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1	DNA sequence variation and methylation in an arsenic tolerant earthworm
2	population
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29 Abstract

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31 Evidence is emerging that earthworms can evolve tolerance to trace element enriched soils. However, few 32 studies have sought to establish whether such tolerance is determined through adaption or plasticity. Here we 33 report results from a combined analysis of mitochondrial (cytochrome oxidase II, COII), nuclear (amplified 34 fragment length polymorphism, AFLP) variation and DNA methylation in populations of the earthworm 35 Lumbricus rubellus from sites across an abandoned arsenic and copper mine. Earthworms from the mine site 36 population demonstrated clear arsenic tolerance in comparison to a naïve strain. COII and AFLP results 37 suggest that L. rubellus from the unexposed and the adapted populations comprises two cryptic lineages 38 (Linages A and B) each of which was present across all of the sites. AFLP analysis by lineage highlighted 39 variations associated with soil metal/metalloid concentrations (most clearly for Lineage A) suggesting a 40 genetic component to the observed tolerance. The methylation sensitive AFLP (Me-AFLP) identified a high 41 genome methylation content (average 13.5%) in both lineages. For Lineage A, Me-AFLP analysis did not 42 identify a strong association with soil arsenic levels. For Lineage B, however, a clear association of 43 methylation patterns with soil arsenic concentrations was found. This suggests that Lineage B earthworms 44 utilise epigenetic mechanisms to adapt to the presence of contamination. These fundamentally different 45 genetic adjustments in the two clades indicate that the two lineages employ distinct adaptive strategies 46 (genetic or epigenetic) in response to arsenic exposure. Mechanisms driving this variation may be founded 47 within the colonisation histories of the lineages.

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54 1. Introduction

55 Many invertebrate species are able to maintain viable populations in polluted soils where total and 56 potentially bioavailable metal/metalloid concentrations greatly exceed toxicity values (e.g. LC_{50} s) for known 57 naïve (and so sensitive) populations (laboratory strains). This suggests that under trace element exposure, 58 some invertebrate populations develop metal tolerance through behaviour or physiological adaptive traits 59 (Posthuma and Van Straalen, 1993; Van Straalen and Roelofs, 2007). Mechanisms underpinning this 60 tolerance have in some cases been shown to involve heritable changes in coding or promoter regions of metal 61 efflux pumps (Callaghan and Denny, 2002) and thiol-rich peptides involved in sequestration (glutathione-S-62 transferases, phytochelatins, and metallothioneins) (Janssens et al., 2007; Vatamaniuk et al., 2005). In other 63 cases, however, the mechanisms underlying tolerance remain unknown and/or unstudied.

64

65 For earthworms, one of the most functionally important of soil taxa (Lavelle et al., 1997), indirect evidence 66 for metal tolerance is provided by the fact that earthworms can be collected from soils containing residue 67 levels that significantly exceed toxic effect concentrations (Spurgeon and Hopkin, 1999a, b). However, 68 difficulties in extrapolating toxicity data between the laboratory and field due to, for example, contaminant 69 aging and speciation (Arnold et al., 2003; Arnold et al., 2007), mean that the mere presence of earthworms in 70 these polluted soils is not alone confirmation that adaptation has occurred (Davies et al., 2003; Donner et al., 71 2010). More directly in relation to tolerance, studies with successive generations of Eisenia fetida selected 72 for tolerance to Zn over two generations found changes in the shape of concentration response relationships 73 for survival that were indicative of tolerance development (Spurgeon and Hopkin, 2000). In the field, 74 Langdon et al. (1999) noted that L. rubellus living in arsenic and copper polluted soil at two abandoned 75 arsenic mines (Devon Great Consols, Carrock Fell) could survive in arsenic-spiked soil that was acutely 76 toxic to earthworms from a clean site. This tolerance was preserved when the mine populations were reared 77 on clean soil over two generations, suggesting a genetic basis for this phenotype (Langdon et al., 2009).

78

79 Despite indications of trace metal and metalloid tolerance in earthworms, the extent to which there is a 80 genetic and/or physiological basis of this trait has not been fully investigated. A study of isozyme specific 81 polymorphisms within *L. rubellus* populations known to be adapted to combined metal and flooding stress failed to identify adaptive variation (Simonsen and Klok, 2010), although the results of this study should be treated with some caution as enzymes known to be related to metal tolerance were not targeted. The only study that has, to date, identified a potential genetic basis for tolerance to adverse soil conditions in earthworms is that for *L. rubellus* living at a lead/zinc mine located at Cwmystwyth, mid Wales. For this population, Andre et al. (2010) used mitochondrial (COII) and amplified fragment length polymorphism (AFLP) genotyping to demonstrate that a mine spoil associated population showed little genetic overlap (in AFLP profile) with individuals within populations at two less polluted sites.

89

90 While the assumption often is that individual/population survival is based on selection for increased 91 tolerance, there is evidence emerging that the plastic responses driven by chemical influences on the 92 epigenome may also be an important mechanism of adaptation (Mirouze and Paszkowski, 2011; Ren et al., 93 2011; Seong et al., 2011). Among the many epigenetic mechanisms, DNA methylation represents a key 94 response. Given that earthworms have been recorded to possess a 13% methylated cytosine content in DNA 95 (Regev et al., 1998), the potential for mediation of adaptive tolerance through epigenetic DNA methylation 96 should be considered. Here, we report a combined toxicological and genetic study, using mitochondrial 97 (COII) and nuclear (AFLP) and DNA methylome analysis, for the earthworm L. rubellus sampled at sites of 98 different metal pollution status within an As-contaminated mine complex - the Devon Great Consols (DGC) 99 site in the UK. For the study, we sampled earthworms from a number of sites within DGC including one 100 (Site 2 in this study) from which tolerant populations previously studied by Langdon et al. (2009; 1999) were 101 collected. Adjacent and distant reference sites were also sampled. That arsenic, a major contaminant at the 102 site, has been reported to induce epigenetic changes including hypo-and hyper-methylation of DNA (Ren et 103 al., 2011; Zhong and Mass, 2001) makes the site particularly suited for the analysis of DNA methylation 104 responses in earthworms. Initially the collected populations were screened to confirm that the tolerance 105 previously reported for populations at study Site 2 was applicable to earthworms inhabiting this and other 106 collection sites located in the mining area. Genetic analyses were then undertaken using these populations. 107 The hypothesis tested was that L. rubellus populations from polluted sites that show evidence of tolerance. 108 would include individuals with mitochondrial or nuclear genotypes and/or DNA methylation patterns that 109 were distinct from those of intolerant earthworms from (adjacent) unpolluted locations.

110 2. Materials and Methods

111 2.1 Site description, sampling and soil characterisation: This study was conducted at the abandoned Devon 112 Great Consols mine complex located in the Tamar Valley, Devon, South-West UK (UK Ordnance Survey. 113 Map coordinates for mine centre: SX426733 – N50:32:52 W4:13:25). This mine was worked for copper and 114 arsenic from 1844-1900 and from 1915-1930. Across the site, the spoil from various extraction processes 115 remain. The soils established on these wastes contain highly elevated concentrations of trace elements, 116 including arsenic and copper. Earthworms (L. rubellus) were sampled from six locations in the region of the 117 Devon Great Consols mine, Four locations (Sites 1-4) were situated on the mine and waste handling area (see 118 Fig. 1). This included a location (Site 2 i.e. close to the area where arsenic was processed using the calciner 119 method) from which the adapted population studied by Langdon et al. (2009; 1999) was collected. Two clean 120 reference site populations were also sampled. These were at a site adjacent to the contaminated area, but 121 which itself was not greatly enriched in arsenic and copper (Site Control - SC) and a site some 20 km distant 122 from DGC which was outside the geological area of arsenic rich soils present in the Tamar Valley (Off-Site 123 Control - OSC) (UK Ordnance Survey. Map coordinates SX 418901 N50:68:89 W4:24:03).

124

125 At each site, approximately 30 fully clitellate adult L. rubellus were collected by digging and hand-sorting 126 over two consecutive days in September 2010. To ensure that genome methylation patterns were not 127 influenced by handling stress, all earthworms were washed and blotted dry on-site and then snap frozen in 128 liquid nitrogen. Triplicate soil samples from surface to 5 cm depth were also collected from each location. 129 These were subsequently oven dried at 80°C and sieved through a 2 mm mesh to remove large roots and 130 stones. Total concentrations of arsenic, barium, calcium, cadmium, chromium, copper, iron, magnesium, 131 nickel, strontium and zinc were determined in a 1 g sample of the processed soil following an aqua regia 132 digestion protocol (Arnold et al., 2008). Digests were analysed on a Perkin Elmer Optima 7300 DV 133 inductively coupled plasma optical emission spectrometry instrument. For quality control, an in house 134 reference traceable to BCR-143R (Commission of the European Communities, Community Bureau of 135 Reference) was included with each batch of digestions. Measured concentrations were always greater than 136 75% of reference values and were above 95% for As. Organic matter content of each soil sample was 137 measured by proxy using loss on ignition following combustion at 500°C (Rowell, 1994) and pH was

138 quantified by electrode from a 1:5 volume soil:water mix (International Standards Organisation, 2005).

139

140 2.2. Toxicity tests to identify putative arsenic tolerance

141 To identify potential tolerance, a 14 day exposure to a single pre-determined arsenic concentration was 142 undertaken to compare survival patterns of earthworms from sites located within and adjacent to the 143 DGC complex to those for a known naïve population. The soil concentration used for this assay was 144 derived from a preliminary study conducted to assess survival of the naïve population at 150 and 145 300 mg/kg arsenic. The earthworms used were taken from a culture established from a field 146 collected population (Lasebo BV, Nijkerkerveen, The Netherlands). At each tested concentration, 147 15 replicate containers, each including 200 g dry weight of a clay loam soil (Broughton Loams, 148 Kettering, UK) (see Spurgeon et al., 2003), were spiked with sodium arsenate solution (Santa Cruz 149 Biotechnology Inc., Santa Cruz, California, US) to give the required metalloid concentration and a 150 soil moisture content of approximately 50% of field capacity. After a one week stabilisation period, 151 one adult L. rubellus was added to each replicate and kept at 13 ± 1 °C under constant light for seven 152 days. Earthworms were observed daily and mortality recorded. Based on these findings, a screening 153 concentration of 300 mg/kg arsenic was selected for the definitive tolerance assay, since this 154 concentration resulted in progressive mortality of the naïve earthworms over the exposure period. 155 Thus the definitive assay was conducted using the 300 mg/kg concentration with 15 earthworms 156 from each of the DGC sites (Sites 1-4) and the SC reference population. The exposure was extended 157 to 14 days to allow the potential to identify survival patterns in more tolerant populations.

158

159 2.3 Mitochondrial cytochrome oxidase II (mtCOII)) sequencing:

DNA was purified from ~10 mg of tissue from the anterior of each individual using the DNAzol reagent
(Life Technologies, Paisley, UK). PCR amplification of the cytochrome oxidase II (COII) mitochondrial
gene made use of forward (TAGCTCACTTAGATGCCA) and reverse (GTATGCGGATTTCTAATTGT)

163 primers and was conducted following Andre et al. (2010). PCR products were assessed electrophoretically prior to purification and sequencing using ABI PRISM[®] BigDye v3.1 Terminator technology (Applied 164 165 Biosystems, USA). Obtained sequences were aligned by ClustalW prior to tree construction using the 166 Maximum Likelihood (ML) and Bayesian methods in Mega v5.01 and MRBAYES v3.2, respectively. ML 167 estimation incorporated the Tajima-Nei model, supported by bootstrap analyses over 1000 iterations. 168 Bayesian analyses were conducted using a General Time Reversible model with a proportion of invariable 169 sites and a gamma-shaped distribution over 2 independent runs. Four Markov Chains were run over 2 million 170 iterations and sampled every 1000 generations, with the first 500 trees discarded as burn-in. Both 171 phylogenetic estimates incorporated outlier sequences from Lumbricus castaneus and Lumbricus terrestris as 172 well as sequences that represent previously recognised L. rubellus clades (Andre et al., 2010).

173

174 2.4 AFLP and methylation sensitive AFLP profiling: A combined AFLP and Me-AFLP protocol was 175 optimised in a pilot methylation study and was based on parallel use of methylation- and non-methylation-176 sensitive restriction enzymes (HapI and MspI) to treat DNA samples prior to primer ligation and 177 amplification (Xiong et al., 1999). Both HapII and MspI recognize a CCGG sequence; however, while MspI 178 is able to cut methylated recognition sites (as well as unmethylated ones), HapII is unable to cut at such 179 locations when they are methylated (i.e. only unmethylated recognition sites are cut). The extent of 180 methylation of restriction sites can therefore be ascertained by recording bands amplified by MspI but not 181 HapII. Such bands can be used to compare individual methylation patterns. AFLP analysis was conducted 182 for individuals from the six collection locations using pre-selective primers and analysis on an Applied 183 Biosystems 3130 x 1 fragment analyser (Andre et al., 2010). Cumulative AFLP fragment profiles were 184 transformed to a binary form and principal coordinates (PCO) analysis used to visualise the genetic 185 relationship between individuals using GenAlEx 6.4.1.

187 **3. Results**

188 Soil analyses highlighted the extent and severity of the arsenic (and copper) contamination at DGC. Arsenic 189 and copper levels were greatly elevated in soils from all sites on the mining area (Sites 1-4), with cobalt also 190 higher than SC and OSC soils by at least a factor of two at the sites (Table 1). The most polluted arsenic soil 191 (Site 4) contained almost 20,000 mg/kg of arsenic, over 900 mg/kg copper and also elevated cobalt, 192 cadmium and lead levels. The remaining three mine spoil contaminated sites each contained over 4000 193 mg/kg As and over 500 mg/kg copper. As expected, the lowest concentrations of arsenic and copper and 194 other trace metals were found at the SC and OSC reference sites. Levels at SC were in the 300 mg/kg As 195 range, a concentration still elevated above background arsenic levels for British soils (Emmett et al., 2010). 196 OSC soils contained arsenic levels consistent with these background concentrations.

197

198 Measured site soil LOI and pH values are presented in Table 1. Whilst the pH of all four sites located on the 199 mine area and the OSC was similar acidic (pH 4.1 - 4.8), the SC site had a pH of 5.6. Overall, there was 200 minimal pH variation between sites, and no correlation with soil arsenic or copper concentration (Pearson 201 correlation, p > 0.05). The absence of a correlation indicates that soil pH influences are unlikely to confound 202 attempts to link genetic variation to soil contaminant levels. For LOI, the lowest values in the mining site 203 soils (4.2 - 17.2%) were found at Sites 1 and 4, while the remaining two soils had higher LOI values (29.7 -204 49.6%). This may reflect the vegetation of the sites: open in the case of Sites 1 and 4, wooded at Sites 2 and 205 3. The two control site soils had %LOI levels intermediate within the range of the two pairs of mine 206 sampling locations.

207

Exposure of the naïve population to 300 mg/kg of arsenic in soil resulted in a progressive mortality, culminating in only 7% survival after 14 days of exposure (Fig. 2). In the SC population progressive mortality over time was also seen. This, however, proceeded at a slower rate than for the naïve earthworms, culminating in 46% survival after 14 days. These variations in mortality rates resulted in different LT₅₀ estimates from Weibull models fits (SigmaPlot 12.0) for the naïve and SC populations; these being 5.3 (95% Confidence Intervals 4.9-5.6) and 12.4 (95% Confidence Intervals 11.6-13.3) days respectively. In the four DGC mine site populations there was observable mortality in the Site 1 population, although 73% survival

after 14 days was higher than for either the naïve or SC earthworms. Populations from the remaining three
DGC mine sites show low mortality, with 100% survival for Site 2 and 4 earthworms and 85% survival for
Site 3 earthworms.

218

219 The mtDNA COII analysis indicated the presence of two distinct lineages (A and B) within the sampled L. 220 rubellus (Fig. 3a). The two cryptic lineages show a 18% and 14% genetic divergence from L. castaneus and 221 L. terrestris respectively. Average difference between lineages was 10.3%. Internal within the lineages, the 222 Lineage A earthworms have a maximum 1.4% genetic difference, while for Lineage B earthworms this was 223 0.06%. This high level of divergence between the two major lineage branches identifies L. rubellus as a 224 complex of cryptic lineages as found previously (Andre et al., 2010). A comparison of the frequency of 225 lineage occurrence at each sampled site identified differences in population COII haplogroup composition. 226 Populations at two sites, Site 4 and Site OSC, included 90% or more of individuals from Lineage B; while in 227 contrast the Site 3 populations included 76% of Lineage A individuals. The remaining three sites each had an 228 approximately equal proportion of each lineage, with Lineage A slightly dominant at Site SC (57%) and Site 229 1 (54%) and Lineage B at Site 2 (64%). That both lineages were found at all sites, often in similar 230 proportions, and also that the two sites showing greatest lineage selection (dominance of Lineage B at both 231 Site 4 and Site OSC) included both the most and least arsenic contaminated soils, is indicative of an absence 232 of a mitochondrial lineage association with soil contamination status.

233

234 Standard AFLP analysis conducted using MspI (which cuts at all recognition sites independent of 235 methylation status) reemphasised the presence of two L. rubellus lineages as indicated by the mitochondrial 236 markers. All mine site earthworms fell clearly into one of the two major lineages, but apparent inter-lineage 237 individuals were observed among SC and OSC earthworms. These hybrids show AFLP genotypes 238 intermediate between the two lineages on PC1 and divergent on PC2 (Fig. 3b). The presence of hybrids is in 239 agreement with previous observation of AFLP profiles in L. rubellus (Andre et al., 2010). The dominance of 240 the lineage effects within a PCO analysis of the AFLP data meant it was not possible to visualise site effects 241 within the complete data-set. Consequently independent lineage-based analyses were conducted (n.b. 242 putative hybrid individuals were excluded from these analyses).

243

Within Lineage A, PCO highlighted a site dependent effect on AFLP marker patterns. Within the PC1 and PC2 scores plot, SC earthworms were clearly separated from earthworms collected from Sites 2 and 3, with the Site 1 individuals intermediate and closer to the SC earthworms (Fig. 4a). Scores for PC2 (and also PC3), but not PC1 within the PCO were significantly correlated with site soil arsenic concentration (Pearson correlation p < 0.01). This significant association suggests that soil arsenic concentrations, as well as possibly the concentration of other co-correlated metals such as Cu, are an important driver of genome structure in Lineage A *L. rubellus* across the DGC site and surrounding area.

251

252 For Lineage B PCO analysis did not identify a separation of populations within a PC1 and PC2 scores plot, 253 although a partial separation of Site 1 and 4 was evident (data not shown). Both of these populations, 254 however, overlap with profiles from the SC and OSC earthworms within this plot. Correlation of PC1 and 255 PC2 scores with soil arsenic concentrations were not significant. Only for the PC3 score was a significant 256 correlation found (Pearson correlation p < 0.02) indicating a weak separation underpinned by the distribution 257 particularly of the Site 2 and Site 4 individuals on this component (Fig. 4b). These results identify that while 258 soil metals such as arsenic and correlated elements are a driver for genome structure in Lineage B, these 259 factors are less important than for Lineage A with effects only observed for the lower contribution PCs.

260

261 To assess the patterns of genome methylation in individual earthworms, a second AFLP analysis was 262 conducted using the MspI methylation sensitive restriction enzyme. Me-AFLP indicated that the genome of 263 L. rubellus had an approximate 13.5% methylated cytosine (m5C) residue content. Across the mine sampling 264 locations, the average extent of genome methylation ranged from 10.6% in earthworms at Site 1 to 22.1% for 265 Site 4. Even though the highest average genome methylation content was at the most arsenic polluted site, 266 the fact that earthworms from the two reference sites had intermediate average methylation levels (SC 267 19.4%, OSC 13.2%) meant there was no clear correlation (Pearson correlation p > 0.05) between methylation 268 level and soil arsenic concentration. This suggests that in the mine soils, arsenic does not have a strong 269 global hyper- or hypo-methylation effect for the resident earthworms.

271 Within the Me-AFLP analysis the presence of two distinct L. rubellus lineages was reconfirmed. 272 Consequently, lineage-specific Me-AFLP profiles were analysed, with the hybrid individuals excluded. For 273 the Lineage A PCO analysis, a segregation of individuals collected from Site 2 and Site 4 was identified 274 within the PC1 and PC2 score plot (Fig. 4c). The remaining sites showed substantial overlap between 275 individual profiles. Correlations of PC1, PC2 and PC3 scores with measured soil arsenic concentration were 276 non-significant in all cases (Pearson correlation p > 0.05). This suggests that soil arsenic was not the 277 principal driver of methylation pattern difference between individuals. For Lineage B, there was a partial 278 separation of profiles of earthworms from the SC and OSC locations from individuals collected from each 279 sampled mine site population (Fig. 4d). Correlation of PC1, PC2 and PC3 scores with measured site soil 280 arsenic concentration indicated a significant correlation for the first principle component (Pearson correlation 281 p < 0.02). This indicates that soil arsenic (and co-correlated trace metals) represents a potentially significant 282 driver of earthworm genome methylation status for lineage B earthworms.

283

285 **4. Discussion**

286 Soil contamination by mineral extraction, fossil fuel consumption, waste disposal and pesticide use is a 287 common problem (Hall et al., 2006). Among trace elements, arsenic represents one of the greatest hazards 288 because of its widespread distribution and toxicity to humans and wildlife (Chen et al., 2009; Thomas et al., 289 2001). The toxicity of arsenic has been established for earthworms. Meharg et al. (1998) determined an LC_{50} 290 of approximately 100 mg/kg As for Lumbricus terrestris after 8 days and Fischer and Koszorus (1992) found 291 that a 25 mg/kg potassium arsenate exposure reduced growth and cocoon production in *Eisenia fetida*. For L. 292 *rubellus*, Langdon et al. (2001) found an LC_{50} of 96 mg/kg As for a clean site population, although 293 populations from mine sites (including DGC) gave higher values (up to 1,510 mg/kg) suggesting tolerance. 294 Building on this work, Langdon et al. (2009) revealed that the adaptation in the mine site earthworms was 295 maintained when earthworms were bred for two generations on clean soil. Cross-tolerance to copper was also 296 found (Langdon et al., 2001).

297

298 In the test to assess the presence of potential tolerance in L. rubellus collected from the DGC line complex 299 area sites, there was a clear indication that the populations inhabiting the DGC site locations substantially 300 enriched in arsenic display a tolerance phenotype. Earthworms sampled from the populations at Site 1-4 all 301 showed low mortality on exposure to a soil arsenic concentration that induced acute toxicity in earthworms 302 from a naïve population and also in the SC reference population. Interestingly the different rates of mortality 303 in naïve and SC earthworms, as highlighted by differences in LT50s for these populations suggest that SC 304 earthworms possess a partial arsenic tolerant phenotype. This may be related to the presence of arsenic 305 concentrations that greatly exceed accepted background concentrations in SC soil (Emmett et al., 2010).

306

Tolerance to chemical exposure can classically take two forms. Most simply, it can be the result of phenotypic plasticity. In this case, exposure to a substance upregulates biochemical pathways (e.g. metal binding proteins, mono-oxygenases and multi-drug resistance transporters), which work to detoxify or eliminate the substance. If the exposure is removed, upregulation of detoxification systems can persist, predisposing individuals to deal with a future chemical challenge. This plasticity has been widely reported in human subjects subjected to long-term drug exposure (Stewart and Badiani, 1993) and also in species

exposed to toxicants in the field (Rajamohan and Sinclair, 2009; Romach et al., 2000). Maintenance of
elevated protein levels and the widely reported effects of stressor exposure on the epigenome (MartinezZamudio and Ha, 2011), including arsenic (Ren et al., 2011), can provide a mechanism through which such
tolerance may be temporally conserved.

317

318 A second mechanism of tolerance development exploits adaptive variation within populations. There is good 319 evidence that this kind of adaptive selection can occur in response to long term chemical exposure. One 320 example is driven by the selection of alleles coding for amino acids associated with active sites of 321 detoxification enzymes. Pesticide resistance is frequently underpinned by this mechanism, with polymorphic 322 cytochrome P450 genes often the selection target (Karunker et al., 2008; Miyo and Oguma, 2010). For 323 metals, selection for metallothionein promoter alleles and other trans-acting genetic factors has been found to 324 underpin cadmium tolerance in the collembolan Orchesella cincta (Janssens et al., 2007; Roelofs et al., 2006; 325 van Straalen et al., 2011).

326

327 Characterisation of metallothionein promoter alleles of earthworms collected from metalliferous and 328 unpolluted soils has so far failed to detect adaptive variation (Stürzenbaum et al., 2004). With evidence for 329 targeted selection absent, a logical next step is to move to genome wide analysis (Baird et al., 2008; 330 Hohenlohe et al., 2010). Using a combined approach applying mitochondrial genotyping and conventional 331 and methylation-sensitive AFLPs, an analysis of both genotypic and epigenetic associations of the confirmed 332 adapted and putative reference populations of L. rubellus with different metal/metalloid exposure histories 333 was conducted. The aim was to assess the basis of the arsenic tolerance observed in the toxicity test. The 334 mitochondrial genotyping and AFLP profiling (using both methylation sensitive and insensitive enzymes) all 335 indicated that L. rubellus comprises two distinct lineages that differ by over 10% in their mitochondrial COII 336 sequence. This reflects the presence of two cryptic lineages within the morphospecies (Andre et al., 2010). 337 Hybrid individuals were found although only at the two reference sites. This prevalence in uncontaminated 338 soils does not support a role of pollution in the breakdown of species boundaries as found by Vonlanthen et 339 al. (2012).

341 As in a previous study with L. rubellus from polluted landscapes (Andre et al., 2010), there was no evidence 342 of a lineage or intra-lineage haplotype association with either polluted or unpolluted sites. This supports the 343 decision to move to a more detailed analysis of population structure. The AFLP analysis conducted for 344 Lineage A indicated a clear separation of earthworms between sites, with the most important principal 345 components associated with soil pollution status. For Lineage B, an influence of soil arsenic on AFLP profile 346 was also found, albeit in this case within one of the more minor principle components (PC3). Such 347 associations that link genetic distance to pollution status have been observed in previous field studies of 348 aquatic invertebrates (Martins et al., 2009) and for both genetic units of the phylogeographically divergent 349 metallophyte Arabidopsis halleri (Pauwels et al., 2012). Such relationships point to a genetic component that 350 may underpin the previous observations of arsenic tolerance in L. rubellus collected at Site 3 by Langdon et 351 al. (2009; 1999), especially given the high frequency of Lineage A individuals at this site.

352

353 Although sequence driven differentiation between populations can clearly be important, there is emerging 354 evidence that epigenetic effects can also play a role in adaptation to local environmental conditions. Known 355 epigenetic mechanisms include DNA methylation, histone modifications, and small interfering (siRNA), and 356 micro RNAs (miRNA). Of these, DNA methylation has so far been most widely studied in animals (Suzuki 357 and Bird, 2008). Studies have identified that metals and metalloids can perturb DNA methylation including 358 hypomethylation by Cd (Takiguchi et al., 2003) and targeted gene silencing via hypermethylation by Ni (Lee 359 et al., 1995). For arsenic, the potential competition with DNA for methyl groups for respectively methyl 360 metabolites and DNA modification can create an interplay between hypomethylation (Arita and Costa, 2009; 361 Zhao et al., 1997) and hypermethylation (Jensen et al., 2008) in arsenic toxicology (Ren et al., 2011).

362

To date relatively little is known about the role of DNA methylation as a component of adaptive variation in invertebrate organisms. Studies on a range of invertebrate species have highlighted extensive variation in the 5-methyl cytosine content of the genome (Regev et al., 1998). Thus, while some species, including the nematode *Caenorhabditis elegans* and fruitfly *Drosophila melanogaster*, have low to negligible 5-methyl cytosine levels (Bird, 2002; Regev et al., 1998), some taxa possess methylation levels in the 10-15% range. Me-AFLP indicated an approximate 13.5% methylated cytosine (m5C) residue content in the *L. rubellus* 369 genome. This represents a high level of DNA methylation for an invertebrate species, but is consistent with 370 previous results for the earthworm *Aporrectodea caliginosa* (Regev et al., 1998). This suggests that DNA 371 methylation may have an important role in annelids, although to date relatively little is known about how 372 such methylation is controlled. For example, a study on the marine annelid species *Chaetopterus* 373 *variopedatus* was able to identify a protein that had a high homology to known invertebrate 374 methyltransferases, but could not confirm a role of this protein in 'de-novo' methylation of double stranded 375 DNA (del Gaudio et al., 1999).

376

377 On exposure to arsenic (and co-contaminant metals), an analysis of methylation patterns using the MeAFLP 378 approach showed a site-specific influence. For Lineage A earthworms, separation between sites for the 379 MeAFLP profiles was seen. This separation could not, however, be significantly associated with soil arsenic 380 concentration as was the case for the standard AFLP analysis for this Lineage. This may indicate that other 381 soil, biotic and local scale climatic factors may instead be modifying the epigenome. For Lineage B 382 earthworms, pattern of DNA methylation could be significantly related to soil arsenic levels, suggesting a 383 potential role of trace element exposure, although it is also feasible that environmental factors (e.g. soil 384 texture, soil moisture, food availability), that are co-correlated to soil pollutant levels, could also be 385 important. Evidence from detailed analysis of stress associated genes, such as metallothionein in the snail 386 Helix pomatia, has already identified the presence of genomic regions that confer a high potential for 387 epigenetic regulation indicating a potential role for epigenetic mechanisms in metal responses (Egg et al., 388 2009). Further, in D. melanogaster stress exposure has been shown to result in epigenetic heterochromatic 389 disruption that is transmissible in a non-Mendelian fashion (Seong et al., 2011). The association of DNA 390 methylation patterns with arsenic exposure observed here suggests a potential role of epigenetic mechanisms 391 in stress adaptation in earthworms that concur with the evidence available for other taxa.

392

To extend the understanding of the role of genetic and epigenetic modification, a fruitful avenue for extending this novel study from a strong associative appreciation to a mechanistic understanding of arsenicmediated molecular-genetic adaptations would entail assaying the transcription levels of specific genes known to be involved in metal/metalloid trafficking and metabolism. Moreover, establishing whether

397 epigenetic marks are preferentially targeted to such genes and their regulatory regions in earthworms 398 exposed to elevated levels of methylation-modifying arsenic in their native field soils is also a matter of 399 priority. Such work clearly has the potential to link genotype to phenotype in adapted populations, so 400 providing insight into the functional basis of adaptive traits in a key soil dwelling taxon.

401

402 The variation in lineage-specific responses observed across the genome and epigenome raises the intriguing 403 prospect that the two L. rubellus cryptic lineages may employ different strategies to response to long-term 404 arsenic exposure. The strong AFLP based separation of Lineage A earthworms in relation to soil arsenic 405 concentrations across major PCs suggests that in this Lineage substantial genome modification has occurred 406 as a result of long-term exposure. In contrast, the evidence for sequence modification is somewhat less 407 compelling in Lineage B and therefore changes in genome methylation status seem to play a role in 408 facilitating plasticity in response to soil arsenic concentration as indicated by the Me-AFLPs. Previous 409 studies have identified differences in sensitivity between closely related lineages or species to chemical 410 exposure. An example is the role of biotransformation capacity for determining the sensitivity of *Capitella* 411 capitata "species" to PAH exposure (Bach et al., 2005). However, to date we are not aware of any studies 412 that have identified such divergent genome responses to chemical exposure within two genetic lineages of a 413 known morphospecies. The detailed basis for the evolution of distinct genetic and/or epigenetic mechanisms 414 that drive arsenic adaptation in the two L. rubellus lineages, thus, emerge as potential models that could be 415 further exploited to understand species plasticity in response to long-term chemical stress.

416

417 The genetic structure evident within the putative L. rubellus lineages is consistent with expectations in 418 relation to survival within glacial refugia and subsequent recolonisation, as has been demonstrated for other 419 species (Hewitt, 1999; Provan and Bennett, 2008). Patterns of recolonisation during the Holocene, including 420 recent human-mediated dispersal, may have resulted in different lineages reaching the DGC area over 421 different timeframes. Andre et al. (2010) inferred that the two L. rubellus lineages have very different 422 evolutionary histories with Lineage A representing a stationary population that has experienced multiple 423 introductions and bottleneck episodes with expansion estimated to have occurred about 250,000 years BP, 424 while Lineage B comprises an unimodal mismatch distribution with an estimated post-glacial population

- 425 expansion time of approximately 17,000 years BP. It is perhaps this differential in the timescale for
- 426 adaptation to local arsenic contamination that has determined the lineage specific balance between adaptive
- 427 variation and plasticity for the two lineages at sites across the DGC mine.
- 428

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660 Table 1. Summary data for trace element concentrations, soil pH and wt% loss on ignition (%LOI) for soil samples collected from sites across the
 661 Devon Great Consols mine complex located in south-west England. For site locations see Fig. 1. Values are means of triplicate subsamples, standard
 662 deviations are given in brackets.

	Al	As	Ва	Cd	Со	Cr	Cu	Fe	Mn	Ni	Pb	Zn	рΗ	%LOI
	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg	mg/kg		
Site 1	9430	4620	47.2	0.7	19.5	10.8	529	48000	430	16.4	61	134	4.5	4.2
	(6720)	(2020)	(32.4)	(0.4)	(11.5)	(9.8)	(266)	(24600)	(335)	(15.8)	(35)	(74)		(0.1)
Site 2	6060	5220	70.0	< 0.2	52.7	6.9	606	99200	802	28.4	191	164	4.1	49.6
	(720)	(470)	(2.5)		(4.8)	(0.3)	(27)	(9300)	(96)	(2.8)	(21)	(22)		(0.4)
Site 3	17300	6270	45.9	0.18	25.7	17.8	2647	79600	630	22.3	225	277	4.8	29.7
	(3900)	(1010)	(8.4)	(0.09)	(5.4)	(3.6)	(606)	(2600)	(135)	(4.0)	(53)	(43)		(0.3)
Site 4	13600	19200	71.2	10.2	< 3.6	20.0	910	65900	262	9.5	148	63	4.6	17
	(2060)	(3470)	(8.9)	(1)		(3.3)	(120)	(10600)	(33)	(1.8)	(15)	(8)		(0.0)
SC	21500	310	45.5	0.41	< 3.6	31.7	107	45800	585	27.5	68	140	5.6	17.2
	1800	(70)	(2.8)	(0.4)		(4.2)	(16)	(6350)	(103)	(9.1)	(8)	(34)		(0.2)
OSC	7840	<50	37.0	< 0.2	< 3.6	10	14	1420	427	3.7	21	69	4.4	14.5
	(4770)		(20.7)			(6.0)	(8)	(8650)	(234)	(2.8)	(12)	(35)		(0.2)

665 FIGURE LEGENDS

666

667 Figure 1: Aerial images showing the location of the Devon Great Consols mine site in the South West UK
668 (top right insert panel) and locations of the 5 sampling locations (Site 1-4 and Site SC) situated in the area
669 on, and immediately adjacent to, the Devon Great Consols mine workings.

670

Figure 2: Temporal patterns of survival of *L. rubellus* collected at five locations of contrasting geochemistry
(4 polluted and 1 site reference) within the Devon Great Consols mine complex and surrounding area and a
known naïve population following exposure to 300 mg/kg of arsenic in a spiked clay loam soil over 14 days.

Figure 3: Mitochondrial and nuclear analysis of *L. rubellus* population structure and corresponding mitochondrial mismatch distributions of collected *L. rubellus*. Panel A: shows a phylogenetic tree of mitochondrial COII genotype showing branching of major lineage (Left and right hand branches of the network are denoted Lineage B & A respectively) and the numbers of individuals from each site within the lineages. Panel B: AFLP multi-locus profiling PCO analysis showing individuals from the six sample stations. Lineage A individuals cluster to the right on PC1, Lineage B to the left. Hybrids (found at SC and OSC only) lie between and above the two Lineage Groups.

682

Figure 4: Nuclear genome analysis of *L. rubellus* collected at six sites (4 polluted and 2 reference) of contrasting geochemistry within the Devon Great Consols mine complex and surrounding area. Panel (i) shows the result of a PCO of AFLP profiles for *L. rubellus* unambiguously ascribed to Lineage A, Panel (ii) shows the result of a PCO of AFLP profiles for *L. rubellus* unambiguously ascribed to Lineage B, Panel (iii) shows the result of a PCO of methylation sensitive AFLP analysis of *L. rubellus* unambiguously ascribed to Lineage B, Panel (iii) shows the result of a PCO of methylation sensitive AFLP analysis of *L. rubellus* unambiguously ascribed to Lineage B, Panel (iii) unambiguously ascribed to Lineage B.

690 Fig. 1.



694 Fig. 2.



698 Fig. 3.



701 Fig. 4.

