

The soil-dwelling earthworm Allolobophora chlorotica modifies its burrowing behaviour in response to carbendazim applications

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1	The soil-dwelling earthworm Allolobophora chlorotica modifies its burrowing
2	behaviour in response to carbendazim applications
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25 Abstract

26 Carbendazim-amended soil was placed above or below unamended soil. Control tests 27 comprised two layers of unamended soil. Allolobophora chlorotica earthworms were 28 added to either the upper or the unamended soil. After 72 hours vertical distributions 29 of earthworms were compared between control and carbendazim-amended 30 experiments. Earthworm distributions in the carbendazim-amended test containers 31 differed significantly to the 'normal' distribution observed in the control tests. In the 32 majority of the experiments earthworms significantly altered their burrowing 33 behaviour to avoid carbendazim. However, when earthworms were added to an upper 34 layer of carbendazim-amended soil they remained in this layer. This non-avoidance is 35 attributed to 1) the earthworms' inability to sense the lower layer of unamended soil 36 and 2) the toxic effect of carbendazim inhibiting burrowing. Earthworms modified 37 their burrowing behaviour in response to carbendazim in the soil. This may explain 38 anomalous results observed in pesticide field trials when carbendazim is used as a 39 control substance.

40

Keywords: earthworm, *Allolobophora chlorotica*, burrowing, avoidance behaviour,
carbendazim, pesticide, field trial

43

1. Introduction

47	The fungicide carbendazim is known to be highly toxic to earthworms and is
48	recommended for use as the reference substance in standardised guidelines for testing
49	the effects of pesticides on earthworms in field situations (ISO, 1999). However,
50	results using carbendazim in field trials have been highly variable (Römbke et al.,
51	2004; Ellis, 2008). This paper reports a study into the behavioural response of
52	Allolobophora chlorotica to carbendazim as part of a wider investigation into this
53	variability.
54	
55	Carbendazim has limited movement in the soil profile and studies have recorded up to
56	97 % of the applied total to remain in the upper 5 cm of the soil profile (Ellis, 2008;
57	Jones et al., 2004; Holmstrup, 2000). The exposure of earthworms to carbendazim in
58	the field will therefore, in part, be determined by their vertical distribution and their
59	ability to detect the chemical and modify their vertical burrowing behaviour as a
60	consequence of this. A field study (Römbke et al., 2004) showed the vulnerability of
61	earthworms to the toxic effects of carbendazim to differ between species. This
62	difference was attributed to the different feeding preferences of the species and their
63	distribution in the soil profile. Species which typically feed on vegetation at the
64	surface of the soil where carbendazim concentration was highest, including Lumbricus
65	terrestris and Lumbricus rubellus had higher mortality than geophageous species
66	including Apporectodea caliginosa which were not dependent on the surface for food
67	and subsequently had lower exposure to the chemical (Römbke et al., 2004). While
68	certain species may be more vulnerable due to their feeding behaviour, earthworms
69	can occupy a range of depths in the soil profile and can adjust their burrowing depth

70	behaviour based on soil conditions (Edwards and Bohlen, 1996). The geophageous
71	species A. chlorotica for example is typically found above a depth of 8 cm when soil
72	conditions are favourable but will burrow to below 8 cm to avoid extremes of
73	temperature or dry soil at the surface (Gerard, 1967). In earthworm avoidance studies,
74	in which earthworms are given a choice between horizontally adjacent soils, (usually
75	a control, contaminant free soil and a contaminant bearing soil, e.g. Yeardley et al.,
76	1996; Natal da Luz et al., 2004; Environment Canada, 2007; ISO, 2008) the
77	earthworm species Eisenia andrei (Loureiro et al., 2005) and Eisenia fetida (Garcia et
78	al., 2008), have been shown to significantly avoid carbendazim and benomyl at
79	concentrations $\geq 1 \text{ mg kg}^{-1}$. However, for chemicals such as carbendazim which have
80	limited mobility through the soil profile, the most significant concentration gradient
81	encountered in the field will be in the vertical plane and a key question is whether or
82	not earthworms are able to modify their behaviour to avoid such chemicals.
83	Horizontal avoidance studies provide useful information on the ability of earthworms
84	to detect and respond to adverse concentrations of chemicals but they do not provide
85	information on whether this avoidance driver is sufficient for earthworms to modify
86	their normal behaviour to avoid such chemicals.
87	
88	The aim of this study was therefore to determine whether the presence of carbendazim
89	led to a modification of the burrowing behaviour of the earthworm A. chlorotica.
90	
91	2. Method
92	
93	2.1. Earthworm species

95	Allolobophora chlorotica is a widely abundant species in the UK. It was selected as a
96	suitable species for the study as it occupies a range of depths in the soil profile, is
97	geophageous, so is not dependent on the soil surface for feeding (Edwards and
98	Bohlen, 1996) and is known to adjust its burrowing depth in response to unfavourable
99	conditions (Gerard, 1967). Earthworms were collected by manual digging and hand
100	sorting soil from a pasture field at the University of Reading farm at Sonning,
101	Berkshire UK and kept in a 3:1 mix of sandy loam soil and sphagnum peat moss at a
102	temperature of 15 °C until the test.
103	
104	2.2. Test substance
105	
106	Delsene 50 Flo, obtained from Nufarm UK Ltd. Belvedere, Kent, UK, was selected as
107	a suitable test substance for the study as it is a commercially available water-based
108	suspension concentrate containing carbendazim at a concentration of 500 g 1^{-1} . The
109	Delsene 50 flo was diluted using deionised water to a concentration of 46 mg l^{-1} .
110	
111	2.3. Test soil
112	
112	
113	Kettering loam, a commercially available sandy loam soil obtained from Broughton
114	Loam, Kettering, UK (Table 1 for soil properties) was used in the avoidance studies.
115	The soil was air dried and sieved to $< 2 \text{ mm}$ prior to use. A carbendazim
116	concentration of 8 mg kg ⁻¹ was used as significant avoidance behaviour was observed
117	in previous studies using similar concentrations (Loureiro et al., 2005; Garcia et al.,

118 2008). Using the relationship of Jänsch et al. (2006) which assumes a soil density of 1

119	500 kg m ⁻³ this concentration is approximately twice that in soil after the typical
120	application rate of 4 kg ha ⁻¹ used in field trials (ISO, 1999). The diluted carbendazim
121	suspension was mixed thoroughly with the soil using a house-hold mixer (Kenwood
122	A907D), to give a soil moisture content of 60 % of the soil water holding capacity.
123	For the control soil, Kettering loam was mixed with deionised water only. The
124	moisture contents of the carbendazim-treated and control soil were the same.
125	
126	2.4. Experimental procedure
127	
128	2.4.1. Arrangement of soils
129	
130	The test containers comprised two sections, one section containing the carbendazim-
131	amended soil and the other the clean unamended soil. The two sections were stacked
132	vertically and earthworms were able to move freely between the two soils. The
133	behavioural response of A. chlorotica to carbendazim was tested with the soils in two
134	arrangements (Figure 1). The first arrangement (Field arrangement) reflected
135	carbendazim application in the field with the carbendazim-amended soil at the top and
136	the unamended soil below. In the second arrangement (Alternative arrangement) the
137	carbendazim-amended soil formed the bottom section. Control tests (with unamended
138	soil in both sections) were used to determine the natural distribution of earthworms
139	without the influence of carbendazim. The test containers were designed to account
140	for the typical burrowing behaviour of A. chlorotica. Allolobophora chlorotica
141	usually form temporary horizontal burrows in the upper 8 cm of the soil profile
142	(Edwards and Bohlen, 1996). The test containers comprised two open-ended,

143 translucent PVC cylinders wrapped in black adhesive tape to exclude light, 8 cm high

144	and with a diameter of 7.5 cm. Four hundred grams (dry weight equivalent) of soil
145	were added to each container which were placed on top of each other. The top of the
146	upper container was covered with mesh (1 mm size) to prevent individuals escaping
147	and to allow light onto the surface of the soil. The bottom of the lower container was
148	closed to prevent earthworm escape. The test containers were kept in a temperature
149	controlled room at 18 °C with a photo period of 12:12 hours (light:dark).
150	
151	2.4.2. Earthworm addition
152	
153	Earthworms were added to the containers in one of 2 ways. In both methods the
154	earthworms were added 24 hours after the soil had been mixed and added to the
155	containers. Five replicates were used per soil arrangement with ten individuals used
156	per replicate. Five replicates were also used for each control. The tests were run for 72
157	hours to ensure that earthworms had sufficient time to burrow into the soil. After 72
158	hours the sections were separated using a card divider and the number of individuals
159	in each section determined by hand sorting.
160	
161	Method 1 (Fig. 1): Earthworms were added to the soil surface at the top of the test
162	container. Thus when the carbendazim-amended soil was in the upper container
163	earthworms were added to the upper surface of the 8 cm thick carbendazim-amended
164	soil. This method allowed us to assess the response of the earthworms when they
165	experienced direct dermal contact with carbendazim-amended soil.
166	
167	Method 2 (Fig. 2): This was intended to be more representative of a field scenario
168	where carbendazim would be sprayed onto the soil surface. Earthworms were initially

169	added to unamended soil and allowed to acclimatise for 24 hours before the
170	carbendazim-amended soil was added, either above or below the unamended soil.
171	This method allowed us to assess whether A. chlorotica would modify its burrowing
172	behaviour in response to either an over-lying or under-lying layer of carbendazim-
173	amended soil. In this method A. chlorotica began the test in two different positions in
174	the test container (either the top or bottom section), dual controls were used for both
175	arrangements. For each arrangement, 5 replicates plus 5 dual controls were used.
176	
177	2.5. Statistical analysis
178	
179	The Fisher exact test in Minitab version 15 was used to determine if earthworms were
180	significantly avoiding the carbendazim-amended soil. This test allows the distribution
181	in the avoidance test to be compared with the normal distribution of earthworms in the
182	controls (Natal da Luz, 2004).
183	
184	3. Results
185	
186	In each arrangement earthworms were observed to burrow rapidly into the soil to
187	which they had been added. For both Method 1 (Fig. 3) and Method 2 (Figs. 4 and 5)
188	in the control experiments there was an uneven distribution of A. chlorotica between
189	the two sections. The greatest proportion of individuals had burrowed to the bottom
190	section, below a depth of 8 cm. Therefore when analysing results from the
191	carbendazim-amended experiments the relative proportion of earthworms in the
192	bottom section was compared to the proportion in the bottom section in the controls.
193	Results indicate that A. chlorotica does indeed modify its natural burrowing behaviour

194 to avoid carbendazim and that exposure to carbendazim inhibits earthworm

195 burrowing.

197	Method 1: Compared to the control earthworms appeared to have modified their
198	burrowing behaviour in response to carbendazim in both the Field and Alternative
199	arrangements. In the Field arrangement with the carbendazim-amended soil at the
200	top, the majority of individuals were found in the carbendazim-amended soil (0.84 \pm
201	s.e 0.05, $n = 5$) and had not burrowed into the unamended soil below (Fig. 3). The
202	proportion in the bottom soil was significantly lower than the control (P < 0.05). In
203	two of the replicates, one earthworm was found dead at the surface of the test soil. In
204	the Alternative arrangement, with the carbendazim-amended soil at the bottom, a
205	significantly lower proportion of A. chlorotica were found in the bottom soil
206	compared to the control (0.42 \pm s.e 0.05, n = 5) (P < 0.05) and had not burrowed into
207	the carbendazim-amended soil below (Fig. 3).
208	
209	Method 2 In the Field arrangement (carbendazim-amended soil at the top) a
210	significantly higher proportion of individuals were found in the bottom section
211	compared to the control (P < 0.05). As this distribution differed significantly from the
212	control, burrowing behaviour appears to have been modified in response to the
213	presence of carbendazim (Figure 4). This was also apparent in the Alternative
214	arrangement in which the carbendazim-amended soil formed the lower section. The
215	majority of individuals were not found in the bottom section but instead remained in

- 216 the unamended soil in the top section (0.78, s.e. \pm 0.07, n = 5) (Figure 5). The
- 217 proportion in the bottom soil was significantly lower than in the control (P < 0.05).
- 218

219 **4. Discussion**

220

221 Although we did not analyse the carbendazim-amended soil used in the experiments, 222 subsamples of the same well-mixed carbendazim-amended soil were used in all the 223 experiments so we can be confident that concentrations of carbendazim were the same 224 in all experiments. The aim of the investigation was to determine whether the 225 presence of carbendazim led to a modification of burrowing behaviour and the lack of 226 precise concentration data does not prevent this. In the current experiments no flow of 227 water occurred through the soil (which had the same moisture content in both the 228 carbendazim-free and carbendazim-amended parts) so it is highly unlikely that the 229 carbendazim would have been redistributed within the soil due to movement of soil 230 solution. Additionally studies by Ellis et al. (In press), Jones et al. (2004) and 231 Holmstrup (2000) indicate that carbendazim is immobile in soils due to very strong 232 partitioning onto the solid phase relative to the solution phase. Thus we can assume 233 that any difference in earthworm behaviour between experiments is due to either 234 exposure to the carbendazim-amended soil (Method 1, Field arrangement) or the 235 detection and consequent avoidance of the carbendazim-amended soil (Method 1, 236 Alternative arrangement and Method 2 Field and Alternative arrangements). 237

We propose two alternate explanations for the modified burrowing behaviour observed in the *Field arrangement* (the majority of individuals remaining in the carebendazim-amended soil held in the top half of the containers compared to the control in which earthworms added to the upper surface burrowed down into the soil in the bottom half of the containers, Fig. 3). The first possible explanation is that earthworms remained in the carbendazim-amended soil because there was no gradient

"leading" them to the unamended soil below, i.e. the earthworms were unaware of the 244 245 less challenging conditions in the bottom half of the test containers. However, as the 246 earthworms in the control experiment clearly showed a preference for burrowing into 247 the bottom half of the test containers this explanation can not be the complete story. 248 Thus it seems more likely that exposure to the carbendazim disrupted the burrowing 249 ability of the earthworms when the earthworms were placed on the upper surface of the carbendazim-amended soil. Carbendazim has been shown to disrupt conduction in 250 251 the giant nerve fibre of earthworms, which is linked with earthworm mobility (Drewes 252 et al., 1987), thus it may be possible that carbendazim reduced the ability of the 253 earthworms to burrow. Unfortunately it is not possible to convert the concentrations 254 used in the filter paper tests by Drewes et al. to equivalent soil concentrations. However, the concentration of carbendazim used in this study (8 mg kg⁻¹) is similar to 255 concentrations at which both acute and chronic toxic effects have been observed in 256 other studies. Van Gestel et al. (1992) reported an LC50 of $4.7 - 6.9 \text{ mg kg}^{-1}$ and 257 sublethal effects on growth at 6.0 mg kg⁻¹ and reproduction at 1.92 mg kg-1 for E. 258 *andrei*. Ellis et al (2007) reported LC50s in the range $2.47 - 16.00 \text{ mg kg}^{-1}$ for E. 259 fetida. Ellis et al. (In press) reported a reduction in surface activity of L. terrestris at 260 surface carbendazim concentrations of c. 2.5 mg kg⁻¹. A third explanation (which we 261 reject as it is contradicted by the avoidance of the carbendazim-amended soil by 262 263 earthworms in the Alternative arrangement) is that the earthworms remained in the 264 carbendazim-amended soil because conditions were preferable to those in the unamended soil. 265

266

267 By adding the earthworms to the unamended soil rather than the amended soil

268 (Method 2), field conditions were more closely represented with the earthworms

269 initially in carbendazim-free soil. The results of Method 2 confirm that the 270 earthworms in the Alternative arrangement of Method 1 modified their burrowing 271 behaviour to avoid the carbendazim-amended soil. In the *Field arrangement* of 272 Method 2 (carbendazim-amended soil in the top half of the containers) significantly 273 more earthworms were found in the bottom half of the containers relative to the 274 control. In the Alternative arrangement (carbendazim-amended soil in the bottom half 275 of the containers) significantly fewer earthworms were found in the bottom half of the 276 containers relative to the control. This indicates that the presence of carbendazim in 277 the soil led to the earthworms altering their burrowing behaviour to avoid burrowing 278 into the carbendazim-amended soil. This finding is consistent with those of Loureiro 279 et al. (2005) and Garcia et al. (2008) who observed avoidance of carbendazim at concentrations $> 1 \text{ mg kg}^{-1}$ for *E. andrei* and *E.* fetida respectively in horizontal 280 281 avoidance tests. The avoidance behaviour by earthworms of potentially toxic 282 chemicals is well documented (e.g. Environment Canada, 2007 and references 283 therein) and is most likely triggered by the detection of chemical substances that 284 render the soil inhospitable by chemoreceptors located on the prosomium or buccal 285 epithelium (Edwards and Bohlen, 1996). Thus earthworms would be able to detect the 286 boundary between the carbendazim-free / carbendazim-amended soils and avoid 287 entering the treated soil. Similar responses resulting in earthworms not burrowing in 288 soils of unsuitable pH have been reported in the literature (e.g. Laverack, 1961). Thus 289 avoidance can occur before an earthworm is in an inhospitable soil and experiments 290 like the ones carried out here are a valid measure of earthworm avoidance behaviour 291 despite, unlike current standardised tests (e.g. Environment Canada, 2007; ISO, 2008) 292 all the earthworms being in the same portion of the test chambers at the start of the experiment. 293

5. Conclusion

297	Carbendazim is used as a reference substance in standardised guidelines for testing
298	the effects of pesticides on earthworms in field situations. If carbendazim application
299	fails to reduce field populations of earthworms to between 40 and 80 % of those in
300	control plots the trial is declared invalid (ISO, 1999). Our results indicate that
301	earthworms may be able to avoid the effects of carbendazim by modifying their
302	burrowing behaviour. This should be borne in mind when determining earthworm
303	population size after application of test chemicals. It is possible that a failure to
304	recover an acceptable number of earthworms from trial plots, which would be
305	interpreted as excess mortality may simply be due to avoidance of treated soil by
306	earthworms. Therefore in field trials when sampling after application of pesticides and
307	control substances care should be taken to sample both outside the treated plot and to
308	sufficient depths so that earthworms exhibiting such behaviour are included in counts
309	of earthworm numbers.

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381

Figure captions

383

Figure 1. Diagrammatic representation of method 1 for assessing vertical avoidance
behaviour of earthworms in which earthworms are added to the upper surface of the
upper soil.

Figure 2. Diagrammatic representation of method 2 for assessing vertical avoidance
behaviour of earthworms in which earthworms are added to the upper surface of the
unamended soil.

390 Figure 3. Mean proportional distribution of *Allolobophora chlorotica* in test

391 containers in the upper and lower soils in the *Field* (carbendazim-amended soil in the

392 upper section) and *Alternative* (carbendazim-amended soil in the bottom section)

393 arrangements with A. chlorotica being added to the upper soil upper surface (Method

394 1). Error bars = standard deviation, n = 5. * = significantly different from the Control.

395 Figure 4. Mean proportional distribution of *Allolobophora chlorotica* in test

396 containers in the upper and lower soils in the *Field arrangement* (carbendazim-

397 amended soil in the upper section) with A. chlorotica being added to the unamended

398 soil (Method 2). Error bars = standard deviation, n = 5. * = significantly different

from the control.

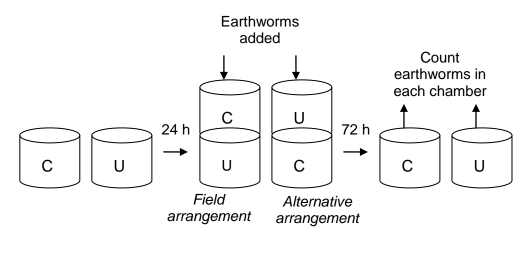
400 Figure 5. Mean proportional distribution of *Allolobophora chlorotica* in test

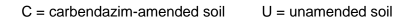
401 containers in the upper and lower soils in the Alternative arrangement (carbendazim-

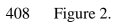
402 amended soil in the bottom section) with A. chlorotica being added to the unamended

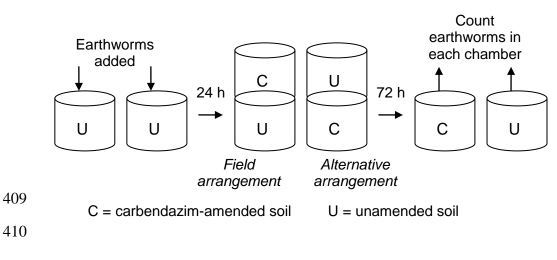
403 soil (Method 2). Error bars = standard deviation, n = 5. * = significantly different

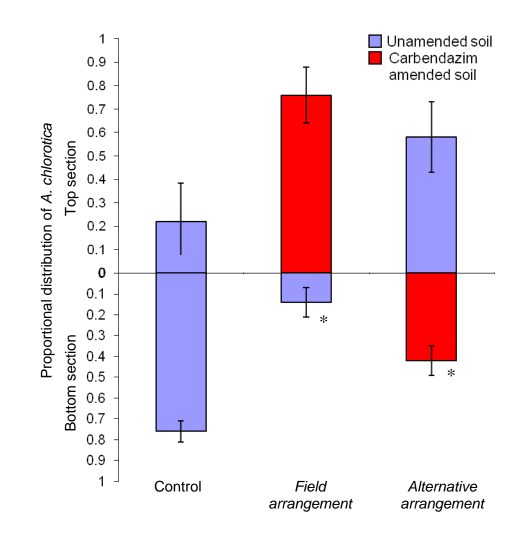
404 from the Control.

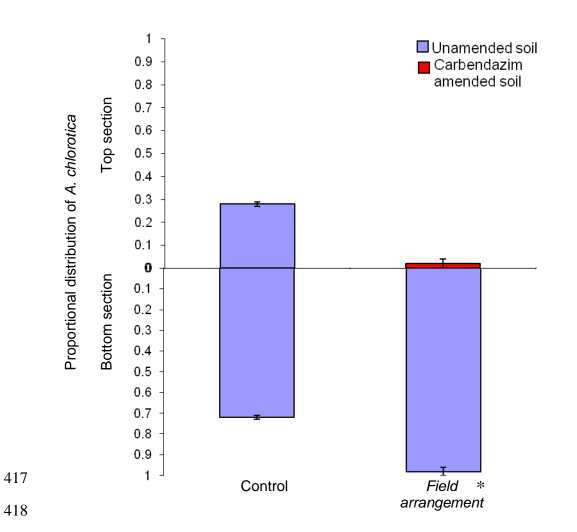


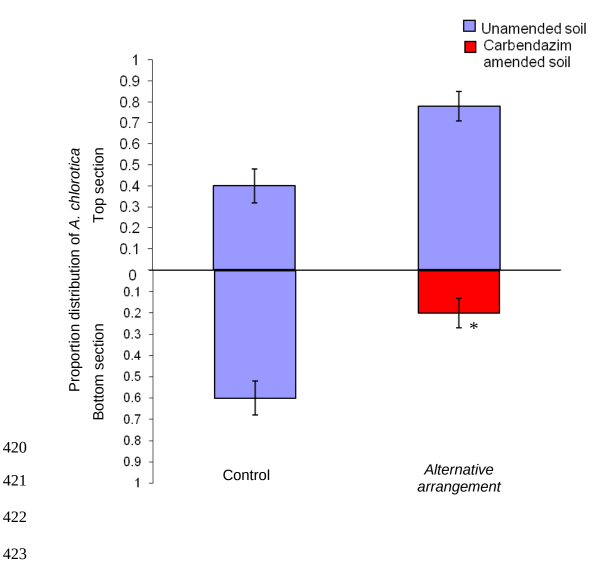












425 Table 1. Selected mean chemical and physical properties of the Kettering loam test

426 soil (n = $3 \pm$ standard error).

Soil property		
рН	6.2 <u>+</u> 0.2	
Organic matter content / %	7.06 ± 0.09	
Texture	11.8 <u>+</u> 1.3 % clay	
	21.7 <u>+</u> 0.3 % silt	
	66.9 ± 1.0 % sand	
Water holding capacity / %	29 <u>+</u> 4	