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1 **Landscape features impact connectivity between soil populations: a comparative study**
2 **of gene flow in earthworms**

3

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14

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17

18 **Running Title** : Landscape genetics of earthworms

19

20

21 **Abstract**

22

23 Landscape features are known to alter the spatial genetic variation of above ground
24 organisms. Here, we tested the hypothesis that the genetic structure of below ground
25 organisms also responds to landscape structure. Microsatellite markers were used to carry out
26 a landscape genetic study of two endogeic earthworm species, *Allolobophora chlorotica* (N =
27 440, 8 microsatellites) and *Aporrectodea icterica* (N = 519, 7 microsatellites), in an
28 agricultural landscape in the North of France, where landscape features were characterised
29 with high accuracy. We found that habitat fragmentation impacted genetic variation of
30 earthworm populations at the local scale. A significant relationship was observed between
31 genetic diversity (H_e , A_r) and several landscape features in *A. icterica* populations and *A.*
32 *chlorotica*. Moreover, a strong genetic differentiation between sites was observed in both
33 species, with a low degree of genetic admixture and high F_{st} values. The landscape
34 connectivity analysis (MLPE) at the regional scale, including Isolation By Distance (IBD),
35 Least Cost Path (LCP) and Cost Weighted Distance (CWD) approaches, showed that genetic
36 distances were linked to landscape connectivity in *A. chlorotica*. This indicates that the
37 fragmentation of natural habitats has shaped their dispersal patterns and local effective
38 population sizes. Landscape connectivity analysis confirmed that *a priori* favourable habitats
39 such as grasslands may constitute dispersal corridors for these species.

40

41 **Introduction**

42

43 A number of studies showed that spatial genetic variations of aboveground organisms
44 respond to changes in their landscape, through mechanisms involving movements of
45 organisms (review in Storfer *et al.* 2010; Manel & Holderegger 2013; Hall & Beissinger
46 2014). It is now well established that landscape features alter aboveground organisms' genetic
47 structure. Comparatively little is known about the impact of landscape-scale habitat
48 heterogeneity on belowground organisms, such as soil invertebrates, whose mobility is more
49 restricted (Vanbergen *et al.* 2007). Despite their importance in ecosystem functioning and in
50 the delivery of many ecosystem services (Lavelle *et al.* 1997; Jouquet *et al.* 2006; Lavelle *et al.*
51 2006; Blouin *et al.* 2013), we still do not have a grasp of even basic information about
52 population genetic structure of soil organisms. For instance, Costa *et al.* (2013) found only
53 sixteen different species among the collembolans, earthworms and isopods groups of soil
54 invertebrates for which a population genetics study was carried out. Some of these papers
55 investigated the spatial genetic structure of soil organisms at a fine-scale (Sullivan *et al.* 2009;
56 Novo *et al.* 2010; Dupont *et al.* 2015), but none addressed the effect of landscape features on
57 genetic variation. However, terrestrial habitat heterogeneity is known to affect the diversity of
58 soil species' assemblages by producing variation in the diversity of plant and litter
59 (Vanbergen *et al.* 2007). It is therefore assumed that aboveground structure and diversity
60 could profoundly impact the population genetic structure of belowground organisms.

61

62 The methodology of landscape genetics allows one to test the influence of the landscape
63 and environmental characteristics on microevolutionary processes and metapopulation
64 dynamics, including gene flow and local adaptation (Manel *et al.* 2003; Storfer *et al.* 2007;

65 Holderegger & Wagner 2008). Landscape connectivity is a twofold parameter made up of
66 structural connectivity and functional connectivity. Structural connectivity refers to the
67 physical relationship between landscape elements while functional connectivity may be
68 defined as the ease with which a landscape can be crossed by an organism (Taylor et al. 2006).
69 Depending on the organisms, the permeability of the landscape will differ and some
70 constituent elements of the landscape can facilitate dispersal (i.e. “corridors”) while others
71 can impede or reduce the passage of dispersers (i.e. “barriers”) (Taylor et al. 1993).
72 Landscape structure can also have an important effect on passive dispersers by altering the
73 abiotic and biotic conditions that affect movement (Matthysen 2012). In order to understand
74 how landscape characteristics influence functional connectivity, resistance surfaces are
75 usually constructed and translated into measures of inter-population connectivity principally
76 using two kinds of models. Least-cost path models (Adriaensen et al. 2003) assume that
77 movement or gene flow rates between each pair of sites is related to the total cumulative
78 resistance or ‘cost’ (sum of per-pixel resistance values) along a single optimal path, while
79 circuit-theory based models (McRae 2006) incorporate all possible pathways across
80 landscapes, and their parameters and predictions can be expressed in terms of random walk
81 probabilities (Cost Weighted Distance "CWD", or "resistance approach"). Spear *et al.* (2010)
82 highlighted that both models provide complementary indices of connectivity with Least-cost-
83 path distances being more informative at local scales and circuit theoretic models being
84 particularly useful for incorporating effects of gene flow over multiple generations.

85

86 Here, we were interested in landscape features impacting genetic variation and functional
87 connectivity in earthworms. Dispersal by passive mechanisms, such as zoochory, wind, water
88 and human activities is believed to be implicated in their long-distance movement (Eijsackers

89 2011; Costa et al. 2013; Dupont et al. 2015), whilst their active dispersal is dependent on
90 habitat quality, conspecific density, and habitat modification by conspecifics in endogeic (i.e.
91 species living in the upper organo-mineral soil layers and forming horizontal non-permanent
92 burrows, Bouché 1977) and anecic (i.e. species forming permanent or semi-permanent
93 vertical burrows in the soil which open at the surface where the earthworm emerges to feed,
94 Bouché 1977) species (Mathieu et al. 2010; Caro et al. 2012; Caro et al. 2013). The
95 distribution of these restricted dispersers is known to be controlled by soil parameters at the
96 field scale and by land use (forest, grassland and agricultural field), soil management, soil
97 type and climatic conditions at scales exceeding the field level; studies at the landscape scale
98 are thus challenging since fine-scale heterogeneity as well as gradients affecting regional-
99 scale patterns have to be accounted for (Palm et al. 2013).

100 In order to investigate whether landscape features impact the genetic structure of
101 earthworm populations, we carried out a regional-scale comparative survey of genetic
102 variation in two species commonly found in European agricultural landscapes, the green
103 morph of *Allolobophora chlorotica* (Savigny, 1826) and *Aporrectodea icterica* (Savigny,
104 1826). Both species are endogeic but present several ecological differences. *A. chlorotica*
105 typically lives between soil surface and the upper 60 mm soil layer (Sims & Gerard 1999) and
106 is theoretically able to travel over 167 m per year in constant suitable conditions (Caro et al.
107 2013). However, in the field, dispersal distances ranging from 6.82 to 7.56m per year were
108 estimated in a recent population genetics study at fine spatial scales (Dupont et al. 2015). *A.*
109 *icterica* is found deeper in soils and is considered to be more mobile, being theoretically able
110 to travel up to 500 m.year⁻¹ under constant artificial conditions (Mathieu et al. 2010; Caro et
111 al. 2013). An extremely low signal of genetic structure was obtained for this species in a fine-
112 scale population genetics study at the within plot scale (100x80m). This result was explained

113 by the great dispersal capacity of the species (Dupont et al. 2015). Moreover, no pattern of
114 isolation by distance (IBD, i.e. decrease in the genetic similarity among populations as the
115 geographic distance between them increases) was observed among six *A. icterica* populations
116 separated by less than 13km (Torres-Leguizamon et al. 2014).

117 We analysed the relationship between landscape features and genetic variation in these
118 two common earthworm species, at fine and regional scales (Fig. 1), in an agricultural
119 landscape in North of France, where both species are common (e.g. Richard et al. 2012). First,
120 we tested the hypothesis that the mosaic of habitats created by anthropogenic drivers alters the
121 genetic diversity in earthworm populations using a buffer approach. This approach consists in
122 assessing the correlation between the genetic variation of the earthworm population and the
123 local landscape structure. Second, we tested the hypothesis that the different elements of the
124 landscape could act either as dispersal barriers or corridors for earthworms with a landscape
125 connectivity analysis at the regional scale (Zeller et al. 2012), encompassing Isolation by
126 Distance (IBD), Least Cost Path (LCP) and resistance (CWD) approaches. The role of three
127 elements, i.e. grasslands, crops and roads, was specifically tested. Grassland represents a
128 suitable habitat that could be easily crossed by endogeic species (Bouché 1972; Decaens et al.
129 2008) while soil tillage and the use of pesticides in cultivated soils are known to have a
130 detrimental effect on earthworms (Bertrand et al. 2015) and roads have been shown to
131 represent dispersal corridors for invasive earthworms (Cameron & Bayne 2009).

132

133 **Material and Methods**

134

135 *Study Area and Sampling*

136 Earthworms were collected in Normandy (northern France, Fig. 2) in 2009 and 2010.
137 The first sampling campaign was carried out in March and April 2009 in two pastures (P_A and
138 P_B) ~ 500m apart at the local agricultural school “Lycée Agricole d’Yvetot”. Details of these
139 sampling sites and methods are given in Dupont *et al.* (2015). The second sampling campaign
140 was carried out in April 2010, during which 39 other pastures were prospected. *A. chlorotica*
141 and *A. icterica* were found in 14 and 19 pastures, respectively (Fig. 2 and Table 1). We
142 selected pastures that had similar management histories, in order to reduce the effect of local
143 environmental variations. The location of the plots was chosen in order to maximise the
144 normality of the pairwise distance between plots. All pastures were at least 5 years old, and
145 the great majority was grazed by cattle. Within each plot of 10 x 10m, 30 individuals were
146 captured by sampling five monoliths of soil (25 x 25cm x 30 cm deep). If a species was
147 present in the samples of a plot, but less than 30 individuals were captured, we sampled other
148 monoliths - less than 10 m apart from the others- until the target of 30 individuals per
149 population was reached. Individuals were preserved in pure ethanol for DNA analysis.

150

151 *DNA extraction, microsatellite genotyping and basic genetic statistics*

152 Total genomic DNA of *A. icterica* was extracted using either the CTAB extraction
153 protocol, as described in Torres-Leguizamon *et al.* (2014) or the DNeasy 96 Blood & Tissue
154 Kit (Quiagen). The latter was also used for *A. chlorotica*.

155 *A. chlorotica* individuals were genotyped at the eight microsatellite loci described in
156 Dupont *et al.* (2011) while *A. icterica* individuals were genotyped at seven microsatellite loci

157 described in Torres-Leguizamon *et al.* (2012) and Dupont *et al.* (2015). Loci were amplified
158 by polymerase chain reaction (PCR) following protocols detailed in Dupont *et al.* (2011),
159 Torres-Leguizamon *et al.* (2012) and Dupont *et al.* (2015). The migration of PCR products
160 was carried out on a 3130xl Genetic Analyser using the LIZ500 size standard, alleles were
161 scored using GENESCAN V3.7 and GENOTYPER V3.7 software (Applied Biosystems,
162 Foster City, CA, USA).

163 Individuals missing 3 or more loci (e.g. failed PCR, poor-quality DNA extract) were
164 excluded from our dataset and ambiguous PCR results were duplicated. Mean genotyping
165 error rates per locus and per allele (Pompanon *et al.* 2005) were estimated from repeat
166 genotyping of 5% of samples (24 individuals per species). The null hypothesis of
167 independence between loci was tested from statistical genotypic disequilibrium analysis using
168 GENEPOP v. 4.4 (Rousset 2008). Evidence of null alleles was examined using the software
169 MICRO-CHECKER (Van Oosterhout *et al.* 2004) and from the frequency of null homozygote
170 within populations. The statistical power to detect genetic divergence was measured for all the
171 samples and markers using POWSIM 4.0 to evaluate the hypothesis of genetic homogeneity
172 under Fisher's exact tests (Ryman & Palm 2006). Microsatellite loci were tested for departure
173 from Hardy–Weinberg equilibrium (HWE) within each sampling population using exact tests
174 implemented in GENEPOP v. 4.4. To adjust for multiple comparisons, the FDR method
175 (Benjamini & Hochberg 1995) as implemented in the software SGoF
176 (<http://webs.uvigo.es/acraaj/SGoF.htm>) was applied.

177

178 *Genetic variation of earthworm populations*

179 For each population, the genetic diversity was analysed by computing allelic richness
180 standardized for sample size (A_r ; $N= 26$ and $N = 9$ for *A. chlorotica* and *A. icterica*

181 respectively) using the program FSTAT v2.9.3.2 (Goudet 2000) and expected heterozygosity
182 (H_e) using Genetix v 4.05 (Belkhir et al. 2004). Weir and Cockerham's (1984) estimator of
183 the inbreeding coefficient F_{is} was calculated using GENEPOP v. 4.4 (Rousset 2008). The
184 distribution of the genetic diversity within populations can diverge from equilibrium models
185 due to demographic changes. We tested whether the populations recently experienced a
186 reduction of their effective size using the approach detailed in Cornuet & Luikart (1996) and
187 implemented in their software BOTTLENECK v. 1.2.02. Using a Wilcoxon test, the observed
188 heterozygosity was compared with the heterozygosity expected under equilibrium,
189 considering a two-phase mutation model (TPM) recommended for microsatellite data (Piry et
190 al. 1999) with 90% single-step mutations and 10% multiple-step mutations (and a variance
191 among multiple step of 12). Populations exhibiting a significant heterozygosity excess would
192 be considered as having experienced a recent genetic bottleneck whereas populations that
193 have been expanding for many generations are characterized by loci exhibiting a
194 heterozygosity deficiency (Cornuet & Luikart 1996).

195 We estimated genetic differentiation between populations by calculating Weir and
196 Cockerham's (1984) estimator of pairwise F_{st} values and carrying out exact tests of allelic
197 differentiation between populations using GENEPOP v.4.0. To adjust for multiple
198 comparisons, the FDR correction was used. Due to the frequent presence of null alleles, we
199 used the program FREENA to calculate pairwise F_{st} estimates corrected for null alleles
200 (F_{st_COR}) using the so-called ENA method (Chapuis & Estoup 2007). This software was also
201 used to estimate the Cavalli-Sforza and Edwards (1967) genetic distance for each pair of
202 populations (D_c) and this distance was also estimated using the INA correction described in
203 Chapuis and Estoup (D_{c_COR} , 2007). Matrices of pairwise genetic distances were compared
204 with Mantel tests (Mantel 1967) using the R program (R Development Core Team 2012).

205 We used the program BAPS v.6 (Corander & Marttinen 2006; Corander *et al.* 2008) to
206 detect clusters of genetically similar populations and to estimate individual coefficients of
207 ancestry with regard to the detected clusters. When testing for population clusters, we ran 5
208 replicates for k = 5, k= 10, k= 15, k= 20, k= 25 and k=30, where k is the maximum number of
209 genetically divergent groups (populations). When estimating individual ancestry coefficients
210 *via* admixture analysis we used recommended values of (i) the number of iterations used to
211 estimate the admixture coefficients for the individuals (100), (ii) the number of reference
212 individuals from each population (200) and (iii) the number of iterations used to estimate the
213 admixture coefficients for the reference individuals (20).

214

215 *Landscape genetics*

216 Landscape elements were mapped at high resolution (precision ~2m) over the whole
217 area. Land use cover and linear elements such as roads and rivers were obtained by merging
218 different sources of data. As background data, we used databases from the French National
219 Geographic Institute (IGN), encompassing shapefiles (BD TOPO, accuracy ~1m), and raster
220 (BD Ortho, resolution = 0.5m) of the year 2010. We crossed this information with field work
221 with a differential GPS with 10 cm real time accuracy, in order to check the boundaries of
222 plots and their management. We also compared our data with Corine Land cover 2006 to
223 identify any inconsistencies. Historical and management information was gained with google
224 maps, from interviews with the farmers, and checked with the different version of Corine
225 Land Cover. The data base and the different geographical layers were built up in ArcGis 10.1
226 (ESRI) in the projection Lambert 93 (EPSG: 2154). Data were stored in a vector format and
227 rasterized at 10m resolution in order to perform the landscape analysis. Polygons and linear
228 elements were rasterized separately and merged in raster format. Linear elements were

229 buffered before rasterization in order to avoid artefact gaps. Landscape structure variables
230 were computed in Fragstats (McGarigal et al. 2012) and were computed at patch scale or at
231 the buffer scale (500m of radius) depending on the metric. Landscape descriptors were then
232 normalized (centred and reduced) and selected for the statistical modelling process based on
233 their Variance Inflation Factor value (VIF), in order to avoid collinearity. There is no
234 theoretical base to choose the threshold of the VIF value to exclude variables, and it is usually
235 recommended to use a predictors with a VIF below ten (Montgomery & Peck 1992; Zuur *et*
236 *al.* 2010; Dormann *et al.* 2013). We used a threshold of six in this study. Landscape structure
237 descriptors were correlated to genetic diversity indices (A_r , H_e) with multiple regression. In
238 order to select the best model, we run all combinations of factors and selected the model with
239 the best weight, based on corrected AIC (AICc) criteria. This approach produces r^2 goodness
240 of fit and avoids over-fitting, thanks to the AIC criteria.

241 Landscape connectivity was performed by defining different scenarios of cost of
242 movements within landscape elements, based on species ecology. Elements were classified in
243 three categories: Barrier, Neutral or Corridors, which corresponded to decreasing movement
244 cost (50, 20, 1 respectively). The result is called a resistance surface map. In all scenarios,
245 urban areas were considered to be barriers; forested areas, hedges and permanent water bodies
246 were considered neutral and temporary water bodies were considered to be corridors. The
247 other elements – grasslands, crops, roads - were considered differently according to the
248 scenario (for details see Table 2). Combining all these possibilities yielded 27 scenarios of
249 resistance surface. In order to test the robustness of our results we also run the analyses by
250 multiplying the costs by 100 in each scenario (giving costs of 5000, 2000, 100). The results
251 were well congruent with initial costs. Connectivity was assessed in three ways. First, simple
252 geographical distance along a straight line between all localities (Euclidian distance) was used

253 to estimate the distance between localities. This scenario makes the assumption that landscape
254 elements do not play a role in dispersal, and is usually referred to as Isolation by Distance
255 (IBD). Second we calculated the least cost path between each pair of site for each of the 27
256 scenarios. This approach makes the assumption that individuals disperse optimally regarding
257 landscape structure, and is usually referred to as Least Cost Path (LCP). Last, we calculated
258 for all the 27 scenarios all paths between each pair of site, weighted by their cumulative cost,
259 to produce 27 corresponding cost weighted distance matrices (CWD), which are usually
260 referred to as resistance distances in circuit theory (McRae & Nürnberger 2006). All these
261 spatial analyses were performed in R with the package {gdistance}. Once all pairwise
262 distances were computed, we looked for the ones that best matched to the (logit transformed)
263 genetic differentiation between populations ($F_{st}/1-F_{st}$ and $F_{st_COR}/1-F_{st_COR}$). This was done
264 using Maximum Likelihood Population Effect (MLPE, Clark et al. 2010; Van Strien et al.
265 2012), a type of linear mixed model that takes into account the non-independence of values
266 within pairwise distance matrices. Marginal r^2 values (R^2_{beta}) were obtained. For this we
267 adapted an R script supplied by Marteen J. Van Strien.

268

269

270 **Results**

271

272 *Microsatellite data*

273 All microsatellite markers were polymorphic across all populations, with 4–19 and 4 -
274 21 alleles per locus for *A. chlorotica* and *A. icterica*, respectively (Supplementary data Tables
275 S1 and S2 respectively). We did not find any evidence of genotypic linkage disequilibrium at
276 any pair of loci in any species. The mean genotyping error rate per locus was 3.12 % and 4.65
277 % in *A. chlorotica* and *A. icterica*, respectively. Genotyping errors were observed in 4 loci
278 over 8 in *A. chlorotica* and they exceeded 5% (i.e. 1 error over 24 genotypes) for the locus
279 Ac419 (8.33%). In *A. icterica*, genotypic errors were observed in 5 loci over 7 and they
280 exceeded 5% for the locus 2PE70 (14.29%). These errors were due to allelic dropout in
281 83% and 71% of the cases in *A. chlorotica* and *A. icterica* respectively. The mean genotyping
282 error rate per allele was 1.56 % and 3.11 % in *A. chlorotica* and *A. icterica*, respectively
283 (ranging from 0% to 4.17% and from 0% to 9.52%, respectively). Significant departures from
284 HWE were observed in 39 of 112 and in 33 of 118 single-locus exact tests after FDR
285 correction in *A. chlorotica* and *A. icterica*, respectively. Across all populations, the presence
286 of null alleles was suggested by MICRO-CHECKER for all *A. chlorotica* loci except Ac 476,
287 with frequencies ranging from 0.08 to 0.34 (Supplementary data Table S1) and for PB10,
288 2PE70 and C4 *A. icterica* loci, with frequencies ranging from 0.13 to 0.41 (Supplementary
289 data Table S2). However, no locus showed null alleles in all populations. A few failures of
290 amplification could be interpreted as null homozygotes that would confirm the presence of
291 null alleles at some loci (Supplementary data Tables S1 and S2). However, amplification
292 failures observed at loci that did not present heterozygote deficit, highlighted that the lack of
293 amplification may be due to causes other than null alleles such as degraded DNA.

294 *Genetic variation within populations*

295 Higher values of genetic diversity were obtained for *A. chlorotica* than for *A. icterica*
296 (Table 1). For example, standardized allelic richness (A_r) ranged from 6.30 to 9.28 and from
297 1.73 to 3.69 in *A. chlorotica* and *A. icterica*, respectively. A significant heterozygosity excess
298 (Wilcoxon test, $P < 0.05$) indicating a recent bottleneck was detected in 1 *A. chlorotica* and 5
299 *A. icterica* populations (Table 1). Significant F_{is} was observed in almost all populations
300 except in P_B and I36 for *A. chlorotica* and in I03, I10 and I31 for *A. icterica* (Table 1).

301

302 *Relationship between genetic diversity and local landscape structure*

303 Landscape features in buffers were correlated to genetic diversity in *A. icterica* and *A.*
304 *chlorotica* (Table 3). In *A. icterica*, the r^2 between A_r and H_e , and landscape features were
305 respectively 0.51 and 0.60 and both were significant ($p < 0.05$). In this species, patch diversity
306 and patch richness were significantly linked to A_r and H_e . In *A. chlorotica*, r^2 was 0.38 and
307 0.42, for A_r and H_e respectively, and both indices were linked to surface in pastures.

308

309 *Genetic structure at regional scale*

310 The statistical power for both sets of microsatellite loci to detect various levels of true
311 population differentiation (F_{st}) between populations is presented Table 4. Both sets of markers
312 will detect a true F_{st} of 0.005 or larger with a probability of 96% or more. The alpha error
313 (corresponding to the probability of obtaining false significances when true $F_{st} = 0$) was close
314 to 5% in all cases.

315 All genetic distances matrices ($F_{st}/1-F_{st}$, $F_{st_COR}/1-F_{st_COR}$, D_c and D_{c_COR}) were
316 significantly correlated with Mantel r value ranging from 0.77 ($p = 0.001$) to 0.99 ($p = 0.001$)

317 in *A. icterica* and from 0.88 ($p = 0.001$) to 0.99 ($p = 0.001$) in *A. chlorotica* (Supplementary
318 data Tables S3, S4 and S5).

319 F_{st} analysis showed significant genetic structure at the level of the whole study for
320 both species ($F_{st} = 0.059$, $F_{st_COR} = 0.055$, $P < 0.001$ and $F_{st} = 0.152$, $F_{st_COR} = 0.138$, $P < 0.001$
321 for *A. chlorotica* and *A. icterica*, respectively). Pairwise F_{st} estimates ranged from 0.008 to
322 0.116 (F_{st_COR} ranged from 0.009 to 0.105) and 0.005 to 0.430 (F_{st_COR} ranged from 0.004 to
323 0.412) for *A. chlorotica* and *A. icterica*, respectively (Supplementary data Tables S3 and S4
324 respectively). All exact tests of allelic differentiation were significant ($P \leq 0.005$). Analyses
325 using BAPS identified 8 and 12 genetic clusters in *A. chlorotica* and *A. icterica*, respectively
326 ($P = 0.99$ and $P = 1$ respectively). For *A. chlorotica*, one cluster was composed of 4
327 populations that were close geographically to one another (P_A , P_B , I32 and I33), another
328 cluster was composed of the I07, I10, I15 and I18 populations and all other populations
329 corresponded to a different cluster. For *A. icterica*, 5 clusters were composed of two
330 geographically close populations (P_A and P_B , I02 and I03, I04 and I32, I07 and I08, I11 and
331 I25) while all other clusters were composed of only one population. Low levels of admixture
332 were observed among the clusters (Supplementary data Fig S1).

333

334 *Relationship between genetic differentiation and landscape connectivity*

335 Genetic differentiation was linked to landscape connectivity mostly in *A. chlorotica*
336 (Table 5, Fig 3). In this species resistance distance (CWD) had the most explanatory power,
337 followed by least cost path (LCP) and finally by isolation by distance (IBD). In *A. chlorotica*,
338 the best scenarios were those in which grasslands were considered to be corridors, whereas
339 crops and roads were considered to be barriers (Table 5). In *A. icterica*, no significant
340 isolation by distance was found except with the non-corrected D_c distance and the only

341 common point between the several most likely scenarios was that crops were considered as
342 barriers (Table 5). The best congruent models between the two species, taking into account
343 the results from the different genetic indices, were scenarios 9, 10 and 11. The most frequent
344 role of the different landscape element in these scenarios was corridor for grasslands and
345 barrier for crops and roads. Despite significant p-value, it is noteworthy that landscape
346 connectivity was poorly linked to genetic differentiation between *A. ictERICA* populations in all
347 cases.
348

349 **Discussion**

350

351 *Microsatellite markers in earthworms*

352 Microsatellites markers have been developed for only a few earthworm species (i.e. 7
353 species so far, review in Torres-Leguizamon et al. 2012; Souleman et al. 2016) and these
354 markers have rarely been used for population genetics studies (but see Velavan et al. 2009;
355 Novo et al. 2010; Dupont et al. 2015). Two different research groups have tried to develop
356 reliable markers for one of the most emblematic European earthworm species, *Lumbricus*
357 *terrestris*. Of the ten markers obtained in this species by Velavan *et al.* (2007), only three
358 were used in a subsequent study (Velavan et al. 2009) suggesting difficulties in genotyping
359 the samples with the other ones. Souleman et al. (2016) couldn't obtain reliable results with
360 these markers. Thus, they developed eight new markers for which they obtained a low
361 amplification success and a significant heterozygote deficit, suggesting null alleles. In our
362 study, null alleles were suspected at seven out of eight loci in *Allobophora chlorotica* and at
363 four out of eight loci in *Aporrectodea icterica*. It is already known that the development of
364 microsatellite molecular markers can be problematic in some taxa (e.g. in molluscs,
365 McInerney et al. 2011 and Lepidoptera, Schmid *et al.* 2016). It was proposed that such
366 methodological difficulties may have been caused by genomic complexities contained within
367 microsatellite flanking regions. In particular, unstable flanking regions may arise when indels
368 or mutations occur at PCR primer binding sites, thereby causing null alleles (McInerney et al.
369 2011). We therefore believe that microsatellite flanking regions are particularly variable in
370 earthworm species. This could be verified by gathering more genomics data on these taxa.
371 Nevertheless, it seemed that the estimation of genetic divergence was not significantly altered
372 by the presence of null alleles in the dataset. Indeed, similar results were obtained with all

373 indices of genetic divergence (F_{st} , F_{st_COR} , D_c , D_{c_COR}), and correction for the presence of null
374 alleles did not change the results.

375

376 *Landscape structure and population genetic diversity*

377 Agriculture and urbanization result in habitat loss and fragmentation that variously
378 impact many animal groups. Anthropogenic landscape fragmentation results in reduced size
379 and increased isolation of habitat patches. Fragmented populations are thus expected to
380 experience increased genetic drift and reduced gene flow, which result in the erosion of
381 genetic diversity and the increase of genetic differentiation among local populations
382 (Keyghobadi 2007). Moreover, small populations isolated by surrounding inhospitable
383 landscapes are more vulnerable to demographic variability, environmental stochasticity and
384 genetic processes including inbreeding depression, the random fixation of deleterious alleles
385 and the loss of adaptive potential (Frankham 1995).

386 In this study, we tested how landscape structure in a man-made environment impacted
387 genetic diversity of earthworm populations by characterizing landscape at the buffer scale. A
388 significant relationship was observed between genetic diversity indices (H_e and A_r) and two
389 features of the immediately surrounding landscape (i.e. patch diversity and patch richness,
390 Table 3) in *A. icterica* while in *A. chlorotica* genetic diversity indices were positively linked
391 to surface in pasture. We thus confirmed that geographic isolation of *A. icterica* populations
392 due to natural and artificial barriers to gene flow probably accentuate the loss of genetic
393 variability through genetic drift, such as already suggested in a previous population genetic
394 study of this species (Torres-Leguizamon et al. 2014). Interestingly, one quarter of the *A.*
395 *icterica* populations seemed recently founded, such as revealed by the heterozygosity excess
396 in these populations. Overall, these results suggest that demographic changes occur more

397 frequently in *A. icterica* than in *A. chlorotica* and that these demographic changes can be
398 explained by the local landscape structure.

399

400 *Genetic differentiation between populations*

401 A local decline of effective population size may be explained by the disruption of
402 historical patterns of gene flow in a fragmented habitat (Keyghobadi 2007). Analyses of
403 spatial patterns of genetic structure showed the presence of a strong genetic differentiation in
404 both species, with a low degree of genetic admixture and high F_{st} values. F_{st} values were
405 higher for *A. icterica*, highlighting that these populations are more genetically isolated than
406 the ones of *A. chlorotica*. This was not expected because *A. icterica* has a higher potential for
407 active dispersal. Caro *et al.* (2013) indeed demonstrated in a mesocosm study that *A. icterica*
408 had a higher dispersal rate than two other endogeic species, namely *A. chlorotica* and
409 *Aporrectodea caliginosa*. Moreover, in a recent population genetic study at very fine scale,
410 Dupont *et al.* (2015) showed a low signal of genetic structure within two *A. icterica*
411 populations sampled in two plots of less than 1 ha separated by ~ 500m while *A. chlorotica*
412 populations showed spatial neighbourhood structure in the same sites. This difference was
413 interpreted as a higher dispersal capacity of *A. icterica*. In the light of the results at very fine
414 scale (Dupont *et al.* 2015) and at landscape scale (this study), we can assume that *A.*
415 *chlorotica* essentially disperse through passive mechanisms over larger distance while passive
416 dispersal might be more restricted for *A. icterica*. *A. chlorotica* is a small bodied species and
417 lives near the soil surface in the upper 60 mm soil layer (Sims & Gerard 1999), two features
418 probably facilitating dispersal *via* various vectors (e.g. zoochory, wind, water and soil transfer
419 *via* human activities) while *A. icterica* is found deeper in the soil and is bigger.

420

421 *Landscape connectivity at regional scale*

422 The landscape connectivity analysis revealed that genetic structure was linked to
423 landscape connectivity in *A. chlorotica*, with resistance distance (cwd) having the most
424 explanatory power. Thus, landscape features better explain genetic structure than Euclidian
425 distances in this species. We specifically tested the hypothesis that linear features such as
426 roads may function as dispersal corridors (see for instance Tyser & Worley 1992; Cameron &
427 Bayne 2009) for these species. It has indeed been shown that European earthworms that are
428 invasive in Canada and the northern USA were introduced and spread along road networks
429 (review in Cameron & Bayne 2009). It was however not clear whether the spread of
430 earthworms along roads is more likely to occur *via* initial transport of earthworms or their
431 cocoons in soil or gravel during road construction or *via* transport by vehicles after the road
432 has been built (Cameron & Bayne 2009). In addition, roads and sidewalks could also function
433 as dispersal corridors when earthworms crawl out of the soil and disperse at night after heavy
434 rain, as is often observed in some species (e.g. Chuang & Chen 2008). Our results rather
435 suggested that roads constitute obstacles for earthworm dispersal. Using MLPE, we indeed
436 showed that the majority of the most likely landscape connectivity scenarios considered roads
437 as barriers (Table 5).

438 The second hypothesis tested was that grasslands represent a suitable habitat that
439 could be easily crossed by endogeic species and thus represent dispersal corridors, while soil
440 under crops has a detrimental effect on earthworms. These expectations were confirmed by
441 the MLPE analysis, the most likely landscape connectivity scenarios generally considered
442 grasslands as corridors and crops as barriers.

443 Thus, this study suggests that landscape configuration influences gene flow in
444 earthworms. However, no distinct dispersal scenario could be clearly identified, which

445 suggests that passive dispersal due to human activities has also probably played a significant
446 role in shaping genetic structure in both species.

447

448 **Conclusion**

449 Simultaneously investigating two ecologically similar species highlighted several
450 common features in the response of each species to the landscape. We showed that functional
451 connectivity was impacted by landscape features and that a favourable habitat could act as a
452 corridor for the dispersal of earthworms. We thus confirmed that the aboveground landscape
453 play a part in dispersal and gene flow of below-ground organisms. However, we also
454 observed some differences between species which could be linked to the dispersal and life
455 history attributes of each species. Indeed, population genetic diversity was significantly
456 influenced by the local landscape structure in both species, but only the genetic differentiation
457 of *A. chlorotica* populations was related to functional connectivity. This result highlights that
458 the exact effect of each habitat type on genetic variation over space and time and of
459 agricultural practices on earthworm dispersal should be studied using specific sampling
460 strategies.

461

462

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469

470

471

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634 **Data Accessibility**

635 The data sets with the microsatellite genotypes of *A. chlorotica* and *A. icterica* are available
636 from the Dryad Digital Repository: <http://dx.doi.org/10.5061/dryad.sc6bb>.

637

638

639 **Author Contributions**

640 J.M initiated the project. L.D. and J.M. designed the study. L.D. drafted the manuscript. L.D.,
641 M.T.L., P. R.-C. and J.M. collected the data. L.D and P. R.-C. performed molecular analyses.
642 J.M. built the GIS database and conducted the landscape genetic analyses (Buffer and
643 connectivity). L.D. and J.M. performed statistical analyses and wrote the manuscript.
644

645

646 **Figure legends**

647

648 **Figure 1**

649 Illustration of the different landscape genetic approaches used in this work. a) Local genetic
650 structure is compared to local landscape features within a buffer around each population; b)
651 genetic difference is linked to geographical distance between populations (IBD); c) genetic
652 difference is linked to the functional distance between populations, calculated as the least cost
653 path between populations (LCP), based on a resistance map; d) genetic difference is linked to
654 the functional distance between populations, calculated as the sum of the cost weighted paths
655 between populations (cwd), based on a resistance map.

656

657 **Figure 2**

658 Geographical distribution of earthworm sampling sites near Yvetot in Normandy (France).
659 “No species” means that neither *A. icterica* nor *A. chlorotica* were found.

660

661 **Figure 3**

662 Best correlation found between genetic distance (D_c distance) and resistance distance scenario
663 (cwd) for *A. chlorotica* (A) and *A. icterica* (B).

664

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667

Table 1. Geographical location*, sample size (N), genetic diversity (A_r : standardized allelic richness; H_e : expected heterozygosity), fixation index F_{is} (significant values are in bold) and Wilcoxon test P value for heterozygote excess compared to expectations at mutation-drift equilibrium (p_{wil}) for each study plot and each species. *Latitude/longitude range in Lambert II étendu.

Plot	Latitude	Longitude	<i>Allolobophora chlorotica</i>					<i>Aporrectodea icterica</i>				
			N	A_r	H_e	F_{is}	p_{wil}	N	A_r	H_e	F_{is}	p_{wil}
P _A	484578.509	2513691.360	42	8.61	0.777	0.128	0.727	22	2.73	0.523	0.352	0.055
P _B	484100.668	2513845.836	30	8.44	0.794	0.091	0.527	30	2.51	0.481	0.094	0.016
I02	484907.071	2511340.636	-	-	-	-	-	12	2.27	0.397	0.274	0.281
I03	484824.797	2511382.199	-	-	-	-	-	28	2.32	0.395	0.115	0.015
I04	483636.894	2511737.970	-	-	-	-	-	12	2.55	0.438	0.103	0.344
I07	483713.045	2511076.294	28	8.60	0.775	0.185	0.371	24	2.95	0.494	0.380	0.281
I08	483823.122	2511110.760	-	-	-	-	-	31	2.71	0.484	0.209	0.406
I10	483411.237	2511337.635	32	7.75	0.781	0.233	0.422	14	2.61	0.495	0.167	0.008
I11	485746.167	2507374.639	27	7.45	0.765	0.195	0.527	29	2.81	0.449	0.172	0.406
I15	481854.533	2514843.110	27	9.28	0.777	0.141	0.727	30	2.54	0.461	0.065	0.008
I18	479823.687	2512611.239	27	8.08	0.742	0.205	0.973	30	3.69	0.575	0.277	0.344
I19	492929.229	2508850.328	-	-	-	-	-	43	3.00	0.493	0.335	0.711
I20	477908.835	2515839.136	29	7.06	0.725	0.293	0.808	33	2.15	0.373	0.184	0.312
I25	480555.744	2508635.999	29	6.30	0.720	0.229	0.527	31	2.84	0.506	0.263	0.078

I27	489902.819	2513148.683	30	8.81	0.788	0.156	0.770	34	2.62	0.485	0.317	0.023
I31	482053.828	2513328.538	-	-	-	-	-	27	1.73	0.226	0.115	0.594
I32	483297.736	2513724.785	29	8.16	0.786	0.179	0.273	30	2.70	0.422	0.305	0.656
I33	481742.786	2511459.498	32	8.61	0.761	0.182	0.902	29	2.52	0.454	0.179	0.148
I34	482430.931	2510886.051	32	7.95	0.747	0.237	0.875	30	2.33	0.366	0.403	0.078
I36	487833.362	2509076.597	46	6.67	0.777	0.067	0.010	-	-	-	-	-

Table 2 Role and cost of landscape elements in the different landscape scenarios tested in this study. The cost of grasslands, crops and roads changed among scenarios. Costs of permanent water bodies (20), temporary water bodies (1), deciduous forest (20), coniferous forest (20), hedges (20) and urban area (50) were fixed in all scenarios.

Scenario	Role of elements			Cost of elements		
	Grasslands	Crops	Roads	Grasslands	Crops	Roads
1	Barrier	Barrier	Barrier	50	50	50
2	Barrier	Barrier	Neutral	50	50	20
3	Barrier	Barrier	Corridor	50	50	1
4	Barrier	Corridor	Barrier	50	1	50
5	Barrier	Corridor	Neutral	50	1	20
6	Barrier	Corridor	Corridor	50	1	1
7	Barrier	Neutral	Barrier	50	20	50
8	Barrier	Neutral	Neutral	50	20	20
9	Barrier	Neutral	Corridor	50	20	1
10	Corridor	Barrier	Barrier	1	50	50
11	Corridor	Barrier	Neutral	1	50	20
12	Corridor	Barrier	Corridor	1	50	1
13	Corridor	Corridor	Barrier	1	1	50
14	Corridor	Corridor	Neutral	1	1	20
15	Corridor	Corridor	Corridor	1	1	1
16	Corridor	Neutral	Barrier	1	20	50
17	Corridor	Neutral	Neutral	1	20	20
18	Corridor	Neutral	Corridor	1	20	1
19	Neutral	Barrier	Barrier	20	50	50
20	Neutral	Barrier	Neutral	20	50	20
21	Neutral	Barrier	Corridor	20	50	1
22	Neutral	Corridor	Barrier	20	1	50
23	Neutral	Corridor	Neutral	20	1	20
24	Neutral	Corridor	Corridor	20	1	1
25	Neutral	Neutral	Barrier	20	20	50
26	Neutral	Neutral	Neutral	20	20	20
27	Neutral	Neutral	Corridor	20	20	1

Table 3 Summary of the best multiple regression between landscape features (predictors) and genetic diversity (A_r : Allele Richness, H_e : expected heterozygosity) of the populations of the two earthworm species *A. icterica* and *A. chlorotica* in the region of Yvetot, Normandy, France, based on AICc weight. A patch represents an element in the landscape and the buffer radius was 500m. Predictors are: Edge Density (length of patch edge/surface of the buffer), Patch Richness (number of different types of patches in the buffer), Patch Diversity (Shannon Diversity of the different Land Use types), Crop surface, Grassland surface and Total length of roads.

	<i>Aporrectodea icterica</i>		<i>Allolobophora chlorotica</i>	
	A_r	H_e	A_r	H_e
Anova table				
F	8.50	11.90	7.50	8.60
p - value	0.03	<10 ⁻⁴	0.02	0.01
adjusted r ²	0.51	0.60	0.38	0.42
AICc	- 20	- 52	- 19	- 66
model weight	0.43	0.32	0.34	0.33
Predictors				
Edge Density	ns	ns	ns	ns
Patch Richness	-0.12	-0.05	ns	ns
Patch Diversity	0.13	0.07	ns	ns
Crop surface	ns	ns	ns	ns
Grassland surface	ns	ns	0.1	0.1
Total length of roads	ns	ns	ns	ns

Table 4 Statistical power for detecting various true levels of population differentiation (F_{st}) by means of Fisher's exact test when using both sets of microsatellite loci, allele frequencies, and sample sizes. The power is expressed as the proportion of simulations that provide statistical significance at the 0.05 level.

True F_{st}	<i>A. chlorotica</i>	<i>A. icterica</i>
0.000	0.064	0.070
0.001	0.417	0.196
0.002	0.852	0.432
0.005	1.000	0.964
0.010	1.000	1.000

Table 5 Summary of the link between landscape connectivity and genetic structure (MLPE), comparing **A.** Euclidian distance and genetic differentiation (pairwise F_{st} , pairwise F_{st} estimates corrected for null alleles, D_c genetic distance and D_c genetic distance corrected for null alleles) between populations (Isolation by distance, IBD).and **B.** cost weighted distances (cwd) and genetic differentiation between populations. The most likely landscape connectivity scenarios are indicated. The roles of the landscape elements in the most likely scenarios are specified (b = barrier, c = corridor and n = neutral); when applicable the most frequent role is in bold. In **C.** the best congruent scenarios are presented. The R^2_{β} statistic described by Van Strien *et al* (2012) is presented. NA = Not Applicable

A. IBD								
	Genetic distance	Geographic distance	scenario	Grasslands	Crops	Roads	R^2_{β}	pvalue
<i>A. chlorotica</i>	F_{st}	euclidian	NA	-	-	-	0.29	<10 ⁻⁵
	F_{st-COR}	euclidian	NA	-	-	-	0.28	<10 ⁻⁵
	D_c	euclidian	NA	-	-	-	0.4	<10 ⁻⁵
	D_{c-COR}	euclidian	NA	-	-	-	0.41	<10 ⁻⁵
<i>A. icterica</i>	F_{st}	euclidian	NA	-	-	-	0.01	0.12
	F_{st-COR}	euclidian	NA	-	-	-	0.02	0.09
	D_c	euclidian	NA	-	-	-	0.03	<10 ⁻⁵
	D_{c-COR}	euclidian	NA	-	-	-	0.02	0.06
B. Best species specific scenario								
	Genetic distance	Geographic distance	scenario	Grasslands	Crops	Roads	R^2_{β}	pvalue
<i>A. chlorotica</i>	F_{st}	CWD	10	c	b	b	0.44	<10 ⁻⁵
	F_{st-COR}	CWD	10	c	b	b	0.43	<10 ⁻⁵

	D_c	CWD	8, 9, 10	b, c	b, n	b, n, c	0.54	$<10^{-5}$
	D_{c-COR}	CWD	8, 9, 14	b, c	n, c	n, c	0.54	$<10^{-5}$
<i>A. icterica</i>	F_{st}	CWD	1, 2, 3, 11, 12, 21	b, n, c	b, c	b, n, c	0.04	$<10^{-4}$
	F_{st-COR}	CWD	1, 2, 3, 11, 12, 21	b, n, c	b, c	b, n, c	0.05	$<10^{-4}$
	D_c	CWD	1, 2, 3, 11, 12, 21	b, n, c	b, c	b, n, c	0.08	$<10^{-5}$
	D_{c-COR}	CWD	1, 11, 12, 21, 25, 26, 27	b, n, c	b, n	b, n, c	0.06	$<10^{-5}$

C. Best congruent scenario

	Genetic distance	Geographic distance	scenario	Grasslands	Crops	Roads	R^2_β	pvalue
<i>A.chlorotica</i>	F_{st}	CWD	10	c	b	b	0.44	$<10^{-5}$
	F_{st-COR}	CWD	10	c	b	b	0.43	$<10^{-5}$
	D_c	CWD	9, 10	b, c	n, b	b, c	0.54	$<10^{-5}$
	D_{c-COR}	CWD	9, 10, 11	b, c	n, b	b, n, c	0.54	$<10^{-5}$
<i>A.icterica</i>	F_{st}	CWD	10	c	b	b	0.04	$<10^{-4}$
	F_{st-COR}	CWD	10	c	b	b	0.04	$<10^{-4}$
	D_c	CWD	9, 10	b, c	n, b	b, c	0.07	$<10^{-5}$
	D_{c-COR}	CWD	9, 10, 11	b, c	n, b	b, n, c	0.06	$<10^{-4}$

Figure 1

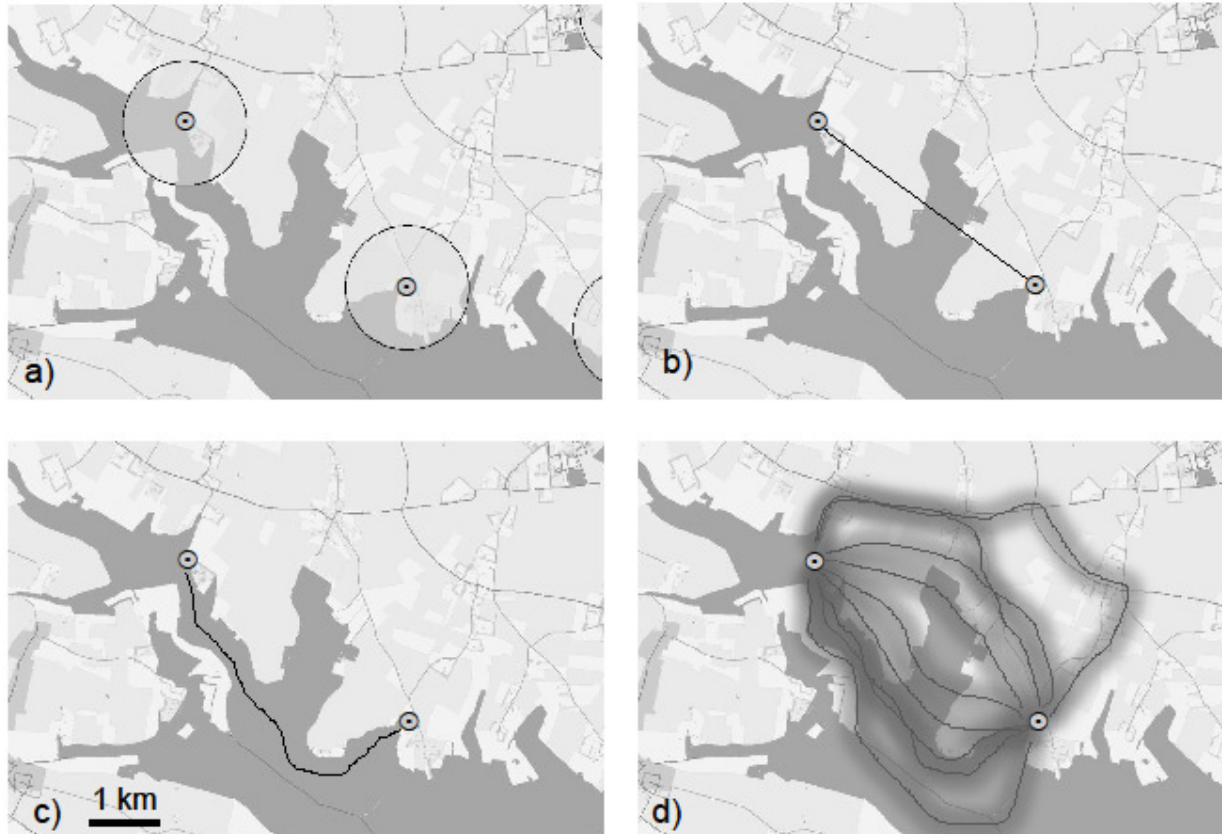


Figure 2

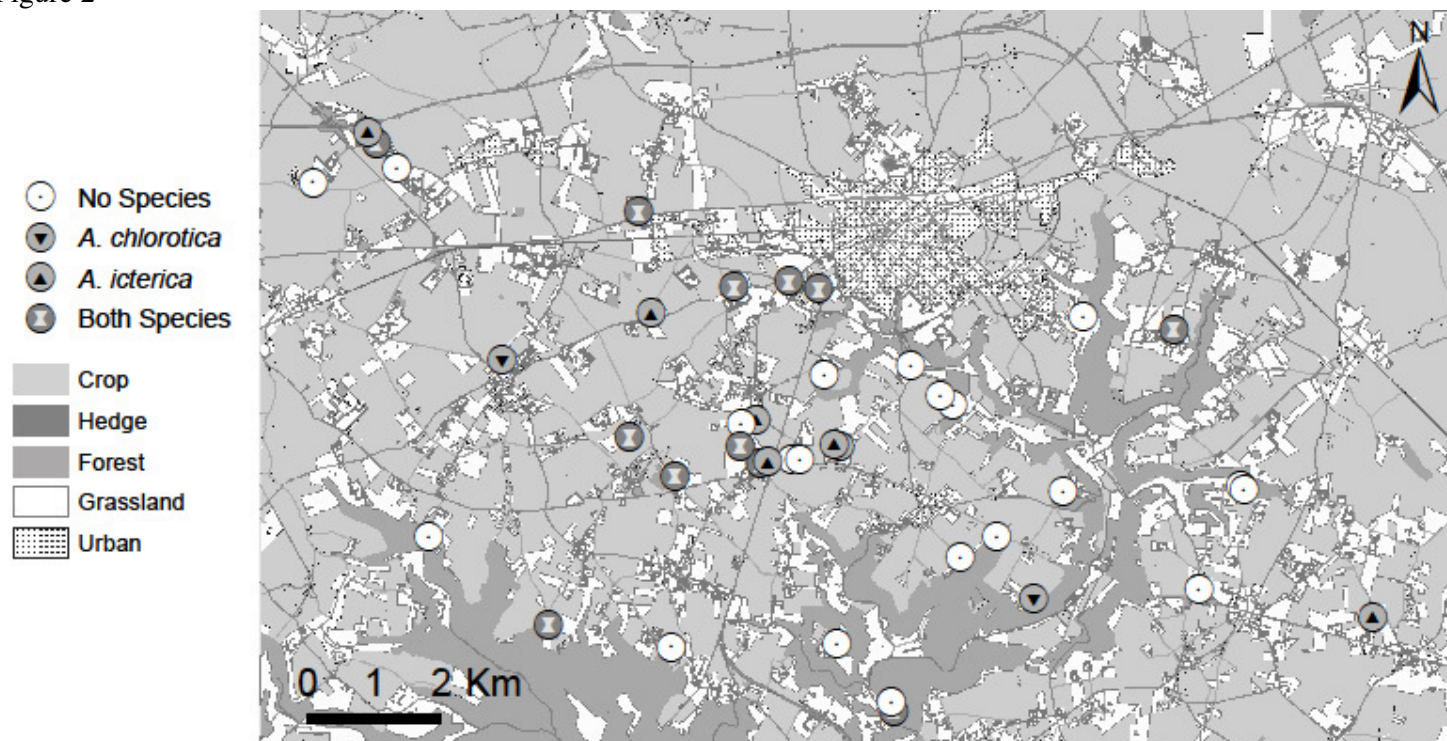


Figure 3

