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1 **An energy-based model to analyze growth data of earthworms exposed to two**  
2 **fungicides.**

3

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26 **Abstract**

27         The pesticide risk assessment for earthworms is currently performed using  
28 standardized tests, the model species *Eisenia fetida*, and the analyses of the data obtained are  
29 performed with *ad hoc* statistical tools. We assessed the impact of two fungicides on the  
30 entire growth pattern of the earthworm species *Aporrectodea caliginosa*, which is highly  
31 representative of agricultural fields. Individuals of three different ages (from hatching to 56  
32 days old) were exposed to Cuprafor micro<sup>®</sup> (copper oxychloride) and Swing<sup>®</sup> Gold  
33 (dimoxystrobin and epoxiconazole). Data were analyzed with an energy-based toxicodynamic  
34 model coupled with a toxicokinetic model. The copper fungicide caused a drastic growth  
35 inhibition once the No Effect Concentration (NEC), estimated at 65 mg kg<sup>-1</sup> of copper, was  
36 exceeded. The Swing<sup>®</sup> Gold negatively affected the growth with NEC values estimated at  
37 0.387 mg kg<sup>-1</sup> and 0.128 mg kg<sup>-1</sup> for the dimoxystrobin and the epoxiconazole in this  
38 fungicide formulation, respectively. The time-profile of the effects on *A. caliginosa*  
39 individuals was fully accounted for by the model, whatever their age of exposure.  
40 Furthermore, toxicity data analyses, supported by measurements of fungicide concentrations  
41 in earthworm at the end of the experiment, allowed bettering understanding of the  
42 mechanisms of action of the fungicides towards earthworm growth.

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49 **Keywords:** Ecotoxicology, *Lumbricidae*, Growth, Risk assessment, Toxicokinetic-  
50 toxicodynamic modelling.

## 51 **Introduction**

52 Pesticides used in agroecosystems can harm biodiversity and biological activity  
53 (Bengtsson et al. 2005; Hole et al. 2005). Among non-target soil organisms that can be  
54 impacted by pesticides, earthworms are commonly used as biological indicators of chemical  
55 stress (OECD, 1984) because they are key soil organisms, involved in nutrient cycling, soil  
56 water regulation and aeration (Blouin et al. 2013; Bertrand et al. 2015; Bart et al., 2019a).  
57 During the last decades, different ecotoxicological laboratory tests have been developed such  
58 as the acute toxicity test (ISO 2012a) or the reproduction tests (ISO 2012b; OECD 2004). In  
59 the ISO and OECD tests, the recommended species is *Eisenia fetida fetida* or *Eisenia fetida*  
60 *andrei* and they are often used to assess the impacts of pesticides or other chemicals on  
61 earthworms. Contrarily, growth tests with earthworms are poorly documented while some  
62 authors reported growth to be a very sensitive endpoint (Springett and Gray 1992; Booth et al.  
63 2000; Booth and O'Halloran 2001). Authors highlighted the need to develop a test system for  
64 measuring key demographic traits in juvenile earthworms, especially growth (Spurgeon et al.  
65 2004). To move towards a more realistic and relevant assessment of the environmental risks  
66 of pesticide use, some issues have to be overcome.

67 First, the model species (i.e. *E. fetida fetida* or *andrei*) do not generally inhabit mineral  
68 soils (Lowe and Butt 2007) and are therefore rarely found in cultivated fields where pesticides  
69 are applied. To complement the use of *E. fetida* in pesticide risk assessment procedures, the  
70 earthworm species *Aporrectodea caliginosa* s.s. (Sims and Gerard 1999) was recently  
71 proposed as a relevant species to be used in soil ecotoxicology tests (Klobucar et al. 2011;  
72 Bart et al. 2018). This species is one of the dominant species in agroecosystems in temperate  
73 areas (Boström and Lofs-Holmin 1996; Boag et al. 1997; Curry et al. 2008; Amossé et al.,  
74 2018) and is found to be more sensitive to pesticides and metabolites than *E. fetida* (Pelosi et  
75 al. 2013). The second issue in currently used risk assessment procedures is data analyses. *Ad*

76 *hoc* statistics are used to test differences between effects measured in polluted and unpolluted  
77 soils or to calculate no observed effect concentrations (NOEC), lowest observed effect  
78 concentrations (LOEC), or effective concentrations (EC<sub>x</sub>). However, these parameters cannot  
79 easily be extrapolated for other exposure durations than the one used for the test, and do not  
80 account for the kinetics of the toxicant in soil. To go towards a better understanding of the  
81 mechanisms of toxicants on life cycle parameters, energy-based models were proposed to  
82 analyze toxicity data (Kooijman and Bedaux 1996). These models are based on the dynamic  
83 energy budget (DEB) theory (Kooijman 1986, 2000, 2010) which partitions the use of energy  
84 between growth, maintenance, and reproduction. Effects models, called DEBtox models,  
85 assume that the use of energy by an organism described in the DEB model can be unbalanced  
86 by a toxicant. The effect is described as an impact on one of the parameters of the energy-  
87 based model and the magnitude of the effects is assumed to be related to the internal  
88 concentration of the toxicant of the organism. The exposure concentration and internal  
89 concentration of the organisms are related throughout time by a toxicokinetic model. The  
90 DEBtox models are toxicokinetic-toxicodynamic models (TK-TD) and have proved their  
91 reliability in the analysis of data from growth and reproduction tests (Péry et al. 2002; Jager et  
92 al. 2004; Goussen et al. 2013). These models also allow estimating a no effect concentration  
93 (NEC) which is a threshold for toxicity that does not depend on the time of exposure. The  
94 NEC can be used to compare ecotoxicity of toxicants avoiding time dependency issues of  
95 classical parameters such as LOEC, NOEC or EC<sub>x</sub> (Baas et al. 2010; Heckmann et al. 2010;  
96 Jager et al. 2014).

97 We here tested the impact of two widely used commercial formulations of fungicides,  
98 that are of interests for the pesticide risk assessment (Bart et al. 2017), on the growth of the  
99 earthworm *A. caliginosa*. The exposure was performed at three different ages to reveal  
100 potential differences in earthworm sensitivity over their development. Data were analyzed

101 with an energy-based model, calibrated for this species (see supplementary material and Bart  
102 et al., 2019b), in order to understand the mechanisms of action and the time-dependence of  
103 the two different fungicides on *A. caliginosa* growth. To support the understanding of the  
104 toxicity mechanisms provided by the model, we performed concentration measurements in  
105 earthworms at the end of the growth experiment.

106

## 107 **Material and methods**

### 108 **Soil, animals and pesticides.**

109 All experiments were performed using a loamy soil texture (Based on the texture  
110 definition of the Food and Agriculture Organization of the United Nations (FAO)), sampled  
111 from a permanent grassland in Versailles (48°48' N, 2°5' E) where no pesticides have been  
112 applied for more than 20 years. The soil was collected from the top 0-20 cm, air-dried and  
113 crushed to pass a 2 mm mesh. Its main physico-chemical characteristics were: pH 7.5, organic  
114 matter 32.6 g kg<sup>-1</sup>, C/N 12.7, 29% sand, 48% silt, 23% clay, and 25.2 mg kg<sup>-1</sup> of copper (see  
115 Bart et al. 2017 for more details).

116 *Aporrectodea caliginosa* s.s used in this experiment were bred in the laboratory from  
117 individuals initially collected from an agricultural field in Estrée-Mons, France (49°52' N, 3°01'  
118 01' E) one year before this study, and determined according to Sims and Gerard (1999). The  
119 earthworms were bred in the same soil described above. To get cohorts of hatchlings, cocoons  
120 were collected in the breeding culture by wet sieving the soil through a 1-mm mesh size (Bart  
121 et al. 2018), and incubated at 20 °C in Petri dishes on wet filter papers (Holmstrup et al.  
122 1991). Cocoons were checked every two days and new hatchlings were collected and stored in  
123 the breeding soil at 4 °C for a maximum of 1 week, to slow their development. This procedure  
124 allowed synchronizing cohorts of individuals to the same level of development (Bart et al.  
125 2018).

126 The first studied fungicide was Swing<sup>®</sup> Gold (BASF Agro SAS, dimoxystrobin 133 g  
127 L<sup>-1</sup>, epoxiconazole 50 g L<sup>-1</sup>), used to protect cereal crops in conventional farming. The  
128 French Recommended Dose (RD) for this product is 1.5 L ha<sup>-1</sup> on wheat (E-phy 2017a). The  
129 RD in laboratory was calculated as 1.16 10<sup>-3</sup> mL kg<sup>-1</sup> of dry soil (corresponding to 150 µg kg<sup>-1</sup>  
130 of dimoxystrobin and to 60 µg kg<sup>-1</sup> of epoxiconazole) with a soil density of 1.29 and  
131 considering that the active compounds of this fungicide are mainly found in the top 10 cm of  
132 soil (McDonald et al. 2013; Chabauty et al. 2016). We tested the following concentrations:  
133 0.33, 1, and 3 times the RD, abbreviated SG0.33, SG1, and SG3, respectively. These  
134 concentrations were assumed to be sub-lethal considering the LC50 estimated at 7.0 10<sup>-3</sup> mL  
135 kg<sup>-1</sup> for *A. caliginosa* (Bart et al. 2017), or 6.03 times the RD.

136 The second studied fungicide was Cuprafor micro<sup>®</sup>, used to prevent spore germination  
137 in organic farming mainly. The French RD for this product is 10 kg ha<sup>-1</sup> for potato crops and  
138 in vineyards (E-phy 2017b). The RD in laboratory was calculated as 15.5 mg kg<sup>-1</sup>  
139 (corresponding to 7.75 mg kg<sup>-1</sup> of copper) of dry soil with a soil density of 1.29 and  
140 considering that copper is mainly found in the top 5 cm of soil (Couto et al. 2015). We tested  
141 the following concentrations: 3.33, 10, and 30 times the RD abbreviated Cu3.33, Cu10 and  
142 Cu30, corresponding respectively to 25.8, 77.5, and 232.5 mg kg<sup>-1</sup> of copper. These  
143 concentrations were assumed to be sublethal (Ma 1984; Spurgeon et al. 2004; Bart et al. 2017;  
144 PPDB 2018).

145 In all experiments, the dry soil was spiked with aqueous solutions of the fungicides,  
146 and the soil water holding capacity was adjusted concomitantly at 70% of the Water Holding  
147 Capacity (WHC).

148

## 149 **Growth experiment**

150

151 In order to monitor the growth, the weight of individuals was measured using an analytical  
152 balance ( $\pm 0.1$  mg). The impact of fungicides was tested exposing earthworm juveniles at  
153 three different ages: just after hatching, after 28 days of growth in a control soil and after 56  
154 days of growth in a control soil (see Fig. S1 in supplementary material). These three ages  
155 were named age 1, age 2 and age 3 respectively (A1, A2 and A3) and individuals weighed  $12$   
156  $\pm 3$  mg,  $90 \pm 15$  mg and  $300 \pm 40$  mg when their exposure began at the three ages,  
157 respectively. Under control condition of the experiment, *A. caliginosa* reach maturity (i.e.  
158 apparition of the clitellum) and are able to reproduce after  $85 \pm 10$  days of growth and at a  
159 weight of  $575 \pm 125$  mg.

160 Earthworms were placed individually in 1 L plastic vessels (15 x 10 x 7 cm) with 400 g of  
161 soil (dry mass). Seven replicates (each replicate corresponded to one individual) were used  
162 per age of exposure and fungicide concentration, including a control without fungicide. All  
163 the vessels were stored in a climate room at  $15 \pm 1$  °C. Individuals were fed with horse dung in  
164 *ad libitum* conditions, as presented in the supplementary material. Individuals were weighed  
165 at least every 14 days and the experiment was stopped when individuals had reached maturity,  
166 characterized by the apparition of a fully developed clitellum. For the individuals who  
167 stopped to grow, without reaching the adult stage, we stopped the experiment after around 35,  
168 70 and 98 days without growth for individuals exposed at Age 3, 2 and 1 respectively. At the  
169 end of the experiment, all individuals were placed in petri dishes on damp filter paper for 48 h  
170 in the dark at  $15 \pm 1$  °C to void gut contents (Hartenstein et al. 1981). Then, they were  
171 weighted and frozen at  $-80^{\circ}\text{C}$  for fungicide analysis.

172

173 For each treatment, the soil was renewed every 28 days to avoid unsuitable conditions for  
174 earthworm growth (e.g. soil compaction). All the soils for a given fungicide treatment were  
175 prepared at the same time (see Fig. S1), to ensure a comparable evolution of pesticide  
176 concentrations and environmental available fraction. Swing<sup>®</sup> Gold fungicide concentration in



177 the soil with earthworms and horse dung and in the new soil was monitored at each soil  
178 renewal (see supplementary material for more details). Moreover, in order to characterize the  
179 exposure concentration of dimoxystrobin (DMX) and epoxiconazole (EPX), the total  
180 concentrations and the environmental available fraction of the active substances were  
181 monitored over the time of the experiment, every 28 days.

182

### 183 **Fungicide analyses.**

184 *The copper fungicide* - The soil was sampled just after the soil preparation soils to verify the  
185 contamination level. The copper concentration in earthworms at the end of the experiment  
186 was also measured in order to quantify the accumulation. Details of the chemicals analyses in  
187 soil and earthworms are available in the supplementary material.

188 *The Swing<sup>®</sup> Gold fungicide* - The total soil concentration and of the environmentally available  
189 fraction of DMX and EPX were measured over time (0, 28, 56 and 84 days after the  
190 contamination) in 4 of the 7 replicates to take into account their dissipation, which changed  
191 the earthworm's exposure. These measured concentrations were used in the toxicokinetic  
192 model. The DMX and EPX concentration in earthworms at the end of the experiment was also  
193 measured in order to quantify the accumulation. Details of the chemicals analyses in soil and  
194 earthworms are available in the supplementary material.

195

### 196 **The energy-based model.**

197

198 We used a growth model shortly presented in the supplementary material and fully  
199 presented in Bart et al., (2019b). This model is based on the DEB theory (Dynamic Energy  
200 Budget) (Kooijman 1986, 2000, 2010). Under *ad libitum* conditions and according to the

201 assumptions of isomorphism and neglected energy costs of maintenance, the growth is  
202 expressed with the following equation:

203 
$$\text{If } l < Cs, \text{ then } \frac{d}{dt}l = a(1 - b) \quad (1)$$

204 
$$\text{If } l > Cs, \text{ then } \frac{d}{dt}l = a \quad (1)$$

205 Where  $l$  is the cubic root of the wet weight of the organisms,  $Cs$  is the Critical size  
206 (below which the individual cannot access all the food)  $a$  is a constant and  $b$  is a food  
207 accessibility factor to take into account that when individuals are too small, they cannot  
208 access the whole food quantity. In our experimental conditions, the parameter values were  
209 optimized with the control treatment as follows  $Cs = 3.99$ ,  $a = 0.075$  and  $b = 0.13$ .

210

### 211 **The toxicokinetic/toxicodynamic (TK/TD) model.**

212

213 The effect model is based on the energy-based model mentioned above. We assumed that  
214 the exposure to the toxicant increases the energy cost of growth. As in the DEBtox model  
215 (Kooijman and Bedaux 1996), we assumed that there is a threshold for effect, a no effect  
216 concentration (NEC), and that the effects are proportional to the difference between the  
217 internal concentration and the NEC value. The toxicokinetic of the internal concentration ( $C_i$ )  
218 was deduced from the exposure concentration with a one compartment model:

219

$$\frac{dC_i}{dt} = Ku \times ce(t) - Ke \times C_i(t) \quad (2)$$

220 where  $Ku$  and  $Ke$  are the uptake and the elimination rate,  $ce$  and  $C_i$  are the external  
221 and internal concentrations of the toxicant, respectively. However, because we do not have

222 access to the internal concentration, it was scaled by the bio-concentration factor as explained  
 223 in a previous study (Péry et al. 2001), leading to the following equation:

$$\frac{dci(t)}{dt} = Ke (ce(t) - ci(t)) \quad (3)$$

$$\text{with } ci = Ci \frac{Ke}{Ku} \quad (4)$$

224 Where  $ci$  is proportional to the internal concentration, but corresponds to an external  
 225 concentration. Moreover, the individuals had a measurable growth during the experiment  
 226 which led to a dilution by growth (the earthworm increased biomass reduced the internal  
 227 concentration). We accounted for this in the toxicokinetic model (Kooijman and Bedaux,  
 228 2010). The elimination rate is assumed to be proportional to the ratio of the surface area to the  
 229 volume, and thus inversely proportional to the length for an isomorphic organism as explained  
 230 theoretically (Kooijman and Bedaux, 2010) and shown experimentally (Sijm and van der  
 231 Linde 1995; Sijm et al. 1995). This is why the elimination rate must be divided by a scaled  
 232 length if the body size changes leading to the following equation:

$$\frac{dci(t)}{dt} = \frac{Ke (ce(t) - ci(t))}{l} - \frac{3a ci}{l} \quad (5)$$

234  
 235 In the case of an increase in the growth costs, we assumed that the costs of building a  
 236 cell are multiplied by a factor  $1 + e (ci(t) - NEC)$ ,  $e$  being a constant and accounting for the  
 237 level of toxicity as soon as the NEC is exceeded by the scaled internal concentration, leading  
 238 to the following equation:

239 If  $l < Cs$ , then  $\frac{d}{dt} l = \frac{a(1-b)}{1 + e (ci - NEC)}$  (6)

240                    If  $l > Cs$ , then  $\frac{d}{dt} l = \frac{a}{1+ e (ci-NEC)}$  (6)

241

242                    All the parameters ( $Ke$ ,  $NEC$  and  $e$ ) were simultaneously calibrated for each fungicide  
243 (including the three different ages of exposure).

244

245

### 246 **Model calibration and statistical analyses.**

247                    The differential equations were implemented in the software R Core Team (2015), and  
248 solved with the package deSolve (Soetaert et al. 2010). The model was fitted to the data, for  
249 all concentrations and ages of exposure for each fungicide, using the least square method.  
250 The bootstrap method<sup>45</sup> was used for the estimation of the confidence intervals of the  
251 parameters. The R script is available on request to the corresponding author.

252

## 253 **Results**

### 254 **The copper fungicide**

255                    The copper contamination led to a drastic growth inhibition in the Cu10 and Cu30  
256 treatments (Fig. 1). We thus chose to simplify the toxicodynamic model as follows, with an  
257 infinite value for  $e$ :

$$\text{If } ci > NEC, \text{ then } \frac{d}{dt} l = 0 \quad (7)$$

258

259 In this situation, there were only two parameters to calibrate in the model:  $Ke$  and the  
260 NEC. The parameter  $Ke$  was estimated at 1.19 and the NEC at 65 mg kg<sup>-1</sup> of copper (Table 1).  
261 These parameter values were common to the three different ages.

262

263

264

265 **Table 1.** Estimated parameter values and confidence intervals (CI 95%) for the copper  
266 fungicide.

Parameters	Value	CI (95%)
$Ke$	1.19	1.17 - 1.67
NEC copper (mg kg <sup>-1</sup> )	65.006	64.79 - 65.01
$e$	infinite	-

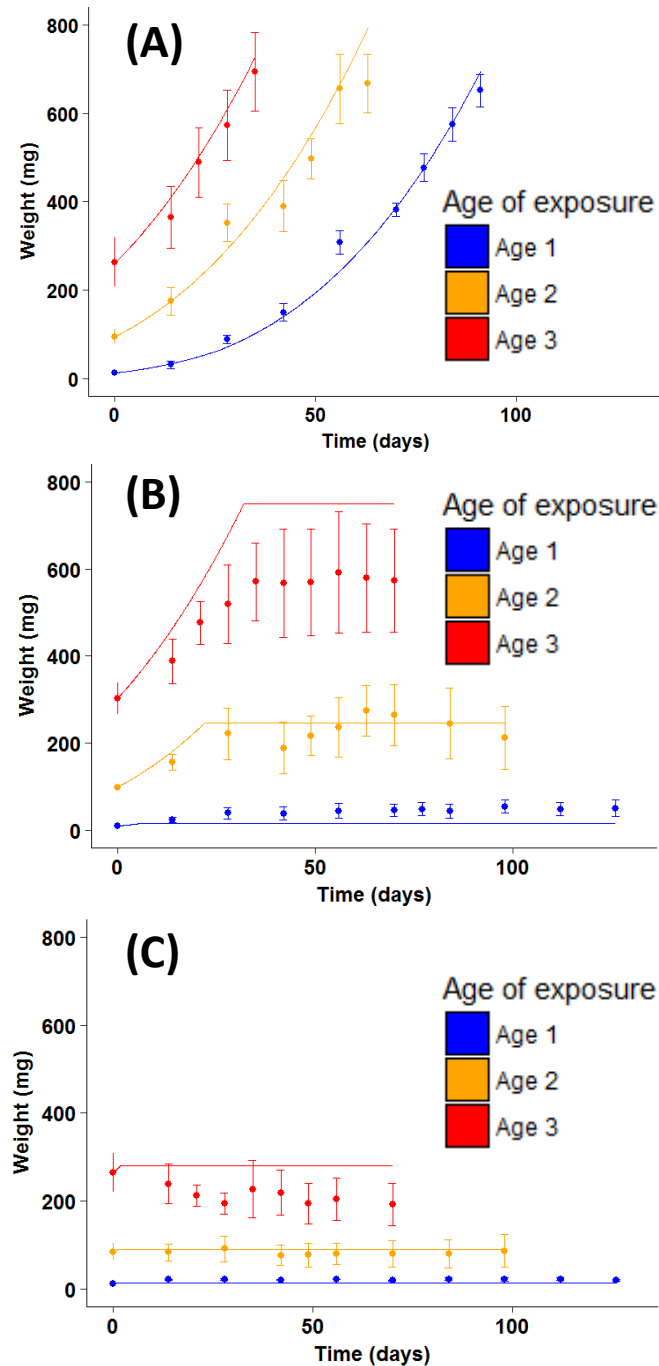
267

268

269 Fig. 1 presents the growth data and the description by our model. The data were not  
270 significantly different from the model description in 59% of the cases ( $P > 0.05$  with  
271 Student's t-tests). The difference was mainly due to over-estimation of the asymptotic mass of  
272 individuals of age 3 exposed to Cu10 and Cu30 treatment, and under-estimation of the  
273 asymptotic mass of individuals of age 1 exposed to the Cu10 treatment.

274 The copper fungicide applied at 25.8 mg kg<sup>-1</sup> of copper did not impact the growth for  
275 the 3 different ages of exposure because the NEC was not exceeded (Fig. 1). The growth  
276 pattern in this treatment thus corresponded to the growth pattern provided by the model in the  
277 control. At 77.5 mg kg<sup>-1</sup> of copper, an inhibition of the growth was observed, appearing at  
278 different times after the beginning of the exposure for the different ages: the effects appeared  
279 immediately after the exposure for new-hatched individuals (Age 1) and after respectively 20  
280 and 30 days after the exposure for individuals of Age 2 and 3. The model accounted for these  
281 differences through the dilution by growth in the toxicokinetic model and provided a good  
282 description of the data although it slightly overestimated the growth of the bigger juveniles

283 (Age 3). At 232.5 mg kg<sup>-1</sup> of copper, the growth was totally inhibited right after the beginning  
284 of the exposure, whatever the age of the earthworms. Our model also accounted for this  
285 absence of difference between ages.

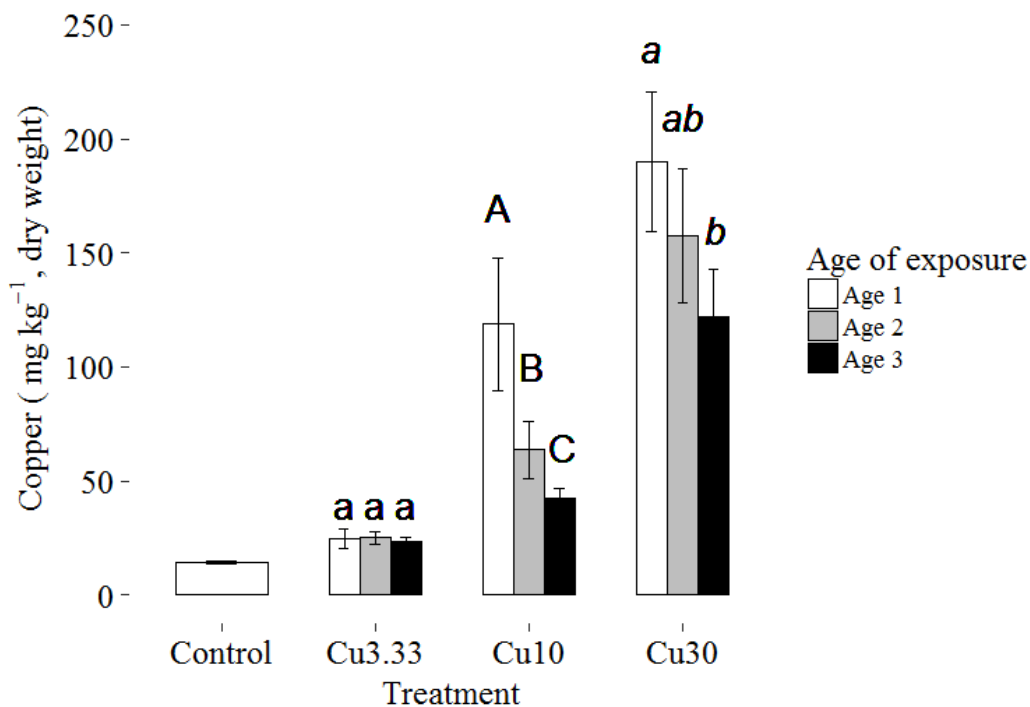


286  
287 **Fig. 1** Growth pattern of *A. caliginosa* juveniles exposed at different ages to the Cuprafor  
288 micro<sup>®</sup> fungicide at (A) 3.33 times the RD (Recommended Dose), corresponding to 25.8 mg  
289 kg<sup>-1</sup> of copper, (B) 10 times the RD, corresponding to 77.5 mg kg<sup>-1</sup> of copper, and (C) 30

290 times the RD, corresponding to 232.5 mg kg<sup>-1</sup> of copper. Full lines represent the description  
291 of the observations (n=7 ± SD) by the model.

292  
293 The copper accumulation in earthworm significantly differed between treatments and  
294 the control (Fig. 2). At 25.8 mg kg<sup>-1</sup> of copper (Cu3.33), the copper internal concentration at  
295 the end of the experiment did not differ between individuals exposed at Age 1, 2, or 3. For the  
296 individuals exposed in the Cu10 and Cu30 treatments, the copper concentration in tissues  
297 significantly decreased with increase of the age of exposed earthworms corresponding to an  
298 increase in copper accumulation with the time of exposure (Fig. 2).

299  
300



301  
302 **Fig. 2** Copper concentration in *A. caliginosa* individuals at the end of the growth experiment  
303 in mg kg<sup>-1</sup> (dry weight). Individuals of Age 1 were exposed for on average 91 days to the  
304 Cu3.33 (25.8 mg kg<sup>-1</sup> of copper) treatment and 126 days to Cu10 (77.5 mg kg<sup>-1</sup> of copper) and  
305 Cu30 (232.5 mg kg<sup>-1</sup> of copper). Individuals of Age 2 were exposed for on average of 63 days

306 to the Cu3.33 treatment and 98 days for the Cu10 and Cu30 treatment. Individuals of Age 3  
 307 were exposed for on average of 35 days to the Cu3.33 treatment and 70 days to the Cu10 and  
 308 Cu30 treatment. Different letters mean significant differences between ages of exposure for  
 309 each copper treatment.

310

### 311 **The Swing<sup>®</sup> Gold fungicide**

312 The effects of Swing<sup>®</sup> Gold on the growth appeared immediately after the start of  
 313 exposure in the SG3 treatment (Fig. 3C). We thus assumed a very fast toxicokinetics and used  
 314 directly the total DMX or EPX soil concentration as internal concentration (*ci*) in the effect  
 315 model. Two parameters were thus calibrated: *e* and the NEC. The parameter *e* was estimated  
 316 at 13.27 and 13.24 for the DMX and the EPX in the Swing<sup>®</sup> Gold formulation respectively  
 317 (Table 2). The NEC was estimated at 0.387 mg kg<sup>-1</sup> (dry soil) and 0.128 mg kg<sup>-1</sup> of DMX and  
 318 EPX in the Swing<sup>®</sup> Gold formulation respectively (Table 2). These parameter values were  
 319 common to the three different ages.

320

321 Table 2. Estimated parameter values and confidence intervals (CI 95%) for the Swing<sup>®</sup> Gold  
 322 fungicide.

Parameters	Value	CI (95%)
<i>Ke</i>	infinite	-
NEC Dimoxystrobin (mg kg <sup>-1</sup> )	0.387	0.375 - 0.402
NEC Epoxiconazole (mg kg <sup>-1</sup> )	0.128	0.123 - 0.143
<i>e</i> Dimoxystrobin	13.27	11.98 - 21.00
<i>e</i> Epoxiconazole	13.24	13.22 - 44.18

323

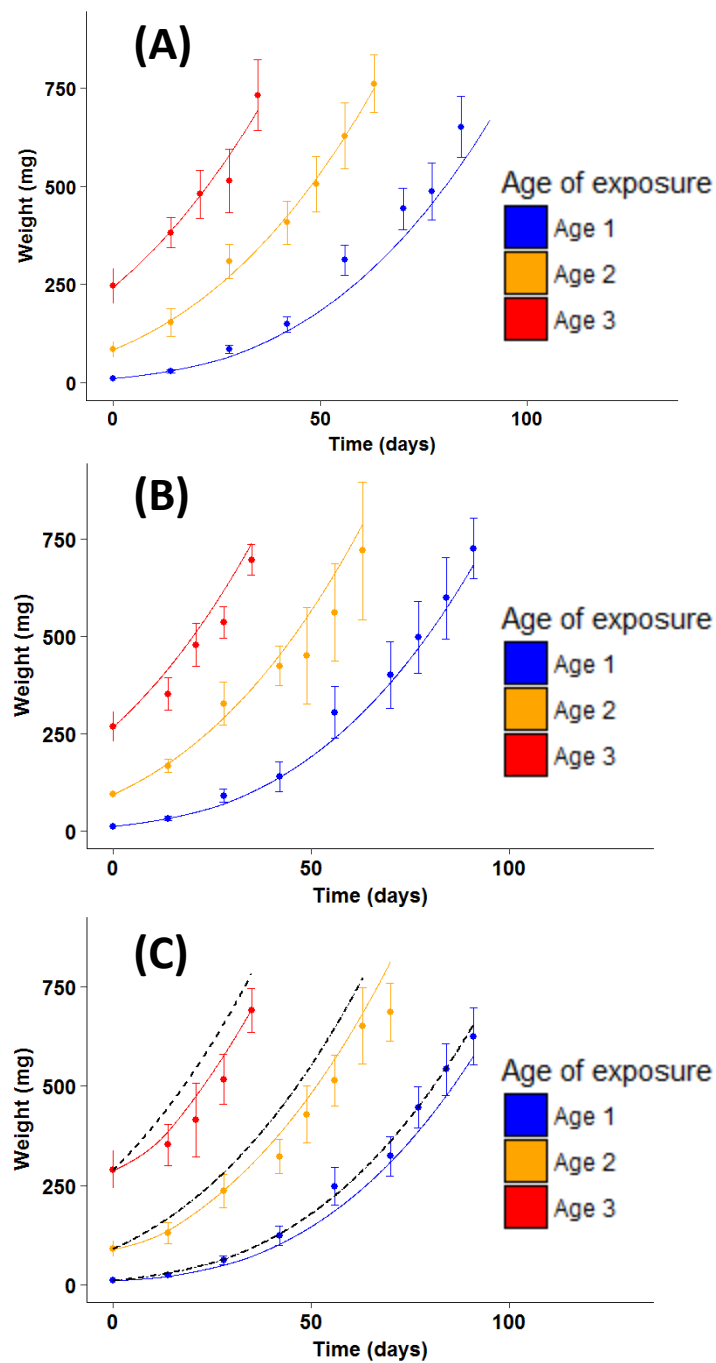
324

325 Fig. 3 presents the growth data and the description by our model. The data were not  
 326 significantly different from the model description in 81% of the cases ( $P > 0.05$  with  
 327 Student's t-tests). The growth of individuals exposed at 0.33 and 1 times the RD (SG0.33 and



328 SG1) was not affected because the NEC was not exceeded. In these treatments, the growth  
329 pattern thus corresponded to the growth pattern provided by the model in the control (Fig. 3A  
330 and 3B). At 3 times the RD (SG3), the growth was negatively affected just after the exposure  
331 whatever the age of exposure (Fig. 3C), and during a period of 15 days corresponding to the  
332 time during which the concentration exceeded the NEC.

333

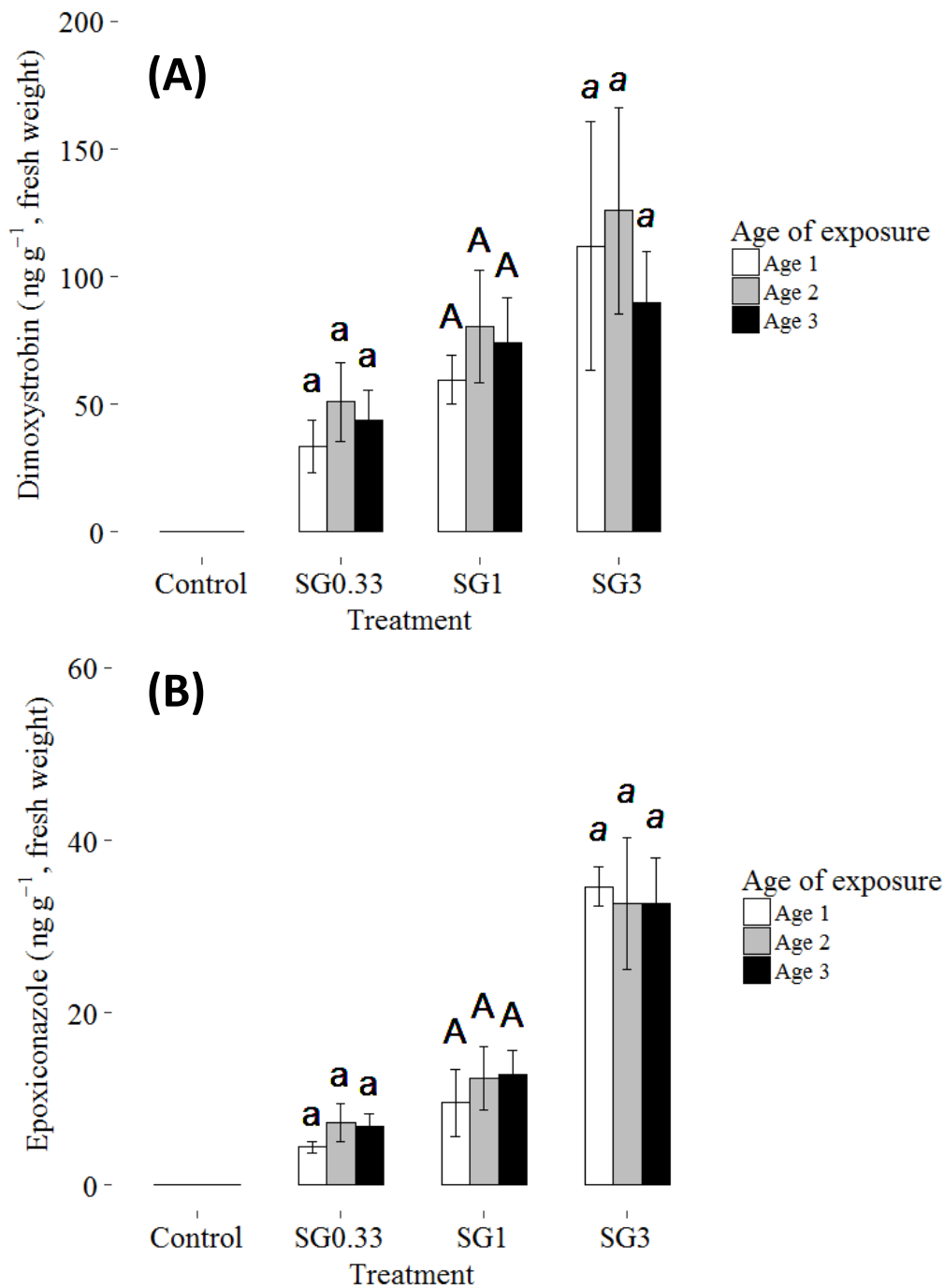


334

335 **Fig. 3** Growth pattern of *A. caliginosa* exposed at different ages to the Swing<sup>®</sup> Gold fungicide  
336 at (A) 0.33 times the RD (Recommended Dose) corresponding to  $5.2 \times 10^{-2}$  mg kg<sup>-1</sup> of DMX  
337 and  $1.94 \times 10^{-2}$  mg kg<sup>-1</sup> of EPX. (B) 1 time the RD corresponding to  $1.55 \times 10^{-1}$  mg kg<sup>-1</sup> of  
338 DMX and  $5.81 \times 10^{-2}$  mg kg<sup>-1</sup> of EPX. (C) 3 times the RD corresponding to  $4.62 \times 10^{-1}$  mg  
339 kg<sup>-1</sup> of DMX and  $1.74 \times 10^{-1}$  mg kg<sup>-1</sup> of EPX. Full lines represent the description of  
340 observations ( $n=7 \pm$  SD) by the model, and the dash lines represent the description of the  
341 observations in the control treatment by the model.

342

343 The DMX and EPX accumulation in earthworm significantly differed between the  
344 treatments and the control (Fig. 4). There was no significant difference in the accumulation of  
345 DMX or EPX between the different ages of exposure corresponding to no difference  
346 accumulation with the time of exposure (Fig. 4).



347

348 **Fig. 4** Concentration of dimoxystrobin (A) and epoxiconazole (B) in *A. caliginosa* individuals  
 349 at the end of the growth experiment in ng g<sup>-1</sup> (fresh weight). Individuals of Age 1 were  
 350 exposed for on average 91 days to the SG 0.33 and SG1 treatment and 95 days to the SG3  
 351 treatment. Individuals of Age 2 were exposed for on average 63 days to the SG 0.33 and SG1  
 352 treatment and 67 days to the SG3 treatment. Individuals of Age 3 were exposed an average 35  
 353 days to the SG 0.33 and SG1 treatment and 38 days to the SG3 treatment. Different letters  
 354 mean significant differences between ages of exposure for each Swing<sup>®</sup> Gold treatment.

355

## 356 **Discussion**

### 357 **Impact of the two tested fungicides on earthworm growth.**

358           We here showed that the growth pattern of one of the most representative species of  
359 earthworms in cultivated fields was highly influenced by the presence of two fungicides at  
360 environmentally relevant doses. Moreover, we highlighted that the magnitude of the effects  
361 depend on the age of the individuals. Finally, we pointed out that the model proposed in the  
362 supplementary material coupled with a TK-TD model can be very useful to understand  
363 impacts and provide threshold value of the ecotoxicity of the tested fungicides.

364           The copper fungicide appeared highly harmful to earthworm growth beyond the NEC,  
365 estimated at 65 mg kg<sup>-1</sup> of copper, corresponding to 8.4 times the RD of the fungicide. This  
366 result is in accordance with the EC50 growth estimated at 81.8 mg kg<sup>-1</sup> of copper in a  
367 previous study (Khalil et al. 1996). Others authors showed that copper affected the growth of  
368 *E. fetida* from 8.92 mg Cu kg substrate<sup>-1</sup> (which was urine-free cattle manure) and that  
369 earthworms exposed to 346.85 mg Cu kg substrate<sup>-1</sup> exhibited hardly any increase in weight  
370 (Helling et al. 2000). It has also been showed that copper inhibited the growth of the  
371 earthworm *Lumbricus rubellus* at a concentration of 370 mg kg<sup>-1</sup> of copper in a sandy soil  
372 (Ma 1984). It thus appears that copper is harmful for earthworm growth but the threshold  
373 concentration inducing impact is highly dependent on the soil characteristics and species  
374 considered (Ma and Rao 1997, EU 2008). Finally, it is important to notice that the NEC value  
375 estimated (i.e. 65 mg kg<sup>-1</sup>) could be reached in agricultural systems because copper  
376 accumulates in soils (Brun et al. 1998). This is the case in vineyards in which copper can  
377 reach more than 100-200 mg kg<sup>-1</sup> and explain the very low density of earthworms in these  
378 agroecosystems (Paoletti et al. 1998).

379 Harmfulness of the Swing<sup>®</sup> Gold fungicide on earthworm growth was estimated by a  
380 NEC value of 2.5 times the RD. Moreover, DMX and EPX have estimated DT50 values (lab  
381 at 20°C) of 210 and 226 days respectively (PPDB, 2018), suggesting that these compound  
382 could accumulate and persist in the environment. The NEC values provided in this study are  
383 valid only in the studied commercial formulation and we could not determine which of the  
384 two substances caused the effect on growth. However, literature suggest that DMX is more  
385 harmful than EPX based on LC50 values (Pelosi et al. 2016; PPDB, 2018).

386 Growth is a key component of the life history parameter that directly influences the  
387 population dynamics in a way comparable to reproduction or survival. In the fields,  
388 earthworms are active between 3 and 7 months per year, generally in spring and autumn  
389 (Baker et al. 1992). In the present study, *A. caliginosa* individuals in the control soil took  
390 three months to become adult in optimal conditions (fed *ad libitum*, soil moisture of 70% of  
391 the WHC, temperature at 15°C, Bart et al. 2018). We can thus assume that there would be no  
392 more than one or two new generations of *A. caliginosa* per year. And it is worthwhile to  
393 underline that *A. caliginosa* grow and reproduce relatively fast (Bart et al. 2018, Bart et al.,  
394 2019c) compared to other species of earthworms such as anecic species (e.g. *Lumbricus*  
395 *terrestris*, Butt 1993; Pelosi et al. 2008. or *Octolasion cyaneum*, Butt 1993). A growth delay of  
396 about ten percent, as we observed for Swing<sup>®</sup> Gold, could have a strong impact on the  
397 population dynamics, with adults appearing significantly later in the year at a period which  
398 could not be optimal for the reproduction. It could be even more problematic with compounds  
399 such as copper that completely inhibit earthworm growth.

400

401

402

403

404 **Relevance of the toxicity analysis with TK-TD model.**

405

406 The two tested fungicides had very different toxicokinetics and toxicodynamics. For  
407 the copper fungicide, we showed a drastic inhibition of growth and a slow kinetics whereas  
408 the toxicokinetics for the Swing<sup>®</sup> Gold fungicide was very fast with moderate effects. These  
409 conclusions came from the analysis of the data with our TK-TD model which indicated the  
410 relevance of assuming either very fast kinetics or very strong effects. The conclusions  
411 regarding the kinetics were confirmed with the measurements of the fungicide internal  
412 concentrations at the end of the experiment.

413 We saw that the effects of the copper fungicide significantly depended on the age of  
414 the exposed individuals and the exposure duration. Indeed, the effects appeared earlier for  
415 small organisms (Age 1) compared to older organisms (Age 2 and 3) and that was accounting  
416 for by the dilution by growth and the influence of weight on the kinetics parameters. Thus, the  
417 difference between ages is fully explained by growth, with same parameters for kinetics and  
418 effects. This also supports that the use of a one compartment-model was satisfactory here. In  
419 some cases, for which, for instance, uptake rate depends on exposure concentrations because  
420 of saturation of the uptake, a model with more compartments could be necessary (Steen  
421 Redeker and Blust 2004). The difference in copper accumulation was certainly due to the  
422 longer exposure of individuals of Age 1 and 2, because low elimination rates mean that the  
423 longer the exposure the higher the accumulation. Still after more than 90 days of exposure, the  
424 plateau for accumulated concentration was thus still not reached. For the Swing<sup>®</sup> Gold  
425 fungicide, EPX and DMX accumulation in earthworms did not differ between the different  
426 ages and times of exposure. This is consistent with rapid kinetics, which implies no  
427 dependence between the accumulated concentration and the exposure duration. However,  
428 further work on the accumulation in earthworm are required to validate our work. The model,

429 which accounts for the exposure throughout time, take into account that the two active  
430 substances of the Swing<sup>®</sup> Gold fungicide degraded over time, and we were able to explain the  
431 toxicodynamics. Indeed, the total concentration in soil became lower than the NEC after 15  
432 days of exposure to the highest nominal concentration (SG3) and the growth was not affected  
433 anymore afterwards.

434 In this study, we hypothesized that the physiological mode of action of the two tested  
435 fungicides was an increase in the growth energy costs. Another possible effect on growth  
436 could have occurred through a decrease in the feeding rate. From a modeling point of view, it  
437 is tricky to assess which model would fit at best the observed data because they are very  
438 similar for small concentrations. To make the difference between these two modes of action,  
439 authors performed experiments in two feeding conditions, *ad libitum* and limited food  
440 conditions (Péry et al. 2003). Here, we only used *ad libitum* conditions. For copper, some  
441 elements in the literature support the assumption of an increase in growth cost. First, this was  
442 the mode of action found for chironomids (Péry et al. 2003). Moreover, the increase in growth  
443 energy costs could be linked to detoxification process. For example, it has been showed for  
444 *Lumbricus rubellus* that the production of metallothionein (MT) proteins increased 5-fold in  
445 soil contaminated with copper compared to a control soil (Stürzenbaum et al. 1998). The same  
446 has been shown with *E. fetida* exposed to cadmium (Brulle et al. 2007). The MT(s) are  
447 responsible for detoxification processes after an exposure to a metal contamination. The  
448 energy could be redirected to the production of such proteins in response to the  
449 contamination.

450

451 **Environmental implications.**

452

453 Earthworm growth appeared as a sensitive endpoint that should be taken into account  
454 in the ecological risk assessment of pesticide. First, the threshold values are, as for the  
455 reproduction (Neuhauser et al. 1985), lower than the LC50 value (based on survival).  
456 Moreover, growth can have a strong impact on population dynamics that determines the  
457 occurrence in the field and the related provided functions. The strength of our study relies on  
458 the ability of the TK-TD model to fit, with the same parameters, the data obtained for three  
459 different ages of exposure, despite apparent differences in the toxicodynamics. The NEC  
460 values provided are common to all ages of exposure and do not depend on the time of  
461 exposure as for ECx values. Finally, TK-TD models are interesting tools that can be used in  
462 the regulatory risk assessment to assess bioaccumulation and effects of pesticides as it is  
463 suggested for aquatic organisms in a recent EFSA report (EFSA, 2018).

#### 464 **Supplementary material**

465 Figure of the experimental design. Description of the growth energy-based model. Chemical  
466 analysis methods, and results.

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473

#### 474 **References**



475 Amosse J, Bart S, Pery ARR, Pelosi C (2018) Short-term effects of two fungicides on  
476 enchytraeid and earthworm communities under field conditions. *Ecotoxicology* 27(3):  
477 300-312.

478 Baas J, Jager T, Kooijman B (2010) Understanding toxicity as processes in time. *Sci Total*  
479 *Environ* 408: 3735–3739

480 Baker GH, Barrett R, Grey-Gardner R, Buckerfield JC (1992) The life history and abundance  
481 of the introduced earthworms *Aporrectodea trapezoides* and *A. caliginosa* (AnneUda:  
482 Lumbricidae) in pasture soils in the Mount Lofty Ranges, South Australia. *Aus J Ecol* 17:  
483 177-188

484 Bart S, Amossé J, Lowe CN, Péry ARR, Mougin C, Pelosi C (2018) *Aporrectodea caliginosa*,  
485 a relevant earthworm species for a posteriori pesticide risk assessment: Current knowledge  
486 and recommendations for culture and experimental design. *Environ Sci Pollut Res* 25:  
487 33867

488 Bart S, Barraud A, Amosse J, Pery ARR, Mougin C, Pelosi C (2019c) Effects of two common  
489 fungicides on the reproduction of *Aporrectodea caliginosa* in natural soil. *Ecotoxicol*  
490 *Environ Saf* 181: 518-524.

491 Bart S, Laurent C, Péry ARR, Mougin C, Pelosi C (2017) Differences in sensitivity between  
492 earthworms and enchytraeids exposed to two commercial fungicides. *Ecotoxicol Environ*  
493 *Saf* 140: 177-184

494 Bart S, Pelosi C, Barraud A, Péry ARR, Cheviron N, Grondin V, Mougin C and  
495 Crouzet O (2019a) Earthworms Mitigate Pesticide Effects on Soil Microbial  
496 Activities. *Front Microbiol* 10:1535

497 Bart S, Pelosi C, Péry ARR (2019b) Towards a better understanding of the life cycle of the  
498 earthworm *Aporrectodea caliginosa*: new data and energy-based modelling. *Pedobiologia*.  
499 *In press*

500 Bengtsson J, Ahnstrom J, Weibull AC (2005) The effects of organic agriculture on  
501 biodiversity and abundance: a meta-analysis. *J Appl Ecol* 42: 261-269

502 Bertrand M, Barot S, Blouin M, Whalen J, de Oliveira T, Roger-Estrade J (2015) Earthworm  
503 services for cropping systems. A review. *Agron Sustain Dev* 35: 553–567

504 Blouin M, Hodson ME, Delgado EA, Baker G, Brussaard L, Butt KR, Dai J, Dendooven L,  
505 Peres G, Tondoh JE, Cluzeau D, Brun JJ (2013) A review of earthworm impact on soil  
506 function and ecosystem services. *Eur J Soil Sci* 64: 161–182

507 Boag B, Palmer LF, Neilson R, Legg R, Chambers SJ (1997). Distribution, prevalence and  
508 intensity of earthworm populations in arable land and grassland in Scotland. *Ann Appl*  
509 *Biol* 130(1): 153-165

510 Booth LH, Heppelthwaite VJ, O'Halloran K (2000) Growth, development and fecundity of the  
511 earthworm *Aporrectodea caliginosa* after exposure to two organophosphates. *N Z Plant*  
512 *Prot* 53: 221-225

513 Booth LH, O'Halloran K (2001) A comparison of biomarker responses in the earthworm  
514 *Aporrectodea caliginosa* to the organophosphorus insecticides diazinon and chlorpyrifos.  
515 *Environ Toxicol Chem* 20(11): 2494-2502

516 Boström U, Lofs-Holmin A (1996) Annual population dynamics of earthworms and cocoon  
517 production by *Aporrectodea caliginosa* in a meadow fescue ley. *Pedobiologia* 40(1): 32-  
518 42

519 Brulle F, Mitta G, Leroux R, Lemiere S, Lepretre A, Vandebulcke F (2007) The strong  
520 induction of metallothionein gene following cadmium exposure transiently affects the  
521 expression of many genes in *Eisenia fetida*: a trade-off mechanism? *Comp Biochem*  
522 *Physiol C Toxicol Pharmacol* 144: 334-41

523 Brun LA, Maillet J, Richarte J, Herrmann P, Remy JC (1998) Relationships between  
524 extractable copper, soil properties and copper uptake by plants in vineyards soils. Environ  
525 Pollut 102: 151–161

526 Butt, K.R., 1993. Reproduction and growth of three deep-burrowing earthworms  
527 (Lumbricidae) in laboratory culture in order to assess production for soil restoration.  
528 Biology and Fertility of Soils, 16(2), pp.135-138.

529 Chabauty F, Pot V, Bourdat-Deschamps M, Bernet N, Labat C, Benoit P (2016) Transport of  
530 organic contaminants in subsoil horizons and effects of dissolved organic matter related to  
531 organic waste recycling practices. Environ Sci Pollut Res Int 23 (7): 6907–6918

532 Couto RR, Benedet L, Comin JJ, Belli Filho P, Martins SR, Gatiboni LC, Radetski M, Valois,  
533 CM, Ambrosini VG, Brunetto G (2015) Accumulation of copper and zinc fractions in  
534 vineyard soil in the mid-western region of Santa Catarina, Brazil. Environ Earth Sci  
535 73(10): 6379–6386

536 Curry JP, Doherty P, Purvis G, Schmidt O (2008) Relationships between earthworm  
537 populations and management intensity in cattle-grazed pastures in Ireland. Appl Soil Ecol  
538 39(1): 58-64

539 E-phy (2017a) <https://ephy.anses.fr/ppp/swing-gold>

540 E-phy (2017b) <https://ephy.anses.fr/ppp/styrocuivre-df>

541 Efron B (1979) Bootstrap methods: another look at the jackknife. Annals stat 7 (1): 1-26

542 EU (2008) European Union Risk Assessment Report. Voluntary risk assessment of copper,  
543 copper II sulphate pentahydrate, copper(I)oxide, copper(II)oxide, dicopper chloride  
544 trihydroxide. Summary of the Terrestrial Effect Chapter. PNEC derivation for copper in  
545 the terrestrial environment.

546 Goussen B, Parisot F, Beaudouin R, Dutilleul M, Buisset-Goussen A, Péry ARR, Bonzom JM  
547 (2013) Consequences of a multi-generation exposure to uranium on *Caenorhabditis*  
548 *elegans* life parameters and sensitivity. *Ecotoxicology* 22: 869–878

549 Hartenstein F, Hartenstein E, Hartenstein R (1981) Gut load and transit time in the earthworm  
550 *Eisenia foetida*. *Pedobiologia* 22: 5–20

551 Heckmann LH, Baas J, Jager T (2010) Time is of the essence. *Environ Toxicol Chem* 29:  
552 1396–1398

553 Helling B, Reinecke SA, and Reinecke AJ (2000) Effects of the fungicide copper oxychloride  
554 on the growth and reproduction of *Eisenia fetida* (Oligochaeta). *Ecotoxicol Environ Saf*  
555 46: 108-116

556 Hole DG, Perkins AJ, Wilson JD, Alexander IH, Grice PV, Evans AD (2005) Does organic  
557 farming benefit biodiversity? *Biol Conserv* 122: 113-130

558 Holmstrup M, Ostergaard IK, Nielsen A, Hansen BT (1991) The relationship between  
559 temperature and cocoon incubation-time for some lumbricoid earthworm species.  
560 *Pedobiologia* 35(3): 179-184

561 ISO (International Organisation for Standardization) (2012a) Soil Quality - Effects of  
562 Pollutants on Earthworms - Part 1: Determination of Acute Toxicity to *Eisenia fetida*/  
563 *Eisenia andrei*. No. 11268-1. Geneva

564 ISO (International Organization for Standardization) (2012b) Effects of pollutants on  
565 earthworms (*Eisenia fetida*). Part 2: determination of effects on reproduction—No. 11268-  
566 2. Geneva

567 Jager T, Crommentuijn T, Van Gestel CAM, Kooijman SALM (2004) Simultaneous  
568 modeling of multiple endpoints in life-cycle toxicity tests. *Environ Sci Technol* 38: 2894–  
569 2900

570 Jager T, Gudmundsdottir EM, Cedergreen N (2014) Dynamic modeling of sublethal mixture  
571 toxicity in the nematode *Caenorhabditis elegans*. Environ Sci Technol 48, 7026–7033

572 Khalil MA, Abdel-Lateif HM, Bayoumi BM, van Straalen NM (1996) Analysis of separate  
573 and combined effects of heavy metals on the growth of *Aporrectodea caliginosa*  
574 (Oligochaeta; Annelida), using the toxic unit approach. Appl Soil Ecol 4: 213-219

575 Klobucar GIV, Stambuk A, Srut M, Husnjak I, Merkas M, Traven L, Cvetkovic Z (2011)  
576 *Aporrectodea caliginosa*, a suitable earthworm species for field based genotoxicity  
577 assessment? Environ Pollut 159: 841–849

578 Kooijman SALM, Bedaux JJM (1996) The Analysis of Aquatic Toxicity Data. Vu University  
579 Press, Amsterdam.

580 Kooijman SALM (1986). Energy budgets can explain body size relations. J. Theor. Biol. 121,  
581 269–282

582 Kooijman SALM (2000) Dynamic energy and mass budgets in biological systems.  
583 Cambridge: Cambridge University Press, 423 pages.

584 Kooijman SALM (2010) Dynamic Energy Budget theory for metabolic organization.  
585 Cambridge University Press, Great Britain ISBN 9780521131919.

586 Lowe CN, Butt KR (2007) Earthworm culture, maintenance and species selection in chronic  
587 ecotoxicological studies: a critical review. Eur J Soil Biol 43: S281–S288

588 Ma W (1984) Sublethal toxic effects of copper on growth, reproduction and litter breakdown  
589 activity in the earthworm *Lumbricus rubellus*, with observations on the influence of  
590 temperature and soil pH. Environ Pollut Ser. A 33, 207219.

591 Ma LQ, Rao GN (1997) Chemical fractionation of cadmium, copper, nickel, and zinc in  
592 contaminated soils. J Environ Qual 26: 259-264

593 McDonald J, Gaston L, Elbana T, Andres K, Crandfield E (2013) Dimoxystrobin sorption and  
594 degradation in sandy loam soil: impact of different landscape positions. *Soil Sci* 178: 662–  
595 670

596 Neuhauser EF, Loehr RC, Malecki MR, Milligan DL, Durkin PR (1985) The toxicity of  
597 selected organic-chemicals to the earthworm *Eisenia fetida*. *J Environ Qual* 14: 383–388

598 OECD (Organization for Economic Co-operation and Development) (2004) Earthworm  
599 Reproduction Test (*Eisenia fetida*/*Eisenia andrei*) (No. 222). OECD Guidelines for the  
600 Testing of Chemicals. OECD, Paris, France.

601 OECD (Organization for Economic Co-operation and Development) (1984) Guideline for the  
602 testing of chemicals. No. 207. Earthworm, acute toxicity tests. OECD Publishing, Paris

603 EFSA PPR Panel (EFSA Panel on Plant Protection Products and their Residues), Ockleford  
604 C, Adriaanse P, Berny P, Brock T, Duquesne S, Grilli S, Hernandez-Jerez AF, Bennekou  
605 SH, Klein M, Kuhl T, Laskowski R, Machera K, Pelkonen O, Pieper S, Smith RH,  
606 Stemmer M, Sundh I, Tiktak A, Topping CJ, Wolterink G, Cedergreen N, Charles S,  
607 Focks A, Reed M, Arena M, Ippolito A, Byers H and Teodorovic I (2018) Scientific  
608 Opinion on the state of the art of Toxicokinetic/Toxicodynamic (TKTD) effect models for  
609 regulatory risk assessment of pesticides for aquatic organisms. *EFSA J* 16(8):e05377

610 Paoletti MG, Sommaggio D, Favretto MR, Petruzzelli G, Pezzarossa B, Barbaferi M (1998)  
611 Earthworms as useful bioindicators of agroecosystem sustainability in orchards and  
612 vineyards with different inputs. *Appl Soil Ecol* 10: 137–150

613 Pelosi C, Bertrand M, Makowski D, Roger-Estrade J (2008) WORMDYN: A model of  
614 *Lumbricus terrestris* population dynamics in agricultural fields. *Ecol Modell* 218: 219-234

615 Pelosi C, Joimel S, Makowski D (2013) Searching for a more sensitive earthworm species to  
616 be used in pesticide homologation tests - a meta-analysis. *Chemosphere* 90: 895–900

617 Pelosi C, Lebrun M, Beaumelle L, Chevion N, Delarue G, Nelieu S (2016) Sublethal effects  
618 of epoxiconazole on the earthworm *Aporrectodea icterica*. Environ Sci Pollut Res 23(4):  
619 3053-3061

620 Péry ARR, Bedaux JJM, Zonneveld C and Kooijman SALM (2001) Analysis of bioassays  
621 with time-varying concentrations. Wat Res 35: 3825-3832

622 Péry ARR, Ducrot V, Mons R, Garric J (2003) Modelling toxicity and mode of action of  
623 chemicals to analyse growth and emergence tests with the midge *Chironomus riparius*.  
624 Aquat Toxicol 65: 281-292

625 Péry ARR, Flammarion P, Vollat B, Bedaux JJM, Kooijman SALM, Garric J (2002) Using a  
626 biology-based model (DEBtox) to analyse bioassays in ecotoxicology: opportunities and  
627 recommendations. Environ Toxicol Chem 21: 459-465

628 PPDB (Pesticide Properties DataBase) (2018)  
629 <https://sitem.herts.ac.uk/aeru/ppdb/en/Reports/246.htm>

630 Sijm DTHM, van der Linde A (1995) Size-dependent bioconcentration kinetics of  
631 hydrophobic organic chemicals in fish based on diffusive mass transfer and allometric  
632 relationships. Environ Sci Technol 29: 2769-2777

633 Sijm DTHM, Verberne ME, de Jonge WJ, Pärt P, Opperhuizen A (1995) Allometry in the  
634 uptake of hydrophobic chemicals determined in vivo and in isolated perfused gills.  
635 Toxicol Appl Pharmacol 131: 130-135

636 Sims RW, Gerard BM (1999) Earthworms. Earthworms: keys and notes for the identification  
637 and study of the species. Synopses of the British fauna. New series; 31. Shrewsbury: Field  
638 Studies Council. p169

639 Soetaert K, Petzoldt T, Setzer RW (2010) Solving Differential Equations in R: Package  
640 deSolve. Journal of Statistical Software, 33(9), 1--25. URL  
641 <http://www.jstatsoft.org/v33/i09/> DOI 10.18637/jss.v033.i09

642 Springett JA, Gray RAJ (1992) Effect of repeated low-doses of biocides on the earthworm  
643 *Aporrectodea caliginosa* in laboratory culture. Soil Biol Biochem 24(12): 1739-1744

644 Spurgeon DJ, Svendsen C, Kille P, Morgan AJ, Weeks JM (2004) Responses of earthworms  
645 (*Lumbricus rubellus*) to copper and cadmium as determined by measurement of juvenile  
646 traits in a specifically designed test system. Ecotox Environ Saf 57: 54-64

647 Steen Redeker E, Blust R (2004) Accumulation and toxicity of cadmium in the aquatic  
648 oligochaete *Tubifex tubifex*: a kinetic modeling approach. Environ Sci Technol 38(2):  
649 537-543.

650 Stürzenbaum SR, Kille P, Morgan AJ (1998) The identification, cloning and characterization  
651 of earthworm metallothionein. FEBS Letters 431-437-442