 days of exposure (uptake); B –
238 using growth measurements for 42 days (28 days of exposure followed by 14 days of recovery).

239 A

[Hg] in food (ppm)	L_{∞}	C.I. 95%	K	C.I. 95%	r^2
0	3.619	3.460 – 3.778	0.029	0.027 – 0.031	0.981
1	3.05	2.906 – 3.194	0.036	0.033 – 0.039	0.958
2	2.917	2.795 – 3.038	0.037	0.035 – 0.040	0.965
3	2.776	2.673 – 2.880	0.041	0.038 – 0.044	0.960
4	2.238	2.191 – 2.285	0.057	0.055 – 0.060	0.960

240

241 B

[Hg] in food (ppm)	L_{∞}	C.I. 95%	K	C.I. 95%	r^2
0	2.862	2.799 – 2.924	0.042	0.04 – 0.044	0.961
1	2.614	2.551– 2.669	0.047	0.045 – 0.049	0.942
2	2.529	2.482 – 2.576	0.048	0.046 – 0.050	0.955
3	2.2458	2.416 – 2.500	0.051	0.049 – 0.054	0.949
4	2.106	2.083 – 2.130	0.065	0.063 – 0.068	0.949

242

243 As expected, collembolans reached their highest values of L_{∞} (maximum length) when
244 exposed to uncontaminated food decreasing its value with the increase of Hg food
245 concentration for the 28 days and 42 days fitting curves, as observed in Figure 1. Also, the
246 pattern was inverse for collembolans' growth rate (k) with values increasing with the increase
247 of Hg in food (28 and 42 days fitting curves). Curves fitted with the 28 days' dataset always
248 showed higher values for L_{∞} and lower values for k when compared to curves fitted with the
249 42 days' data set. In addition, the statistical analysis showed significant differences in the
250 growth rate within the same treatment when comparing the 28 and 42 days fitting curves
251 (Table SD1; $p < 0.05$). Using the same approach for collembolans exposed to for both 28 or 42
252 days curves, the growth rate was statistically different for all treatments comparing to the
253 control curve (Table SD1; $p < 0.05$). Due to those differences, an analysis per day of exposure of
254 collembolans' length was performed for the 28 days of exposure. As expected, statistical
255 differences were found regarding growth for the 4-ppm treatment already at day 12 and
256 onward, comparing to the control (Dunnett's test, $p < 0.05$). For collembolans exposed to
257 contaminated food with 3 and 2 ppm, statistical differences were found at day 14 and onward,
258 while for the 1ppm treatment only at day 20 and onward differences were attained, always
259 comparing with the control animals supplied with non-contaminated yeast (Dunnett's test,
260 $p < 0.05$). These significant differences were maintained during the recovery period thus

261 showing that organisms were not completely recovering from Hg exposure even when clean
262 food was provided (Dunnett's test, $p < 0.05$).

263 The time of hatched eggs was also evaluated in our study. Analyzing the hatching day of
264 juveniles, a significant delay was observed for organisms exposed to 4 ppm Hg in food ($H_4 =$
265 15.747; $p = 0.003$). The mean number of days for juveniles to hatch (\pm SD) in control, 1, 2, 3 and
266 4 ppm was respectively 22.1 (± 0.78); 21.6 (± 0.70); 22.3 (± 0.71), 22.2 (± 0.73) and 23.78 (± 1.30)
267 days. These results are in accordance with the ones found by Fountain and Hopkins (2011)
268 where reproduction (eggs laid in time) was delayed, due to retarded growth at high metal
269 concentrations. This species of collembolan has an unpigmented body and a gut that crosses
270 the organisms' body, and when they avoid food and start eating the black mixture of plaster of
271 Paris and activated charcoal, their gut acquires a visible black color. With the increase of Hg in
272 food the number of organisms showing a black ribbon along the body also increased (personal
273 observation/data not shown) thus supporting a possible avoiding food behavior. Avoidance
274 behavior is well explained by Fountain and Hopkin (2001), extrapolating that the reduced
275 growth of collembolans, when exposed to higher concentrations of metals, could be explained
276 to presumably use of the substrate as an alternative food source to springtails on an attempt
277 to acquire nutrition from the graphite.

278 In the study of Marigomez et al. (1986), the terrestrial slug *Arion ater* was exposed to
279 concentrations of mercury chloride ranging 10 to 100 ppm, and significant food consumption
280 and growth reductions were observed for concentrations higher than 10 ppm in a dose-related
281 manner. These results are in accordance with ours since an increase in Hg concentrations leads
282 to lower growth. Another study performed by Abassi and Soni (1983) kept adult earthworms
283 (*Octochaetus pattoni*) in cement tanks for 60 days, at the same density as the one found in the
284 wild, in a mixture of soil and animal dung contaminated with mercuric chloride ranging from
285 0.5 - 5.0 ppm Hg. The calculated LC_{50} was 2.39 ppm after 10 days' decreasing to 0.79 ppm at

286 the end of the 60 days' exposure period. Differences in the Hg-induced toxicity could be
287 explained by different exposure routes, since earthworms would ingest the contaminated
288 food, but would also be exposed to the contaminated soil by dermal contact and soil ingestion.
289 Lock and Janssen (2001) studied the effects of Hg in three different representative soil
290 invertebrates: the enchytraeid *Enchytraeus albidus*, the collembolan *Folsomia candida* and the
291 earthworm *Eisenia fetida*. An EC₅₀ of 9.16 ppm was calculated for *E. fetida* (21 days of
292 exposure), while an EC₅₀ of 22 ppm for *E. albidus* (42 days of exposure) and an EC₅₀ of 3.26
293 ppm for *F. candida* (28 days of exposure) were derived, based on data from reproduction
294 output for all organisms. A similar study conducted by Liu et al. (2010), using only the species
295 *F. candida* found a higher EC₅₀ for reproduction of 9.29 ppm and an EC₅₀ value for the
296 avoidance of 3.88 ppm in the presence of an agricultural fluvoaquic sandy loam soil (Cambisol,
297 9.0% clay, 21.8% silt, 69.2% sand) spiked with HgCl₂. Also, a more recent study from Buch et al.
298 (2016) assessed the effects of natural and artificial soils contaminated with Hg in two
299 collembolan species (*Folsomia candida* and *Proisotoma minuta*). For *F. candida* exposed to
300 natural soil, an AC₅₀ (avoidance behavior) of 5.44 (CI 4.13–6.75) ppm, an EC₅₀ for reproduction
301 of 3.40 (3.18–3.62) ppm and an LC₅₀ of 6.12 (3.74–8.50) ppm were attained. Despite the
302 apparent differences between both studies (different exposure routes and different endpoints
303 analyzed), the range of concentrations used in our study is lower than the LC₅₀ and AC₅₀, and
304 the highest concentration used similarly to the EC₅₀ derived by Buch et al. (2016). It should also
305 be noticed that in that study *F. candida* was more sensitive to mercury contamination than *P.*
306 *minuta*. Even knowing that different exposure routes could lead to differences regarding
307 toxicity, our results reveal that ingestion of contaminated food could be a very sensitive
308 endpoint and growth could be a relevant sensitive endpoint to look at.

309

310 **3.3 - Bioaccumulation test with *Folsomia candida***

311 Before Hg analysis, 20 collembolans per replicate from each treatment and control were
312 sampled and weighted at each sampling time. The results of the one-way ANOVA in each
313 sampling time revealed that until day 28 (complete uptake period), no statistical differences
314 were found in the weight of collembolans exposed to different Hg concentrations (ANOVA,
315 $p>0.05$). Collembolans sampled at day 35 (first sampling time in the recovery period) from the
316 2 and 3 ppm food exposure presented statistically higher weights than those from the control
317 (Dunnett's test, $p<0.05$). Surprisingly, no statistical differences were observed for the ones
318 exposed to 4 ppm Hg in food compared to the control. Regarding the collembolans used in day
319 42, statistical differences in their weight were also observed, with collembola from the 4-ppm
320 exposure showing higher weights (Dunnett's test, $p<0.05$), on a possible attempt to ingest as
321 much food as possible to compensate the lack of quality feeding when previously
322 contaminated food was supplied, as suggested above. Analyzing the results with a two-way
323 ANOVA, as expected, the time of exposure was the factor that explained the majority of the
324 collembola weight differences (two-way ANOVA, $F_{5,88} = 106.007$, $p<0.05$) with an R^2 of 0.82.
325 Regarding the factor for Hg concentration no statistical effect in collembolans' weight was
326 derived (two-way ANOVA, $F_{4,88} = 1.062$, $p>0.05$) with an R^2 of 0.014 with a slight interaction
327 with time of exposure (two-way ANOVA, $F_{20,88} = 2.23$, $p<0.05$; R^2 of 0.07).

328 The fit of the one-compartment kinetics model and the corresponding uptake and elimination
329 rate constants are shown in Figure 2 and Table 2. For 1 and 3 ppm treatments, Hg body
330 concentrations did not reach equilibrium after the 28 days of exposure. Mean Hg body
331 concentration ($n=3$) at the end of the uptake phase (28 days) was 0.02, 0.03, 0.07 and 0.07
332 ppm, corresponding to a total body burden of 0.07, 0.13, 0.25 and 0.26 ng of Hg at exposure
333 concentrations of 1, 2, 3 and 4 ppm, respectively. Within this, Hg values in collembola at day
334 28 of exposure increased with the increase in Hg concentrations in food, with the two highest
335 concentrations of exposure (3 and 4 ppm) producing similar body burden values.

336 Considering the size of collembolans decreased with increasing Hg concentrations, and their
 337 weight did not follow the same pattern, the hypotheses that an extra source of food occurred
 338 can be raised. This was somehow supported by observations of the presence of black material
 339 (possibly charcoal) in the digestive system of collembolans during the exposure to the highest
 340 Hg concentration. The hypothesis of extra ingestion of plaster of Paris and charcoal to
 341 compensate for the lack or low quality of the provided food source can be raised and
 342 corroborated with these findings. By looking at the kinetics curves patterns, collembolans
 343 exposed to 2 and 4 ppm were those deriving data that formed a (close to) plateau in the
 344 uptake phase, showing a potential equilibrium between uptake and elimination. This was also
 345 shown by the DT_{50} values at 2 and 4 ppm where Hg half-life in collembola was much lower
 346 than those from 1 and 3 ppm, revealing a faster elimination of Hg. The same pattern was also
 347 seen when comparing BAF values calculated with the kinetic parameters or through the
 348 concentrations measured.

349

350 Table 2 - Hg uptake and elimination rate constants (k_1 and k_2), bioaccumulation factor ($BAF_{kinetics}$ and
 351 $BAF_{org/food}$ and time that takes for Hg to be reduced by half inside the organisms (DT_{50}) for the
 352 accumulation of Hg in *Folsomia candida* exposed for 28 days to Hg spiked food, followed by an
 353 elimination phase of 14 days. The 95 % confidence intervals are presented in brackets.

[Hg] in food (ppm)	K_1 ($kg_{food}/kg_{org}/day$)	K_2 (day^{-1})	$BAF_{kinetics}$ (kg_{food}/kg_{org})	$BAF_{[Hg]org/[Hg]food}$ (kg_{food}/kg_{org})	DT_{50} (days)
1	0.0011 (0.001 - 0.002)	0.0307 (-0.001 - 0.062)	0.0359	0.018	22.6
2	0.0020 (0.001 - 0.003)	0.1038 (0.057 - 0.150)	0.0198	0.0165	6.7
3	0.0014 (0.001 - 0.002)	0.0456 (0.031 - 0.061)	0.0312	0.022	15.2
4	0.0016 (0.001 - 0.002)	0.0972 (0.068 - 0.126)	0.0173	0.017	7.1

354

355 Uptake rate constants (k_1) were compared using the likelihood-ratio test. The statistical
 356 analysis did not show any significant difference between treatments, thus showing that Hg

357 uptake and accumulation was concentration independent. The same procedure was conducted
358 for the elimination rate constant (k_2) and statistical differences were observed between
359 concentrations of 1 and 2 ppm ($\chi^2_{(1)} = 5.766$ $p < 0.05$), 1 and 4 ppm ($\chi^2_{(1)} = 5.34$ $p < 0.05$), 2 and 3
360 pp ($\chi^2_{(1)} = 8.10$ $p < 0.05$), and between 3 and 4 ppm ($\chi^2_{(1)} = 13.04$ $p < 0.05$). These differences in k_2
361 revealed that collembolans are eliminating Hg faster as Hg exposure increases.

362 Collembolans exposed to the lowest concentration of Hg (1 ppm) in food showed a low Hg
363 uptake (< 0.02 ng/g), which is in accordance with the no effects observed in their growth during
364 the test. At the highest concentrations (3 and 4 ppm) a maximum of approximately 0.07 ng/g
365 was attained, which could be close to the maximum sub-lethal concentrations that
366 collembolans can tolerate. Organisms exposed to 4 ppm had to compensate food intake with
367 extra food available (as mentioned above) as possibly they could not tolerate any further
368 increase of Hg in their body. Although this avoidance behavior towards the contaminated food
369 along with the ingestion of plaster led to an increase in their weight, no energy could be
370 allocated from the plaster towards their growth (expressed as body size).

371 The comparison of these data with literature is difficult as studies on mercury bioaccumulation
372 in soil-dwelling organisms are scarce, and toxicity studies are based on soil exposure using
373 reproduction as a preferred endpoint. However, as stated before, Fountain and Hopkin found
374 an avoidance behavior to Cd, Cu, Pb, and Zn exposures in food, at high concentrations.
375 Although they did not assess metal bioaccumulation, graphite was found in the digestive tract
376 which supports our data regarding food compensation. In the study of Zhang et al. (2009)
377 earthworms showed to accumulate high levels of Hg from a polluted soil with total and
378 methylmercury concentrations differing between earthworms' species. The comparison
379 between earthworms (or other species) and collembolans should be analyzed carefully since
380 the routes of exposure may be different. Also, the exposure route may be constant or
381 avoidable, like is the case of food, which can be avoided, or soil under a reproduction test

382 where avoidance is not an available option. Fountain and Hopkin (2001) also discussed the
383 potential higher tolerance of Collembolans regarding metal exposure through diet since they
384 can avoid contaminated food. When in contact with contaminated media they excrete metals
385 by molting, exfoliating the midgut epithelium where the elements are retained as part of a
386 storage detoxification system, in order to decrease metal accumulation (Köhler, 2002).

387 In soil ecotoxicology, the presence of soil as exposure matrix or as substrate can provide some
388 difficulties when assessing some crucial endpoints, like growth. In the present case study, the
389 dietary exposure was assessed in plaster of Paris/charcoal, providing continuous monitoring
390 during the exposure period, and assessing the effects in time. This approach would not have
391 been possible in the soil, due to the difficulty of spotting each collembola and following its
392 growth (in time).

393

394 **4 - Conclusions**

395 This study provides a new approach to understand Hg-induced effects in *F. candida*, by
396 combining an ecotoxicological endpoint (growth) and with Hg bioaccumulation following a
397 timeline. For the first time, it is reported the Hg-induced effects on growth and
398 bioaccumulation from a dietary source to the springtail *F. candida*. Even at low and
399 environmental relevant Hg concentrations, collembola growth rate was impaired. Despite a
400 clear difference in the growth of collembolans with increasing Hg concentrations, collembolans
401 did not reach a complete steady state in all concentrations of the uptake phase, being
402 independent of the Hg exposure concentration.

403 Regarding depuration, after exposure to the higher Hg concentration, collembolans showed an
404 increase in the k_2 , with a faster elimination and decreasing Hg residence time. Also, by
405 comparing collembolans' length and weight at different Hg concentration, along with their
406 bioaccumulation patterns, it could be highlighted a potential food avoidance and exploring a

407 different food source available (plaster with charcoal). This was perceived by the dark color of
408 collembola exposed to the highest concentrations. It was also highlighted a possible maximum
409 sub-lethal concentration of 0.07 ng Hg/g of collembolan. The present study can be considered
410 as a step forward in the assessment of the potential risks of mercury or other metals in
411 terrestrial environments, also providing a mechanistic approach looking simultaneously at
412 toxicity and bioaccumulation.

413

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421

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Figure
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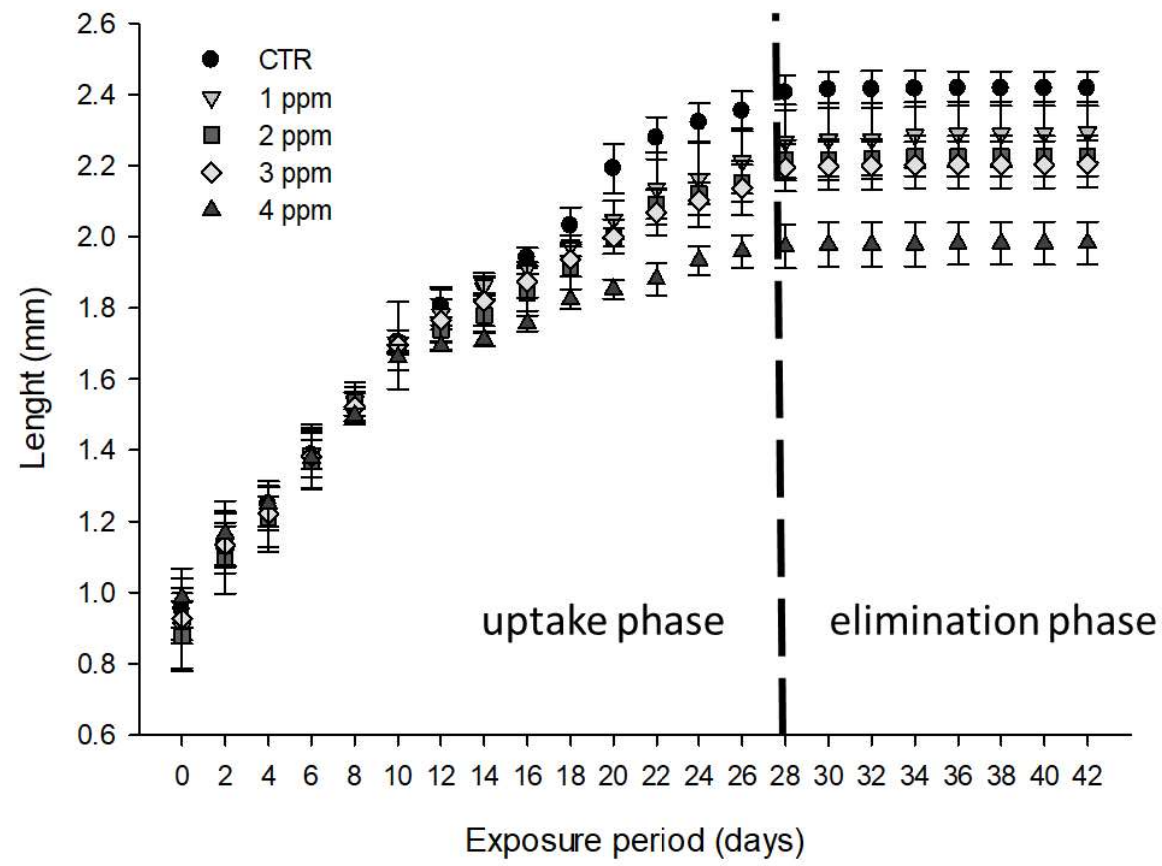


Figure 1 – Body length of *Folsomia candida* recorded during a 28 day exposure to Hg contaminated yeast (uptake phase), followed by a 14 day exposure to clean yeast (elimination phase). The negative control was clean yeast (diamond) and the vertical dashed line separates the uptake phase from the elimination phase. Data is expressed as average \pm standard error.

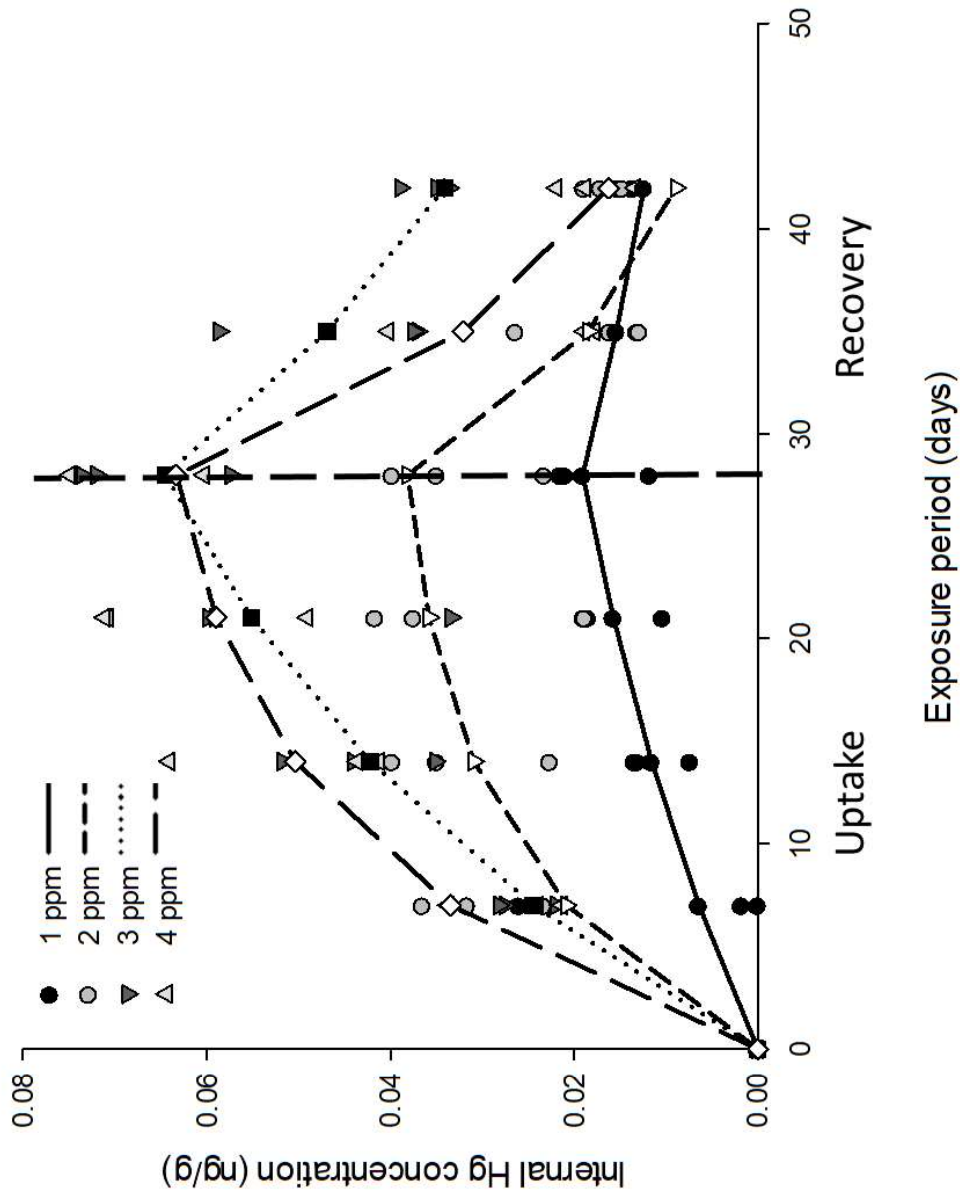


Figure 2 - Uptake and elimination kinetics of Hg in *Folsomia candida* exposed to nominal concentrations ranging from 1 to 4 ppm in contaminated dry yeast. Uptake and elimination phases lasted for 28 and 14 days, respectively. The vertical dashed line separates the uptake phase from the elimination phase. Lines represent the modeled Hg body concentration, using the one compartment model.

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