

# Genetic, epigenetic and microbiome characterisation of an earthworm species (Octolasion lacteum) along a radiation exposure gradient at Chernobyl

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#### 29 Abstract

30 The effects of exposure to different levels of ionising radiation were assessed on the genetic, 31 epigenetic and microbiome characteristics of the "hologenome" of earthworms collected at sites 32 within the Chernobyl exclusion zone (CEZ). The earthworms Aporrectodea caliginosa (Savigny, 33 1826) and Octolasion lacteum (Örley, 1881) were the two species that were most frequently found 34 at visited sites, however, only O. lacteum was present at sufficient number across different 35 exposure levels to enable comparative hologenome analysis. The identification of morphotype O. 36 lacteum as a probable single clade was established using a combination of mitochondrial 37 (cytochrome oxidase I) and nuclear genome (Amplified Fragment Length Polymorphism (AFLP) 38 using Mspl loci). No clear site associated differences in population genetic structure was found 39 between populations using the AFLP marker loci. Further, no relationship between ionising 40 radiation exposure levels and the percentage of methylated loci or pattern of distribution of DNA 41 methylation marks was found. Microbiome structure was clearly site dependent, with gut 42 microbiome community structure and diversity being systematically associated with calculated 43 site-specific earthworm dose rates. There was, however, also co-correlation between earthworm 44 dose rates and other soil properties, notably soil pH; a property known to affect soil bacterial 45 community structure. Such co-correlation means that it is not possible to attribute microbiome 46 changes unequivocally to radionuclide exposure. A better understanding of the relationship 47 between radionuclide exposure soil properties and their interactions on bacterial microbiome 48 community response is, therefore, needed to establish whether these the observed microbiome 49 changes are attributed directly to radiation exposure, other soil properties or to an interaction 50 between multiple variables at sites within the CEZ.

51

52 *Capsule:* Selected earthworm hologenome traits (e.g. gut microbiome) are potentially influenced
 53 by radiation exposure, but also confounding soil property variables

54

55 *Keywords:* Radioecology, <sup>137</sup>Caesium, Earthworm, DNA methylation, Microbiome,

#### 56 Introduction

The Chernobyl nuclear accident on April 26th 1986 was responsible for a large-scale release of 57 58 radionuclides into the surrounding ecosystem, resulting in long-term exposure of the local and 59 wider environment. Exposure to radioactivity in controlled, high dose laboratory studies has been 60 shown to have effects on soil invertebrate species including earthworms (Hertel-Aas et al., 2007; 61 Rusin et al., 2019; Sowmithra et al., 2015), springtails (Nakamori et al., 2008) and nematodes 62 (Dubois et al., 2019; Lecomte-Pradines et al., 2017). Further, research on the long-term effects of 63 ionising radiation in the surrounds of the former power station, now known as the Chernobyl 64 Exclusion Zone or CEZ, have provided information on the physiological and ecological responses 65 of individuals, populations and ecosystems to chronic radiation exposure (see Beresford et al., 66 2016; Geras'kin et al., 2008; Hinton et al., 2007; Lourenco et al., 2016). Despite these studies, 67 there is a lack of consensus on the long-term effects of radiation on wildlife in the CEZ, with some 68 studies suggesting wide-scale impacts and others limited effects (Beresford et al., 2019b). Further 69 work is, therefore, needed to understand how radiation exposure affected soil organisms, 70 especially under long-term chronic exposure scenarios.

71

72 The soil was one of the main sinks for radionuclides released by the Chernobyl accident. The 73 initially high levels in CEZ soils were shown to alter soil invertebrate population sizes and 74 community structure (Krivolutsky, 1996). For example, reduced recruitment rates of the 75 earthworms Aporrectodea caliginosa and Dendrobaena octaedra (Savigny, 1826) resulted in a 76 population decline in contaminated forest sites (Krivolutzkii et al., 1992). Over time, however, 77 earthworm populations at sites in the CEZ showed some recovery, such that within 3 years of the 78 accident, immature/adult ratios and overall earthworm biomass in the previously impacted sites 79 no longer differed from that in uncontaminated areas (Krivolutsky, 1996). Despite this rebound, 80 species diversity has, however, remained lower at some sites (Geras'kin et al., 2008). This initial 81 decline and recovery of earthworm populations in the CEZ in response to acute exposure may 82 provide only a limited picture of potential response trajectory of earthworms to exposure which 83 may also encompass chronic effects on traits relevant for long-term individual physiology and

population dynamics. For example, in a review of the impacts of the 1957 Kyshtym (Russian Urals) accident, reports indicate that soil invertebrate communities had not been restored at a contaminated site (the main contaminant being 90Sr) c. 30 years after the accident (see Zaitsev et al., 2014). Given the low dispersal rates (e.g. of the order of 5-10 m a-1 for earthworms which may limit recolonization potential (Marinissen and Vandenbosch, 1992), such a long-term impact of an acute radiation event would seem plausible.

90

91 The decline and potentially ongoing change for earthworm populations observed at some CEZ 92 sites in the aftermath of the accident raises fundamental questions regarding the potentially 93 adaptive mechanisms by which these organisms may respond to ongoing chronic exposure (as 94 reviewed by Geras'kin et al., 2008). One possibility may lie in population recovery from a low 95 starting density as exposure abates through radioactive decay and radionuclide transport to 96 deeper soil layers. However, even within this scenario, the selection of tolerant individuals leading 97 to the development of an adapted population may also play a role. At Chernobyl, both plant and 98 fungal species have been shown to develop tolerance to radioisotope exposures (Boubriak et al., 99 2008; Egorova et al., 2015). However, despite evidence for the development of tolerance in 100 earthworms to other pollutants (Fisker et al., 2013; Kille et al., 2013; Langdon et al., 2003), genetic 101 adaptation to radionuclide exposure in this taxa has yet to be clearly established. Adaptation to 102 adverse conditions can result both from phenotypic plastic responses, such as changes in gene 103 expression mediated through modifications in the epigenetic gene regulation, as well as more 104 classically through the selection of traits associated with tolerance. Additionally, changes in the 105 structure and associated function of the host-microbiome can also play a role (Pass et al., 2015).

106

107 The concept of the "hologenome" recognises that changes in the regulation/activity of the genes 108 of the organism itself *and* changes in the community and functional activity of mutualistic 109 (microbial) communities can in combination contribute to the overall development of adapted 110 phenotypes (Bordenstein and Theis, 2015). To assess how different components of the 111 hologenome respond to radiological exposure (and other environmental co-variables), we here

112 measure genetic, epigenetic and gut microbiome changes occurring in earthworms (predominantly 113 Octolasion lacteum) collected from multiple sites along a contamination gradient in the CEZ. 114 Genetic trait analysis was undertaken using cytochrome oxidase 1 (COI) sequencing and the 115 measurement of nuclear sequence polymorphisms using AFLP analysis using one restriction 116 enzyme, epigenetic traits were assessed by DNA methylation profiling using methylation sensitive 117 AFLP (meAFLP) conducted by comparing the first AFLP profile to those obtained using a second 118 DNA methylation sensitive restriction enzyme. Gut microbiome structure was quantified using 16S 119 rRNA loci sequencing. These analyses allow us to test the hypothesis that different genetic, 120 epigenetic and microbiome traits of the hologenome show radiation exposure level dependent 121 changes.

#### 123 Materials and methods

124 An pilot field trip to the Chernobyl Exclusion Zone (CEZ) was carried out in October 2014 to identify 125 the earthworm species present within the zone. Twenty sites representing dominant habitats 126 (grasslands, woodlands, wetlands) were visited. Qualitative sampling collected earthworms from 127 a 50 m x 50 m area over a 2 h search period. This initial survey indicated that earthworms are 128 relatively uncommon in the CEZ. Indeed, many sites lack this taxon. This limited distribution is 129 likely due mainly to the sandy, low pH, low organic matter nature of the soils present, which are 130 known to be unfavourable for earthworms (Edwards, 2004; Ouellet et al., 2008). Where 131 earthworms were found, they were usually associated with the margins of water bodies, 132 marshlands or in improved soils, such as the gardens of abandoned communities. Based on this 133 information, a second field trip was carried out in October 2016 to collect earthworms of one or 134 more species systematically from sites ranging in expected radionuclide exposure from 135 background to relatively high, with wetlands identified as the main habitats for sampling.

136

137 During the second field campaign, six earthworm species, *Eiseniella tetraedra* (Savigny, 1826), 138 Eisenia fetida (Savigny, 1826), O. lacteum, Lumbricus rubellus (Hoffmeister, 1843), A. caliginosa 139 and Aporrectodea rosea (Savigny, 1826) were found at nine sampled sites. Three of these species 140 (E. fetida, L. rubellus and A. rosea) were collected from a limited number of sites giving insufficient 141 variation in radioactivity exposure for the collected specimens. Although *E. tetraedra* was collected 142 from five sites that covered different radioactivity exposure levels, densities were very low limiting 143 statistical power. Only, O. lacteum was found in sufficient, although sometimes at low, numbers 144 across multiple sites with measured and modeled radiation exposure levels ranging from 145 background to high. Therefore, this species has been selected for the genetic, epigenetic and 146 microbiome trait analyses. It should, however, be noted that due to challenges in distinguishing 147 between species based only upon morphology, additional genetic marker analysis was applied to 148 a further earthworm species (A. caliginosa) in order to corroborate in field taxonomic identity. At 149 each site, effort was made to collect multiple individuals per site within the restricted timeframe 150 (as Governed by access permissions) that was available for collection. This was often challenging

as even at sites where earthworms were present, densities were often low. Each individual
 collected was treated thereafter as a separate biological replicate for that sample location.

153

154 Live earthworms were collected from the soil by digging and hand sorting. Individuals were 155 maintained on own site soil for transfer to a local working area for identification and subsequent 156 dissection for tissue samples. Due to the unique conditions presented by the sample site (remote 157 location, sampling period limitations, and minimal facilities) an on-site protocol was used for 158 sampling and tissue preservation. All earthworms were initially weighed, surface cleaned in 159 phosphate-buffered saline to remove any adhering soil particles. The position of the worm's 160 clitellum was used to divide the tissue into anterior (including the clitellum) and posterior section 161 samples. To preserve sample integrity, tissues intended for later DNA extraction were cut into 162 small pieces and placed in 1:10 ratio of sample weight to DNA/RNA shield<sup>™</sup> lysis and preservation 163 buffer (Zymo Research, Irvine, USA). Samples were lysed in the field using the super FastPrep™ 164 handheld homogenizer and Lysing Matrix E (both from MP Biomedicals, Eschwege, Germany). 165 Lysed samples and sampled soils (to 10 cm depth) were transported back to the main laboratory 166 and stored at -20°C prior to DNA extraction. The soils sampled at each site were also used for 167 radionuclide (Cs-137, Sr-90, Am-241, Pu-total isotope) activity measurements for dosimetry 168 calculation. This augmented the simple measurement of air dose rate made *in-situ* at each site.

169

#### 170 Radionuclide Measurement and dosimetry approach

Soil and earthworm <sup>137</sup>Cs concentrations were measured using gamma spectrometry (Ge and Nal scintillation detectors, Canberra). Whole earthworm and earthworm soil gut measurements of <sup>137</sup>Cs were carried out when samples were available. Soil <sup>90</sup>Sr concentrations were determined using beta spectrometry of the daughter nuclide <sup>90</sup>Y, and <sup>241</sup>Am using low energy gamma spectrometry. Pu concentrations (total Pu-isotopes: <sup>238</sup>Pu, <sup>239</sup>Pu, <sup>240</sup>Pu) were determined after sample dissolution (65% HNO<sub>3</sub>), anion exchange separation (BioRad AG 1 × 8, 100–200 mesh) and co-precipitation using <sup>242</sup>Pu added as a yield tracer. These samples were then counted using

- a planar ion implanted silicon detector. All methods were calibrated against standards and counting errors were typically <3% for  ${}^{90}$ Sr, <7% for  ${}^{137}$ Cs and <20% for the Pu isotopes.
- 180

181 Controversies in interpreting results from field-based radioecology studies can arise due to an 182 incomplete assessment of actual organism exposure (Bonzom et al., 2016). Thus, while external 183 dose rates can provide an indication of exposure at sites, they usually underestimate actual doses 184 received by an organism, since they do not include the contribution from internalised 185 radionuclides. Therefore, we chose to estimate exposure as weight absorbed dose rate, which 186 includes the contribution of all radiation types (alpha, beta and gamma emitters) from all exposure 187 pathways (internal and external), as opposed to just ambient external dose rates measured with 188 dosimeters (e.g. Moller and Mousseau, 2009; Uematsu et al., 2015). The approach to dosimetry 189 assessment for all sites used a stepwise approach that: 1) recorded air dose rates and ranges 190 (mGy/hr) at the soil surface and 1 m above ground at each collection sites; and 2) used the 191 measured activity levels in soils to estimate the relative internal and external dose rate 192 contributions of each radionuclide to earthworm exposure. Dosimetry modelling was carried out 193 within a tier 3 assessment using the ERICA tool, a software system that has a structure based 194 upon the tiered approach to assessing the radiological risk to terrestrial, freshwater and marine 195 biota (Brown et al., 2008), using the creation of organisms (different earthworm sizes) and site-196 specific data. Measured tissue concentrations of <sup>137</sup>Cs were used to calculate internal dose rates, 197 while the ERICA default transfer ratios for earthworms were used for the other radionuclides (Popic 198 et al., 2012). Weighting factors of 1, 3 and 10 were used for gamma and beta > 10 keV, beta < 10 199 keV and alpha radiation, respectively. Based on the radionuclide concentration in soil and the 200 sampled organism, external and internal dose rates ( $\mu$ Gy h<sup>-1</sup>) were calculated and used to 201 determine total weight absorbed dose rates.

202

203 DNA extraction

Following sample return to the main research laboratory, a protocol was developed which enabled the co-extraction of the earthworm and microbial DNA from the tissue and gut microbiome

206 samples. Preserved tissue lysate was extracted using the Zymo research ZR-96 soil microbe DNA 207 kit™, with the following modifications. Prior to DNA extraction samples were thawed, then 208 incubated with 2 µl proteinase K (Promega) for 1 hour at 20°C. All samples underwent a further 209 mechanical lysis step on a fast prep 24 (MP Biomedicals) at 5 k for 30 sec. Lysed samples were 210 centrifuged at 2100 g for 5 minutes and 250 µl of lysate combined with 750 µl soil binding buffer. 211 All further steps were carried out according to manufacturer's protocol. Anterior (head) and 212 posterior (tail) samples were extracted separately for analysis of earthworm genomic and 213 microbiome traits, respectively. Large worms often contained more tail tissue than optimal for each 214 lysis tube, where this was the case, the available tissue was spread across a number of tubes, 215 from all of which DNA was extracted. To ensure that the entire microbiome was represented 216 accurately, the results from all assessment for each individual worm were pooled.

217

#### 218 Genotyping

219 DNA samples extracted from the body wall from the anterior proportion of the earthworm were 220 used for the corroboration of field species identification. Approximately 20 ng of DNA template 221 was used to amplify ~710 bp region of the mitochondrial cytochrome oxidase subunit I (COI) gene, 222 in a 50  $\mu$ l total reaction, following the addition of; 0.4  $\mu$ l of tag polymerase (Sigma), 1  $\mu$ l of 10  $\mu$ M 223 dNTP mix (Bioline), 0.5 µl bovine serum albumin (New England Biolabs), and 0.5 µl of 100 µM 224 primers LCOI1490F and HC02198R (Folmer et al., 1994). Amplification conditions were as 225 follows: 95 °C for 2 min, then 35 cycles of 95 °C for 1 min., 40 °C for 1 min, and 72 °C for 1:30 min, 226 with a final extension at 72 °C of 10 min. PCR products were electrophoretically analysed and then 227 purified using the ZR-96 DNA Clean-up kit™ (Zymo research). Purified PCR product was diluted 228 at a 1:10 ratio and 1 µl used separate F and R sequencing reaction carried out using 1 µl BigDye™ 229 terminator V 3.1 reaction mix (Applied Biosystems, USA), 1.5 µl BigDye™ terminator V 3.1 buffer, 230 1 µl of 3.2 µM primer solution, on an Applied Biosystems 3730 genetic analyser. Because of 231 inconsistent amplification using LCOI1490F and HC02198R, a custom additional internal 232 sequencing primer was designed and included for this study COI lumbricF seq (5'-233 TACAGCCACGCATTCGTTA-3'). Sequencing reactions were purified using Big Dye®

234 Xterminator<sup>™</sup> purification kit (Applied Biosystems, USA) and sequenced using ABI PRISM<sup>®</sup> 235 BigDye v3.1 Terminator technology (Applied Biosystems). Resultant sequences were base called, 236 quality checked and assembled using Sequencher V. To act as suitable phylogenetic outgroups 237 resultant contigs were aligned using ClustalW with reference COI sequences sourced from the 238 barcode of life (BOLD) database, NCBI genebank and unpublished reference sequences from 239 native worms sourced in UK (OS1). Optimal likelihood settings were determined to be TVM+G 240 through the implementation of the Akaike Information Criterion in JModelTest V2.17 (Darriba et 241 al., 2012; Posada, 2008). PAUP4b8 was used to generate a Neighbour Joining (NJ) tree using the 242 likelihood criterion (with optimal settings), and bootstrap support values for 1000 replicates. The 243 resultant tree (OS2) was used to determine the taxonomic affiliation of each earthworm and 244 identities corroborated via inclusion in the nearest supported cluster. For O. lacteum, a set of 245 successfully amplified product sequences clustered with O. lacteum COI sequences available 246 from BOLD within the constructed phylogenetic tree, supporting the morphological identification 247 for these individuals. For a remaining set of O. lacteum individuals (38 of 72 individuals), 248 amplification of the COI gene failed, despite best efforts to optimize template, primers and reaction 249 conditions for PCR. Hence, support for the morphospecies identification for these individuals was 250 not achieved through COI sequencing. For these samples a combination of onsite morphological 251 taxonomy and distinct species separation through AFLP profile (see Supplementary Fig. 1) was 252 used to verify species identity.

253

#### 254 Microbiome characterisation

Approximately 20-30 ng of template DNA collected from samples was amplified using Q5 High Fidelity Polymerase (New England Biolabs, Hitchin, UK), each sample reaction include a unique barcode-primer combination to allow separation of sequences associated with the different individuals (Kozich et al., 2013). Amplification conditions consisted of 25 cycles of an initial 30s, a 98°C denaturation step, followed by annealing phase of 30s at 53°C, and a final extension step lasting 90 sec at 72°C. Primer sequence was based on the universal bacterial primer sequence combination 341F and 806R, producing amplicons of ~550 bp spanning the V3-V4 hypervariable

regions of the 16S small subunit ribosomal RNA gene (herein, 16S rRNA gene). PCR Products were normailsed using Sequalprep normalisation plates (Invitrogen, Carlsbad, CA, USA) and the resultant amplicon library was sequenced at a concentration of 5.4 pM with a 0.6 pM addition of Illumina generated PhiX control library. Sequencing was performed on an Illumina MiSeq platform using V3 chemistry (Illumina Inc., San Diego, CA, USA).

267

268 Sequenced paired-end reads were joined using VSEARCH (Rognes et al., 2016), quality filtered 269 using FASTX tools (hannonlab.cshl.edu), length filtered with the minimum length of 300 bp, 270 presence of PhiX and adapters were checked and removed with BBTools (jgi.doe.gov/data-and-271 tools/bbtools/), and chimeras were identified and removed with VSEARCH\_UCHIME\_REF 272 (Rognes et al., 2016) using Greengenes Release 13\_5 (at 97%) (DeSantis et al., 2006). Singletons 273 were removed and the resulting sequences were clustered into operational taxonomic units 274 (OTUs) with VSEARCH\_CLUSTER (Rognes et al., 2016) at 97% sequence identity (Tindall et al., 275 2010). Representative sequences for each OTU were taxonomically assigned by RDP Classifier 276 with the bootstrap threshold of 0.8 or greater (Wang et al., 2007) using Greengenes Release 13 5 277 (full) (DeSantis et al., 2006) as the reference. Unless stated otherwise, default parameters were 278 used for the steps listed. The raw sequence data reported in this study have been deposited in 279 the European Nucleotide Archive under accession number ERS3594341-ERS3594509.

280

281 After guality filtering a total of 13725096 sequences were rarefied to an even depth within the 282 phyloseq package (McMurdie and Holmes, 2013). To visualise the relationship between Miseq 283 community profiles from differing sample sites, nonmetric multidimensional scaling (NMDS) was 284 performed using the 'ordinate' function, based on dissimilarities calculated using the Bray-Curtis 285 index. Changes in bacterial diversity related to mean dosage were assessed using Fishers log 286 series [alpha], as this is largely unaffected by sample size and independent if individuals (N) 287 >1000. Diversity was analysed in relation to potential environmental drivers relevant for sites in 288 the CEZ including soil pH, loss on ignition and the calculated dose rates.

289

290 Genetic and epigenetic characterisation using dual restriction enzyme AFLP

291 A dual restriction enzyme based meAFLP protocol was optimised to assess genetic markers and 292 DNA methylation at these sites. The protocol used is based on parallel use of methylation- and 293 non-methylation-sensitive restriction enzymes (Hpall and Mspl) to treat DNA samples prior to 294 primer ligation and amplification (Xiong et al., 1999). Initially extracted genomic DNA is digested 295 with EcoRI and then there is a second digestion step conducted for an aliquot of this sample using 296 either the restriction enzymes Mspl or Hpall. Both Hpall and Mspl recognize a CCGG sequence. 297 Mspl is able to cut methylated recognition sites (as well as unmethylated ones) thereby providing 298 an assessment of relevant genetic polymorphism. In contrast, Hpall is unable to cut at such 299 locations when they are methylated (i.e. only unmethylated recognition sites are cut) which by 300 comparing to Mspl profiles provides an assessment of epigenetic DNA methylation status.

301

302 The meAFLP analysis was conducted in duplicate for all O. lacteum and A. caliginosa using 303 selective primers for the PCR reactions and analysis on an Applied Biosystems 3130 analyser 304 (Andre et al., 2010). The presence or absence of fragments was scored on chromatograms using 305 GeneMapper Genotyping Software 1.5. Profile bin widths were checked and manually adjusted to 306 encompass all detected peaks. To differentiate signal from background, a Fluorescence Unit (FU) 307 threshold of 40 units was used for a presence/absence binary matrix. All peaks were manually 308 checked for inclusion in analysis. Rare alleles differing only in one individual were initially deleted 309 from the data set. The total number of loci included in the analysis was 202. Fragments were 310 scored as follows: non-methylated state if present in both EcoRI-Hpall and EcoRI-Mspl products 311 (1/1); methylated state if present in either EcoRI-Hpall (1/0) or EcoRI-Mspl (0/1) products (either 312 internal cytosine methylation (0/1) or hemimethylation (1/0)) or absent (0/0). Fragments were 313 classified as "methylation-susceptible loci" if the observed proportion of methylated scores (1/0, 314 0/1 and 0/0) and "non-methylation fragments". Principal coordinates (PCO) analysis was also used 315 to visualise the genetic relationship between individuals using GenAlEx ver 6.4.1 (Srut et al., 316 2017). Site and also potential earthworm strain effects were assessed using the site principal 317 component scores (PCoA1 and PCoA 2) after verification for normality using the Kolmogorov-

- 318 Smirnov test. Tests were conducted using ANOVA with site and potential strain differences as
- 319 factors.

#### 320 **Results**

321 Earthworm distribution, collection and identification by COI barcoding

Morphotype *O. lacteum* were collected in sufficient numbers ( $\geq$ 3 individuals per site) across five sites covering measured surface dose rates from 0.12 - 12 µGy/h. All sampled locations yielding *O. lacteum* were associated with wetland or marshland habitats (Table 1). Morphotype *A. caliginosa* were collected from three sites covering dose rates from 0.12 - 8 µGy/h. However, all but one earthworm came from sites with exposure levels <0.2 µGy/h. This one *A. caliginosa* collected at site M2: Forest Lake 2 was the only individual to co-occur with *O. lacteum*, indicating an almost completed segregation of the two species across visited sites in the CEZ (Table 1).

329

330 To confirm the field based morphological identifications, COI gene sequencing was conducted for 331 all individuals to provide a mitochondrial genotype for comparison with previously published 332 sequences for the two species. For the morphotype A. caliginosa, the phylogenetic analysis 333 indicated that all amplified sequences clustered with the A. caliginosa COI sequences available in 334 the BOLD database. This correspondence supports the morphotype identification of the 335 individuals as A. caliginosa. For O. lacteum, a set of successfully amplified product sequences 336 clustered with O. lacteum COI sequences available from BOLD within the constructed 337 phylogenetic tree, supporting the morphological identification for these individuals (Supplementary 338 Fig. 2). For a remaining set of O. lacteum individuals (38 of 72 individuals), amplification of the 339 COI gene failed, despite best efforts to optimize primers and reaction conditions for PCR. Hence, 340 support for the morphospecies identification for these individuals was not achieved through COI 341 sequencing (see methods). Fortunately, the nuclear genomic loci analysed using the meAFLP 342 method could, however, be used to confirm species identity for these individuals (see below).

343

344 Soil analysis and dosimetry modeling

Measured soil <sup>137</sup>Cs, <sup>90</sup>Sr, <sup>241</sup>Am and Pu-total, and earthworm gut <sup>137</sup>Cs were close to or below detection limits in all control sites and were highest at the H1 Glubokya Marsh site (Table 2). For modeling the transfer of radionuclides to earthworm tissues at CEZ sites, the external irradiation

348 from <sup>137</sup>Cs dominates the exposure profile independent of site (Fig. 1), with internal <sup>241</sup>Am 349 representing the next largest dose rate contributor (assuming a radiation weighting factor of 10). 350 The contribution of radionuclides to dose rates were similar across sites (e.g. Fig. 1a/b). Total 351 median modeled dose rates projected for earthworms across the gradient of sites ranged from 352 0.09 to 45 µGy/h, these values being correlated the with ambient dose rates, albeit at a factor of 353 2-3 times higher. The quoted internal dose rates represent only the contribution from actual 354 transfer across the gut or skin to the tissues, although the soil in the gut will act as an additional 355 source of exposure. For <sup>137</sup>Cs, this consideration makes little difference to the total dose rate 356 experienced. There could be a localised dose from <sup>90</sup>Sr to the gut surface, but within one order of 357 magnitude of the estimated internal dose rate, although localised dose may be similar. The dominance of external exposure due to <sup>137</sup>Cs and the relative contributions of the different 358 359 radionuclides to internal dose are in agreement with dose estimates based upon radionuclide 360 activity concentrations measured in earthworms (Lumbricidae) and soil sampled from the Red 361 Forest (CEZ) in 2014 (Beresford et al., 2019a).

362

Earthworm size has a relatively small influence on modeled dose rates, with slightly lower <sup>90</sup>Sr dose rates in small earthworms, which is expected from differences in absorption and attenuation. At the most highly contaminated sites (H1 Glubokya Marsh), measured activity concentrations in the gut soil of individual earthworms (29 - 53 Bq/g) showed good agreement with the levels measured in the top 10 cm of soil (49 - 86 Bq/kg). These observations support the modeling predictions that individuals at this site are more highly exposed than earthworms from the other sampled locations.

370

371 Site soil pH range from mildly acidic (4.67) at the H1 Glubokya Marsh site with the highest 372 earthworm dose rate to close to neutral (6.23 and 6.66) at two of the low dose rate sites (C1 373 Glinka, C3 Chernobyl Garden). Soils contained between 4.5 and 15.6% organic matter as 374 measured by loss on ignition, the site with the highest dose rate having the highest value which 375 was almost double that any other site (Table 2). The calculated dose rate was significantly

negatively correlated with soil pH (Pearson correlation -0.86, p<0.02) and positively correlated with soil loss on ignition (Pearson correlation 0.781, p<0.05), with soil pH and loss on ignition also significantly negatively correlated (Pearson correlation -0.78, p<0.05). When the highest dose rate site was removed from the data-set, there was no longer a significant correlation with either soil pH or loss on ignition (p>0.05) indicating the importance of the low pH and high loss on ignition values for the H1 Glubokya Marsh site in driving the significant relationships seen between the soil properties.

383

#### 384 Genetic and epigenetic characterisation of earthworms

385 Genetic structure within the collected earthworms was assessed using the AFLP analysis for 386 samples at the Mspl cleavage site for all sampled individuals. Two distinct groupings were 387 identified based on the obtained profiles. One group (orange squares in Supplementary Fig. 1) 388 comprised confirmed morphotype A. caliginosa. These individuals included earthworms collected 389 predominantly from two low exposure sites - C2: Zamozhnya and C3: Chernobyl gardens and a 390 single individual from medium exposure site M2: Forest Lake 2. Given the similarly low levels of 391 exposure experienced by all but one of the A. caliginosa, we focussed our further analysis on O. 392 lacteum for which individuals were available from a number of locations with different measured 393 and modeled exposure dose rates. Nonetheless, the A. caliginosa data was still valuable for 394 taxonomic assignment.

395

396 The remaining AFLP profiles (red circles and green triangles in Supplementary Fig. 1) comprise a 397 single and distinct group within the PCoA plot separated from the A. caliginosa samples along 398 PCoA1. This second set of profiles contains all individuals from morphotype O. lacteum for which 399 COI sequencing gave both a best BLAST hit against an existing O. lacteum COI sequence and 400 for which clustered with a known O. lacteum COI reference sequences (green triangle, 401 Supplementary Fig. 1) and also those morphotype O. lacteum from which the COI loci could not 402 be amplified and sequenced (red circles, Supplementary Fig. 1). Based on morphological similarity 403 of the individuals to the COI loci confirmed O. lacteum and the clustering of the AFLP profiles with

404 those of the COI confirmed individuals in the PCoA plot, we conclude that all the amplified 405 individuals correspond to one or more clades of *O. lacteum* for which the used primer set does not 406 amplify the COI gene. As likely *O. lacteum*, we concluded that these individuals can be analysed 407 alongside the COI confirmed *O. lacteum* for the assessment of the genetic, epigenetic and 408 microbiome response to ionising radiation at CEZ sites. However, when assessing driver of 409 differences between profiles we used the two potential mitochondrial lineages as fixed factors 410 within ANOVAs of the PCoA scores.

411

The AFLP analysis based on *Msp*l cleavage site markers for all *O. lacteum* showed overlap of profiles between replicate earthworms (Fig. 2). Analysis of PCoA 1 scores indicated a significant site effect (GLM F=7.95, p<0.001), but not of *O. lacteum* mitochondrial strain (GLM, F=0.02, P>0.01). Between site differences were found for earthworms from the 6.8  $\mu$ Gy/h site and all other locations, with the exception of the 0.12  $\mu$ Gy/h site. The differences in AFLP profiles of earthworms from intermediate dose rate sites compared to those from the higher and lower dose rate sites suggests that exposure to ionising radiation is not the main driver of AFLP profile differences.

419

To assess DNA methylation patterns in sampled O. lacteum, a second AFLP analysis was 420 421 conducted using the methylation sensitive Hpall restriction enzyme. The meAFLP indicated an 422 average of 20.1% of methylation sensitive loci. DNA methylation levels showed no trend of hyper-423 or hypo-methylation across sites (lowest average 16.7% at an exposure level of 0.12  $\mu$ Gy/h, 424 highest average 23.4% at an exposure level of 9.1 µGy/h, Fig. 3a). Principle coordinate analysis 425 of obtained methylation patterns indicated no site-specific structure among the profiles. Sampled 426 individuals showed no segregation within the PCoA on either of the first two principal coordinate 427 axes, which together account for only 20.6% of variance (Fig. 3b). No significant differences in 428 PCoA 1 or PCoA 2 scores were evident (PCoA 1 ANOVA F=2.09, p>0.05; PCoA 2 ANOVA F=1.08, 429 p>0.05). Consistent methylation levels and the overlap of methylation patterns among earthworms 430 sampled from sites with different radiation dose rates, suggests no effects of radiation exposure 431 on global DNA methylation patterns at CEZ sites for O. lacteum.

433 Sequencing of the 16S rRNA loci amplified from O. lacteum gut DNA identified 18870 OTUs at the 434 97% similarity threshold. An NMDS analysis using data for sequences allocated to these OTUs 435 for each earthworm microbiome indicated a clear segregation of all individuals from each site 436 within an NMDS analysis plot (Fig. 4). NMDS1 and NMDS2 scores were significantly correlated 437 with median dose rate, soil pH and soil loss on ignition in all cases (p<0.001 all cases). However, 438 due to the small number of sites, such correlations with average dose rate may be a potential 439 artifact due to co-correlation effects. For this reason, to determine whether a given environmental 440 variable may be a driver of microbiome change, we assessed the structure or this relationship -441 notably the monotonic nature of the response (i.e. score increase or decrease progressively in 442 relation to environmental variable level). This site based separation by OTU profiles, did not show 443 a monotonic relationship with radiation exposure level along the NMDS 1 axis, with low exposure 444 level and high exposure levels sites having high NMDS 1 scores and medium exposure sites low 445 values (Supplementary Fig. 3A). There was also a non-monotonic relationship of NMDS 1 score 446 with site soil pH, the highest axis values found for the sites with the most acidic and alkaline soils 447 (Supplementary Fig. 3B). There was an approximate monotonic relationship of NMDS 1 score 448 with site soil loss on ignition (Supplementary Fig. 3C), indicating that soil organic matter 449 composition may act as a critical driver of soil bacterial community structure either directly or 450 through indirect effects on other soil properties such as water holding capacity which may affect 451 soil responses to drought or frost. NMDS 2 score was close to monotonic related to calculated 452 dose rate (Supplementary Fig. 3D). However, as there is co-correlation of the environmental 453 drivers, both soil pH and loss on ignition also show a monotonic relationship with NMDS 2 score 454 (Supplementary Fig. 3E and 3F).

455

Diversity expressed as Fishers log alpha was highest in the bacterial gut microbiome samples in *O. lacteum* earthworms from the two lowest exposure sites (C1 Glinka, C4 Marsh) and lowest in the two highest exposure sites (H1 Glubokya Marsh, M2 Forest Lake). Diversity increased systematically with increasing median earthworm dose rate (Fig. 5A). As pH is strongly co-

- 460 correlated with earthworm dose rate, there is also a relationship of increasing diversity at high soil
- 461 pH values. Diversity was not systematically related to soil loss on ignition, with high diversity at
- 462 both intermediate and high soil loss on ignition sites (Fig. 5B).

#### 464 **Discussion**

465 The scale and devastating nature of the Chernobyl accident has created a legacy of exposure for 466 wildlife living around the site of the former power plant. Radioecologists have studied the long-467 term effects of the resulting exposure, both to support local management and to better understand 468 any long-term effects on individuals, populations, and ecosystems. Comprehensive reviews of the 469 phenotypic effects observed in wildlife experiencing prolonged exposure to elevated radionuclide 470 levels in the CEZ have highlighted the range of studies on the structure and function of exposure 471 populations and communities from soil microbial communities to charismatic vertebrate wildlife 472 (e.g. Beresford et al., 2016; Geras'kin et al., 2008; Hinton et al., 2007; Lourenco et al., 2016; 473 Steinhauser et al., 2014). Within the range of published studies, there is a lack of consensus on 474 the long-term effects of chronic radiation exposure for wildlife in the CEZ (Beresford et al., 2019b). 475 The impacts of radiation exposure on the hologenome and subsequent impacts on exposure and 476 subsequent generations have been identified as one factor that may contribute to the impacts of 477 long-term exposure in the CEZ (Horemans et al., 2019). Understanding such effects may benefit 478 from addressing uncertainties relating to the chronic effects of radiation exposure.

479

480 The prolonged exposure of soil invertebrates to long-lived contaminants has been shown to 481 change population genetic and epigenetic traits that underpin the development of tolerance 482 (Horemans et al., 2019). The relatively high reproductive rate of many invertebrates enables rare 483 or novel gene variants associated with adaptive traits to spread relatively quickly throughout 484 populations under selection pressure. Many invertebrates have been shown to develop adaptation 485 to contaminants (Klerks & Weis 1987; Posthuma & Van Straalen 1993). For earthworms, tolerance 486 to metal exposure has been shown in populations inhabiting sites polluted with a number of 487 different metals. Andre et al. (2010) used genetic marker analysis to identify an 'ecological island' 488 with little genetic overlap in L. rubellus populations inhabiting lead polluted and clean soils under 489 different pH conditions. Mutations in the Ca-transport gene SERCA were found between 490 populations, identifying this locus as a possible mechanism for adaptation. Langdon et al. (1999) 491 also working with L. rubellus found that individuals inhabiting two mining areas heavily polluted

492 with arsenic and copper (Devon Great Consols, Carrock Fell) could survive in arsenic-spiked soil 493 that was acutely toxic to earthworms from an uncontaminated location. This tolerance was 494 preserved following culturing over two generations, suggesting a genetic basis for the adapted 495 phenotype (Langdon et al., 2009). Kille et al. (2013) used mitochondrial (COI) marker analysis and 496 meAFLP to assess the genetic and epigenetic traits of *L. rubellus* from the same Devon Great 497 Consols mine site studied by Langdon et al. (2009). Both the COI analysis and AFLP markers 498 identified two genetic lineages of L. rubellus at the mine site consistent with previous findings 499 (Anderson et al., 2017; Giska et al., 2015; Spurgeon et al., 2016). For the most genetically diverse 500 lineage (A), segregation of local collection site populations by AFLP profiles was seen. For the 501 second, less diverse, lineage (B), this genetic segregation by AFLP profile was not seen, however, 502 site segregation by meAFLP profiles was observed. This indicates a contribution of the epigenome 503 to the observed adaptive phenotype in Lineage B L. rubellus. This unique finding indicates the 504 need to assess how both genetic sequence changes and DNA methylation can play a role in the 505 development of adaptive phenotypes in earthworms.

506

507 An initial survey of Chernobyl earthworm communities identified that species are patchily 508 distributed across the exclusion zone. The dominant soil type in the region is sandy in nature with 509 characteristically low pH and organic matter content. Such free draining acid soils have been found 510 to be poorly suited for earthworms, which maintain larger and more diverse communities in areas 511 with high clay or silt content, high water holding capacity and high electrical conductivity (Nuutinen 512 et al., 1998: Valckx et al., 2009). The main locations from which earthworms were collected were 513 associated either with water bodies (lake or river margins, marshland) or improved soils managed 514 as domestic gardens in areas of past human habitation. This indicates that soil hydrological status 515 acts as a major driver of earthworm distributions within the CEZ.

516

517 Two species of earthworms were found in sufficient numbers to provide potential samples for 518 analysis. Both *O. lacteum* and *A. caliginosa* are endogeic species that live in the mineral soil 519 layers. Exposure in this habitat would be lower than to epigeic species living in the top-soil, since

520 almost 20 years after the accident, radionuclides remain largely in the upper 0-20 cm soil layers 521 (Almgren and Isaksson, 2006; Ivanov and Kashparov, 2003). However, at the soil sampling depth 522 used (10 cm), calculated dose rates would be generally representative for species living within the 523 mineral soil layer. The individual measurements of relevant external and internal radionuclide 524 concentrations for dosimetry assessment confirmed this exposure. Among the earthworms 525 species found during the preliminary survey and main field sampling campaign, only O. lacteum 526 was found in sufficient number across CEZ sites with comparatively low, medium and high 527 exposure. O. lacteum is an endogeic earthworms species that is able to undertake both sexual 528 and parthenogenetic reproduction, which may be relevant both for its relative prevalence in the 529 CEZ and also potentially for its adaptation to local environmental conditions. A. caliginosa, another 530 albeit larger endogeic earthworm species, was in contrast collected only from sites with near 531 background levels. Too few sites were sampled to attribute the absence of A. caliginosa to the 532 prevailing exposure levels. Absence is likely to be explained simply by the lack of suitable habitat 533 for this species at the more contaminated sites.

534

535 Our analysis of the O. lacteum hologenome included measurement of genetic, epigenetic and 536 microbiome characteristics. However, while covering key aspects, not all hologenome traits were 537 included. For example in assessing epigenome responses, we focussed on an established method 538 to measure coarse DNA methylation coverage. These DNA methylation marks were not mapped 539 to genomic locations to provide insights in their functional role in gene regulation. Further, other 540 potential epigenetic changes linked to the modification of histone protein and expression of 541 regulatory microRNAs were not measured. These epigenome components have previously been 542 found to be active in earthworms (Gong et al., 2010; Novo et al., 2015). Hence, their future 543 inclusion within a more complete assessment of hologenome responses to ionising radiation 544 exposures may be warranted.

545

546 The genetic analysis of *O. lacteum* using AFLP markers at *Msp*l restriction sites indicated that all 547 morphotype earthworm from this species exist as a single group. Using the COI primers,

548 sequences from 53% of all collected morphotype O. lacteum failed to amplify despite best efforts. 549 The reasons for this failure are currently not clear, but may be related to the presence of specific 550 point sequence variations in the primer binding region of the COI sequence in these individuals. 551 The presence of these polymorphisms may be indicative of the cryptic mitochondrial lineages in 552 this species, as has been found for O. lacteum (Klarica et al., 2012) and also commonly for other 553 earthworm including L. rubellus (Anderson et al., 2017; Andre et al., 2010), Eisenia fetida 554 (PerezLosada et al., 2009) and Allolobophora chlorotica (King et al., 2008). The more 555 comprehensive AFLP data, however, refutes the presence of cryptic O. lacteum species, as the 556 PCoA analysis of the AFLP data shows that CEZ O. lacteum comprise a single clade.

557

558 Site-specific structure was observed within the Mspl restriction site AFLP analysis. This was not, 559 however, correlated with modeled earthworm dose rate. This suggests that there has been no 560 selection relating to the ionising radiation exposure. Similarly, we found no difference in DNA 561 methylation levels or structure within the meAFLP analysis, indicating that there is no site, and 562 hence ionising radiation exposure level, influence on the distribution of DNA methylation within the 563 O. lacteum genome. Many contaminants have been shown to alter gene methylation patterns in 564 invertebrates (Santoyo et al., 2011) and plants (Chinnusamy and Zhu, 2009). In response to 565 ionising radiation exposure in the CEZ, hypermethylation of DNA has been observed in pine trees 566 (Pinus sp.) and Arabidopsis thaliana (Kovalchuk et al., 2004; Kovalchuk et al., 2003) and it has 567 been suggested that individuals of adult pale blue grass butterfly Zizeeria maha at Fukahima have 568 shown inherited abnormalities from the F1 to the F2 generation (Hiyama et al., 2012). Epigenetic 569 mechanisms were proposed to play a role in both cases. Absence of a clear indication of changes 570 in patterns of DNA methylation measured using meAFLP for O. lacteum may indicate that adaptive 571 phenotypes may be regulated through other epigenome components. Alternatively, it is possible 572 that exposure at the sampled CEZ sites is not sufficient to cause significant epigenetic changes in 573 this species. However, in this context, it is important also to consider that meAFPL provides only 574 a very course assessment of one aspect of the response to the epigenome to perturbations. Other 575 methods for quantifying DNA methylation targeting either specific genes through methylationspecific PCR or genome wide methylation analysis using high throughput sequencing approaches
may better resolve any specific DNA methylation changes resulting from radiation exposure.
Further, there are other epigenetic mechanisms that may also play a role.

579

580 The analysis of the structure of the CEZ O. lacteum microbiome indicated clear site structure in 581 the composition of gut bacteria as analysed using 16S RNA gene loci amplification and 582 sequencing. The earthworm microbiome has previously been found to be shifted in response to 583 the presence of arsenic in contaminated soils at a mine site (Pass et al., 2015). In this study, the 584 earthworm-associated microbiome was found to differ from the surrounding environment. Several 585 taxa observed in uncontaminated control microbiomes were found to be suppressed by 586 metal/metalloid field exposure, including the eradication of the hereto ubiquitously associated 587 Verminephrobacter symbiont, which raises implications to its functional role in the earthworm 588 microbiome (Pass et al., 2015).

589

590 It has been shown that the earthworm gut bacterial microbiome is a combination of soil associated 591 species passing through the gut and also a range of strongly host related core species present 592 across different soils (Liu et al., 2018; Pass et al., 2015). As the earthworms sampled in this survey 593 were not starved or washed to remove soil in the gut, both core O. lacteum and also soil associated 594 species would be present. Microbiome structure and diversity was systematically related to site 595 soil properties. Analyses through 16s rRNA sequencing have indicated that soil properties can 596 significantly affect the bacterial community (George et al., 2019; Plassart et al., 2019), Griffiths et 597 al. (2011) showed that pH was a dominant driver of national patterns of soil bacterial species 598 distribution, with other environmental variables and climatic variables and spatial correlations at 599 local scale also influencing community structure. Given that the soil bacterial community 600 contributes a significant proportion of earthworm microbiome species, the potential exists for soil 601 properties to be responsible for differences in the microbiome.

602

603 O. lacteum gut bacterial microbiome structure was site specific and was weakly correlated with 604 dose rate. However, across sites there was also co-correlation between earthworm dose rates 605 and other soil properties, notably pH. Subtle shifts in soil bacterial community structure have 606 previously been shown to respond to radiation exposure, although at much higher dose rates than 607 those guantified here (Jones et al., 2004; McNamara et al., 2007; Niedree et al., 2013). Hence, 608 changes in earthworm gut microbiome structure and diversity may be related to soil dose rates. 609 However, the widely known role of soil pH as a driver of bacterial community structural difference 610 makes the attribution of the observed difference in microbiome structure to radiation exposure 611 challenging. Further, soil pH has been shown to be a key determinant of the speciation of metals 612 and radionuclides in soil (Lofts and Tipping, 2011; Vandenhove et al., 2007), with resulting effects 613 on bioavailability (Hegazy et al., 2013; Spurgeon et al., 2006). Hence, as well as the potential 614 direct effects of radionuclide and pH, there may be interactive pH effects on radionuclide 615 bioavailability which may complicate the attribution of O. lacteum gut bacterial community variation 616 to any single driver.

617

#### 618 Conclusions

619 Prevailing habitat, soil and climatic conditions mean that the distribution of earthworms in the CEZ 620 is spatially limited. Our survey suggests that earthworms are restricted to sites associated with 621 water bodies and in abandoned urban gardens. Our hologenome analyses indicate that there is 622 no major impact of the prolonged exposure of earthworms to radionuclides on the measured 623 genetic or epigenetic characteristics of earthworms collected across our studied sites. Earthworm 624 gut bacterial microbiome community did show strong site specific structure that could be related 625 to the prevailing dose rate predicted for earthworms living at each site. Co-correlation of other soil 626 properties with dose rate and potential interactions, however, means that it is not possible to 627 attribution of these changes specifically to radionuclide exposure. A wider survey is needed to 628 more robustly address questions relating to the impacts of radionuclide exposure on the 629 microbiome. Due to their limited distribution in the CEZ earthworm may not be the best study group 630 for such an extended survey. Robust wide-scale microbiome community structure is a potentially

- 631 effective approach, although it is acknowledged that microbe may be among some of the most
- radio insensitive groups of organisms, the interplay between host health and microbiome structure
- 633 may indicate a wider interaction at the level of the hologenome.

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930 Table 1. Site, habitat, number of earthworm species present and count of *A. caliginosa* and *O. lacteum* collected and thereafter treated as separate

931 biological replicates from each collection location.

Site Number	Northing	Easting	Distance to Power Plant (km)	Habitat	Sample date	Species present	Aporrectdea caliginosa	Octolasion lacteum
C1 Glinka (4)	51.241295	29.9908887	18.3	Wetland margin	06/10/2016	5	0	3
C2 Zamozhnya (7)	51.236371	29.898201	20.7	Wetland margin	07/10/2016	4	13	0
C3 Chernobyl Garden (6)	51.279219	30.213228	14.5	Abandoned Garden	07/10/2016	3	18	0
C4 Marsh (9)	51.474772	29.633966	33.5	Marshland	10/10/2016	2	0	11
M1 Forest lake 1 (2)	51.41190	30.100092	2.6	Wetland margin	06/10/2016	2	0	27
M2 Forest lake 2 (3)	51.410225	30.100816	2.2	Wetland margin	06/10/2016	3	1	16
H1 Glubokya Marsh) (8)	51.445025	30.063611	6.5	Marshland	08/10/2016	2	0	13

Table 2. Site soil properties, measured radionuclide concentrations, modeled total earthworm and gut dose rates and median dose rate values (used for site labels) for all collection locations.

Site Number	Soil pH	Soil loss on ignition (%)	Measured surface dose rate (μGy/h)	Soil Bq/g <sup>137</sup> CS	Soil Bq/g <sup>90</sup> Sr	Soil Bq/g <sup>241</sup> Am	Soil Bq/g <sup>total</sup> Pu	Earthworm dose rate range (μGy/h)	Worm gut activty Bq/g <sup>137</sup> Cs	Median worm dose rate for site (µGy/h)
C1 Glinka (4)	6.23	4.8	0.05 - 0.11	0.3	<0.48	0.003	<0.02	0.096 - 0.144	<2	0.12
C2 Zamozhnya (7)	5.87	8.5	0.1	0.2	<0.48	0.004	<0.02	0.072 - 0.108	<2	0.09
C3 Chernobyl Garden (6)	6.66	5.1	0.2	0.66	<0.48	0.02	<0.02	0.27 - 0.41	<2	0.34
C4 Marsh (9)	5.78	10.3	0.15 - 0.2	0.54	<0.48	0.01	<0.02	0.19 - 0.29	2.2+/-0.9	0.24
M1 Forest lake 1 (2)	5.73	4.5	2.8 - 3.5	10.1	20.3	0.54	0.18	6.12 - 7.48	NA	6.8
M2 Forest lake 2 (3)	5.49	6.1	7 - 8	16.9	7.1	0.59	0.33	8.2 - 10	NA	9.1
H1 Glubokya Marsh (8)	4.67	15.6	8 - 12	49 - 86	18 - 28	2.3 - 4.1	0.6 - 2.3	29 - 53	20 - 76	45

#### Legends to figures

Figure 1. Relative proportionate contribution of different measured radionuclides at sites with median ambient dose rates of 0.24 (Left pie chart) and 45  $\mu$ Gy/h (Right pie chart B) for earthworms sampled in the CEZ.

Figure 2. Principle coordinate analysis of *O. lacteum Mspl* AFLP multi-locus profiling collected at five sites in the CEZ with different soil total earthworm dose rates.

Figure 3. (Panel A) Percentage of methylated susceptible loci for *O. lacteum* collected at five sites in the CEZ with different soil total earthworm dose rates ( $\mu$ Gy/h); and, (Panel B) principle coordinate analysis for *Mspl/Hpall* meAFLP analysis for methylation loci for *O. lacteum Mspl* AFLP multi-locus profiling collected at five sites in the CEZ with different total soil earthworm dose rates.

Figure 4. Non-metric multidimensional scaling plot of 16S rRNA gene sequence analysis of gut microbiomes of *O. lacteum* collected at five sites in the CEZ with different total soil earthworm dose rates and the relationships with environmental vectors oriented in the direction of greatest increase with length is proportional to their correlation with the two NMDS axes.

Figure 5. Diversity of bacterial sequenced OTUs within the earthworm microbiome for *O. lacteum* collected from five locations within the CEZ expressed as Fischer's alpha in relation to soil (A) pH and (B) loss on ignition, bars are shaded according to earthworm dose rate ( $\mu$ Gy/h).













#### Fig. 5

Supplementary Figure 1. Principle coordinate analysis of AFLP *Mspl* multi-locus profiles for morphotype *O. lacteum* and *A. caliginosa*: morphotype *A. caliginosa* confirmed by COI sequencing cluster to the right; morphotype *O. lacteum* both confirmed by COI sequencing and which also fail to amplify for the COI locus cluster to the left as a single group.

Supplementary Figure 2. Phylogenetic gene tree showing the relationships between individual collected earthworms successfully amplified and sequenced for the Cytochrome oxidase I loci; branch distance indicates the degree of base substitutions between individual; all samples labeled starting P1 or P2 correspond to study samples with all other samples corresponding to reference specimen sequences.

Supplementary Figure 3. Relationship between NMDS axis 1 score and soil (A) median earthworm dose rate ( $\mu$ Gy/h), (B) pH, (C) loss on ignition and NMDS axis 2 score and soil (D) median earthworm dose rate, (E) pH, (F) loss on ignition.