



The role of dispersal and local environment in urban land snail assemblages: an example of three cities in Central Italy

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(Article begins on next page)

1 **The role of dispersal and local environment in urban land snail assemblages: an example of three cities in central**
2 **Italy**

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19 *constructive comments and criticisms.*

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22 **Abstract** Ecologists increasingly appreciate the central role that urban biodiversity plays in ecosystems, however much
23 urban biodiversity is neglected, especially some very diverse groups of invertebrates. For the first time in southern
24 Europe, land snail communities are analysed in four urban habitats along a geographical gradient of three cities, using
25 quantitative methods and assessing the relative roles of local environmental conditions (“distance from sea”, “distance
26 from city centre”, “vegetation cover”) and spatial effects by principal coordinate analysis of neighbour matrices,
27 redundancy analysis and variation partitioning. A total of 53 species was recorded, a richness similar to that of natural
28 areas. At habitat level, species richness did not show a clear increasing trend from more to less urbanized habitats, but
29 rather a homogeneous pattern. At city level, study areas hosted rather heterogeneous species assemblages and biotic
30 homogenization did not seem to have any impact; thus, only three species could be considered alien. Variation
31 partitioning showed that land snail communities were mostly structured by environmental factors, even when spatial
32 structures independent of measured environmental variables were included: “vegetation cover” and “distance from city
33 centre” were the environmental variables that explained most of the variation in species composition. The lack of strong
34 spatial structure also unexpectedly suggested that transport by humans aids dispersal of organisms with low mobility,
35 which are usually limited by spatial constraints in natural environments. These results provide ecological and
36 conservation implications for other invertebrate groups, suggesting to set priorities in management strategies that
37 include habitat conservation at local scale.

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39 **Introduction**

40

41 Nowadays more than half the world's population lives in large urban settlements. This percentage is predicted to
42 increase to 66% by 2050 (United Nations 2014). Although it is the most common human habitat (Grimm et al. 2008),
43 the urban environment has been neglected by researchers (Maurer et al. 2000; Beninde et al. 2015; Hartop et al. 2015)
44 as unnatural and poor in biodiversity (Grimm et al. 2008; Paul and Meyer 2008; Güneralp and Seto 2013), besides
45 being subject to biotic homogenization due to the increasing number of non-native species (Lockwood 2004; McKinney
46 2006; La Sorte et al. 2007; Williams et al. 2009) and habitat fragmentation (McKinney 2002; Cadenasso et al. 2007).
47 While urbanization is certainly a major cause of biodiversity and habitat loss (Czech et al. 2000; McKinney 2006;
48 Grimm et al. 2008; Hahs et al. 2009), the influence of urban land use on biodiversity is more complex than expected
49 (McKinney 2008).

50 In recent years, the belief that cities only affect the surrounding environment negatively has been challenged. An
51 increasing body of literature has investigated urban biodiversity (Beninde et al. 2015), covering groups such as birds,
52 invertebrates, plants and even zooplankton (Mimouni et al. 2015). Surprisingly, researchers are discovering that cities
53 can protect and control vulnerable and particular ecosystems (UNEP 2005), as well as high levels of biodiversity
54 (Aronson et al. 2014) including native species (Barratt et al. 2015), sometimes showing the same dynamic interactions
55 as in nature (Baldock et al. 2015; Beninde et al. 2015). Studies have revealed the extreme ecological importance of
56 particular urban habitats for species dispersal (Angold et al. 2006). Moreover, urban green areas can have a positive
57 influence on human quality of life (Fuller et al. 2007; Mitchell and Popham 2008; Carrus et al. 2015; Taylor and
58 Hochuli 2015) in terms of recreation and public participation in scientific activities (Beumer and Martens 2015).

59 Although interest in urban ecology is increasing, not much is yet known about less charismatic invertebrates such as
60 land snails. This is surprising because according to the IUCN Red List, molluscs are the group most subject to
61 extinctions, which often go completely unnoticed (Lydeard et al. 2004; Régnier et al. 2009). The current lack of
62 scientific documentation and their low mobility make land snails susceptible to anthropogenic disturbance and habitat
63 fragmentation (Douglas 2011). Malacofauna currently includes several rare and endangered species, and constitutes
64 20% of all threatened animals and 37% of known animal extinctions since the year 1600 (Seddon 1998). Land snails are
65 numerous and diverse; they live in almost all terrestrial environments and play a key role in worldwide trophic webs
66 (Baur and Baur 1993; Douglas et al. 2013). Snails are preyed on by a wide range of animals from insects to salamanders,
67 toads, lizards, snakes, birds and mammals. Some species of fireflies (i.e. *Pyrocoelia pectoralis*, *Cratomorphus* spp.)
68 feed mainly on land snails (Viviani 2001; Wang et al. 2007). Many birds require a huge quantity of calcium carbonate
69 for the formation of eggshells, relying on land snails for Ca supplementation (Mänd et al. 2000). Their low mobility
70 makes snails, who also have good bioaccumulation capacity (Pauget et al. 2013), perfect bioindicators of environmental
71 quality (Cuttelod et al. 2011; Rota et al. 2016).

72 Studies on urban ecology are often descriptive checklists of traditional biodiversity components, such as richness
73 and diversity measures. Since interactions between communities and their physical environment, and between
74 organisms, occur at precise spatial and temporal scales (Borcard et al. 2004), discovering spatial structures at each scale
75 and the processes involved in their creation is important for understanding the ecological patterns of natural
76 communities (Borcard et al. 2004). In ecology, space has a key role in shaping the distribution of species assemblages,
77 even if it is often neglected. Spatial heterogeneity is often driven by a wide range of factors that interact with each other
78 in different ways (Borcard et al. 1992). The traditional niche-based model assumes that the local environment itself
79 controls the spatial structure of species assemblages: in other words the interaction between a species and its physical

80 environment may mould its distribution through adaptation to and interaction with different habitat features (Borcard et
81 al. 2004). According to this model, it is the environment itself, through resource types and habitat conditions, that
82 determines how similar/dissimilar communities are, irrespective of spatial proximity or distance between assemblages
83 (Caruso et al. 2012). Sometimes spatial structures may be generated by species assemblages themselves and can persist
84 despite environmental changes (neutral model) (Hubbell 2001). In this case the space effect is described directly by
85 community dynamics through dispersal limitations, demographic stochasticity or competitive/predatory processes
86 without any environmental influence (Legendre et al. 2009).

87 Advanced and nowadays well established multivariate techniques such as principal coordinate analysis of neighbour
88 matrices (PCNM) and variation partitioning (ter Braak 1986; Borcard et al. 1992; Legendre and Legendre 1998; Borcard
89 and Legendre 2002; Borcard et al. 2004) have recently made it possible to disentangle and quantify the relative and pure
90 effects of environment and space in shaping variations in community composition (β diversity), as well as the fraction
91 of species variation explained by space and environment together. These techniques provided a useful tool for
92 quantifying the amount of variation uniquely attributable to measured environmental factors, separating it from spatial
93 structures potentially created by unmeasured environmental factors but also population and dispersal dynamics
94 independent of environmental factors (e.g Legendre et al. 2009).

95 In southern Europe, there has been a complete lack of any quantitative and spatially explicit research into any aspect
96 of urban land snail biodiversity, whereas in central Europe Horsák et al. (2009) and Lososová et al. (2011) have been
97 the only authors to compare plant and land snail diversity in different urban habitats across several cities, using a
98 standardized sampling protocol. Chytrý et al. (2012) subsequently highlighted the importance not only of the
99 environmental but also of the space effect in shaping urban community structure across groups of different sizes and
100 with different dispersal abilities (i.e. subaerial cyanobacteria and algae, vascular plants, land snails, grass, and animal-
101 dispersed or wind-dispersed trees and shrubs) in central European cities. Thus we have a major gap in scientific
102 knowledge, since the Mediterranean is one of the world's richest areas in terms of species diversity, but also one of the
103 most threatened, mainly by human impact, especially habitat loss and degradation (Cuttelod et al. 2008). In this
104 geographical context, the extremely complex biogeographical framework of Italy makes it a huge source of
105 biodiversity: with regard to molluscs, the second in Europe, ranking just after Greece in terms of species and subspecies
106 richness (Cuttelod et al. 2011).

107 Here for the first time, the spatial variation component of urban land snail communities is studied in different
108 habitats along a geographical gradient of three cities in Tuscany (Italy). The aims of the study were: 1) to analyse and
109 compare land snail biodiversity in terms of richness and potential differences in species composition (alpha α , beta β
110 and gamma γ diversity) in four different human habitats along a gradient of three cities in central Italy; 2) to evaluate
111 the presence of species of European Conservation Concern (ECC) and the impact of alien species, if any; 3) to detect
112 factors affecting community structure by partitioning the pure effect of "environment" (measured environmental factors)
113 from that of "space" (unmeasured environmental variation, biotic interactions that structure population spatially even
114 within an environmentally homogeneous habitat and also dispersal limitations), as well as the combined effect, on the
115 distribution of urban land snail communities by means of multivariate variation partitioning.

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118 **Materials and methods**

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120 **Study area**

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We investigated three central Italian cities along a geographical transect about 100 km long, extending from the Tyrrhenian coast to the Apennine mountain chain (Fig. 1). The cities differ for example in altitude, population and foundation date, but all have old city walls.

Grosseto (altitude 10 m, population 82,284, area 474 km²) is located in southwestern Tuscany, 10 km from the Tyrrhenian coast. Situated in the alluvial plain of the Ombrone river, the largest plain in southern Tuscany, Grosseto was founded in the High Middle Ages. Its Medicean walls were begun in 1574, replacing those from the 12th-14th centuries. Climate is Mediterranean with mild wet winters and hot dry summers.

Siena (altitude 322 m, population 52,774, area 118 km²) is located in central Tuscany and is the highest of the three cities and the one with the smallest population. Founded in Roman times, it has well preserved green areas within its perimeter. The historical centre of Siena, a UNESCO World Heritage Site, is surrounded by medieval walls, which include the Medici fort (1561-1563). Climate is sub Mediterranean with precipitation sufficiently distributed throughout the year, so there is no real dry season.

Arezzo (altitude 296 m, population 99,232, area 386 km²) is further east, near the Apennines. It is the oldest of the three, believed to have been one of twelve major Etruscan cities. Climate is continental due to its location straddling the Arno and Chiana valleys at the foot of the Apennines. In the high part of the town the Medicean walls enclose the cathedral, town hall and Medici Fortress.

Sampling design and data collection

Sampling design included four urban habitat types over a gradient of human impact from heavily affected and disturbed sites to quite well preserved naturalized habitats (Lososová et al. 2011). Google Earth and city maps were used to choose two spatially independent sites for each habitat type in the three areas, making a total of 24 sites. Habitat types were chosen according to the standardized protocol of Lososová et al. (2011) with modifications: 1. WALLS: historical walls of the city centre; 2. PARK: city parks and public gardens with old deciduous trees (tree cover 10–50%) and frequently mown lawns; 3. E-SUC: early successional sites, strongly disturbed in the last 1–3 years, with a prevalence of bare ground and sparse vegetation cover, usually in or around construction and industrial sites; 4. WOOD: quite well preserved wooded patches, with well-structured vegetation (old and young trees, especially holm oak with scattered shrubs).

A qualitative visual search (presence/absence) was conducted for a standard time of 2 h at each site. The time-constrained sampling strategy made it possible to maximize species numbers by including a huge number of heterogeneous microhabitats and environmental conditions. It also avoided logistic problems related to sampling activity on vertical surfaces (city walls), where a common standard-sized plot can be hard to arrange.

In each site, 5 litres of debris and leaf litter was also collected to detect small and very small species (Cameron and Pokryszko 2005). Despite the huge investment in effort and time, this “volume method” combined with visual search produces more information on small species (Menez 2007; Benocci et al. 2015).

The litter was sieved with a 10 mm mesh, then dried and sieved again with meshes of decreasing size down to 0.5 mm. Specimens were determined at species rank and recorded in a database. The nomenclature follows Manganelli et al. (1995) and subsequent updates. Voucher specimens were deposited in Manganelli collection, Dipartimento di Scienze Fische, della Terra e dell’Ambiente, Università di Siena (Italy).

162 **Explanatory variables**

163

164 Two groups of explanatory variables (environmental and spatial) were calculated for each site and used to model land
165 snail species richness and composition. The environmental properties of each site were quantified by three variables: 1)
166 distance from city centre (“*Dist_center*”) measured as linear distance from the city cathedral to the sampling site
167 (Horsák et al. 2009); 2) distance from sea (“*Dist_sea*”) assessed as minimum linear distance from the site to the nearest
168 point on the coastline; 3) vegetation cover (“*Tree_cover*”), ranging from 0 to 100% of total area of sampling site.
169 Spatial variables were XY geographical coordinates and all the spatial structures at different scales detected by PCNM.

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171 **Statistical analysis**

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173 Observed and estimated species richness

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175 α , β and γ diversity were used to assess richness and diversity patterns at habitat and city level. α diversity was defined
176 as species richness in each of the 24 sites and γ diversity as species richness in each urban habitat or city. β diversity,
177 concerning species composition turnover within and between habitats and cities, was examined by the Simpson
178 dissimilarity index (β_{sim} ; Baselga et al. 2007), which only considers dissimilarity due to spatial turnover, irrespective
179 of species richness (Baselga 2010). Analysis of Variance (ANOVA) and Bonferroni p-value adjustment for multiple
180 comparisons were used to test for statistically significant differences. The *Chao2* richness estimator and its standard
181 error *Chao.se* (Chao 1987) were also used to estimate the minimum asymptote of species richness for habitats and cities
182 in order to detect the number of unseen or undetected species in relation to observed species richness and sampling
183 efficiency.

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185 Environmental and spatial effects on land snail assemblages: PCNM and Variation Partitioning

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187 To evaluate the effects of environment and space on species composition, principal coordinate analysis of neighbour
188 matrices (PCNM, Borcard et al. 2004) was used to detect spatial structure in the data. Compared to traditional
189 multivariate trend-surface analysis, PCNM detects, decomposes and models any type of hidden spatial structure in the
190 data, within the limits of sampling design (Borcard and Legendre 2002). It makes it possible to select the best linear
191 combination of eigenvectors, maximizing correlation with the data and minimizing the number of vectors (Dray et al.
192 2006). We first computed a matrix of Euclidean distances among the 24 sites using the latitudes and longitudes of
193 sampling sites. Before running PCNM, the data was detrended by regressing all variables against the geographical
194 coordinates (X-Y), retaining residuals. The Euclidean distances were then truncated: PCNM is based on the assumption
195 that the only meaningful spatial signals for plots are those circumscribed by neighbourhood sites (Dray et al. 2006). The
196 truncation distance, which defines neighbourhood thresholds, is considered the maximum distance connecting all plots
197 based on a minimum spanning tree criterion (Rangel et al. 2006). The *give.thresh* function in the *spacemakerR* package
198 was used.

199 PCNM was performed on the modified (i.e. truncated) distance matrix to extract eigenvectors (PCNM variables) to
200 use as new spatial explanatory variables. Since PCNM are orthogonal, each PCNM variable represents an independent
201 spatial gradient. The higher the eigenvalue of a PCNM variable, the broader the spatial scale represented, while PCNM
202 variables with low eigenvalues only explain small fractions of the total spatial structure and are related to finer spatial

203 gradients (Duarte et al. 2012). The model selection procedure was based on multivariate extension of the AIC criterion
204 using the *ortho.AIC* function in the SpacemakeR package. Six PCNM vectors obtained from the X-Y coordinates of
205 each sampling site were then entered as explanatory variables for Redundancy Analysis (RDA) to account for the space
206 effect on the urban land snail matrix. To reveal the urban gradient effect after removing the effect of space (i.e. pure
207 spatial patterns that cannot be related to any measured variable + unmeasured spatially structured environmental
208 variation), we computed the relative contribution of the three environmental variables (distance from city centre,
209 distance from sea, % vegetation cover) by partial RDA (pRDA) that excluded spatial influence from the community
210 matrix.

211 Variation partitioning was later conducted to detect the fractions of total variance explained exclusively by the
212 environmental predictors [a], by environment and space [b], exclusively by spatial variables [c] and finally by residual
213 variation (Borcard et al. 1992; Peres-Neto et al. 2006), which can be ascribed to internal biological variation, survey
214 error, or variables not included in the analysis (Legendre and Legendre 1998). Quantification of the variance
215 components was based on the *varpart* function of the Vegan R package (Oksanen et al. 2016).

216 All multivariate statistical analysis based on Hellinger-transformed land snail presence/absence data was performed
217 with RStudio version 0.99.473 (RStudio Team 2015).

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220 **Results**

221

222 Observed and estimated species richness

223

224 Total land snail richness was 53 species (Table 1). Communities were dominated by *Cornu aspersum* and *Xerotracha*
225 *conspurcata* present in 19 and 18 out of 24 sites, respectively, while rare species such as *Howenwartiana howenwartii*,
226 *Marmorana serpentina*, *Cepaea nemoralis* and *Vitrea etrusca* occurred in 2 or 3 out of 24 sites. Some species are
227 ranked as LC (Least Concern) in the IUCN Red List, while only three are alien (the Australasian *Paralaoma servilis*,
228 the Sardo-Corsican *Marmorana serpentina* and the Mediterranean *Deroceras invadens*).

229 Numbers of land snail species found in sites ranged from 5 to 29 with a mean of 14 (Lososová et al. 2011: 0-21,
230 mean 7.5). The lowest mean number of land snail species per site (α diversity) was found in early successional sites (E-
231 SUC) and old city walls (WALLS) (9 and 13, respectively) while the highest mean α diversity was in wooded sites
232 (WOOD) and urban gardens and parks (PARK) (17 and 16, respectively). The total number of species found in each
233 habitat type (γ diversity) was lowest in early successional sites (E-SUC: 30) and increased in old city walls (WALLS:
234 31), reaching its highest values in wooded suburban areas (WOOD: 37) and parks (PARK: 38) (Fig. 2a). Across
235 habitats, γ diversity was correlated with mean α diversity ($r=0.94$, $p=0.05$). One-way ANOVA (df 3; F-value 2.793; P-
236 value 0.067) revealed no significant difference among habitat types. β diversity was highest in E-SUC, lowest in all the
237 other habitats. This agrees with the findings of Lososová et al. (2011) (Fig. 3a). One-way ANOVA (df 3; F-value 12.25;
238 P-value 2.87e-06) and Bonferroni p-value adjustment for multiple comparisons showed that E-SUC significantly
239 differed with respect to all the other habitats in β diversity values (E-SUC-PARK $p=9.5e-05$; E-SUC-WALLS $p=3.7e-05$;
240 E-SUC-WOOD $p=2.5e-05$). Pairwise β diversity values between habitat types were: WALLS-PARK 0.16;
241 WALLS-E-SUC 0.37; WALLS-WOOD 0.32; PARK-E-SUC 0.20; PARK-WOOD 0.19; E-SUC-WOOD 0.17.

242 The lowest mean α diversity in the cities was recorded in Grosseto (GR) and Arezzo (AR) (11 in both cases) while
243 the highest was recorded in Siena (SI) (19). The total number of species found in each city (γ diversity) was lowest in

244 Grosseto (29), higher in Arezzo (35) and highest in Siena (44) (Fig. 2b). Across cities, γ diversity was not correlated
245 with mean α diversity ($r=0.84$, $p>0.05$). After one-way ANOVA (df 2; F-value 6.439; P-value 0.007) and Bonferroni
246 correction for multiple comparisons, the SI-GR and SI-AR pairs proved to be significantly different in α diversity (SI-
247 GR $p=0.014$; SI-AR $p=0.018$). Arezzo showed the highest internal β diversity, while Grosseto and Siena showed similar
248 values (Fig. 3b). After one-way ANOVA (df 2; F-value 9.739; P-value 0.0002) and Bonferroni p-value adjustment, the
249 AR-GR and AR-SI pairs resulted to be significantly different in β diversity values (AR-GR $p=0.001$; AR-SI $p=0.001$).
250 Pairwise β diversity values between cities were: SI-GR 0.17; SI-AR 0.14; AR-GR 0.38.

251 The minimum estimated *Chao2* richness and its standard error *Chao.se* differed among habitat types and cities (Fig.
252 4a,b). However, a few species are missing: more or less all species were detected with a reasonable degree of certainty
253 through sampling effort.

254

255 Environmental and spatial effects on land snail assemblages: PCNM and Variation Partitioning

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257 Total species matrix variation (Table 2) can be portioned in four parts: (a) pure effects of space, (b) combined variation
258 i.e. spatially structured environmental variation, (c) pure effects of environment, (d) residual variation (Borcard et al.
259 1992). In our data matrix, the total contribution of spatial structures (a + b) accounted for 16% of community structure
260 whereas the variation attributed solely to pure spatial patterns, i.e. patterns that cannot be related to any measured
261 environmental variable, was 13% (a). Environmental factors (c + b) are invoked to explain a higher portion of species
262 matrix variation (22%) than explained by spatial structures. Furthermore, almost all of this rather high variation (c)
263 (19%) is ascribed to the pure effect of environment, i.e. species-environment relationship associated with local
264 environment, irrespective of space. The percentage of residual variation (d) was large: more than half the total variation
265 remained unexplained, while only 3% of the explained variation was shared (b) between the two factors. All the
266 variation components had a significant effect ($p<0.05$). Redundancy analysis (Fig. 5) detected significant effect in
267 shaping urban land snail composition for two out of three local environmental variables, namely “distance from the city
268 centre” and “vegetation cover”. *Papillifera papillaris* was clearly negatively associated with *Dist_center* and was the
269 most characteristic species of WALLS assemblages followed by *Mediterranea hydatina*, which was also negatively
270 associated with *Tree_cover*; *Hohenwartiana hohenwartii* was most linked to E-SUC habitat type, *Monacha*
271 *parumcincta* was clearly positively associated with *Dist_center* and *Tree_cover*, proving the most characteristic species
272 of WOOD assemblages, while *Deroceras invades* and *Cernuella virgata* were associated with PARK habitat type.

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275 Discussion

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277 This study is the first to consider the influence of environment and spatial factors on urban land snail communities in
278 southern Europe, and the insights it provides are unexpected. In terms of land snail total species richness, we recovered
279 53 species over our 100 km transect, compared to the 87 species recorded over a much larger transect (>1200 km) in
280 central Europe by Chytrý et al. (2012) and the 54 forest entities found in three well preserved forest areas of Tuscany
281 (Benocci et al. 2015). These numbers suggest that urban environments may actually offer a range of different habitat
282 and environmental conditions to land snails, thereby creating a sort of heterogeneous melting pot of unlimited
283 ecological situations. Perhaps, the many species found in the assemblage possess a wide variety of ecological
284 requirements, which even in urban environments allow them to find optimal conditions to adapt, flourish and spread,

285 just as happens in naturalized areas. The various results collected in this study support this general interpretation and we
286 discuss these results below in terms of our three main aims.

287

288 α , β and γ diversity

289

290 Concerning habitat γ diversity, we did not find a clear decreasing trend from more (WALLS and PARK) to less
291 urbanized habitats (E-SUC and WOOD), but rather a homogeneous pattern with just a dip in proximity to early
292 successional sites. These open sub-urban habitats are typically in city peripheries; many are highly disturbed, often
293 close to busy roads or industrial settlements, and function as ecological corridors (or barriers?) to the open countryside
294 or woods. On the other hand, city parks, public gardens and wooded patches proved to be genuine islands of
295 biodiversity in the urban matrix, showing the highest values of both α and γ diversity. This finding agrees with other
296 research from which it emerged that parks were the most diverse urban habitat for several groups of organisms
297 (Lososová et al. 2011). Pairwise species composition differences between habitats showed major β diversity values, the
298 highest difference being between WALLS and E-SUC habitats: PARK and WOOD shared a common more or less
299 forested matrix, whereas WALLS and E-SUC showed the strongest differences in terms of ecological conditions and
300 environmental heterogeneity. At city level, Siena had the richest α and γ diversity: it is the greenest of the three cities,
301 characterized by many well preserved wooded and green patches inside the urban core, also relatively less impacted by
302 vehicular traffic and the effects of urbanization. These results overall imply the great diversity of conditions in urban
303 environments, which is reflected by diversity patterns in the land snail assemblage.

304

305 Conservation status

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307 The checklist did not include species in danger of extinction or threatened; xenodiversity was apparently very low and
308 only three species (5.7% of the total) were true aliens: *Paralaoma servilis* is a worldwide litter- and humus-dwelling
309 invasive species, probably of Australasian origin (Manganelli et al. 2015); *Deroceras invadens*, probably of central
310 Mediterranean origin, is an invasive species worldwide (Reise et al. 2011) and *Marmorana serpentina*, first reported
311 from the city of Grosseto by the present research, occurs in Sardinia and Corsica and has been introduced into certain
312 Tuscan cities (Livorno, Pisa, Siena) (Fiorentino et al. 2009). As regards the remaining species, at least two (*Lehmannia*
313 *melitensis*, *Papillifera papillaris*) may also have been introduced by anthropochory. They are nearly always found in
314 habitats at the edge of urban areas or used by man at some time in the past. Finally, vast open and built-up areas have
315 probably facilitated colonization by potentially indigenous but originally less widespread synanthropic species, such as
316 the slug *Limacus flavus* and land snails *Xerotricha conspurcata*, *Eobania vermiculata* and *Cornu aspersum* (Manganelli
317 et al. 2015). Biotic homogenization therefore does not seem to have had remarkable ecological impact: the small
318 Tuscan cities still maintain considerable β diversity, both within and between study areas, despite the restricted
319 geographical distance gradient. Thus they differ from other cosmopolitan urban areas in Europe where the introduction
320 of invasive species, together with human-mediated biotic interchange, resulted in widespread floristic and faunistic
321 homogenization (La Sorte et al. 2007; Ferenc et al. 2014).

322

323 Factors affecting community structure

324

325 Our attempt to discover the processes underlying land snail community structure by partitioning the pure effects of
326 “environment” and that of unmeasured factors (dispersal, biotic interactions, unmeasured environmental variables) that
327 create spatial structure (called “space”), provided the most interesting findings. Environment and space can be equally
328 important in shaping communities but their function should always be considered in relation to the dispersal capacity of
329 the group in question (Heino 2013). The relative role of deterministic and stochastic aspects driving ecological
330 metacommunities has been examined in different organisms in natural environments (e.g. oribatid mites, Caruso et al.
331 2012; oribatid mites and collembola, Ingimarsdóttir et al. 2012; microbial communities, Caruso et al. 2011; several
332 groups of aquatic organisms, Padial et al. 2014, Rádková et al. 2014, Zhai et al. 2015) but few studies have concerned
333 urban areas (spiders, bees and birds, Sattler et al. 2010; subaerial cyanobacteria and algae, vascular plants and land
334 snails, Chytrý et al. 2012). According to general belief, high dispersers should be less influenced by spatial effects since
335 they move more readily to suitable habitat than low dispersers and species are consequently sorted according to their
336 ecological requirements (Martiny et al. 2006; Heino 2013). Conversely, weak dispersers are expected to show
337 significant spatial structuring as a direct consequence of their low mobility while their distribution is often due to
338 stochastic processes mediated by limited dispersal (Bell 2000; Hubbell 2001). Paradoxically, although land snails are
339 widely recognised as having low mobility (Baur and Baur 1993), they proved to be most significantly shaped by pure
340 and spatially independent environmental processes, which often turn out to be irrelevant since the environment is
341 usually spatially structured by various ecological processes (Legendre and Fortin 1989). In other words, after removing
342 the effect of space (i.e. pure spatial patterns that cannot be related to any measured variables + unmeasured spatially
343 structured environmental variation), urban gradient turned out to be the main factor modelling land snail community
344 structure.

345 Some exceptions to the rule were also found by Hájek et al. (2011), who described much stronger environmental
346 than spatial structure in bryophytes but not in high dispersers such as diatoms in Western Carpathian fens, and similarly
347 Astorga et al. (2012) confirmed a preponderant influence of niche-related factors rather than spatially limited dispersal
348 in bryophytes and macroinvertebrates in Finnish streams. Sattler et al. (2010) first took urban environments seriously
349 into account, observing higher environmental influence and weak spatial structure for spiders over a gradient of three
350 Swiss cities. According to Sattler et al. (2010), this combination with traditionally limited dispersers may be a feature
351 typical of urban areas, where high disturbance affecting the heterogeneous mosaic of urban habitat could inhibit
352 establishment of well-defined spatial structures, selecting only the species able to adapt and coexist with human
353 activities. Recent studies on land snail dispersal confirmed the possibility of long distance passive dispersal (Ožgo et al.
354 2016) so that human-mediated transport in urban environments may be the most reasonable explanation for the high
355 pure environmental constraint detected in urban land snail data: molluscs can be passively dispersed, but only if
356 environmental conditions are suitable do they establish and develop well defined assemblages. Diverse microhabitat
357 conditions and small-scale habitat structures are in fact necessary conditions for mollusc diversification, assuming a
358 suitable geological substratum (Cuttelod et al. 2011). The pure environmental contribution therefore suggests selection
359 of species well adapted to man-made environments, while other species show random distributions, probably as a
360 consequence of occasional dispersal events.

361 The hypothesis of stronger environmental effect was also confirmed by RDA analysis that showed four well
362 structured and differentiated species assemblages at habitat level, as demonstrated by 95% confidence ellipses (Fig. 5).

363 “Distance from the city centre” and “Vegetation cover” proved to be the environmental variables most significantly
364 involved in explaining variation in species composition. With the exception of parks and gardens within the city limits,
365 which can be considered genuine islands of richness and biodiversity inside the urban core (Dedov and Penev 2004),

366 the greater the distance from the urban centre, the higher the vegetation density. It is well known that forest cover
367 provides shelter for land snails under unfavourable conditions (Dedov and Penev 2004). This has strong implications
368 for species richness since the richest urban habitats are generally those with the highest tree and vegetation cover values
369 (PARK and WOOD in our study). At the same time different forest cover values work as ecological filters, selecting
370 species that depend on different environmental conditions. It is true that poor assemblages with few species are usually
371 more frequent close to city centres, due to the reduced frequency of natural habitat (Horsák et al. 2009), but although
372 poor in terms of richness, these completely anthropogenic assemblages often consist of surprisingly unique entities:
373 WALLS harbour extremely specialist species (e.g. *Marmorana serpentina*, *Papillifera papillaris*, *Papillifera solida*),
374 that shelter in rock clefts and cracks, instead of in litter and vegetation. Furthermore, subterranean species living in the
375 ground behind walls, such as *Mediterranean hydatina*, which proved to be typical species in WALLS assemblages, can
376 be included in this habitat.

377 The response of local land snail assemblages to urbanisation cannot be described as a gradual and linear decline in
378 species richness with proximity to the city centre (Horsák et al. 2009), but rather as a clear change in species
379 composition through selection/adaptation to different ecological conditions, presumably after passive human-mediated
380 dispersal.

381 This significant environmental effect, reflecting niche processes as well as environmental filtering, is an interesting
382 result since a strong spatially independent environmental constraint has never previously been detected for organisms
383 limited by low dispersal within the urban matrix. This finding suggests the effect of human-mediated transport in
384 enhancing the dispersal capacity of organisms normally considered to have low mobility. However, the snails would not
385 become established in the absence of suitable microhabitat conditions. In particular, two urban environmental variables
386 were the main constraints shaping urban land snail communities, namely vegetation cover and distance from the city
387 centre. The detection of a pure spatially independent environmental effect has important conservation implications
388 which go beyond the analysis of a single taxonomic group: irrespective of study area position, sampling in different
389 cities would presumably show similar species assemblages explained by exactly the same dataset of environmental
390 variables.

391

392

393 **Conclusion**

394

395 In conclusion, urban environments can show high land snail richness, like natural areas; species richness does not show
396 a clear inverse trend in relation to urbanization of habitats, but rather a homogeneous pattern. While parks prove to be
397 real islands of biodiversity inside the urban patchwork in terms of species composition, only urban environments such
398 as old city walls provide ideal environments for many specialist species with particular ecological requirements (i.e.
399 carbonate rocks, shaded walls rich in calcium and moist refuges).

400 Although the transect we studied was only 100 km, the gradient explained an higher amount of variation to that
401 obtained by Lososová et al. (2011) and Chytrý et al. (2012) with transects of more than 1200 km in central Europe. In
402 addition to demonstrate the wealth of diversity patterns offered by the Mediterranean area and especially the Italian
403 peninsula, this finding furnishes a proof of the necessity of combining traditional visual search with debris and leaf litter
404 collection. Indeed, leaf litter collections allowed quantification of small and micro-snails.

405 City study areas host rather heterogeneous species assemblages where biotic homogenization seems very low: only
406 three out of 53 species were true alien species (*Paralaoma servilis*, *Marmorana serpentina* and *Deroceras invadens*).

407 After detecting and removing the influence of spatial structures, the effect of urban gradient turned out to be the
408 principal component structuring urban land snail assemblages. In particular, “Distance from the city centre” and
409 “Vegetation cover” were the environmental variables that explained most of the variation in species composition. This
410 finding shows the importance of environmental factors, possibly because of a combination of niche and environmental
411 filtering at local level for organisms with low mobility, which are usually structured by spatial constraints. It also
412 suggests new urban scenarios with major ecological and conservation implications for other invertebrate groups: inside
413 this complex urban matrix, even intensively managed habitats could support particular assemblages as results of unique
414 interactions between species and environment at local scale. This provides guidance for managers to set priorities in
415 management strategies that include habitat conservation at local scale.
416

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601

602 **Table captions**

603 **Table 1** Land snail species in three cities (Grosseto, Siena, Arezzo) and total number of species in each urban area

604
605 **Table 2** Summary of variation partitioning constituents for urban land snails obtained by partial RDAs with environmental and
606 spatial data as predictors

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608

609 **Figure captions**

610 **Fig. 1** Map of study area and location of sampling sites (red dots) in each city

611

612 **Fig. 2** α diversity (species richness per site, box and whiskers) and γ diversity (total number of species, numbers outside box and
613 whiskers) in habitat types (a, on the left) and cities (b, on the right). Different letters denote significant differences between groups
614 (ANOVA with Bonferroni p-value adjustment, $p < 0.05$)

615

616 **Fig. 3** Box plots of Simpson β diversity in habitat types (a, on the left) and cities (b, on the right). Different letters denote significant
617 differences between groups (ANOVA with Bonferroni p-value adjustment, $p < 0.05$)

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619 **Fig. 4** Bar plots of γ diversity in habitat types (a, on the left) and cities (b, on the right) showing the minimum estimated richness
620 Chao2 and its standard error Chao.se

621

622 **Fig. 5** Ordination plots of RDA showing the pure effect of environmental variables (arrows in blue) shaping land snail communities.
623 Above the entire ordination diagram while below we zoomed in to focus on a section of the ordination plot. The species numbers (s1,
624 s2, ..., s53) match the numerical order in Table 1, while row numbers (row1, row2, ...row24) match the 24 sampling sites. Ellipses
625 show the 95% confidence intervals associated with each habitat type

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